

PURIFICATION
OF
LABORATORY CHEMICALS

Eighth Edition

Purification of Laboratory Chemicals

Eighth Edition

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AUSTRALIA



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DEDICATION

From the very first to all later editions of this book, I have received every encouragement, support, and assistance through discussions and proofreading from my devoted wife Dr Pauline Marjorie Armarego. Her efforts and contributions during the preparation of this eighth edition were very considerable. She was responsible for locating and inserting the DOI and PMID numbers after thousands of references to the original literature. In so doing she amended all errors in faulty references, checked that the science corresponded with the respective reference, and that the numbers indeed allowed the download of the respective original references. In addition to this she proofread the whole manuscript four times on a 24 inch screen on a 'word by word' basis, correcting the typos while keeping an 'eagle eye' on the formatting and the chemistry. Her efforts have saved me more than twelve months of hard work, which are deserving of co-authorship. However, her modesty would not allow me this, but finally conceded me the pleasure of dedicating this edition to her.

WLFA

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Preface to the Eighth Edition

THE INCENTIVE to write the eighth edition came from Ms Fiona Geraghty, Acquisitions Editor, S & T Books, Elsevier Limited who approached me to write an 8th edition of *Purification of Laboratory Chemicals* and proposed an increase in the size of the book by about 200 pages. I accepted because I had many ideas in mind that would improve the book's usefulness, and this was a good opportunity to carry them through. These ideas were conveyed to Ms Geraghty who solicited the views of a few reviewers. Their responses, as well as being uniformly favourable, were also very constructive.

The following propositions were thus implemented in this edition.

- (a) All Chapters and sections were updated, new information and new substances were inserted in keeping with the original theme of '*purification of commercially available substances*'. In several instances brief synthetic procedures were included because they gave insights into purifying these substances.
- (b) The empirical formulae and new physical data were inserted for each entry.
- (c) Almost all substances now have one or more references to the original literature with their corresponding DOI's (digital object identifiers), PMID (unique PubMed system identifiers), ISBN (International Standard Book Number), and ISSN (International Standard Serial Number) numbers. These will allow users to download the original publications directly from a laptop computer connected to the internet without leaving the laboratory or requiring a comprehensive library.
- (d) Chapter 3 of the 7th edition was deleted and replaced by sections on *Recent Advances in Physical and in Chemical Techniques, used in Purification* at the end of Chapters 1 and 2 respectively. The 'Miscellaneous Compounds' section in Chapter 6 (formerly Chapter 7) was renamed 'Physiologically Active Compounds' and the more important active compounds, and many new ones, were located here.
- (e) Every endeavour was made to state briefly the more common applications where possible, e.g. industrial, agricultural, pharmaceutical, physiological, medical, catalytic etc.
- (f) The General Index was made more comprehensive.
- (g) A 20% increase in the Chemical Abstracts Service Registry numbers (CASRNs), and consequently of new substances, was made, providing a total of over 8000 chemical and biochemical substances. It does not include those that were not allocated CASRNs, e.g. allotropic and polymorphic forms, polymer bound reagents, enzymes, some catalysts and some nanomaterials.

The large variety and number of examples of purification procedures described in this book should make it possible to devise appropriate methods for the purification of any number of new substances in hand.

I should like to make the following acknowledgements:

I am gratefully indebted to Professor Jill E. Gready (John Curtin School of Medical Research, ANU) for her continued encouragement and for strongly supporting my Visiting Fellowships in her research group over many years. I am also grateful to Professor David Tremethick (Head of the of Genome Biology Department, John Curtin School of Medical Research, ANU) for his continued encouragement and for supporting my Visiting Fellowship. I thank Professor Martin Banwell FAA (Research School of Chemistry, ANU) for pointing out to me that specific optical rotations are now designated without a degree sign. I have now deleted the degree sign from the thousands of specific rotations in this edition.

I am indebted to Professor Hiroyasu Taguchi (Department of Medical Chemistry, Molecular Neuroscience Research Centre, Shiga University of Medical Science, Japan) for his encouragement and support for this and previous editions of this book, and particularly for allowing me to include data on curcumin and its derivatives, and their use in the diagnostics of Alzheimer's disease. The continued help from Joe Papa BS MS (EXAXOL in Clearwater, Florida, USA) with the preparation and purification of several inorganic compounds, particularly of beryllium and cerium, and trace metal analyses of a large number of commercially available inorganic compounds included in this book, is much appreciated and gratefully acknowledged.

I wish to thank my wife Dr Pauline M. Armarego for her continued help (see Dedication), my daughter Dr Sarah H. Armarego FANZCA, CCP (Aust) (Anaesthetist and Perfusionist, Senior Staff Specialist) for providing and discussing with me information on many physiologically active substances and their medical aspects, and my grand daughter Ms Gemma M. M. Armarego (University student in Nursing and Paramedics) for providing me with a long list of drugs that she uses in her profession, many of which have been included in this book. Finally, I thank the ANU library and its staff, both on campus and at the ANU Print Repository, for their prompt and unfailing assistance.

W.L.F. Armarego September 2016

PREFACE

Preface to the First Edition

WE BELIEVE that a need exists for a book to help the chemist or biochemist who wishes to purify the reagents she or he uses. This need is emphasised by the previous lack of any satisfactory central source of references dealing with individual substances. Such a lack must undoubtedly have been a great deterrent to many busy research workers who have been left to decide whether to purify at all, to improvise possible methods, or to take a chance on finding, somewhere in the chemical literature, methods used by some previous investigators.

Although commercially available laboratory chemicals are usually satisfactory as supplied, for most purposes in scientific and technological work, it is also true that for many applications further purification is essential.

With this thought in mind, the present volume sets out, first, to tabulate methods, taken from the literature, for purifying some thousands of individual commercially available chemicals. To help in applying this information, two chapters describe the more common processes currently used for purification in chemical laboratories and give fuller details of new methods which appear likely to find increasing application for the same purpose. Finally, for dealing with substances not separately listed, a chapter is included setting out the usual methods for purifying specific classes of compounds.

To keep this book to a convenient size, and bearing in mind that its most likely users will be laboratory-trained, we have omitted manipulative details with which they can be assumed to be familiar, and also detailed theoretical discussion. Both are readily available elsewhere, for example in Vogel's very useful book **Practical Organic Chemistry** (Longmans, London, 3rd ed., 1956), or Fieser's **Experiments in Organic Chemistry** (Heath, Boston, 3rd ed., 1957).

For the same reason, only limited mention is made of the kinds of impurities likely to be present, and of the tests for detecting them. In many cases, this information can be obtained readily from existing monographs.

By its nature, the present treatment is not exhaustive, nor do we claim that any of the methods taken from the literature are the best possible. Nevertheless, we feel that the information contained in this book is likely to be helpful to a wide range of laboratory workers, including physical and inorganic chemists, research students, biochemists, and biologists. We hope that it will also be of use, although perhaps to only a limited extent, to experienced organic chemists.

We are grateful to Professor A. Albert and Dr D.J. Brown for helpful comments on the manuscript.

D.D.P., W.L.F.A. & D.R.P.
1966

Preface to the Second Edition

SINCE the publication of the first edition of this book, there have been major advances in purification procedures. Sensitive methods have been developed for the detection and elimination of progressively lower levels of impurities. Increasingly stringent requirements for reagent purity have gone hand-in-hand with developments in semiconductor technology, in the preparation of special alloys and in the isolation of highly biologically active substances. The need to eliminate trace impurities at the micro- and nanogram levels has placed greater emphasis on ultrapurification technique. To meet these demands the range of purities of laboratory chemicals has become correspondingly extended. Purification of individual chemicals thus depends more and more critically on the answers to two questions—Purification from what, and to what permissible level of contamination. Where these questions can be specifically answered, suitable methods of purification can usually be devised.

Several periodicals devoted to ultrapurification and separations have been started. These include "Progress in Separation and Purification" (vol. 1) Ed. E.S. Perry, Wiley-Interscience, New York, vols. 1-4, 1968-1971, and **Separation and Purification Methods**, Ed. E.S. Perry and C.J. van Oss, Marcel Dekker, New York, vol. 1, 1973. Nevertheless, there still remains a broad area in which a general improvement in the level of purity of many compounds can be achieved by applying more or less conventional procedures. The need for a convenient source of information on methods of purifying available laboratory chemicals was indicated by the continuing demand for copies of this book even though it had been out of print for several years.

We have sought to revise and update this volume, deleting sections that have become more familiar or less important, and incorporating more topical material. The number of compounds in Chapters 3 and 4 have been increased appreciably. Also, further details in purification and physical constants are given for many compounds that were listed in the first edition.

We take this opportunity to thank users of the first edition who pointed out errors and omissions, or otherwise suggested improvements or additional material that should be included. We are indebted to Mrs S. Schenk who emerged from retirement to type this manuscript.

D.D.P., W.L.F.A. & D.R.P.
1980

Preface to the Third Edition

THE CONTINUING demand for this monograph and the publisher's request that we prepare a new edition are an indication that **Purification of Laboratory Chemicals** fills a gap in many chemists' reference libraries and laboratory shelves. The present volume is an updated edition that contains significantly more detail than the previous editions, as well as an increase in the number of individual entries and a new chapter.

Additions have been made to Chapters 1 and 2 in order to include more recent developments in techniques (e.g. Schlenk-type, *cf* p. 10), and chromatographic methods and materials. Chapter 3 still remains the core of the book, and lists in alphabetical order relevant information on *ca* 4000 organic compounds. Chapter 4 gives a smaller listing of *ca* 750 inorganic and metal-organic substances, and makes a total increase of *ca* 13% of individual entries in these two chapters. Some additions have also been made to Chapter 5.

We are currently witnessing a major development in the use of physical methods for purifying large molecules and macromolecules, especially of biological origin. Considerable developments in molecular biology are apparent in techniques for the isolation and purification of key biochemicals and substances of high molecular weight. In many cases something approaching homogeneity has been achieved, as evidenced by electrophoresis, immunological and other independent criteria. We have consequently included a new section, Chapter 6, where we list upwards of 100 biological substances to illustrate their current methods of purification. In this chapter the details have been kept to a minimum, but the relevant references have been included.

The lists of individual entries in Chapters 3 and 4 range in length from single-line entries to *ca* one page or more for solvents such as acetonitrile, benzene, ethanol and methanol. Some entries include information such as likely contaminants and storage conditions. More data referring to physical properties have been inserted for most entries [i.e. melting and boiling points, refractive indexes, densities, specific optical rotations (where applicable) and UV absorption data]. Inclusion of molecular weights should be useful when deciding on the quantities of reagents needed to carry out relevant synthetic reactions, or preparing analytical solutions. The Chemical Abstracts registry numbers have also been inserted for almost all entries and should assist in the precise identification of the substances.

In the past ten years laboratory workers have become increasingly conscious of safety in the laboratory environment. We have therefore in three places in Chapter 1 (pp. 3 and 33, and bibliography p. 52) stressed more strongly the importance of safety in the laboratory. Also, where possible, in Chapters 3 and 4 we draw attention to the dangers involved with the manipulation of some hazardous substances.

The worldwide facilities for retrieving chemical information provided by the Chemical Abstract Service (CAS on-line) have made it a relatively easy matter to obtain CAS registry numbers of substances, and most of the numbers in this monograph were obtained *via* CAS on-line. We should point out that two other available useful files are CSCHEM and CSCORP, which provide, respectively, information on chemicals (and chemical products) and addresses and telephone numbers of the main branch offices of chemical suppliers.

The present edition has been produced on an IBM PC and a Laser Jet printer using the **Microsoft Word (4.0)** word-processing program with a set style sheet. This has allowed the use of a variety of fonts and font sizes which has made the presentation more attractive than in the previous edition. Also, by altering the format and increasing slightly the sizes of the pages, the length of the monograph has been reduced from 568 to 391 pages. The reduction in the number of pages has been achieved in spite of the increase of *ca* 15% of total text.

PREFACE

We extend our gratitude to the readers whose suggestions have helped to improve the monograph, and to those who have told us of their experiences with some of the purifications stated in the previous editions, and in particular with the hazards that they have encountered. We are deeply indebted to Dr M.D. Fenn for the several hours that he has spent on the terminal to provide us with a large number of CAS registry numbers.

This monograph could not have been produced without the expert assistance of Mr David Clarke who has spent many hours loading the necessary fonts in the computer, and for advising one of the authors (W.L.F.A.) on how to use them together with the idiosyncrasies of Microsoft Word.

D.D.P. & W.L.F.A.
1988

Preface to the Fourth Edition

THE AIMS of the first three editions, to provide purification procedures of commercially available chemicals and biochemicals from published literature data, are continued in this fourth edition. Since the third edition in 1988 the number of new chemicals and biochemicals that have been added to most chemical and biochemical catalogues have increased enormously. Accordingly there is a need to increase the number of entries with more recent useful reagents and chemical and biochemical intermediates. With this in mind, together with the need to reorganise and update general purification procedures, particularly in the area of biological macromolecules, as well as the time lapse since the previous publication, this fourth edition of **Purification of Laboratory Chemicals** has been produced. Chapter 1 has been reorganised with some updating, and by using a smaller font it was kept to a reasonable number of pages. Chapters 2 and 5 were similarly altered and have been combined into one chapter. Eight hundred and three hundred and fifty entries have been added to Chapters 3 (25% increase) and 4 (44% increase), respectively, and four hundred entries (310% increase) were added to Chapter 5 (Chapter 6 in the Third Edition), making a total of 5700 entries-all resulting in an increase from 391 to 529 pages, i.e., by ca 35%.

Many references to the original literature have been included remembering that some of the best references happened to be in the older literature. Every effort has been made to provide the best references, but this may not have been achieved in all cases. Standard abbreviations, listed on page 1, have been used throughout this edition to optimise space, except where no space advantage was achieved, in which cases the complete words have been written down to improve the flow of the sentences.

With the increasing facilities for information exchange, chemical, biochemical and equipment suppliers are making their catalogue information available on the Internet; e.g., Aldrich-Fluka-Sigma catalogue information is available on the World Wide Web by using the address <http://www.sigma.sial.com>, and GIBCO BRL catalogue information from <http://www.lifetech.com>, as well as on CD-ROMS which are regularly updated. Facility for enquiring about, ordering and paying for items is available *via* the Internet. CAS on-line can be accessed on the Internet, and CAS data is available now on CD-ROM. Also biosafety bill boards can similarly be obtained by sending SUBSCRIBE SAFETY John Doe at the address "listserv@uvmvm.uvm.edu", SUBSCRIBE BIOSAFETY at the address "listserv@mitvma.mit.edu", and SUBSCRIBE RADSAF at the address "listserv@romulus.ehs.uiuc.edu"; and the Occupational, Health and Safety information (Australia) is available at the address "<http://www.worksafe.gov.au/~wsa1>". Sigma-Aldrich provided Material Safety data sheets on CD-ROMs.

It is with much sadness that Dr Douglas D. Perrin was unable to participate in the preparation of the present edition due to illness. His contributions towards the previous editions have been substantial, and his drive and tenacity have been greatly missed.

The Third Edition was prepared on an IBM-PC, and the previous IBM files were converted into Macintosh files. These have now been reformatted on a Macintosh LC575 computer, and all further data to complete the Fourth Edition were added to these files. The text was printed with a Hewlett-Packard 4MV -600dpi Laser Jet printer, which gives a clearer resolution.

I thank my wife Dr Pauline M. Armarego, also an organic chemist, for the arduous and painstaking task of entering the new data into the respective files, and for the numerous hours of proofreading as well as the corrections of typographic errors in the files. I should be grateful to my readers for any comments, suggestions, amendments and criticisms which could, perhaps, be inserted in the second printing of this edition.

W.L.F. Armarego, 30 June 1996

Preface to the Fifth Edition

THE DEMAND for **Purification of Laboratory Chemicals** has not abated since the publication of the fourth edition as evidenced by the number of printings and the sales. The request by the Editor for a fifth edition offered an opportunity to increase the usefulness of this book for laboratory purposes. It is with deep regret that mention should be made that Dr Douglas D. Perrin had passed away soon after the fourth edition was published. His input in the first three editions was considerable, and his presence has been greatly missed. A fresh, new and young outlook was required in order to increase the utility of this book, and it is with great pleasure that Dr Christina L.L. Chai, a Reader in Chemistry and leader of a research group in organic and bio-organic chemistry, has agreed to coauthor this edition. The new features of the fifth edition have been detailed below.

Chapters 1 and 2 have been reorganised and updated in line with recent developments. A new chapter on the Future of Purification has been added. It outlines developments in syntheses on solid supports, combinatorial chemistry as well as the use of ionic liquids for chemical reactions and reactions in fluorous media. These technologies are becoming increasingly useful and popular, so much so that many future commercially available substances will most probably be prepared using these procedures. Consequently, knowledge of their basic principles will be helpful in many purification methods of the future.

Chapters 4, 5 and 6 (3, 4 and 5 in the 4th ed.) form the bulk of the book. The number of entries has been increased to include the purification of many recent commercially available reagents that have become more and more popular in the syntheses of organic, inorganic and bio-organic compounds. Several purification procedures for commonly used liquids, e.g., solvents, had been entered with excessive thoroughness, but in many cases the laboratory worker only requires a simple, rapid but effective purification procedure for immediate use. In such cases a **rapid purification** procedure has been inserted at the end of the respective entry, and should be satisfactory for most purposes. With the increased use of solid phase synthesis, even for small molecules, and the use of reagents on solid support (e.g., on polystyrene) for reactions in liquid media, compounds on solid support have become increasingly commercially available. These have been inserted at the end of the respective entry and have been listed in the General Index together with the above rapid purification entries.

A large number of substances are ionisable in aqueous solutions, and knowledge of their ionisation constants, stated as pK (pKa) values, can be of importance not only in their purification but also in their reactivity. Literature values of the pK's have been inserted for ionisable substances, and where values could not be found they were estimated (pK_{Est}). The estimates are usually so close to the true values as not to affect the purification process or the reactivity seriously. The book will thus be a good compilation of pK values for ionisable substances.

Almost all the entries in Chapters 4, 5 and 6 have CAS (Chemical Abstract Service) Registry Numbers to identify them, and these have been entered for each substance. Unlike chemical names which may have more than one synonymous name, there is only one CAS Registry Number for each substance (with only a few exceptions, e.g., where a substance may have another number before purification, or before determination of absolute configuration). To simplify the method for locating the purification of a substance, a CAS Registry Number Index with the respective page numbers has been included after the General Index at the end of the book. This will also provide the reader with a rapid way to see if the purification of a particular substance has been reported in the book. The brief General Index includes page references to procedures and equipment, page references to abbreviations of compounds, e.g., TRIS, as well as the names of substances for which a Registry Number was not found.

Website references for distributors of substances or/and of equipment have been included in the text. However, since these may be changed in the future we must rely on the suppliers to inform users of their change in website references.

We wish to thank readers who have provided advice, constructive criticism and new information for inclusion in this book. We should be grateful to our readers for any further comments, suggestions, amendments and criticisms which could, perhaps, be inserted in a second printing of this edition. In particular, we thank Professor Ken-chi Sugiura (Graduate School of Science, Tokyo Metropolitan University, Japan) who has provided us with information on the purification of several organic compounds from his own experiences, and Joe Papa BS MS (EXAXOL in Clearwater, Florida, USA) who has provided us not only with his experiences in the purification of many inorganic substances in this book, but also gave us his analytical results on the amounts of other metal impurities at various stages of purification of several salts. We thank them graciously for permission to include their reports in this work. We express our gratitude to Dr William B. Cowden for his generous advice on computer hardware and software over many years and for providing an Apple LaserWriter (16/600PS) which we used to produce the master copy of this book. We also extend our sincere thanks to Dr Bart Eschler for advice on computer hardware and software and for assistance in setting up the computers (iMac and eMac) used to produce this book.

We thank Dr Pauline M. Armarego for assistance in the painstaking task of entering data into respective files, for many hours of proofreading, correcting typographical errors and checking CAS Registry Numbers against their respective entries.

One of us (W.L.F.A) owes a debt of gratitude to Dr Desmond (Des) J. Brown of the Research School of Chemistry, ANU, for unfailing support and advice over several decades and for providing data that was difficult to acquire not only for this edition but also for the previous four editions of this book.

PREFACE

One of us (C.L.L.C) would especially like to thank her many research students (past and present) for their unwavering support, friendship and loyalty, which enabled her to achieve what she now has. She wishes also to thank her family for their love, and would particularly like to dedicate her contribution towards this book to the memory of her brother Andrew who had said that he should have been a scientist.

We thank Mrs. Joan Smith, librarian of the Research School of Chemistry, ANU, for her generous help in many library matters, which has made the tedious task of checking references more enduring.

W.L.F. Armarego & C.L.L. Chai
November 2002

Preface to the Sixth Edition

THERE IS a continuing demand for the **Purification of Laboratory Chemicals** book, to the extent that the 5th edition which was published in early 2003 was carefully translated into Chinese (ISBN 978-7-5025-94367) by Ying-Jie Lin, Wei Liu, Hui-Ping Wang, Xiao-Bo Sun, Qing-Shan Li and Jun-Gang Cao from Jilin University (People's Republic of China) in 2007. In response to the demand, it was timely to update the 5th edition to include the more recently developed purification procedures, as well as add to the list of compounds for purification. The latter comprise some commercially available compounds that have gained usefulness and popularity in the past few years.

The first two chapters have been updated, sections of current interest have been expanded and new sections added. Chapter 3 has been rewritten so that areas of work that have lost popularity have been reduced in size or deleted and sections on recent, and now commonly adopted, technologies have been inserted. Chapters 4, 5 and 6 are now completely reorganized, and each is subdivided into several sections which will make it easier for the reader to locate compounds of similar classification. Chapter 4 is subdivided into aliphatic, alicyclic, aromatic and heterocyclic compounds, Chapter 5 has been subdivided into inorganic and metal-organic compounds, and Chapter 6 has been subdivided into amino acids and peptides, proteins, enzymes, DNA and RNA, carotenoids, carbohydrates, steroids and a miscellaneous section which includes small biologically active substances such as antibiotics, coenzymes, co-factors, lipids, phospholipids, polynucleotides and vitamins. Some useful compounds that have been added recently to commercial catalogues have been included in these three chapters. A large number of derivatives of previous entries with their physical properties and purifications have been inserted together with extensive referencing to the original literature including *Beilstein* references. This resulted in an increase in size of the 5th edition, in text and number of compounds, by over 20%. The purifications of some 7400 substances are described. As in the 5th edition, substance entries are in alphabetical order within subsections and each substance is defined by its Chemical Abstracts Service (CAS) Registry Number. An index of these numbers with their respective page numbers at the end of the book will make it possible to locate the purification of a desired substance readily and to check if the substance is contained in the book. For this purpose we thank Rodney Armarego for setting up a *Macro* on the MacBook Pro computer used for collating the CAS Registry Numbers for the index. There is also a General Index of Contents.

Website references of distributors of substances and/or of equipment have been included in the text. However, since these may change in the future, users should check for current websites of suppliers. The bibliographies have been updated, and websites of a few publishers and book suppliers have been included. Several texts with publication dates older than fifteen years have been deleted except for a few very useful textbooks which are out of print and where recent editions have not been produced. In these cases it is usually possible to obtain used copies from good suppliers of old books, for which there are several websites, e.g. visit Google under "old books suppliers"; also visit websites such as <<http://www.abebooks.com>>, <<http://www.betterworld.com/usedbooks>>, <<http://www.booksandcollectibles.com.au/index>>, <<http://www.ebay.com.au>>. Further information for almost every entry in Chapters 4, 5 and 6 of the 6th edition can be obtained from the references to the original literature, which are cited under each entry together with their respective *Beilstein* reference(s).

We thank readers who have provided advice, constructive criticism and new information. We are grateful for any further comments, suggestions, amendments and criticisms which could, perhaps, be inserted in a second printing of this edition. We thank Joe Papa BS MS (EXAXOL in Clearwater, Florida, USA) in particular for sharing his experiences on the purification of several inorganic substances in this and previous editions, and also for allowing us to use his analytical results on the amounts of metal impurities at various stages of purification of several salts.

We thank Dr Pauline M. Armarego for assistance in the painstaking task of entering data into respective files, for many hours of proofreading, correcting typographical errors and checking CAS Registry Numbers against their respective entries.

One of us (W.L.F.A) owes a debt of gratitude to Dr Desmond (Des) J. Brown of the Research School of Chemistry, ANU, for unfailing support and advice over several decades and for providing data that was difficult to acquire not only for this edition but also for the previous five editions of this book.

One of us (C.L.L.C) would like to acknowledge the support and friendship of her many research staff and students (past and present at ANU and A*STAR). She especially thanks Drs Paul Huleatt, Paul Bernardo, Felicity Moore and Brendan Burkett for their unfailing faith in her, through chemical and personal journeys both in Singapore and Australia. The legacy of this book is for Kimberley and Victoria Tse because it is cool to be a scientist!

We thank Mrs Joan Smith, librarian of the Research School of Chemistry, ANU, for her generous help in many library matters which made the tedious task of checking references more enduring.

W.L.F. Armarego & C.L.L. Chai
November 2008

Preface to the Seventh Edition

The sales of the sixth edition, which appeared in April 2009, were high by about October 2009, and one of us (WLFA) was approached by Ms Melanie Benson, Editorial Project Manager of Elsevier Science & Technology Books (who was mainly responsible for the production of the 6th edition), about writing a 7th edition within 2-3 years. In the past, 6-9 years were allowed to lapse between editions. However, the attraction this time, was that we were allowed to increase the size of the work by up to 249 pages. This has given us the opportunity to update all the previous chapters in the light of current thinking on safety (personal and environmental), and to introduce two new chapters. The award of five Nobel Prizes in the past ten years or so in Chemistry and one in Physics, of which three were awarded for work on *Catalysis* and the *catalytic process* (2001, 2005 and 2010), and two for work on *Nanomaterials and Nanotechnology* (1996, 2010) have prompted us to write a new chapter on *Catalysts* and a new chapter on *Nanomaterials and Nanotechnology*. Chemical suppliers have now made commercially available a large number of catalysts as well as many nanomaterials of various sorts. Since the number of commercially available catalysts are currently considerably larger than that of nanomaterials, the chapter on Catalysts is larger than that on Nanomaterials and Nanotechnology, and had to be divided into two parts. The availability, preparation and purification of a large range of these, are presented in these chapters. The other chapters have been updated and expanded, also in keeping with the purpose of all previous editions which is to provide information for the purification of commercially available laboratory materials. Of course, the General Subjects Index and the Chemical Abstracts Registry Numbers (CASRNs) Index increased in size accordingly. Much of the cross referencing is done *via* CASRNs and a page of how to use this book through these is included before Chapter 1 to assist the reader, not only to locate the pages where the required CASRNs are to be found, but also to let the reader know whether a particular substance is included in this work. CASRNs can be readily obtained from chemical catalogues or from SciFinder.

We would like to acknowledge Professor Martin Banwell FAA (Director, Research School of Chemistry, ANU) for his generosity in allowing the use of IT services, and to Dr Emil Mittag (Research School of Chemistry, ANU) for editing and updating the *Macro* program used for making the CASRNs Index.

We are greatly indebted to many people who have made valuable and constructive comments, and indicated errors, in previous editions. The continued help from Joe Papa of BS MS (EXAXOL in Clearwater, Florida, USA) with the preparation and purification of several inorganic compounds, particularly beryllium and cerium, is appreciated and gratefully acknowledged.

One of us (WLFA) owes a debt of gratitude to Professor Jill E. Gready (John Curtin School of Medical Research, ANU) for her continued encouragement and for strongly supporting a Visiting Fellowship over a period of many years.

We thank Dr Pauline M. Armarego for assistance in the painstaking task of proofreading and correcting typographical errors as well as checking the General Index and the CASRNs Index.

We thank the ANU library and its staff, both on campus and at the ANU Print Repository, for their prompt and unfailing assistance.

W.L.F. Armarego & C.L.L. Chai
December 2011

HOW THIS BOOK SHOULD BE USED

Substances have been entered in their respective chapters, sections and subsections. In these sections, compounds are listed in alphabetical order according to their more commonly used names. Because compounds can be named in various ways, some alternate names, as well as the commonly used abbreviations and commercial synonyms, have been included in brackets after their entries. Sometimes it may be difficult to find a particular substance; in which case *it is advisable to obtain the page number of the entry from its Chemical Abstracts Registry Number in the 'CASRNs Index' at the end of the book.* This index is the **gateway** to the book. CASRNs of substances are readily obtained from 'SciFinder', or better, from any commercial catalogue that sells these compounds, as almost all of these have CASRNs inserted after the names of their products. Some substances that could be formally included in more than one section of the book are entered in the preferred section but are cross-referenced by inserting 'see CASRN' or just the 'CASRN'. Thus the cross-reference is its CASRN. The CASRNs Index provides the page numbers in bold/italic type. If the CASRN of a desired substance is *not* in the CASRN Index, or its full or abbreviated name is *not* in the General Index, then it will not be present in this book. CASRNs are unique for each substance and are internally consistent. They are set up according to a specific formula. Refer to the first page of the CASRNs Index to calculate the formula in order to check whether or not the number is a valid CAS number.

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WLFA

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WLFA

ABOUT THE AUTHOR

Wilf Armarego was a British Subject born in Alexandria, Egypt, on 23 April 1931, and was educated at British Boys' School (Alexandria), where he matriculated externally (University of London). In 1949 he enrolled as an external free auditor at Farouk 1st University (later, University of Alexandria). He attended some lectures and all practical classes while studying mostly at home. In June 1951 he obtained the *Intermediate BSc* qualification (external, University of London) in Chemistry, Physics and Biology, and two years later in June 1953 succeeded in the University of London External BSc Special Honours degree in Chemistry. In September 1953 he left Egypt for London. He began research training in stereochemistry at Bedford College (University of London) in November 1953 under the tutelage of Professor E. E. Turner FRS, and subsequently submitted a PhD thesis in December 1955. In September 1956 he went to Melbourne, Australia, and was appointed Research Officer at the new ICIANZ Central Research Laboratories where he worked on plant growth substances for 2 years. This was followed by a year as Senior Demonstrator in Organic Chemistry at the University of Melbourne. He joined the Department of Medical Chemistry, John Curtin School of Medical Research (JCSMR), ANU, in January 1960 where he worked until his retirement in December 1996. He was awarded a DSc of the University of London for his work in heterocyclic chemistry at the early age of 36 years. In 1972 he changed his research field to studies of pteridine requiring enzymes associated with a variety of inborn errors of metabolism in children. This involved work on enzyme purification, cloning, mutation, gene expression, kinetics and enzyme mechanisms. He became a naturalised Australian in 1969. Since retirement he has been a Visiting Fellow in the JCSMR. He has written four books,¹⁻⁴ in addition to over 140 refereed publications. Awarded FRSC (FRIC 1963), C.Chem, FRACI (1972), A.D. Olle Prize (1968) and Worldwide WHO'sWHO 2014 Professional of the Year, Representing Medical Research.

1. W.L.F. Armarego, *Fused Pyrimidines Part 1 – Quinazolines* **1967** 537pp (Wiley-Interscience: NY).
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CHAPTER 1

COMMON PHYSICAL TECHNIQUES USED IN PURIFICATION

INTRODUCTION

Purity is a matter of degree. Other than contaminants such as dust, paper fibres, wax, cork, etc., that may have been inadvertently introduced into the sample during manufacture, all commercially available chemical substances are in some measure impure. Any amounts of unreacted starting material, intermediates, by-products, isomers and related compounds may be present depending on the synthetic or isolation procedures used for preparing the substances. Inorganic reagents may deteriorate because of defective packaging (glued liners affected by sulfuric acid, zinc extracted from white rubber stoppers by ammonia), corrosion or prolonged storage. Organic molecules may undergo changes on storage. In extreme cases the container may be incorrectly labeled or, where compositions are given, they may be misleading or inaccurate for the proposed use. Where any doubt exists, it is usual to check for impurities by appropriate spot tests, or by recourse to tables of physical or spectral properties such as the extensive infrared and NMR libraries published by the Sigma Aldrich Chemical Co.

The important question, then, is not whether a substance is pure but whether a given sample is sufficiently pure for some intended purpose. That is, are the contaminants likely to interfere in the process or measurement that is to be studied. By suitable manipulation it is often possible to reduce levels of impurities to acceptable limits, but absolute purity is an ideal which, no matter how closely approached, can never be attained. A *negative* physical or chemical test indicates only that the amount of an impurity in a substance lies below a certain sensitivity level; no test can demonstrate that a likely impurity is entirely absent.

When setting out to purify a laboratory chemical, it is desirable that the starting material is of the best grade commercially available. Particularly among organic solvents there is a range of qualities varying from *laboratory chemical* to *spectroscopic* and *chromatographic* grades. Many of these are suitable for use as received. With the more common reagents it is usually possible to obtain from the current literature some indications of likely impurities, their probable concentrations and methods for detecting them. However, in many cases complete analyses are not given so that significant concentrations of unspecified impurities may be present.

THE QUESTION OF PURITY

Solvents and substances that are specified as *pure* for a particular purpose may, in fact, be quite impure for other uses. Absolute ethanol may contain traces of benzene, which makes it unsuitable for ultraviolet spectroscopy, or plasticisers which make it unsuitable for use in solvent extraction. See also the section on 'Criteria of Purity' in Chapter 2.

Irrespective of the grade of material to be purified, it is essential that some criteria exist for assessing the degree of purity of the final product. The methods used are preferably those that do not destroy the material, and if they do, denoted with a hash #, it is important that as little material is used as possible since it is not recoverable. The more common methods include:

1. Examination of physical properties such as:
 - (a) Melting point, freezing point, boiling point, and the freezing curve (i.e. the variation, with time, in the freezing point of a substance that is being slowly and continuously frozen).

- (b) Density at a specified temperature, and whether it is relative to the density of some other standard, e.g. water at 0.
 - (c) Refractive index at a specified temperature and wavelength. The sodium D line at 589.26 nm (weighted mean of the D₁ and D₂ lines) is the usual wavelength used but the refractive index values at other wavelengths can often be interpolated from a plot of refractive index versus $1/(\text{wavelength})^2$.
 - (d) Specific conductivity. (This can be used to detect, for example, water, salts, inorganic and organic acids and bases, in non-electrolytes).
 - (e) Optical rotation, optical rotatory dispersion and circular dichroism.
2. Empirical analysis, for C, H, N, S, metals, ash, etc.#
3. Chemical tests for particular types of impurities, e.g. for peroxides in aliphatic ethers (with acidified KI), or for water in solvents (quantitatively by the Karl Fischer method, see Fieser and Fieser, *Reagents for Organic Synthesis*, J. Wiley & Sons, NY, Vol 1 pp. 353, 528 1967, Library of Congress Catalog Card No 66-27894, also see Karl Fischer titrant or Hydranal –Titrant type 5E [64-17-5]. A number of other formulations for general and specific purposes, in aqueous and non-aqueous solutions, are commercially available [see Sigma-Aldrich Catalogue].
4. Physical tests for particular types of impurities:
- Emission and atomic absorption spectroscopy for detecting organic impurities and determining metal ions.
 - Chromatography, including paper, thin layer, liquid (high, medium and normal pressure), flash and vapour phase.
 - Electron spin resonance for detecting free radicals.
 - Other spectroscopic methods (see 5 below).
5. Examination of spectroscopic properties
- Nuclear Magnetic Resonance (¹H, ¹³C, ³¹P, ¹⁹F, ¹¹B NMR etc)
 - Infrared spectroscopy (IR, NIR and Fourier Transform IR)
 - Ultraviolet (UV), visible and fluorescence spectroscopy
 - X-ray photoelectron spectroscopy (XPS)
 - Atomic absorption spectroscopy (AAA)
 - Mass spectroscopy [electron ionisation (EI), chemical ionisation (CI), electrospray ionisation (ESI), fast atom bombardment (FAB), matrix-associated laser desorption ionisation (MALDI), inductively coupled plasma-mass spectrometry (ICP-MS), etc.#
6. Electrochemical methods (see 'Introduction' in Chapter 6 for macromolecules).
7. Nuclear methods which include a variety of radioactive elements as in organic reagents, complexes or salts.

A substance is usually taken to be of an acceptable purity when the measured property is unchanged by further treatment (especially if it agrees with a recorded value). In general, at least two different methods, such as recrystallisation and distillation, should be used in order to ensure maximum purity. Crystallisation may be repeated (from the same solvent or better from different solvents) until the substance has a constant melting point, and until it distils repeatedly within a narrow specified temperature range. The purified product should have spectroscopic properties which indicate that the traces of impurities left in the sample are of acceptable levels for the intended purpose.

With liquids, the refractive index at a specified temperature and wavelength is a sensitive test of purity. Note however that this is sensitive to dissolved gases such as O₂, N₂ or CO₂. Under favourable conditions, freezing curve studies are sensitive to impurity levels of as little as 0.001 moles percent. Analogous fusion curves or heat capacity measurements can be up to ten times as sensitive as this. With these exceptions, most of the above methods are rather insensitive, especially if the impurities and the substances in which they occur are chemically similar. In some cases, even an impurity comprising many parts per million of a sample may escape detection.

The common methods of purification, discussed below, comprise distillation (including fractional distillation, distillation under reduced pressure, sublimation and steam distillation), crystallisation, extraction, chromatographic, electrophoresis and other methods. In some cases, volatile and other impurities can be removed

simply by heating or application of a vacuum. Impurities can also sometimes be eliminated by the formation of derivatives from which the purified material is regenerated (see Chapter 2).

SOURCES OF IMPURITIES

Some of the more obvious sources of contamination of solvents arise from storage in metal drums and plastic containers, and from contact with grease and screw caps. Many solvents contain water. Others have traces of acidic materials such as hydrochloric acid in chloroform. In both cases this leads to corrosion of the drum and contamination of the solvent by traces of metal ions, especially Fe^{3+} . Grease, for example on stopcocks of separating funnels and other apparatus, e.g. greased ground joints, is also likely to contaminate solvents during extractions and chemical manipulation. Oxygen from the air is a source of contamination by virtue of its ability to produce small or large amounts of oxidation products (see section on the *Solubility of gases in liquids* below).

A much more general source of contamination that has not received the consideration it merits comes from the use of plastics for tubing and containers. Plasticisers can readily be extracted by organic solvents from PVC and other plastics, so that most solvents, irrespective of their grade (including spectrograde and ultrapure), have been reported to contain 0.1 to 5ppm of plasticiser [de Zeeuw et al. *Anal Biochem* **67** 339 1975, DOI: 10.1016/0003-2697(75)90303-6]. Where large quantities of solvent are used for extraction followed by evaporation, this can introduce significant amounts of impurity, even exceeding the weight of the genuine extract and giving rise to spurious peaks in gas chromatography, for example of fatty acid methyl esters [Pascaud, *Anal Biochem* **18** 570 1967, DOI: 10.1016/0003-2697(67)90116-9]. Likely contaminants are di(2-ethylhexyl)phthalate and dibutyl phthalate, but upwards of 20 different phthalate esters are listed as plasticisers as well as adipates, azelates, phosphates, epoxides, polyesters and various heterocyclic compounds. These plasticisers would enter the solvent during passage through plastic tubing or from storage in containers or from plastic coatings used in cap liners for bottles. Such contamination could arise at any point in the manufacture or distribution of a solvent. The problem with cap liners is avoidable by using corks wrapped in aluminium foil, although even in this case care should be taken because aluminium foil can dissolve in some liquids e.g. benzylamine and propionic acid. Polycarbonate containers invariably leach out the 'estrogenic chemical' **Bisphenol A** (see 'Aromatic Compounds' in Chapter 3) into the liquid in the container [Fiona Case *Chemistry World* **5** (No. 4) 12 2008, Rebecca Trager *Chemistry World* **5** (No. 5) 8 2008]. Also see issues of the SAGE journal *Toxicology and Industrial Health* first published in 1985. Solutions in contact with polyvinyl chloride can become contaminated with trace amounts of lead, titanium, tin, zinc, iron, magnesium or cadmium from additives used in the manufacture and moulding of PVC.

N-Phenyl-2-naphthylamine is a contaminant of solvents and biological materials that have been in contact with black rubber or neoprene (in which it is used as an antioxidant). Although this potential carcinogenic naphthylamine was only an artifact of the isolation procedures, it was at first thought to be a genuine component of vitamin K preparations, and of extracts of plant lipids, algae, butter, animal livers, eye tissue and kidney tissue [Brown *Chem Br* **3** 524 1967, Wang et al. *Cancer Res* **44** 3098 1984, PMID: 6327034].

Most of the above impurities can be removed by prior distillation of the solvent, and care should be taken to avoid further contact with plastic or black rubber materials. When using large volumes of solvents for extraction and chromatographic purification of substances it is imperative that the solvents are of the highest purity because impurities in them could severely contaminate the desired substance after solvent evaporation. This is particularly serious when working with small quantities of material (see above).

PRACTICES TO AVOID IMPURITIES

Cleaning practices

Laboratory glassware and Teflon equipment can be cleaned satisfactorily for most purposes by careful immersion into a solution of sodium dichromate in concentrated sulfuric acid, followed by draining, and rinsing copiously with distilled water. This is an exothermic reaction and should be carried out **very** cautiously in an efficient fume cupboard. [To prepare the **chromic acid bath**, dissolve 5 g of sodium dichromate (CARE: cancer suspect agent) in 5 ml of water. The dichromate solution is then cooled and stirred while 100 ml of concentrated sulfuric acid is added slowly. Store it in a glass bottle.] Where traces of chromium (adsorbed on the glass) must be avoided, a 1:1 mixture of concentrated sulfuric and nitric acid is a useful alternative. (*Use in a fumehood to remove vapour and with adequate face and body protection.*) Acid washing is also suitable for polyethylene ware, but prolonged contact (some weeks) leads to severe deterioration of the plastic. Alternatively, an alcoholic solution of sodium

hydroxide (*alkaline base bath*) can be used. This strongly corrosive solution (CAUTION: alkali causes serious burns) can be made by dissolving 120g of NaOH in 120 ml of water, followed by dilution to 1 L with 95% ethanol. This solution is conveniently stored in suitable alkali-resistant containers (e.g. Nalgene heavy duty rectangular tanks) with lids. Glassware can be soaked overnight in the base bath and rinsed thoroughly after soaking until the pH of the wash is close to 7. For much glassware, washing with hot detergent solution, using tap water, followed by rinsing well with distilled water and acetone, and heating at 200-300° overnight, is adequate. (Volumetric apparatus should not be heated: after washing it is rinsed with acetone, then pure diethyl ether, and air-dried. Prior to use, equipment can be rinsed with acetone, then with petroleum ether or pure diethyl ether, to remove the last traces of contaminants.) Teflon equipment should be soaked, first in acetone, then in petroleum ether or pure diethyl ether for ten minutes, then dried in a vacuum or flushed with dry nitrogen prior to use. For trace metal analyses, prolonged soaking of equipment in 1M nitric acid may be needed to remove adsorbed metal ions.

Soxhlet thimbles and filter papers may contain traces of lipid-like materials. For manipulations with highly pure materials, as in trace-pesticide analysis, thimbles and filter papers should be thoroughly extracted with the solvent used for the extractions, followed by pure diethyl ether before use.

Trace impurities in silica gel for TLC can be removed by heating at 300° for 16 hours or by Soxhlet extraction for 3 hours with distilled chloroform, followed by 4 hours extraction with distilled pure diethyl ether and drying in a vacuum.

Silylation of glassware and plasticware

Silylation of apparatus makes it repellant to water and hydrophilic materials. It minimises loss of solute by adsorption onto the walls of the container. The glassware is placed in a desiccator containing dichloromethyl silane (1ml) in a small beaker and evacuated for 5 minutes. The vacuum is turned off and air is introduced into the desiccator, which allows the silylating agent to coat the glassware uniformly. The desiccator is then evacuated, closed and set aside for 2 hours. The glassware is removed from the desiccator and baked at 180° for 2 hours before use.

Silylating mixtures for coating glassware, as well as for silylation reactions, can be readily prepared, e.g. (a) ~5% dimethyldichlorosilane [DMDCS, 75-78-5] in hexane, (b) ~2% dimethyldichlorosilane in 1,1,1-trichloroethane for coating micro electrodes, (c) 10% hexamethyldisilazane [HMDS, 999-97-3] and 6% trimethylchlorosilane [TMCS, 75-77-4] in 1-chloronaphthalene, (d) ~4% trimethylchlorosilane in *o*-xylene, (e) ~5% dimethyldichlorosilane in *o*-xylene, and (f) ~3% tributylchlorosilane [995-45-9] in 1-chloronaphthalene for coating micropipette electrodes by the dip-and-bake method. A more powerful general silylating mixture consists of *N*-tert-butyltrimethylsilyl-*N*-methyltrifluoroacetamide [MTBSTFA, 77377-52-7] containing 1% of *tert*-butyldimethylchlorosilane [TBDMSCl, 18162-48-6], which is also available commercially.

Plasticware is treated similarly except that it is rinsed well with water before use instead of baking. Note that dichloromethylsilane is highly **TOXIC** and **VOLATILE**, and the whole operation should be carried out in an efficient fume cupboard.

An alternative procedure used for large apparatus is to rinse the apparatus with a 5% solution of dichloromethyl silane in chloroform, followed by several rinses with water before baking the apparatus at 180°/2 hours (for glass) or drying in air (for plasticware). A solution of 2% w/v of dichloromethylsilane in octamethyl cyclooctasilane (Repel-silane ES, Lim et al. *J Nanomater* **2015** 1 2015, article ID 561586 DOI: [org/10.1155/2015/561586](https://doi.org/10.1155/2015/561586)) or octmethylcyclotetrasiloxane [556-67-2] is used to prevent the sticking of polyacrylamide gels, agarose gels and nucleic acids to glass surfaces and these chemicals are available commercially.

SAFETY PRECAUTIONS ASSOCIATED WITH THE PURIFICATION OF LABORATORY CHEMICALS

Although most of the manipulations involved in purifying laboratory chemicals are inherently safe, care is necessary if hazards are to be avoided in the chemical laboratory. In particular there are dangers inherent in the inhalation of vapours and absorption of liquids and low melting solids through the skin. In addition to the toxicity

of solvents there is also the risk of their flammability and the possibility of eye damage. Chemicals, particularly in admixture, may be explosive. Compounds may be carcinogenic or otherwise deleterious to health. Present-day chemical catalogues specifically indicate the particular dangerous properties of the individual chemicals they list, and these should be consulted whenever the use of commercially available chemicals is contemplated. Radioisotopic labeled compounds pose special problems of human exposure and of disposal of laboratory waste. Hazardous purchased chemicals are accompanied by detailed MSDS (Material Safety Data Sheets), which contain information regarding their toxicity, safety handling procedures and the necessary precautions to be taken. These should be read carefully and filed for future reference. In addition, chemical management systems such as Chem-ChemWatch, which include information and videos on hazards, handling and storage of chemicals, are commercially available. There are a number of websites which provide selected safety information: these include the Sigma-Aldrich website <www.sigmaaldrich.com> and other chemical websites, e.g. <www.ilpi.com/msds>.

The most common hazards are:

- (1) Explosions due to the presence of peroxides formed by aerial oxidation of ethers and tetrahydrofuran, decahydronaphthalene, acrylonitrile, styrene and related compounds.
- (2) Compounds with low flash points (below room temperature). Examples are acetaldehyde, acetone, acetonitrile, benzene, carbon disulfide, cyclohexane, diethyl ether, ethyl acetate and *n*-hexane.
- (3) Contact of oxidising agents (KMnO_4 , HClO_4 , H_2O_2) with organic liquids.
- (4) Toxic reactions with tissues (Me_2SO_4 , H_2CrO_4).

The laboratory should at least be well ventilated and safety glasses should be worn, particularly during distillations and manipulations carried out under reduced pressure or at elevated temperatures. With this in mind we have endeavoured to warn users of this book whenever greater than usual care is needed in handling chemicals. As a general rule, however, **all chemicals which users are unfamiliar with should be treated with extreme care and assumed to be highly flammable and toxic.** The safety of others in a laboratory should always be foremost in mind, with ample warning to others whenever a potentially hazardous operation is in progress. Also, unwanted solutions or solvents should never be disposed *via* the laboratory sink. The operator should be aware of the usual means for disposal of chemicals in her/his laboratories, and she/he should remove unwanted chemicals accordingly. **Organic liquids for disposal should be temporarily stored, as is practically possible, in respective containers. Avoid placing all organic liquids in the same container particularly if they contain small amounts of reagents which could react with each other. Halogenated waste solvents should be kept separate from other organic liquids. These containers should be stored in cupboards, preferably made of metal, and separated from containers of common laboratory solvents.**

Laboratory coats, disposable aprons, caps, sleeves, dust/mist respirators and foot protection, hearing protection as well as a variety of safety glasses, goggles, face and body shields should be used when the demand arises and are available commercially (see e.g. the Sigma-Aldrich Labware catalogue).

SOME HAZARDS OF CHEMICAL MANIPULATION IN PURIFICATION AND RECOVERY OF RESIDUES

Performing chemical manipulations calls for some practical knowledge if danger is to be avoided. However, with care, hazards can be kept to an acceptable minimum. A good general approach is to consider every operation as potentially perilous and then to adjust one's attitude as the operation proceeds. A few of the most common dangers are set out below. For a larger coverage of the following sections, and of the literature, the bibliography at the end of this chapter should be consulted.

Perchlorates and perchloric acid. At 160° perchloric acid is an exceedingly strong oxidising acid and a strong dehydrating agent. Organic perchlorates, such as methyl and ethyl perchlorates, are unstable and are violently explosive compounds. A number of heavy-metal perchlorates are extremely prone to explode. The use of anhydrous magnesium perchlorate, *Anhydron*, *Dehydrite*, as a drying agent for organic vapours is **not** recommended. Desiccators which contain this drying agent should be adequately shielded at all times, preferably in a metal basket, and kept in a cool place, i.e. **never** on a window sill where sunlight can fall on it.

No attempt should be made to purify perchlorates, except for ammonium, alkali metal and alkaline earth salts which, in water or aqueous alcoholic solutions are insensitive to heat or shock. Note that perchlorates react relatively slowly in aqueous organic solvents, but as the water is removed there is the increased possibility of an

explosion. Perchlorate salts, often used in non-aqueous solvents, are explosive in the presence of even small amounts of organic compounds when heated. Hence stringent care should be taken when purifying perchlorates, and direct flame and infrared lamps should be avoided. Tetra-alkylammonium perchlorates should be dried below 50° under vacuum (and protection). Only very small amounts of such materials should be prepared, and stored, at any one time.

Peroxides. These are formed by aerial oxidation or by autoxidation of a wide range of organic compounds, including diethyl ether, allyl ethyl ether, allyl phenyl ether, dibenzyl ether, benzyl butyl ether, *n*-butyl ether, *iso*-butyl ether, *t*-butyl ether, dioxane, tetrahydrofuran, olefins, and aromatic and saturated aliphatic hydrocarbons. They accumulate during distillation and can detonate violently on evaporation or distillation when their concentration becomes high. If peroxides are likely to be present materials should be tested for peroxides before distillation (for tests see entry under 'Ethers' in Chapter 2). Also, distillation should be discontinued when at least one quarter of the residue is left in the distilling flask.

Heavy-metal-containing explosives. Ammoniacal silver nitrate, on storage or treatment, will eventually deposit the highly explosive silver nitride *fulminating silver*. Silver nitrate and ethanol may give silver fulminate (see 'Inorganic Compounds' in Chapter 4), and in contact with azides or hydrazine and hydrazides may form silver azide. Mercury can also form such compounds. Similarly, ammonia or ammonium ions can react with gold salts to form '*fulminating gold*'. Metal fulminates of cadmium, copper, mercury and thallium are powerfully explosive, and some are detonators [Matyáš & Pachman *Fulminates in Primary Explosives* Springer Verlag, pp 37-70 2013, ISBN 978-3-642-28436-6_3, DOI: 10.1007/978-3-642-28436-6_3]. Heavy-metal-containing solutions, particularly when organic material is present, should be treated with great respect, and precautions towards possible explosion should be taken.

Strong acids. In addition to perchloric acid (see above), extra care should be taken when using strong mineral acids. Although the effects of concentrated sulfuric acid are well known, these cannot be stressed strongly enough. Contact with tissues will leave irreparable damage. **Always dilute the concentrated acid (Oil of Vitriol) by carefully adding the acid down the side of the flask which contains the water, and the process should be carried out under cooling. This solution is not safe to handle until the acid has been thoroughly mixed (care) with the water. Protective face, and body coverage should be used at all times.** Fuming sulfuric acid and chlorosulfonic acid are even more dangerous than concentrated sulfuric acid, and adequate precautions should be taken. Chromic acid cleaning mixture contains strong sulfuric acid and should be treated in the same way; and in addition this mixture is potentially *carcinogenic*.

Concentrated and fuming nitric acids are also dangerous because of their severe deleterious effects on tissues.

Picric acid. This acid and related nitro compounds, e.g. styphnic acid, are explosive and should **not** be allowed to dry. The acid is generally stored wet by covering the crystals with water. Solutions in ethanol and benzene are used occasionally. They should be stored in the cold (to minimise evaporation), and a rubber or plastic stopper (**not a ground glass stopper, because twisting the stopper could cause any dry substance on the side of the stopper to explode due to heat produced by friction**) should be used. **Note** that picric acid and picrates stain skin proteins in a yellow colour that is not readily washed off. This can be avoided by wearing rubber gloves.

Reactive halides and anhydrides. Substances like acid chlorides, low-molecular-weight anhydrides and some inorganic halides (e.g. PCl₃) can be **highly toxic and lachrymatory, affecting mucous membranes and lung tissues. Utmost care should be taken when working with these materials. Work should be carried out in a very efficient fume cupboard.**

Salts and organic esters of some inorganic acids. In addition to the dangers of perchlorate salts, other salts such as nitrates, azides, diazo salts, organic nitrates, organic azides and picrates (see above) can be hazardous, and due care should be taken when these are dried. Organic nitrites are dangerous as they affect the heart rate. Large quantities should never be prepared or stored for long periods.

Solvents. The flammability of low-boiling organic liquids cannot be emphasised strongly enough. These invariably have very low flash points and can ignite spontaneously. Special precautions against explosive flammability should be taken when recovering such liquids. Care should be taken with small volumes (*ca* 250ml) as well as large volumes (> 1L), and the location of all the fire extinguishers, and fire blankets, in the immediate

vicinity of the apparatus should be checked. Fire extinguishers should be operational and checked on a regular basis and the dates noted. The following flammable liquids (in alphabetical order) are common fire hazards in the laboratory: acetaldehyde, acetone, acrylonitrile, acetonitrile, benzene, carbon disulfide, cyclohexane, diethyl ether, ethyl acetate, hexane, low-boiling petroleum ether, tetrahydrofuran and toluene. Toluene should always be used in place of benzene wherever possible due to the potential *carcinogenic* effects of the liquid and vapour of the latter. The drying of flammable solvents with sodium or potassium metal and metal hydrides poses serious potential fire hazards, and adequate precautions should be stressed.

Safety is an issue that goes hand in hand with ‘Green Science’, and cannot be emphasised enough. It should be the first consideration when planning a scientific operation. Safety issues should never be considered a *nuisance*, or an aspect which hampers procedures. This is no longer a *local* or a *regional* issue, but is becoming more and more a *global* problem. It is true that it means more effort should be put in planning an operation, but this is something that the worker should get used to. It is totally *unacceptable* today, and certainly will be in the future, not to take **safety** very seriously.

Questions to be addressed include:

Could this procedure be carried out in a safer way with respect not only to the operator but also to those around her/him?

How does it affect the immediate as well as the extended environment around the operation?

Could the operation be performed in an alternative way that can better satisfy these criteria?

Safety now goes well beyond the laboratory and all should be consciously aware of the consequences of a potentially unsafe situation.

SAFETY DISCLAIMER

Experimental chemistry is a very dangerous occupation, and extreme care and adequate safety precautions should be taken at all times. Although we have stated the safety measures that have to be taken under specific entries, these are by no means exhaustive and some may have been unknowingly or accidentally omitted. The experimenter without prior knowledge or experience must seek further safety advice on reagents and procedures from experts in the field before undertaking the purification of any material. Chemical Suppliers should provide detailed safety documentation for every chemical they list. If for some reason this has not been provided, the purchaser should demand that the respective safety details be forwarded by the Suppliers. ***We take no responsibility whatsoever if any mishaps occur when using any of the procedures described in this book.***

METHODS OF PURIFICATION OF REAGENTS AND SOLVENTS

Many methods exist for the purification of reagents and solvents. A number of these methods are routinely used in synthetic as well as analytical chemistry and biochemistry. These techniques, outlined below, will be discussed in greater detail in the respective sections in this chapter. It is important to note that more than one method of purification may need to be implemented in order to obtain compounds of highest purity.

Common methods of purification are:

Solvent Extraction and Distribution

Distillation

Recrystallisation

Sublimation

Electrophoresis

Chromatography

For substances contaminated with water or solvents, drying with appropriate absorbents and desiccants may be sufficient.

SOLVENT EXTRACTION AND DISTRIBUTION

Extraction of a substance from suspension or solution into another solvent can sometimes be used as a purification process. Thus, organic substances can often be separated from inorganic impurities by shaking an aqueous solution

or suspension with suitable immiscible solvents such as benzene, carbon tetrachloride, chloroform, diethyl ether, diisopropyl ether or petroleum ether. After several such extractions, the combined organic phase is dried and the solvent is evaporated. Grease from the glass taps of conventional separating funnels is invariably soluble in the solvents used. Contamination with grease can be very troublesome particularly when the amounts of material to be extracted are very small. Instead, the glass taps should be lubricated with the extraction solvent; or better, the taps of the extraction funnels should be made of the more expensive material *Teflon*. Immiscible solvents suitable for extractions are given in Table 1. Addition of electrolytes (such as ammonium sulfate, calcium chloride or sodium chloride) to the aqueous phase helps to ensure that the organic layer separates cleanly and also decreases the extent of extraction of water into the latter. Emulsions can also be broken up by filtration (with suction) through Celite, or by adding a little diethyl ether, octyl alcohol or some other paraffinic alcohol. The main factor in selecting a suitable immiscible solvent is to find one in which the material to be extracted is readily soluble, whereas the substance from which it is being extracted is not. The same considerations apply irrespective of whether it is the substance being purified, or one of its contaminants, that is taken into the new phase. (The second of these processes is described as washing.)

Common examples of washing with aqueous solutions include the following:

Removal of acids from water-immiscible solvents by washing with aqueous alkali, sodium carbonate or sodium bicarbonate.

Removal of phenols from similar solutions by washing with aqueous alkali.

Removal of organic bases by washing with dilute hydrochloric or sulfuric acids.

Removal of unsaturated hydrocarbons, of alcohols and of ethers from saturated hydrocarbons or alkyl halides by washing with cold concentrated sulfuric acid.

This process can also be applied to purification of the substance if it is an acid, a phenol or a base, by extracting into the appropriate aqueous solution to form the salt which, after washing with pure solvent, is again converted to the free species and re-extracted. Paraffin hydrocarbons can be purified by extracting them with phenol (in which aromatic hydrocarbons are highly soluble) prior to fractional distillation.

For extraction of solid materials with a solvent, a *Soxhlet* extractor is commonly used. This technique is applied, for example, in the alcohol extraction of dyes to free them from insoluble contaminants such as sodium chloride or sodium sulfate.

Acids, bases and amphoteric substances can be purified by taking advantage of their ionisation constants (see below).

The recovery of some fifty more commonly used solvents from water, other solvents, residues etc. have been discussed, together with information on their behaviour before and after use, by I.M. Smallwood in the *Solvent Recovery Handbook*, Blackwood Science Publ Ltd, 2001, ISBN 9780632056477.

DISTILLATION

One of the most widely applicable and most commonly used methods of purification of liquids or low melting solids (especially of organic chemicals) is fractional distillation at atmospheric, or some lower, pressure. Almost without exception, this method can be assumed to be suitable for all organic liquids and most of the low-melting organic solids. For this reason it has been possible, e.g. in Chapter 3, to omit many procedures for purification of organic chemicals when only a simple fractional distillation is involved—the suitability of such a procedure is implied from the boiling point.

The boiling point of a liquid varies with the 'atmospheric' pressure to which it is exposed. A liquid boils when its vapour pressure is the same as the external pressure on its surface, its normal boiling point being the temperature at which its vapour pressure is equal to that of a standard atmosphere (760.000 mm Hg = 101.325 kPa). Lowering the external pressure lowers the boiling point. For most substances, boiling point and vapour pressure are related by an equation of the form,

$$\log p = A + B/(t + 273),$$

where p is the pressure in mmHg, t is in °C, and A and B are constants. Hence, if the boiling points at two different pressures are known, the boiling point at another pressure can be calculated from a simple plot of $\log p$ versus $1/(t + 273)$. For organic molecules that are not strongly associated, this equation can be written in the form,

$$\log p = 8.586 - 5.703 (T + 273)/(t + 273)$$

where T is the boiling point in °C at 760mm Hg. Tables 2A and 2B give computed boiling points over a range of pressures. Some examples illustrate its application. Ethyl acetoacetate, b 180° (with decomposition) at 760mm Hg has a predicted b of 79° at 16mm (where it is stable); the experimental value is 78°. Similarly 2,4-diaminotoluene, b 292° at 760mm, has a predicted b of 147° at 8mm; the experimental value is 148-150°. For self-associated molecules the predicted b are lower than the experimental values. Thus, glycerol, b 290° at 760mm, has a predicted b of 146° at 8mm: the experimental value is 182°.

Similarly an estimate of the boiling points of liquids at reduced pressure can be obtained using a nomogram (see Fig. 1).

For pressures near 760mm, the change in boiling point is given approximately by

$$\delta t = a(760 - p)(t + 273)$$

where $a = 0.00012$ for most substances, but $a = 0.00010$ for water, alcohols, carboxylic acids and other associated liquids, and $a = 0.00014$ for very low-boiling substances such as nitrogen or ammonia [Crafts *Chem Ber* **20** 709 1887, DOI: 10.1002/cber.188702001162]. When all the impurities are non-volatile, simple distillation is adequate purification. The observed boiling point remains almost constant and approximately equal to that of the pure material. Usually, however, some of the impurities are appreciably volatile, so that the boiling point progressively rises during the distillation because of the progressive enrichment of the higher-boiling components in the distillation flask. In such cases, separation is effected by fractional distillation using an efficient column. [For further reading see section on 'Variation of Boiling Points with Pressure' in *CRC—Handbook of Chemistry and Physics 96th Edition, 2015-2016*, William M. Haynes (Editor-in-Chief) CRC Press, Taylor & Francis Publishing Group, Boca Raton, Florida, USA, ISBN-10: 1482260964; ISBN-13: 978-1482260960.]

Techniques

The distillation apparatus consists basically of a distillation flask, usually fitted with a vertical fractionating column (which may be empty, or packed with suitable materials such as glass helices or stainless-steel wool) to which is attached a condenser leading to a receiving flask. The bulb of a thermometer projects into the vapour phase just below the region where the condenser joins the column. The distilling flask is heated so that its contents are steadily vaporised by boiling. The vapour passes up into the column where, initially, it condenses and runs back into the flask. The resulting heat transfer gradually warms the column so that there is a progressive movement of the vapour phase-liquid boundary up the column, with increasing enrichment of the more volatile component. Because of this fractionation, the vapour finally passing into the condenser (where it condenses and flows into the receiver) is commonly that of the lowest-boiling components in the system. The conditions apply until all of the low-boiling material has been distilled, whereupon distillation ceases until the column temperature is high enough to permit the next component to distil. This usually results in a temporary fall in the temperature indicated by the thermometer.

Distillation of liquid mixtures

The principles involved in fractional distillation of liquid mixtures are complex but can be seen by considering a system which approximately obeys *Raoult's law*. (This law states that the vapour pressure of a solution at any given temperature is the sum of the vapour pressures of each component multiplied by its mole fraction in the solution.) If two substances, A and B, having vapour pressures of 600mm Hg and 360mm Hg, respectively, were mixed in a molar ratio of 2:1 (i.e. 0.666:0.333 mole ratio), the mixture would have (ideally) a vapour pressure of 520mm Hg (i.e. $600 \times 0.666 + 360 \times 0.333$, or $399.6 + 119.88$ mm Hg) and the vapour phase would contain 77% ($399.6 \times 100/520$) of A and 23% ($119.88 \times 100/520$) of B. If this phase was now condensed, the new liquid phase would, therefore, be richer in the volatile component A. Similarly, the vapour in equilibrium with this phase is still further enriched in A. Each such liquid-vapour equilibrium constitutes a 'theoretical plate'. The efficiency of a fractionating column is commonly expressed as the number of such plates to which it corresponds in operation. Alternatively, this information may be given in the form of the height equivalent to a theoretical plate, or HETP. The number of theoretical plates and equilibria between liquids and vapours are affected by the factors listed to achieve maximum separation by fractional distillation in the section below on techniques.

In most cases, systems deviate to a greater or lesser extent from Raoult's law, and vapour pressures may be greater

or less than the values calculated. In extreme cases (e.g. azeotropes), vapour pressure-composition curves pass through maxima or minima, so that attempts at fractional distillation lead finally to the separation of a constant-boiling (azeotropic) mixture and one (but not both) of the pure species if either of the latter is present in excess.

Elevation of the boiling point by dissolved solids. Organic substances dissolved in organic solvents cause a rise in boiling point which is proportional to the concentration of the substance, and the extent of rise in temperature is characteristic of the solvent. The following equation applies for dilute solutions and non-associating substances:

$$\frac{M \Delta t}{c} = K$$

where M is the molecular weight of the solute, Δt is the elevation of boiling point in $^{\circ}\text{C}$, c is the concentration of solute in grams for 1000gm of solvent, and K is the *Ebullioscopic Constant* (molecular elevation of the boiling point) for the solvent. K is a fixed property (constant) for the particular solvent. This has been very useful for the determination of the molecular weights of organic substances in solution.

The efficiency of a distillation apparatus used for purification of liquids depends on the difference in boiling points of the pure material and its impurities. For example, if two components of an ideal mixture have vapour pressures in the ratio 2:1, it would be necessary to have a still with an efficiency of at least seven plates (giving an enrichment of $2^7 = 128$) if the concentration of the higher-boiling component in the distillate was to be reduced to less than 1% of its initial value. For a vapour pressure ratio of 5:1, three plates would achieve as much separation. In a fractional distillation, it is usual to reject the initial and final fractions, which are likely to be richer in the lower-boiling and higher-boiling impurities respectively. The centre fraction can be further purified by repeated fractional distillation.

To achieve maximum separation by fractional distillation:

1. The column must be flooded initially to wet the packing. For this reason it is customary to operate a still at reflux for some time before beginning the distillation.
2. The reflux ratio should be high (i.e. the ratio of drops of liquid which return to the distilling flask and the drops which distil over), so that the distillation proceeds slowly and with minimum disturbance of the equilibria along the column.
3. The hold-up of the column should not exceed one-tenth of the volume of any one component to be separated.
4. Heat loss from the column should be prevented, but if the column is heated to offset this, its temperature must not exceed that of the distillate in the column.
5. Heat input to the still-pot should remain constant.
6. For distillation under reduced pressure there must be careful control of the pressure to avoid flooding or cessation of reflux.

Types of distillation

The distilling flask. To minimise superheating of the liquid (due to the absence of minute air bubbles or other suitable nuclei for forming bubbles of vapour), and to prevent bumping, one or more of the following precautions should be taken:

(a) The flask is heated uniformly over a large part of its surface, either by using an electrical heating mantle or, by partial immersion in a bath above the boiling point of the liquid to be distilled (Table 3).

(b) Before heating begins, small pieces of unglazed fireclay or porcelain (porous pot, boiling chips), pumice, diatomaceous earth, or platinum wire are added to the flask. These act as sources of air bubbles.

(c) The flask may contain glass siphons or boiling tubes. The former are inverted J-shaped tubes, the end of the shorter arm being just above the surface of the liquid. The latter comprise long capillary tubes sealed above the lower end.

(d) A steady slow stream of inert gas (e.g. N₂, Ar or He) is passed through the liquid.

(e) The liquid in the flask is stirred mechanically. This is especially necessary when suspended insoluble material is present.

For simple distillations a Claisen flask is often used. This flask is, essentially, a round-bottomed flask to the neck of which is joined another neck carrying a side arm. This second neck is sometimes extended so as to form a Vigreux column [a glass tube in which have been made a number of pairs of indentations which almost touch each other and which slope slightly downwards. The pairs of indentations are arranged to form a spiral of glass inside the tube].

For heating baths, see Table 3. For distillation apparatus on a macro, semi-micro or micro scale see Aldrich and other glassware catalogues. Alternatively, visit some useful websites for suppliers of laboratory glassware, e.g. <www.wheatonsci.com>; <www.sigmaaldrich.com> and <www.kimble-kontes.com>.

Types of columns and packings. A slow distillation rate is necessary to ensure that equilibrium conditions operate and also that the vapour does not become superheated so that the temperature rises above the boiling point. Efficiency is improved if the column is heat insulated (either by vacuum jacketing or by lagging) and, if necessary, heated to just below the boiling point of the most volatile component. Efficiency of separation also improves with increase in the heat of vaporisation of the liquids concerned (because fractionation depends on heat equilibration at multiple liquid-gas boundaries). Water and alcohols are more easily purified by distillation for this reason.

Columns used in distillation vary in their shapes and types of packing. Packed columns are intended to give efficient separation by maintaining a large surface of contact between liquid and vapour. Efficiency of separation is further increased by operation under conditions approaching total reflux, i.e. under a high reflux ratio. However, great care must be taken to avoid flooding of the column during distillation. The minimum number of theoretical plates for satisfactory separation of two liquids differing in boiling point by δt is approximately $(273 + t)/3\delta t$, where t is the average boiling point in °C.

Some of the commonly used *columns* are:

Bruun column. A type of all-glass bubble-cap column.

Bubble-cap column. A type of plate column in which inverted cups (bubble caps) deflect ascending vapour through reflux liquid lying on each plate. Excess liquid from any plate overflows to the plate lying below it and ultimately returns to the flask. (For further details, see Bruun & West *Ind Eng Chem Anal Ed* **9** 247 1937, DOI: 10.1021/ac50109a023). Like most plate columns, it has a high through-put, but a relatively low number of theoretical plates for a given height.

Dufton column. A plain tube, into which fits closely (preferably ground to fit) a solid glass spiral wound round a central rod. It tends to choke at temperatures above 100° unless it is lagged (Dufton *J Soc Chem Ind* (London) **38** 45T 1919, DOI: 10.1002/jetb.5000380406).

Hempel column. A plain tube (fitted near the top with a side arm) which is almost filled with a suitable packing, which may be of rings or helices.

Oldershaw column. An all-glass perforated-plate column. The plates are sealed into a tube, each plate being equipped with a baffle to direct the flow of reflux liquid, and a raised outlet which maintains a definite liquid level on the plate and also serves as a drain on to the next lower plate [see Oldershaw *Ind Eng Chem (Anal Ed)* **11** 265 1941, DOI: 10.1021/i560092a026].

Podbielniak column. A plain tube containing 'Heli-Grid' Nichrome or Inconel wire packing. This packing provides a number of passage-ways for the reflux liquid, while the capillary spaces ensure very even spreading of the liquid, so that there is a very large area of contact between liquid and vapour while, at the same time, channelling and flooding are minimised. A column 1m high has been stated to have an efficiency of 200-400 theoretical plates (for further details, see Podbielniak *Ind Eng Chem (Anal Ed)* **13** 639 1941, DOI: 10.1021/i560097a020; Mitchell & O'Gorman *Anal Chem* **20** 315 1948, DOI: 10.1021/ac60016a012).

Stedman column. A plain tube containing a series of wire-gauze discs stamped into flat, truncated cones and welded together, alternatively base-to-base and edge-to-edge, with a flat disc across each base. Each cone has a hole, alternately arranged, near its base, vapour and liquid being brought into intimate contact on the gauze surfaces (Stedman *Can J Research B* **15** 383 1937, DOI: 10.1139/cjr37b-044).

Todd column. A column (which may be a Dufton type, fitted with a Monel metal rod and spiral, or a Hempel type, fitted with glass helices) which is surrounded by an open heating jacket so that the temperature can be adjusted to be close to the distillation temperature (Todd *Ind Eng Chem (Anal Ed)* **17** 175 1945, DOI: 10.1021/i560139a016).

Vigreux column. A glass tube in which have been made a number of pairs of indentations which almost touch each

other and which slope slightly downwards. The pairs of indentations are arranged to form a spiral of glass inside the tube.

Widmer column. A Dufton column, modified by enclosing within two concentric tubes the portion containing the glass spiral. Vapour passes up the outer tube and down the inner tube before entering the centre portion. Thus flooding of the column, especially at high temperatures, is greatly reduced (Widmer *Helv Chim Acta* **7** 59 1924, DOI: 10.1002/hlca.19240070107).

The packing of a column greatly increases the surface of liquid films in contact with the vapour phase, thereby increasing the efficiency of the column, but reducing its capacity (the quantities of vapour and liquid able to flow in opposite directions in a column without causing flooding). Material for packing should be of uniform size, symmetrical shape, and have a unit diameter less than one-eighth that of the column. (Rectification efficiency increases sharply as the size of the packing is reduced but so, also, does the hold-up in the column.) It should also be capable of uniform, reproducible packing.

The usual *packings* are:

(a) **Rings.** These may be hollow glass or porcelain (Raschig rings), of stainless steel gauze (Dixon rings), or hollow rings with a central partition (Lessing rings) which may be of porcelain, aluminium, copper or nickel.

(b) **Helices.** These may be of metal or glass (Fenske rings), the latter being used where resistance to chemical attack is important (e.g. in distilling acids, organic halides, some sulphur compounds, and phenols). Metal single-turn helices are available in aluminium, nickel or stainless steel. Glass helices are less efficient, because they cannot be tamped to ensure uniform packing.

(c) **Balls.** These are usually glass of uniform diameter, but can be made of stainless steel.

(d) **Wire packing.** For use of 'Heli-Grid' and 'Heli-Pak' packings. see references given for the Podbielniak column. For Stedman packing, see entry under Stedman column.

Types of condensers:

Air condenser. A glass tube such as the inner part of a Liebig condenser. Used for liquids with boiling points above 90°. Can be of any length.

Allihn condenser. The inner tube of a Liebig condenser is modified by having a series of bulbs to increase the condensing surface. Further modifications of the bubble shapes give the Julian and Allihn-Kronbitter (glass-jacketed) condensers.

Bailey-Walker condenser. A type of all-metal condenser fitting into the neck of extraction apparatus and being supported by the rim. Used for high-boiling liquids.

Coil condenser. An open tube, into which is sealed a glass coil or spiral through which water circulates. The tube is sometimes also surrounded by an outer cooling jacket.

Double surface condenser. A tube in which the vapour is condensed between an outer and inner water-cooled jacket after impinging on the latter. Very useful for liquids boiling below 40°.

Friedrichs condenser. A 'cold-finger' type of condenser sealed into a glass jacket open at the bottom and near the top. The cold finger is formed into glass screw threads.

Graham condenser. A type of coil condenser.

Hopkins condenser. A cold-finger type of condenser resembling that of Friedrichs.

Liebig condenser. An inner glass tube surrounded by a glass jacket through which water is circulated.

Othmer condenser. A large-capacity condenser which has two coils of relatively large bore glass tubing inside it, through which the water flows. The two coils join at their top and bottom.

West condenser. A Liebig condenser with a light-walled inner tube and a heavy-walled outer tube, with only a narrow space between them.

Wiley condenser. A condenser resembling the Bailey-Walker type.

[For more information see: [https://en.wikipedia.org/wiki/Condenser_\(laboratory\)](https://en.wikipedia.org/wiki/Condenser_(laboratory))]

Vacuum distillation. This expression is commonly used to denote a distillation under reduced pressure lower than that of the normal atmosphere. As the boiling point of a substance depends on the pressure, it is often possible to distil materials at a temperature low enough to avoid partial or complete decomposition by lowering the pressure, even if they are unstable when boiled at atmospheric pressure.

Sensitive or high-boiling liquids should invariably be distilled or fractionally distilled under reduced pressure. The apparatus is essentially as described for distillation except that ground joints connecting the different parts of the apparatus should be air tight by using grease, or better *Teflon sleeves*. For low, moderately high, and very high temperatures Apiezon L, M and T greases, respectively, are very satisfactory. Alternatively, it is often preferable to avoid grease and to use thin Teflon sleeves in the joints. The distillation flask, must be supplied with a capillary bleed (which allows a fine stream of air, nitrogen or argon into the flask), and the receiver should be of the fraction collector type. When distilling under vacuum it is very important to place a loose packing of glass wool above the liquid to buffer sudden boiling of the liquid. The flask should **not** be more than two-thirds full of liquid. The vacuum must have attained a steady state, i.e. the liquid has been completely degassed, before the heating

source is applied, and the temperature of the heat source must be raised *very slowly* until boiling is achieved.

If the pump is a filter pump off a high-pressure water supply, its performance will be limited by the temperature of the water because the vapour pressure of water at 10°, 15°, 20° and 25° is 9.2, 12.8, 17.5 and 23.8 mm Hg, respectively. The pressure can be measured with an ordinary manometer. For vacuums in the range of 10⁻² mm Hg to 10 mm Hg, rotary mechanical pumps (oil pumps) are used and the pressure can be measured with a Vacustat McLeod-type gauge. If still higher vacuums are required, for example for high vacuum sublimations, a mercury diffusion pump is suitable. Such a pump can provide a vacuum up to 10⁻⁶ mm Hg. For better efficiencies, the diffusion pump can be backed up by a mechanical pump. In all cases, the mercury pump is connected to the distillation apparatus through several traps to remove mercury vapours. These traps may operate by chemical action, for example the use of sodium hydroxide pellets to react with acid vapours, or by condensation, in which case empty tubes cooled in solid carbon dioxide-ethanol or liquid nitrogen (contained in wide-mouthed Dewar flasks) are used.

Special oil or mercury traps are available commercially, and a liquid-nitrogen (**b** -209.9°C) trap is the most satisfactory one to use between these and the apparatus. It has an advantage over liquid air or oxygen in that it is non-explosive if it becomes contaminated with organic matter. Air should **not** be sucked through the apparatus before starting a distillation because this will cause liquid oxygen (**b** -183°C) to condense in the liquid nitrogen trap, and this is potentially explosive (especially in mixtures with organic materials). Due to the potential lethal consequences of liquid oxygen/organic material mixtures, care must be exercised when handling liquid nitrogen. Hence, it is advisable to degas the system for a short period before the trap is immersed into the liquid nitrogen (which is kept in a Dewar flask).

Spinning-band distillation. Factors which limit the performance of distillation columns include the tendency to flood (which occurs when the returning liquid blocks the pathway taken by the vapour through the column) and the increased hold-up (which decreases the attainable efficiency) in the column that should, theoretically, be highly efficient. To overcome these difficulties, especially for distillation under high vacuum of heat sensitive or high-boiling highly viscous fluids, spinning band columns are commercially available. In such units, the distillation columns contain a rapidly rotating, motor-driven, spiral band, which may be of polymer-coated metal, stainless steel or platinum. The rapid rotation of the band in contact with the walls of the still gives intimate mixing of descending liquid with ascending vapour while the screw-like motion of the band drives the liquid towards the still-pot, helping to reduce hold-up. There is very little pressure drop in such a system, and very high throughputs are possible, with high efficiency. For example, a 765-mm long 10-mm diameter commercial spinning-band column is reported to have an efficiency of 28 plates and a pressure drop of 0.2 mm Hg for a throughput of 330ml/hour. The columns may be either vacuum jacketed or heated externally. The stills can be operated down to 10⁻⁵ mm Hg. The principle, which was first used commercially in the Podbielniak Centrifugal Superfractionator, has also been embodied in descending-film molecular distillation apparatus.

Steam distillation. When two immiscible liquids distil, the sum of their (independent) partial pressures is equal to the atmospheric pressure. Hence in steam distillation, the distillate has the composition

$$\frac{\text{Moles of substance}}{\text{Moles of water}} = \frac{P_{\text{substance}}}{P_{\text{water}}} = \frac{760 - P_{\text{water}}}{P_{\text{water}}}$$

where the *P*'s are vapour pressures (in mm Hg) in the boiling mixture.

The customary technique consists of heating the substance and water in a flask (to boiling), usually with the passage of steam, followed by condensation and separation of the aqueous and non-aqueous phases in the distillate. Its advantages are those of selectivity (because only some water-insoluble substances, such as naphthalene, nitrobenzene, phenol and aniline are volatile in steam) and of ability to distil certain high-boiling substances well below their boiling point. It also facilitates the recovery of a non-steam-volatile solid at a relatively low temperature from a high-boiling solvent such as nitrobenzene. The efficiency of steam distillation is increased if superheated steam is used (because the vapour pressure of the organic component is increased relative to water). In this case the flask containing the material is heated (without water) in an oil bath and the steam passing through it is superheated by prior passage through a suitable heating device (such as a copper coil heated electrically or an oil bath).

Azeotropic distillation. In some cases two or more liquids form constant-boiling mixtures, or azeotropes. Azeotropic mixtures are most likely to be found with components which readily form hydrogen bonds or are otherwise highly associated, especially when the components are dissimilar, for example an alcohol and an aromatic hydrocarbon, but have similar boiling points.

Examples where the boiling point of the distillate is a minimum (less than either pure component) include:

Water with ethanol, *n*-propanol and isopropanol, *tert*-butanol, propionic acid, butyric acid, pyridine,
methanol with methyl iodide, methyl acetate, chloroform,
ethanol with ethyl iodide, ethyl acetate, chloroform, benzene, toluene, methyl ethyl ketone,
benzene with cyclohexane,
acetic acid with toluene.

Although less common, azeotropic mixtures are known which have higher boiling points than their components. These include water with most of the mineral acids (hydrofluoric, hydrochloric, hydrobromic, perchloric, nitric and sulfuric) and formic acid. Other examples are acetic acid-pyridine, acetone-chloroform, aniline-phenol, and chloroform-methyl acetate.

The following azeotropes are important commercially for drying ethanol:

ethanol 95.5% (by weight) - water 4.5%	b 78.1°
ethanol 32.4% - benzene 67.6%	b 68.2°
ethanol 18.5% - benzene 74.1% - water 7.4%	b 64.9°

Materials are sometimes added to form an azeotropic mixture with the substance to be purified. Because the azeotrope boils at a different temperature, this facilitates separation from substances distilling in the same range as the pure material. (Conversely, the impurity might form the azeotrope and be removed in this way.) This method is often convenient, especially where the impurities are isomers or are otherwise closely related to the desired substance. Formation of low-boiling azeotropes also facilitates distillation.

One or more of the following methods can generally be used for separating the components of an azeotropic mixture:

1. By using a chemical method to remove most of one species prior to distillation. (For example, water can be removed by suitable drying agents; aromatic and unsaturated hydrocarbons can be removed by sulfonation).
2. By redistillation with an additional substance which can form a ternary azeotropic mixture (as in the ethanol-water-benzene example given above).
3. By selective adsorption of one of the components. (For example, of water on to silica gel or molecular sieves, or of unsaturated hydrocarbons onto alumina).
4. By fractional crystallisation of the mixture, either by direct freezing or by dissolving in a suitable solvent.

Kügelrohr distillation. The Aldrich Kügelrohr Distillation Apparatus (see Aldrich-Sigma Labware catalogue) is made up of small glass bulbs (*ca* 4-5cm diameter) that are joined together *via* Quickfit joints at each pole of the bulbs. The liquid (or low melting solid) to be purified is placed in the first bulb of a series of bulbs joined end to end, and the system can be evacuated. The first bulb is heated in a furnace (e.g. Büchi Kügelrohr micro distillation oven from Sigma-Aldrich Labware catalogue) at a high temperature whereby most of the material distils into the second bulb (which is outside of the furnace). The second bulb is then moved into the furnace and the furnace temperature is reduced by *ca* 5° whereby the liquid in the second bulb distils into the third bulb (at this stage the first bulb is now out at the back of the furnace, and the third and subsequent bulbs are outside the front of the furnace). The furnace temperature is lowered by a further *ca* 5°, and the third bulb is moved into the furnace. The lower boiling material will distil into the fourth bulb. The process is continued until no more material distils into the subsequent bulb. The vacuum (if applied) and the furnace are removed, the bulbs are separated and the various fractions of distillates are collected from the individual bulbs. For volatile liquids, it may be necessary to cool the receiving bulb with solid CO₂ held in a suitable container (a Kügelrohr distillation apparatus with an integrated cooling system is available). This procedure is used for preliminary purification and the distillates are then redistilled or recrystallised.

Isopiestic or isothermal distillation. This technique can be useful for the preparation of metal-free solutions of volatile acids and bases for use in trace metal studies. The procedure involves placing two beakers, one of distilled water and the other of a solution of the material to be purified, in a desiccator. The desic-

cator is sealed and left to stand at room temperature for several days. The volatile components distribute themselves between the two beakers whereas the non-volatile contaminants remain in the original beaker. This technique has afforded metal-free pure solutions of ammonia, hydrochloric acid and hydrogen fluoride.

RECRYSTALLISATION

Techniques

The most commonly used procedure for the purification of a solid material by recrystallisation from a solution involves the following steps:

- (a) The impure material is dissolved in a suitable solvent, by shaking or vigorous stirring, at or near the boiling point, to form a near-saturated solution.
- (b) The hot solution is filtered to remove any insoluble particles. To prevent crystallisation during this filtration, a heated filter funnel can be used, or the solution can be diluted with more of the solvent.
- (c) The solution is then allowed to cool so that the dissolved substance crystallises out.
- (d) The crystals are separated from the mother liquor, either by centrifuging or by filtering, under suction, through a sintered glass, a Hirsch or a Büchner, funnel. Usually, centrifugation is preferred because of the greater ease and efficiency of separating crystals and mother liquor, and also because of the saving of time and effort, particularly when very small crystals are formed or when there is entrainment of solvent.
- (e) The crystals are washed free from mother liquor with a little fresh cold solvent, then dried.

If the solution contains extraneous coloured material likely to contaminate the crystals, this can often be removed by adding some activated charcoal (decolourising carbon) to the hot, but not boiling, solution which is then shaken frequently for several minutes before being filtered. (The large active surface of the carbon makes it a good adsorbent for this purpose.) In general, the cooling and crystallisation steps should be rapid so as to give small crystals which occlude less of the mother liquor. This is usually satisfactory with inorganic material, so that commonly the filtrate is cooled in an ice-water bath while being vigorously stirred. In many cases, however, organic molecules crystallise much more slowly, so that the filtrate must be set aside to cool to room temperature or left in the refrigerator. It is often desirable to subject material that is very impure to preliminary purification, such as steam distillation, Soxhlet extraction, or sublimation, before recrystallising it. A greater degree of purity is also to be expected if the crystallisation process is repeated several times, especially if different solvents are used. The advantage of several crystallisations from different solvents lies in the fact that the material sought, and its impurities, are unlikely to have similar solubilities because solvents and temperatures are varied.

For the final separation of solid material, sintered-glass discs are preferable to filter paper. Sintered glass is unaffected by strongly acidic solutions or by oxidising agents. Also, with filter paper, cellulose fibres are likely to become included in the sample. The sintered-glass discs or funnels can be readily cleaned by washing in freshly prepared *chromic acid cleaning mixture*. This mixture is made by adding 100ml of concentrated sulfuric acid slowly with stirring to a solution of 5g of sodium dichromate (CARE: cancer suspect) in 5ml of water. (The mixture warms to about 70°, and sulfuric acid becomes hot when water is added to it; see p 3).

For materials with very low melting points it is sometimes convenient to use dilute solutions in acetone, methanol, pentane, diethyl ether or CHCl₃/CCl₄. The solutions are cooled to -78° in a dry-ice/acetone bath, to give a slurry which is filtered off through a precooled Büchner funnel. Experimental details, as applied to the purification of nitromethane, are given by Parrett and Sun [*J Chem Educ* **54** 448 1977, DOI: 10.1021/ed054p448].

Where substances vary little in solubility with temperature, *isothermal crystallisation* may sometimes be employed. This usually takes the form of a partial evaporation of a saturated solution at room temperature by leaving it under reduced pressure in a desiccator.

However, in rare cases, crystallisation is not a satisfactory method of purification, especially if the impurity forms crystals that are *isomorphous* with the material being purified. In fact, the impurity content may even be greater in such recrystallised material. For this reason, it still remains necessary to test for impurities and to remove or adequately lower their concentrations by suitable chemical manipulation prior to recrystallisation.

Filtration. Filtration removes particulate impurities rapidly from liquids and is also used to collect insoluble or crystalline solids which separate or crystallise from solution. The usual technique is to pass the solution, cold or hot, through a fluted filter paper in a conical glass funnel.

If a solution is hot and needs to be filtered rapidly, a Büchner funnel and flask are used and filtration is performed

under a slight vacuum (water pump), the filter medium being a circular cellulose filter paper wet with solvent. If filtration is slow, even under high vacuum, a pile of about twenty filter papers, wet as before, are placed in the Büchner funnel and, as the flow of solution slows down, the upper layers of the filter paper are progressively removed. *Alternatively*, a filter aid, e.g. Celite, Florisil or Hyflo-supercel, is placed on top of a filter paper in the funnel. When the flow of the solution (under suction) slows down, the upper surface of the filter aid is scratched gently. Filter papers with various pore sizes are available covering a range of filtration rates. Hardened filter papers are slow filtering, but they can withstand acidic and alkaline solutions without appreciable hydrolysis of the cellulose (see Table 4). When using strong acids it is preferable to use glass micro fibre filters, which are commercially available (see Tables 4 and 5).

Freeing a solution from extremely small particles [e.g. for optical rotatory dispersion (ORD) or circular dichroism (CD) measurements] requires filters with very small pore size. Commercially available (Millipore, Gelman, Nucleopore) filters other than cellulose or glass include nylon, Teflon, and polyvinyl chloride, and the pore diameter may be as small as 0.01micron (see Table 5). Special containers are used to hold the filters, through which the solution is pressed by applying pressure, e.g. from a syringe. Some of these filters can be used to clear strong sulfuric acid solutions (and remove bacteria from contaminated water).

As an alternative to the Büchner funnel for collecting crystalline solids, a funnel with a sintered glass-plate under suction may be used. Sintered-glass funnels with various porosities are commercially available and can be easily cleaned with warm chromic or nitric acid (see above).

When the solid particles are too fine to be collected on a filter funnel because filtration is extremely slow, separation by **centrifugation** should be used. Bench-type centrifuges are most convenient for this purpose. The solid is placed in the centrifuge tube, the tubes containing the solutions on opposite sides of the rotor should be balanced accurately (at least within 0.05 to 0.1g), and the solutions are spun at maximum speed for as long as it takes to settle the solid (usually *ca* 3-5minutes). The solid is washed (by shaking) with cold solvent by centrifugation, and finally twice with a pure volatile solvent in which the solid is insoluble, also by centrifugation. After decanting the supernatant, the residue is dried in a vacuum, at elevated temperatures if necessary. In order to avoid 'spitting' and contamination with dust while the solid in the centrifuge tube is dried, the mouth of the tube is covered with aluminium foil and held fast with a tight rubber band near the lip. The flat surface of the aluminium foil is then perforated in several places with a pin, and the tube and contents are dried in a vacuum desiccator over a desiccant.

Solvents

Choice of solvents. The best solvents for recrystallisation have the following properties:

- (a) The material is much more soluble at higher temperatures than it is at room temperature or below.
- (b) Well-formed (but not large) crystals are produced.
- (c) Impurities are either very soluble or only sparingly soluble.
- (d) The solvent must be readily removed from the purified material.
- (e) There must be no reaction between the solvent and the substance being purified.
- (f) The solvent must not be inconveniently volatile or too highly flammable. (These are reasons why diethyl ether and carbon disulfide are not commonly used in this way.)

The following generalisations provide a rough guide to the selection of a suitable solvent:

- (a) Substances usually dissolve best in solvents to which they are most closely related in chemical and physical characteristics. Thus, hydroxylic compounds are likely to be most soluble in water, methanol, ethanol, acetic acid or acetone. Similarly, petroleum ether might be used with water-insoluble substances. However, if the resemblance is too close, solubilities may become excessive.
- (b) Higher members of homologous series approximate more and more closely to their parent hydrocarbon.
- (c) Polar substances are more soluble in polar than in non-polar solvents.

Although Chapters 3 to 7 provide details of the solvents used for recrystallising a large portion of commercially available laboratory chemicals, they cannot hope to be exhaustive, nor are they necessarily the best choice, but they are the solvents reported in the literature. In other cases where it is desirable to use this process, it is necess-

ary to establish whether a given solvent is suitable. This is usually done by taking only a small amount of material in a small test-tube and adding enough solvent to cover it. If it dissolves readily in the cold or on gentle warming, the solvent is unsuitable. Conversely, if it remains insoluble when the solvent is heated to boiling (adding more solvent if necessary), the solvent is again unsuitable. If the material dissolves in the hot solvent but does not crystallise readily within several minutes of cooling in an ice-salt mixture, another solvent should be tried.

Water

The properties and purification of water are described in the 'Inorganic Compounds' section of Chapter 4. Fluka (Riedel-de Haën) supply purified water prepared specifically for a variety of uses, e.g. LC-MS, HPLC, gradient elution, for cell biology which is freed from *enterotoxins* by *ultrafiltration* and autoclaving, for organic and for inorganic trace analysis, for residue analysis and other analytical purposes. Some of these have been prepared by *reverse osmosis*, or ultrafiltration, under clean room conditions and filtered through 0.2 μm membranes into bottles of high purity glass under inert gas. They have a limited shelf life once opened most probably because O_2 from the air dissolves readily in the water. The solubility of O_2 in 100ml of water is $\sim 1.02\text{ml}$ (0.455mM) at 0° , 0.68ml (0.282mM) at 20° , 0.63ml (0.258mM) at 25° , 0.63ml (0.237mM) at 30° , and 0.12ml (0.033mM) at 100° , all at $\sim 760\text{mmHg}$ in equilibrium with air (see Tables 25-28). This is in comparison with the concentration of O_2 of 0.23mM in 0.1M Tris HCl buffer at pH 7.2 and 25° in equilibrium with air at 760mmHg. Routinely, water is best purified by redistilling it twice in an all glass apparatus, storing it under N_2 or He in stoppered glass containers and, if necessary, preferably subjected to ultrafiltration through a single or multistage 0.2 μm membrane system or to reverse osmosis (visit <www.millipore.com>). If oxygen-free water is required, N_2 or argon should be bubbled through a sintered glass frit in the highly purified water for 2-3hours, and stoppered immediately. It is best to use a glass container from which the water can be withdrawn without it coming into contact with air. Note that boiling and distilling water, and condensing it in an inert atmosphere should de-gas it.

Petroleum ethers are commercially available fractions of refined petroleum and are sold in fractions of about 20° boiling ranges. This ensures that little of the hydrocarbon ingredients boiling below the range is lost during standing or boiling when recrystallising a substance. Petroleum ethers with boiling ranges (at 760mm pressure) of $35-60^\circ$, $40-60^\circ$, $60-80^\circ$, $80-100^\circ$, and $100-120^\circ$ are generally free from unsaturated and aromatic hydrocarbons. The lowest boiling petroleum ether commercially available has **b** $30-40^\circ/760\text{mm}$ and is mostly *n*-pentane. The purer spectroscopic grades are almost completely free from olefinic and aromatic hydrocarbons. **Petroleum spirit** (which is sometimes used synonymously with petroleum ether or light petroleum) is usually less refined petroleum, and **ligroin** is used for fractions boiling above 100° . The lower boiling fractions consist of mixtures of *n*-pentane (**b** 36°), *n*-hexane (**b** 68.5°) and *n*-heptane (**b** 98°), and some of their isomers in varying proportions. For purification see petroleum ether **b** $35-60^\circ$ in 'Aliphatic Compounds', Chapter 3, which is typical.

Solvents commonly used for recrystallisation, and their boiling points, are given in Table 6.

For comments on the toxicity and use of **benzene** see the 'Introduction' pages of Chapters 3, 4 and 6.

Mixed Solvents. Where a substance is too soluble in one solvent and too insoluble in another, for either to be used for recrystallisation, it is often possible (provided the solvents are miscible) to use them as a mixed solvent system. (In general, however, it is preferable to use a single solvent if this is practicable.) Table 7 contains many of the common pairs of miscible solvents.

Several procedures with mixed solvents have been used successfully for crystallisation. These include the following:

(a) The material is dissolved in the solvent in which it is more soluble at room temperature, then the second solvent (heated to near boiling) is added cautiously to the cold solution until a slight turbidity persists or crystallisation begins. The turbidity is cleared by warming or by adding several drops of the first solvent, and the clear solution is allowed to cool slowly for crystallisation to occur. The supernatant is decanted off carefully (do not disturb the crystals unduly) and more of the second solvent is added to the clear decanted supernatant until turbidity begins again, and is set aside for further crystals to form. The procedure is repeated until no more crystals separate.

(b) A variation of the procedure in (a) is simply to precipitate the material in a microcrystalline form from solution in one solvent at room temperature, by adding a little more of the second solvent also at room temperature, filtering off the crystals, adding a little more of the second solvent and repeating the process. This ensures, at least in the first or last precipitation, a material which contains as little as possible of the impurities, which may also be precipitated in this way. With inorganic salts or metal salts of organic acids, the first solvent is commonly water

and the second solvent is alcohol or acetone. With salts of organic bases and inorganic acids, e.g. hydrochloride, or salts of organic acids and organic bases, the first solvent is usually an alcohol or acetone, in which the salt is very soluble and the second solvent is dry diethyl ether.

(c) A very concentrated solution of the compound in the first solvent in one beaker, and a second beaker containing the second solvent in which the compound is insoluble are placed in a desiccator. As the vapours of the two solvents equilibrate in the desiccator, and crystals separate in the first beaker that contains the compound.

(d) This procedure is best carried out in a cold room (at *ca* 4°). A strong solution of the solid in the solvent in which it is very soluble is *layered* carefully with the second solvent. As the second solvent diffuses and dissolves into the solution, crystals begin to form at the 'interface'. When separation of crystals is complete and the solvent mixture is homogeneous, another layer of the second solvent is applied and the process is repeated.

Seeding is well known to initiate the crystallisation process. A good way to procure seed crystals is to dissolve the crystals in the minimum amount of solvent, place the solution in a watch glass, then blow a fine stream of dry N₂ or argon gently over the surface of the solution until seed crystals are formed. *Alternatively*, the inert gas is allowed to evaporate all the solvent, and the residual crystals or fine powder are used for seeding. The seeds are applied in the above procedures at the appropriate time, e.g. when first turbidity appears, or placed onto the tip of a glass rod which is then rubbed against the sides of the container of the solution until crystallisation begins.

Recrystallisation from the melt. A crystalline solid melts when its temperature is raised sufficiently for the thermal agitation of its molecules or ions to overcome the restraints imposed by the crystal lattice. Usually, impurities weaken crystal structures, and hence lower the melting points of solids (or the freezing points of liquids). If an impure material is melted and cooled slowly (with the addition, if necessary, of a trace of solid material near the freezing point to avoid supercooling), the first crystals that form will usually contain less of the impurity, so that fractional solidification by partial freezing can be used as a purification process for solids with melting points lying in a convenient temperature range (or for more readily frozen liquids). Some examples of cooling baths that are useful in recrystallisation are summarised in Table 8. In some cases, impurities form higher melting eutectics with substances to be purified, so that the first material to solidify is less pure than the melt. For this reason, it is often desirable to discard the first crystals and also the final portions of the melt. Substances having similar boiling points often differ much more in melting points, so that fractional solidification can offer real advantages, especially where ultrapurity is sought. For further information on this method of recrystallisation, consult the earlier editions of this book as well as references by Schwab and Wichers (*J Res Nat Bur Stand* **25** 747 1940, DOI: 10.6028/jres.025.001). This method works best if the material is already nearly pure, and hence tends to be a final purification step.

Zone refining. Zone refining (or zone melting) is a particular development for fractional solidification and is applicable to all crystalline substances that show differences in the concentrations of impurities in liquid and solid states at solidification. The apparatus used in this technique consists essentially of a device in which the crystalline solid to be purified is placed in a glass tube (set vertically) which is made mechanically to move slowly upwards while it passes through a fixed coil (one or two turns) of heated wire. A narrow zone of molten crystals is formed when the tube is close to the heated coil. As the zone moves away from the coil the liquid crystallises, and a fresh molten zone is formed below it at the coil position. The machine can be set to recycle repeatedly. At its advancing side, the zone has a melting interface with the impure material whereas on the upper surface of the zone there is a constantly growing face of higher-melting, resolidified purer material. This leads to a progressive increase in impurity in the liquid phase which, at the end of the run, is discarded from the bottom of the tube. Also, because of the progressive increase in impurity in the liquid phase, the resolidified material contains correspondingly less of the impurities. For this reason, it is usually necessary to make several zone-melting runs before a sample is satisfactorily purified. This is also why the method works most successfully if the material is already fairly pure. In all these operations the zone must travel slowly enough to enable impurities to diffuse or be convected away from the area where resolidification is occurring.

The technique finds commercial application in the production of metals of extremely high purity (tubes other than glass are used in these cases, and impurities can be reduced down to 10⁻⁹ ppm), in purifying refractory oxides, and in purifying organic compounds, using commercially available equipment. Criteria for indicating that definite purification is achieved include elevation of melting point, removal of colour, fluorescence or smell, and a lowering of electrical conductivity. Difficulties likely to be found with organic compounds, especially those with low melting points and low rates of crystallisation, are supercooling and, because of surface tension and contraction, the tendency of the molten zone to seep back into the recrystallised areas. The method is likely to be

useful in cases where fractional distillation is not practicable, either because of unfavourable vapour pressures or ease of decomposition, or where super-pure materials are required. The method has been used for the latter purpose for purifying anthracene, benzoic acid, chrysene, morphine, 1,8-naphthyridine and pyrene to name a few. [See E.F.G.Herington, *Zone Melting of Organic Compounds*, Wiley & Sons, NY, 1963; W.Pfann, *Zone Melting*, 2nd edn, Wiley, NY, 1966; H.Schildknecht, *Zonenschmelzen*, Verlag Chemie, Weinheim, 1964; W.R.Wilcox, R.Friedenberg and Back *Chem Rev* **64** 187 1964, DOI: 10.1021/cr60228a006; M.Zief and W.R.Wilcox (Eds), *Fractional Solidification*, Vol I, M Dekker Inc. NY, 1967.]

SUBLIMATION

Sublimation differs from ordinary distillation because the vapour condenses to a solid instead of a liquid. Usually, the pressure in the heated system is diminished by pumping, and the vapour is condensed (after travelling a relatively short distance) onto a cold finger or some other cooled surface. This technique, which is applicable to many organic solids, can also be used with inorganic solids such as aluminium chloride, ammonium chloride, arsenious oxide and iodine to name a few. In some cases, passage of a stream of inert gas over the heated substance secures adequate vaporisation and reduces oxidation. This procedure has the added advantage of removing occluded solvent used for recrystallising the solid.

CHROMATOGRAPHY

Chromatography is often used with advantage for the purification of small, and large, amounts of complex organic mixtures. Chromatography techniques all rely on the differential distribution of the various components in the solution, between the mobile phase and the stationary phase. The mobile phase can either be a gas or a liquid, whereas the stationary phase can either be a solid or a non-volatile liquid adsorbed on a solid surface.

The major chromatographic techniques can also be categorised according to the nature of the mobile phase used - vapour phase chromatography for when a gas is the mobile phase and liquid chromatography for when a liquid is the mobile phase.

The suppliers of chromatography equipment for every need are too numerous to list here but can be viewed on the internet under 'Chromatography products'. Details and orders can be obtained from the respective websites listed at the end of the section on HPLC below.

Vapour phase chromatography (GC or gas-liquid chromatography)

The mobile phase in vapour phase chromatography is a gas (e.g. hydrogen, helium, nitrogen or argon), and the stationary phase is a non-volatile liquid impregnated onto a porous material. The mixture to be purified is injected into a heated inlet whereby it is vaporised and taken into the column by the carrier gas. It is separated into its components by partition between the liquid on the porous support and the gas. For this reason vapour-phase chromatography is sometimes referred to as gas-liquid chromatography (g.l.c). Vapour phase chromatography is very useful for the resolution of a mixture of volatile compounds. This type of chromatography uses either packed or capillary columns. Packed columns have internal diameters of 3-5 mm with lengths of 2-6 metres. These columns can be packed with a range of materials including firebrick derived materials (chromasorb P, for separation of non-polar hydrocarbons) or diatomaceous earth (chromasorb W, for separation of more polar molecules such as acids, amines). Capillary columns have stationary phase bonded to the walls of long capillary tubes. The diameters of capillary columns are less than 0.5 mm, and the lengths of these columns can go up to 50 metres! These columns have much superior separating capabilities than the packed columns. Elution times for equivalent resolutions with packed columns can be up to ten times shorter. It is believed that almost any mixture of compounds can be separated using one of the four stationary phases, OV-101, SE-30, OV-17 and Carbowax-20M. Capillary columns for analysis in gas chromatography are now routinely used. An extensive range of packed and capillary columns is available from chromatographic specialists such as Supelco, Alltech, Hewlett-Packard, Phenomenex (for stainless steel capillary columns see <phenomenex.com>, etc. (see above and at the end of the section on HPLC below). Some typical liquids used for stationary phases in gas chromatography are listed in Table 9.

Although gas chromatography is routinely used for the analysis of mixtures, this form of chromatography can also be used for separation/purification of substances. This is known as preparative GC. In preparative GC, suitably packed columns are used, and as substances emerge from the column, they are collected by condensing the vapour

of these separated substances in suitable traps. The carrier gas blows the vapour through these traps; hence these traps have to be very efficient. Improved collection of the effluent vaporised fractions in preparative work is attained by strong cooling, increasing the surface of the traps by packing them with glass wool, and/or by applying an electrical potential which neutralises the charged vapour and causes it to condense.

When the gas chromatograph is attached to a mass spectrometer, a very powerful analytical tool (*gas chromatography-mass spectrometry*; **GC-MS**) is produced. Gas chromatography allows the separation of mixtures but does not allow the definitive identification of unknown substances, whereas mass spectrometry is good for the identification of individual compounds of the mixtures of compounds. This means that with GC-MS, both separation *and* identification of substances in mixtures can be achieved. The spectrometer can be connected to a computer that has a library from which the mass peaks can be compared and is a very powerful analytical tool. Because of the relatively small amounts of material required for mass spectrometry, a splitting system is inserted between the column and the mass spectrometer. This enables only a small fraction of the effluent to enter the spectrometer; the rest of the effluent is usually collected or vented to the air. See p 29 below for **GC-MS-MS**.

For more detail on apparatus and chromatographic columns see <http://www.sigmaaldrich.com/analytical-chromatography/gas-chromatography.html> and websites at the end of the section on HPLC below.

Liquid chromatography

In contrast to vapour phase chromatography, the mobile phase in liquid chromatography is a liquid. In general, there are four main types of liquid chromatography: *adsorption*, *partition*, *ion-chromatography*, and *gel filtration*.

Adsorption chromatography is based on the difference in the extent to which substances in solution are adsorbed onto a suitable surface. The main techniques in adsorption chromatography are TLC (thin layer chromatography), paper and column chromatography.

Thin layer chromatography (TLC). In thin layer chromatography, the mobile phase, i.e. the solvent, creeps up the stationary phase (the adsorbent) by capillary action. The adsorbent (e.g. silica, alumina, cellulose) is spread on a rectangular glass plate (or solid inert plastic sheet or aluminium foil). Some adsorbents (e.g. silica) are mixed with a setting material (e.g. CaSO₄) by the manufacturers which causes the film to set hard on drying. The adsorbent can be activated by heating at 100-110° for a few hours. Other adsorbents (e.g. celluloses) adhere on glass plates without a setting agent. Thus some grades of adsorbents have prefixes; e.g. prefix G means that the adsorbent can cling to a glass plate and is used for TLC (e.g. silica gel GF₂₅₄ is for TLC plates which have a dye that fluoresces under 254nm UV light). Those lacking this binder have the letter H after any coding and is suitable for column chromatography e.g. silica gel 60H. The materials to be purified or separated are spotted in a solvent close to the lower end of the plate and allowed to dry. The spots will need to be placed at such a distance so as to ensure that when the lower end of the plate is immersed in the solvent, the spots are a few mm above the eluting solvent. The plate is placed upright in a tank containing the eluting solvent. Elution is carried out in a closed tank to ensure equilibrium. Good separations can be achieved with square plates if a second elution is performed at right angles to the first using a second solvent system. For rapid work, plates of the size of microscopic slides or even smaller are used which can decrease the elution time and cost without loss of resolution. The advantage of plastic backed and aluminium foil backed plates is that the size of the plate can be made as required by cutting the sheet with scissors or a sharp guillotine. Visualisation of substances on TLC can be carried out using UV light if they are UV absorbing or fluorescing substances or by spraying or dipping the plate with a reagent that gives coloured products with the substance (e.g. iodine solution or vapour gives brown colours with amines), or with dilute sulfuric acid (organic compounds become coloured or black when the plates are heated at 100° if the plates are of alumina or silica, but not cellulose). (see Table 10 for some methods of visualisation.) Some alumina and silica powders are available with fluorescent materials in them, in which case the whole plate fluoresces under UV light. Non-fluorescing spots are thus clearly visible, and fluorescent spots invariably fluoresce with a different colour. The colour of the spots can be different under UV light at 254nm from light at 365nm. Another useful way of showing up non-UV absorbing spots is to spray the plate with a 1-2% solution of Rhodamine 6G in acetone. Under UV light the dye fluoresces and reveals the non-fluorescing spots. For preparative work, if the material in the spot or fraction is soluble in ether or petroleum ether, the desired substance can be extracted from the adsorbent with these solvents which leave the water soluble dye behind.

TLC can be used as an analytical technique, or as a guide to establishing the progress of a reaction, conditions for column chromatography or as a preparative technique in its own right.

The thickness of the adsorbent on the TLC plates can be between 0.2mm to 2mm or more. In preparative work, the thicker plates are used and hundreds of milligrams of mixtures can be purified conveniently and quickly. The spots or areas are easily scraped off the plates and the desired substances extracted from the adsorbent with the

required solvent. For preparative TLC, non-destructive methods for visualising spots and fractions are required. As such, the use of UV light is very useful. If substances are not UV active, then a small section of the plate (usually the right or left edge of the plate) is sprayed with a visualising agent while the remainder of the plate is kept covered.

Thin layer chromatography has been used successfully with ion-exchange celluloses as stationary phases and various aqueous buffers as mobile phases. Also, gels (e.g. Sephadex G-50 to G-200 superfine) have been adsorbed on glass plates and are good for fractionating substances of high molecular weights (1500 to 250,000). With this technique, which is called *thin layer gel filtration (TLG)*, molecular weights of proteins can be determined when suitable markers of known molecular weights are run alongside (see Chapter 6).

Commercially available pre-coated plates with a variety of adsorbents are generally very good for quantitative work because they are of a standard quality. Plates of a standardised silica gel 60 (as medium porosity silica gel with a mean porosity of 6mm) released by Merck have a specific surface of 500 m²/g and a specific pore volume of 0.75 ml/g. They are so efficient that they have been called *high performance thin layer chromatography (HPTLC)* plates (Ripphahn & Halpaap *J Chromatogr* **112** 81 1975, DOI: 10.1016/S0021-9673(00)99944-8). In another variant of thin layer chromatography the adsorbent is coated with an oil as in gas chromatography thus producing *reverse-phase thin layer chromatography (R-P TLC)*. R-P TLC plates e.g. silica gel RP-18 are available from Fluka and Merck.

A very efficient form of chromatography makes use of a circular glass plate (rotor) coated with an adsorbent (silica, alumina or cellulose). As binding to a rotor is needed, the sorbents used may be of a special quality and/or binders are added to the sorbent mixtures. For example, when silica gel is required as the adsorbent, silica gel 60 PF-254 with calcium sulfate (Merck catalog 7749) is used. The thickness of the adsorbent (1, 2 or 4 mm) can vary depending on the amount of material to be separated. The apparatus used is called a **Chromatotron** (available from Harrison Research, USA). The glass plate is rotated by a motor, and the sample followed by the eluting solvent is allowed to drip onto a central position on the plate. As the plate rotates the solvent elutes the mixture, centrifugally, while separating the components in the form of circular bands radiating from the central point. The separated bands are usually visualised conveniently by UV light and as the bands approach the edge of the plate, the eluent is collected. The plate with the adsorbent can be re-used many times if care is employed in the usage, and hence this form of chromatography utilises less adsorbents as well as solvents.

Recipes and instructions for coating the rotors are available from the Harrison website <<http://www.harrisonresearch.com/chromatotron/>>. In addition, information on how to regenerate the sorbents and binders is also included.

Paper chromatography. This is the technique from which thin layer chromatography was developed. It uses cellulose paper (filter paper) instead of the TLC adsorbent and does not require a backing like the plastic sheet in TLC. It is used in the **ascending procedure** (like in TLC) whereby a sheet of paper is hung in a jar, and the materials to be separated are spotted (after dissolving in a suitable solvent and drying) near the bottom of the sheet which dips into the eluting solvent just below the spots. As the solvent rises up the paper the spots are separated according to their adsorption properties. A variety of solvents can be used, the sheet is then dried in air (fume cupboard), and can then be run again with the solvent running at right angles to the first run to give a two-dimensional separation. The spots can then be visualised as in TLC or can be cut out and analysed as required. A **descending procedure** had also been developed where the material to be separated is spotted near the top of the paper and the top end is made to dip into a tray containing the eluting solvent. The whole paper is placed in a glass jar, and the solvent then runs down the paper causing the materials in the spots to separate also according to their adsorption properties and to the eluting ability of the solvent. This technique is much cheaper than TLC and is still used (albeit with thicker cellulose paper) with considerable success for the separation of protein hydrolysates for sequencing analysis and/or protein identification. However, modern and more efficient technologies are available for analysing proteins and their hydrolysates although the equipment is expensive. (Whatman papers for chromatography and electrophoresis are available also from Sigma-Aldrich Labware.)

Column Chromatography. The substances to be purified are usually placed on the top of the column and the solvent is run down the column. Fractions are collected and checked for compounds using TLC (UV and/or other means of visualisation). The adsorbent for chromatography can be packed dry and solvents to be used for chromatography are used to equilibrate the adsorbent by flushing the column several times until equilibration is achieved. *Alternatively*, the column containing the adsorbent is packed wet (slurry method), and pressure is applied at the top of the column until the column is well packed (i.e. the adsorbent is settled).

Graded Adsorbents and Solvents. Some materials used in columns for adsorption chromatography are grouped in Table 11 in an approximate order of effectiveness. Other adsorbents sometimes used include barium carbonate, calcium sulfate, calcium phosphate, charcoal (usually mixed with Kieselguhr or

other forms of diatomaceous earth, for example, the filter aid Celite) and cellulose. The alumina can be prepared in several grades of activity (see below).

In most cases, adsorption takes place most readily from non-polar solvents such as petroleum ether and least readily from polar solvents such as alcohols, esters, and acetic acid. Common solvents, arranged in approximate order of increasing eluting ability are also given in Table 11. Eluting power roughly parallels the dielectric constants of solvents. The series also reflects the extent to which the solvent binds to the column material, thereby displacing the substances that are already adsorbed. This preference of alumina and silica gel for polar molecules explains, for example, the use of percolation through a column of silica gel for the following purposes:— drying of ethylbenzene, removal of aromatics from 2,4-dimethylpentane and of ultraviolet absorbing substances from cyclohexane. Mixed solvents are intermediate in strength, and so provide a finely graded series. In choosing a solvent for use as an eluent it is necessary to consider the solubility of the substance in it and the ease with which it can subsequently be removed.

Preparation and Standardisation of Alumina. The activity of alumina depends inversely on its water content, and a sample of poorly active material can be rendered more active by leaving for some time in a round bottomed flask heated up to about 200° in an oil bath or a heating mantle while a slow stream of a dry inert gas is passed through it. *Alternatively*, it is heated to red heat (380-400°) in an open vessel for 4-6 hours with occasional stirring and then cooled in a vacuum desiccator: this material is then of grade I activity. Conversely, alumina can be rendered less active by adding small amounts of water and thoroughly mixing for several hours. Addition of about 3% (w/w) of water converts grade I alumina to grade II.

Used alumina can be regenerated by repeated extraction, first with boiling methanol, then with boiling water, followed by drying and heating. The degree of activity of the material can be expressed conveniently in terms of the scale due to Brockmann and Schodder (*Chem Ber* **74** 73 1941, DOI: 10.1002/cber.19410740113).

Alumina is normally slightly alkaline. A (less strongly adsorbing) neutral alumina can be prepared by making a slurry in water and adding 2M hydrochloric acid until the solution is acid to Congo red. The alumina is then filtered off, washed with distilled water until the wash water gives only a weak violet colour with Congo red paper, and dried.

Alumina used in TLC can be recovered by washing in ethanol for 48 hours with occasional stirring, to remove binder material and then washed with successive portions of ethyl acetate, acetone and finally with deionised water. Fine particles are removed by siphoning. The alumina is first suspended in 0.04M acetic acid (30min), then in distilled water (2 x 30min) and siphoning off after each wash. The process is repeated 7-8 times. It is then dried and activated at 200° [Vogh & Thomson *Anal Chem* **53** 1345 1981, DOI: 10.1021/ac00232a010].

Preparation of other adsorbents

Silica gel can be prepared from commercial water-glass by diluting it with water to a density of 1.19 and, while keeping it cooled to 5°, adding concentrated hydrochloric acid with stirring until the solution is acid to thymol blue. After standing for 3 hours, the precipitate is filtered off, washed on a Büchner funnel with distilled water, then suspended in 0.2M hydrochloric acid. The suspension is set aside for 2-3 days, with occasional stirring, then filtered, washed well with water and dried at 110°. It can be activated by heating up to about 200° as described for alumina.

Powdered commercial silica gel can be purified by suspending and standing overnight in concentrated hydrochloric acid (6ml/g), decanting the supernatant and repeating with fresh acid until the latter remains colourless. After filtering with suction on a sintered-glass funnel, the residue is suspended in water and washed by decantation until free of chloride ions. It is then filtered, suspended in 95% ethanol, filtered again and washed on the filter with 95% ethanol. The process is repeated with anhydrous diethyl ether before the gel is heated for 24 hours at 100° and stored for another 24 hours in a vacuum desiccator over phosphorus pentoxide.

To buffer silica gel for flash chromatography (see later), 200g of silica is stirred in 1L of 0.2M NaH₂PO₄ for 30 minutes. The slurry is then filtered with suction using a sintered glass funnel. The silica gel is then activated at 110°C for 16 hours. The pH of the resulting silica gel is ~4. Similar procedures can be utilized to buffer the pH of the silica gel at various pHs (up to pH ~8: pH higher than this causes degradation of silica) using appropriate phosphate buffers.

Commercial silica gel has also been purified by suspension of 200g in 2L of 0.04M ammonia, and stood for 5 minutes before siphoning off the supernatant. The procedure was repeated 3-4 times, before rinsing with distilled water and drying, and activating the silica gel in an oven at 110° [Vogh & Thomson, *Anal Chem* **53** 1345 1981,

DOI: 10.1021/ac00232a010]. Although silica gel is not routinely recycled after use (due to fear of contamination as well as the possibility of reduced activity), the costs of using new silica gel for purification may be prohibitive. In these cases, recycling may be achieved by stirring the used silica gel (1 kg) in a mixture of methanol and water (2L MeOH/4L water) for 30-40 minutes. The silica gel is filtered (as described above) and reactivated at 110°C for 16 hours.

Diatomaceous earth (Celite 535 or 545, Hyflo Super-cel, Dicalite, Kieselguhr) is purified before use by washing with 3M hydrochloric acid, then water, or it is made into a slurry with hot water, filtered at the pump and washed with water at 50° until the filtrate is no longer alkaline to litmus. Organic materials can be removed by repeated extraction at 50° with methanol or chloroform, followed by washing with methanol, filtering and drying at 90-100°.

Charcoal is generally satisfactorily activated by heating gently to red heat in a crucible or quartz beaker in a muffle furnace, finally allowing to cool under an inert atmosphere in a desiccator. Good commercial activated charcoal is made from wood, e.g. *Norit* (from Birch wood), *Darco* and *Nuchar*. If the cost is important, then the cheaper *animal charcoal* (bone charcoal) can be used. However, this charcoal contains calcium phosphate and other calcium salts and cannot be used with acidic materials. In this case the charcoal is boiled with dilute hydrochloric acid (1:1 by volume) for 2-3 hours, diluted with distilled water and filtered through a fine grade paper on a Büchner flask, washed with distilled water until the filtrate is almost neutral, and dried first in air, then in a vacuum, and activated as above. To improve the porosity, charcoal columns are usually prepared in admixture with diatomaceous earth.

Cellulose for chromatography is purified by sequential washing with chloroform, ethanol, water, ethanol, chloroform and acetone. More extensive purification uses aqueous ammonia, water, hydrochloric acid, water, acetone and diethyl ether, followed by drying in a vacuum. Trace metals can be removed from filter papers by washing for several hours with 0.1M oxalic, citric acid, or 0.1M EDTA solution, followed by repeated washing with distilled water.

Supelco supply a variety of 'solvent desorption tubes', which are cartridges that remove specific impurities (e.g. LpDNPH cartridges which contain a high purity silica adsorbent coated with 2,4-dinitrophenylhydrazine and remove carbonyl compounds; ozone scrubbers which eliminate ozone). Other cartridges such as the 'ORBO charcoal' cartridges contain various beds such as *activated coconut charcoal*, *activated petroleum charcoal*, *HBr on petroleum charcoal* or *4-tert-butyl catechol on charcoal* and are used for specific or for general purposes. Other ORBO cartridges contain activated silica gel and coated silica gel, Florisil, Carboxen, Carbosieve and carbon-coated traps, as well as a variety of ORBO porous polymers, polyurethane, and glass fibre coated with 1-(2-pyridyl)piperazine (which is specific for sampling diisocyanates). They also supply filter cartridges for trapping aerosols and particulate forms of semivolatiles.

Flash Chromatography (FC and HPFC)

A faster method of separating components of a mixture is *flash chromatography* (see Still et al. *J Org Chem* **43** 2923 1978, DOI: 10.1021/jo00408a041). Flash chromatography has become an extremely useful and popular means of purification of small as well as large quantities of compounds. In flash chromatography the eluent flows through the column under a pressure of *ca* 1 to 4 atmospheres. The lower end of the chromatographic column has a relatively long taper closed with a tap. The upper end of the column is connected through a ball joint to a tap. *Alternatively*, a specially designed chromatographic column with a solvent reservoir can also be used (for an example, see the Aldrich Chemical Catalog-glassware section). The tapered portion is plugged with cotton, or quartz wool and *ca* 1 cm length of fine washed sand (the latter is optional). The adsorbent is then placed in the column as a dry powder or as a slurry in a solvent and allowed to fill to about one-third of the column. A fine grade of adsorbent is required in order to slow the flow rate at the higher pressure, e.g. Silica 60, 230 to 400 mesh with particle size 0.040-0.063mm (e.g. from Merck). The top of the adsorbent is layered with *ca* 1 cm length of fine washed sand. The mixture in the smallest volume of solvent is applied at the top of the column and allowed to flow into the adsorbent under gravity by opening the lower tap momentarily. The top of the column is filled with eluent, the upper tap is connected by a tube to a nitrogen supply from a cylinder, or to compressed air, and turned on to the desired pressure (monitor with a gauge). The lower tap is turned on and fractions are collected rapidly until the level of eluent has reached the top of the adsorbent (do not allow the column to run dry). If further elution is desired then both taps are turned off, the column is filled with more eluting solvent and the process repeated. The top of the column can be modified so that gradient elution can be performed. *Alternatively*, an apparatus for producing the gradient is connected to the upper tap by a long tube and placed high above the column in order to produce the required hydrostatic pressure. Much better resolution is obtained by dry loading

the sample for purification rather than loading the sample as a solution. Flash chromatography is more efficient and gives higher resolution than conventional chromatography at atmospheric pressure and is completed in a relatively shorter time. A successful separation of components of a mixture by TLC using the same adsorbent is a good indication that flash chromatography will give the desired separation on a larger scale.

Very elaborate equipment is now available for FC and HPFC (high-performance flash chromatography), which may include a pump, facility for gradient elution, UV detection and fraction collection of effluent. A large variety of columns (disposable cartridges) with packings such as silicate, carbon, reverse phases for a wide range of applications are commercially available. In addition a plethora of cartridges are available for preliminary purification, prior to FC or HPFC, packed with adsorbents which can remove specific impurities, e.g. unwanted reaction products such as aldehydes or ketone which may be suspected by-products and/or starting materials. [see Supelco online catalog <http://sigma-aldrich.dirxion.com/WebProject.asp?BookCode=chr09flx#>; Sigma Aldrich: <http://www.sigmaaldrich.com/analytical-chromatography/analytical-chromatography-catalog.html>; Biotage: *Synthesis and Purification Catalogue* and the *Analytical Sample Preparation Catalogue* contain details on available FC and HPFC equipment, accessories and consumables, as well as means of optimising purification, see <www.biotage.com>.]

Paired-ion Chromatography (PIC)

Mixtures containing ionic compounds (e.g. acids and/or bases), non-ionisable compounds, and zwitterions can be separated successfully by paired-ion chromatography (PIC). It utilises the 'reverse-phase' technique (Eksborg & Schill *Anal Chem* **45** 2092 1973, DOI: 10.1021/ac60334a019). The stationary phase is lipophilic, such as μ -BONDAPAK C₁₈ or any other adsorbent that is compatible with water. The mobile phase is water or aqueous methanol containing the acidic or basic counter ion. Thus the mobile phase consists of dilute solutions of strong acids (e.g. 5mM 1-heptanesulfonic acid) or strong bases (e.g. 5 mM tetrabutylammonium phosphate) that are completely ionised at the operating pH values which are usually between 2 and 8. An equilibrium is set up between the neutral species of a mixture in the stationary phase and the respective ionised (anion or cation) species which dissolve in the mobile phase containing the counter ions. The extent of the equilibrium will depend on the ionisation constants of the respective components of the mixture, and the solubility of the unionised species in the stationary phase. Since the ionisation constants and the solubility in the stationary phase will vary with the water-methanol ratio of the mobile phase, the separation may be improved by altering this ratio gradually (gradient elution) or stepwise. If the compounds are eluted too rapidly, the water content of the mobile phase should be increased, e.g. by steps of 10%. Conversely, if components do not move, or move slowly, the methanol content of the mobile phase should be increased by steps of 10%.

The application of pressure to the liquid phase in liquid chromatography generally increases the separation (see HPLC). In PIC also, improved efficiency of the column is observed if pressure is applied to the mobile phase (Wittmer et al. *Anal Chem* **47** 1422 1975, DOI: 10.1021/ac60358a072). [See the Fluka (Riedel-deHaën) catalogue and Supelco catalogue, <<http://sigma-aldrich.dirxion.com/WebProject.asp?BookCode=chr09flx#>> for IPC reagents for the separation of cations and anions.]

Ion-exchange Chromatography

Ion-exchange chromatography involves an electrostatic process which depends on the relative affinities of various types of ions for an immobilised assembly of ions of opposite charge. The mobile phase is an aqueous buffer with a fixed pH or an aqueous mixture of buffers in which the pH is continuously increased or decreased as the separation may require. This form of liquid chromatography can also be performed at high inlet pressures of liquid with increased column performances.

Ion-exchange Resins. An ion-exchange resin is made up of particles of an insoluble elastic hydrocarbon network to which is attached a large number of ionisable groups. Materials commonly used comprise synthetic ion-exchange resins made, for example, by crosslinking polystyrene to which has been attached non-diffusible ionised or ionisable groups. Resins with relatively high crosslinkage (8-12%) are suitable for the chromatography of small ions, whereas those with low cross linkage (2-4%) are suitable for larger molecules. Applications to hydrophobic systems are possible using aqueous gels with phenyl groups bound to the rigid matrix (Phenyl-Superose/Sepharose, Pharmacia-Amersham or merged companies) or neopentyl chains (Alkyl-Superose, Biosciences merged companies suppliers). (Superose is a cross-linked agarose-based medium with an almost uniform bead size.) These groups are further distinguishable as strong [-SO₂OH, -NR₃⁺] or weak [-OH, -CO₂H, -PO(OH)₂, -NH₂]. Their charges are counterbalanced by diffusible ions, and the operation of a column depends on its ability and selectivity to replace these ions. The exchange that takes place is primarily an electrostatic process

but adsorptive forces and hydrogen bonding can also be important. A typical sequence for the relative affinities of some common anions (and hence the inverse order in which they pass through such a column) is the following, obtained using a quaternary ammonium (strong base) anion-exchange column:

Fluoride < acetate < bicarbonate < hydroxide < formate < chloride < bromate < nitrite < cyanide < bromide < chromate < nitrate < iodide < thiocyanate < oxalate < sulfate < citrate.

For an amine (weak base) anion-exchange column in its chloride form, the following order has been observed:

Fluoride < chloride < bromide = iodide = acetate < molybdate < phosphate < arsenate < nitrate < tartrate < citrate < chromate < sulfate < hydroxide.

With strong cation-exchangers (e.g. with SO_3H groups), the usual sequence is that polyvalent ions bind more firmly than mono- or di- valent ones, a typical series being as follows:

$\text{Th}^{4+} > \text{Fe}^{3+} > \text{Al}^{3+} > \text{Ba}^{2+} > \text{Pb}^{2+} > \text{Sr}^{2+} > \text{Ca}^{2+} > \text{Co}^{2+} > \text{Ni}^{2+} = \text{Cu}^{2+} > \text{Zn}^{2+} = \text{Mg}^{2+} > \text{UO}_2^{+} = \text{Mn}^{2+} > \text{Ag}^{+} > \text{Tl}^{+} > \text{Cs}^{+} > \text{Rb}^{+} > \text{NH}_4^{+} = \text{K}^{+} > \text{Na}^{+} > \text{H}^{+} > \text{Li}^{+}.$

Thus, if an aqueous solution of a sodium salt contaminated with heavy metals is passed through the sodium form of such a column, the heavy metal ions will be removed from the solution and will be replaced by sodium ions from the column. This effect is greatest in dilute solution. Passage of sufficiently strong solutions of alkali metal salts or mineral acids readily displaces all other cations from ion-exchange columns. (The regeneration of columns depends on this property.) However, when the cations lie well to the left in the above series it is often advantageous to use a complex-forming species to facilitate removal. For example, iron can be displaced from ion-exchange columns by passage of sodium citrate or sodium ethylenediaminetetraacetate. Some of the more common commercially available resins are listed in Table 12.

Ion-exchange resins swell in water to an extent which depends on the amount of crosslinking in the polymer, so that columns should be prepared from the wet material by adding it as a suspension in water to a tube already partially filled with water. (This also avoids trapping air bubbles.) The exchange capacity of a resin is commonly expressed as mg equiv/ml of wet resin. This quantity is pH-dependent for weak-acid or weak-base resins but is constant at about 0.6-2.0 for most strong-acid or strong-base types.

Apart from their obvious applications to inorganic species, sulfonic acid resins have been used in purifying amino acids, aminosugars, organic acids, peptides, purines, pyrimidines, nucleosides, nucleotides and polynucleotides. Thus, organic bases can be applied to the H^{+} form of such resins by adsorbing them from neutral solution and, after washing with water, they are eluted sequentially with suitable buffer solutions or dilute acids. *Alternatively*, by passing alkali solution through the column, the bases will be displaced in an order that is governed by their pK values. Similarly, strong-base anion exchangers have been used for aldehydes and ketones (as bisulfite addition compounds), carbohydrates (as their borate complexes), nucleosides, nucleotides, organic acids, phosphate esters and uronic acids. Weakly acidic and weakly basic exchange resins have also found extensive applications, mainly in resolving weakly basic and acidic species. For demineralisation of solutions without large changes in pH, mixed-bed resins can be prepared by mixing a cation-exchange resin in its H^{+} form with an anion-exchange resin in its OH^{-} form. Commercial examples include Amberlite MB-1 (IR-120 + IRA-400) and Bio-Deminrolit (Zeo-Karb 225 and Zerolit FF). The latter is also available in a self-indicating form.

Ion-exchange Celluloses and Sephadex. A different type of ion-exchange column that finds extensive application in biochemistry for the purification of proteins, nucleic acids and acidic polysaccharides derives from cellulose by incorporating acidic and basic groups to give ion-exchangers of controlled acid and basic strengths. Commercially available cellulose-type resins are listed in Tables 13 and 14. AG 501 x 8 (Bio-Rad) is a mixed-bed resin containing equivalents of AG 50W-x8 H^{+} form and AG 1-x8 HO^{-} form, and Bio-Rex MSZ 501 resin. A dye marker indicates when the resin is exhausted. Removal of unwanted cations, particularly of the transition metals, from amino acids and buffer can be achieved by passage of the solution through a column of Chelex 20 or Chelex 100. The metal-chelating abilities of the resin reside in the bonded iminodiacetate groups. Chelex can be regenerated by washing in two bed volumes of 1M HCl, two bed volumes of 1M NaOH and five bed volumes of water.

Ion-exchange celluloses are available in different particle sizes. It is important that the amounts of 'fines' are kept to a minimum otherwise the flow of liquid through the column can be extremely slow to the point of no liquid flow. Celluloses with a large range of particle sizes should be freed from 'fines' before use. This is done by suspending the powder in the required buffer and allowing it to settle for one hour and then decanting the 'fines'. This separation appears to be wasteful, but it is necessary for reasonable flow rates without applying high pressure

at the top of the column. Good flow rates can be obtained if the cellulose column is packed dry whereby the 'fines' are evenly distributed throughout the column. Wet packing causes the 'fines' to rise to the top of the column, which thus becomes clogged.

Several ion-exchange celluloses require recycling before use, a process that must be applied for recovered celluloses. Recycling is done by stirring the cellulose with 0.1M aqueous sodium hydroxide, washing with water until neutral, then suspending in 0.1M hydrochloric acid and finally washing with water until neutral. When regenerating a column it is advisable to wash with a salt solution (containing the required counter ions) of increasing ionic strength up to 2M. The cellulose is then washed with water and recycled if necessary. Recycling can be carried out more than once if there are doubts about the purity of the cellulose and when the cellulose had been used previously for a different purification procedure than the one to be used. The basic matrix of these ion-exchangers is cellulose and it is important not to subject them to strong acid ($> 1\text{M}$) and strongly basic ($> 1\text{M}$) solutions.

When storing ion-exchange celluloses, or during prolonged usage, it is important to avoid growth of microorganisms or moulds which slowly destroy the cellulose. Good inhibitors of microorganisms are phenyl mercuric salts (0.001%, effective in weakly alkaline solutions), chlorohexidine (Hibitane at 0.002% for anion exchangers), 0.02% aqueous sodium azide or 0.005% of ethyl mercuric thiosalicylate (Merthiolate); these are most effective in weakly acidic solutions for cation exchangers. Trichlorobutanol (Chloreton, at 0.05% is only effective in weakly acidic solutions) can be used for both anion and cation exchangers. Most organic solvents (e.g. methanol) are effective antimicrobial agents but only at high concentrations. These inhibitors must be removed by washing the columns thoroughly before use because they may have adverse effects on the material to be purified (e.g. inactivation of enzymes or other active preparations).

Sephadex. Other carbohydrate matrices such as *Sephadex* are a bead form of cross-linked gels (based on dextran) which have more uniform particle sizes. Their advantages over the celluloses include faster and more reproducible flow rates and they can be used directly without removal of 'fines'. *Sephadex*, which can also be obtained in a variety of ion-exchange forms (see Table 14) consists of beads of a cross-linked dextran gel which swells in water and aqueous salt solutions. The smaller the bead size, the higher the resolution that is possible but the slower the flow rate. Typical applications of *Sephadex* gels are the fractionation of mixtures of polypeptides, proteins, nucleic acids, polysaccharides and for desalting solutions.

Sephadex ion-exchangers, unlike celluloses, are available in narrow ranges of particle sizes. These are of two medium types, the G-25 and G-50, and their dry bead diameter sizes are *ca* 50 to 150 microns. They are available as cation and anion exchange *Sephadex*. One of the disadvantages of using *Sephadex* ion-exchangers is that the bed volume can change considerably with alteration of pH. *Ultragels* also suffer from this disadvantage to a varying extent, but ion-exchangers of the bead type have been developed e.g. *Fractogels*, *Toyopearl*, which do not suffer from this disadvantage.

Sepharose (e.g. *Sepharose CL* and *Bio-Gel A*) is a bead form of agarose gel which is useful for the fractionation of high molecular weight substances, for molecular weight determinations of large molecules (molecular weight > 5000), and for the immobilisation of enzymes, antibodies, hormones and receptors usually for affinity chromatography applications. In preparing any of the above for use in columns, the dry powder is evacuated, then mixed under reduced pressure with water or the appropriate buffer solution. *Alternatively*, it is stirred gently with the solution until all air bubbles are removed. Because some of the wet powders change volumes reversibly with alteration of pH or ionic strength (see above), it is imperative to make allowances when packing columns (see above) in order to avoid overflowing of packing when the pH or salt concentrations are altered.

Cellex CM ion-exchange cellulose can be purified by treatment of 30-40g (dry weight) with 500ml of 1mM cysteine hydrochloride. It is then filtered through a Büchner funnel and the filter cake is suspended in 500ml of 0.05M NaCl/0.5M NaOH. This is filtered and the filter cake is resuspended in 500ml of distilled water and filtered again. The process is repeated until the washings are free from chloride ions. The filter cake is again suspended in 500ml of 0.01M buffer at the desired pH for chromatography, filtered, and the last step repeated several times.

Cellex D and other anionic celluloses are washed with 0.25M NaCl/0.25M NaOH solution, then twice with deionised water. This is followed with 0.25M NaCl and then washed with water until chloride-free. The *Cellex* is then equilibrated with the desired buffer as above.

Crystalline Hydroxylapatite is a structurally organised, highly polar material which, in aqueous solution (in buffers) strongly adsorbs macromolecules such as proteins and nucleic acids, permitting their separation by virtue of the interaction with charged phosphate groups and calcium ions, as well by physical adsor-

ption. The procedure therefore is not entirely ion-exchange in nature. Chromatographic separations of singly and doubly stranded DNA are readily achievable, whereas there is negligible adsorption of low-molecular-weight species.

Gel Filtration

The gel-like, bead nature of wet Sephadex enables small molecules such as inorganic salts to diffuse freely into it while, at the same time, protein molecules are unable to do so. Hence, passage through a Sephadex column can be used for complete removal of salts from protein solutions. Polysaccharides can be freed from monosaccharides and other small molecules because of their differential retardation. Similarly, amino acids can be separated from proteins and large peptides.

Gel filtration using Sephadex G-types (50 to 200) is essentially useful for fractionation of large molecules with molecular weights above 1000. For Superose, the range is given as 5000 to 5×10^6 . Fractionation of lower molecular weight solutes (e.g. ethylene glycols, benzyl alcohols) can now be achieved with Sephadex G-10 (up to Mol.Wt 700) and G-25 (up to Mol.Wt 1500). These dextrans are used only in aqueous solutions. In contrast, Sephadex LH-20 and LH-60 (prepared by hydroxypropylation of Sephadex) are used for the separation of small molecules (Mol.Wt less than 500) using most of the common organic solvents as well as water.

Sephacorb HP (ultrafine, prepared by hydroxypropylation of crossed-linked dextran) can also be used for the separation of small molecules in organic solvents and water, and in addition it can withstand pressures up to 1400 psi making it useful in HPLC. These gels are best operated at pH values between 2 and 12, because solutions with high and low pH values slowly decompose them (see further in Chapter 6).

Supelco (see catalogue) supply a variety of SUPELCOGEL columns (for small molecule separations), TSK-GEL columns (for large molecules separation) and guard columns for gel permeation chromatography. They have columns of the latter type (e.g. TSK-GEL column G4000SW) which can separate globular proteins of $20-10,000 \times 10^3$ Daltons in molecular weight. They also supply 'Ascentis HPLC Applications CDs' containing a comprehensive library of their columns and possible applications.

High Performance Liquid Chromatography (HPLC)

When pressure is applied at the inlet of a liquid chromatographic column the performance of the column can be increased by several orders of magnitude. This is partly because of the increased speed at which the liquid flows through the column and partly because fine column packings which have larger surface areas can be used. Because of the improved efficiency of the columns, this technique has been referred to as high performance, high pressure, or high speed liquid chromatography and has found great importance in chemistry and biochemistry.

The equipment consists of a hydraulic system to provide the pressure at the inlet of the column, a column, a detector, data storage and output, usually in the form of a computer. The pressures used in HPLC vary from a few psi to 4000-5000 psi. The most convenient pressures are, however, between 500 and 1800psi. The plumbing is made of stainless steel or non-corrosive metal tubing to withstand high pressures. Plastic tubing and connectors are used for low pressures, e.g. up to ~500psi. Increase of temperature has a very small effect on the performance of a column in liquid chromatography. Small variations in temperatures, however, do upset the equilibrium of the column, hence it is advisable to place the column in an oven at ambient temperature in order to achieve reproducibility. The packing (stationary phase) is specially prepared for withstanding high pressures. It may be an adsorbent (for adsorption or solid-liquid HPLC), a material impregnated with a high boiling liquid (e.g. octadecyl sulfate, in *reverse-phase* or *liquid-liquid* or *paired-ion* HPLC), an ion-exchange material (in *ion-exchange* HPLC), or a highly porous non-ionic gel (for high performance *gel filtration* or *permeation*). The mobile phase is water, aqueous buffers, salt solutions, organic solvents or mixtures of these.

Detectors. The more commonly used detectors for column chromatography in general have UV, visible, diode array or fluorescence monitoring for light absorbing substances in the effluent, and refractive index monitoring and evaporative light scattering for transparent compounds in the effluent. UV detection is not useful when molecules do not have UV absorbing chromophores, and solvents for elution should be carefully selected when UV monitoring is used so as to ensure the lack of background interference in detection. The sensitivity of the refractive index monitoring is usually lower than the light absorbing monitoring by a factor of ten or more. It is also difficult to use a refractive index monitoring system with gradient elution of solvents. When substances have readily oxidised and reduced forms, e.g. phenols, nitro compounds, heterocyclic compounds etc. then electrochemical detectors are useful. These detectors oxidise and/or reduce these substances and make use of this process to provide a peak on the recorder. The cells of the monitoring devices are very small (*ca* 5 μ l) and the detection is very good. The volumes of the analytical columns are quite small (*ca* 2ml for a 1 metre column)

hence the result of an analysis is achieved very quickly. Larger columns have been used for preparative work and can be used with the same equipment. Most machines have solvent mixing chambers for solvent gradient or ion gradient elution. The solvent gradient (for two solvents) or pH or ion gradient can be adjusted in a linear, increasing or decreasing exponential manner. Splitters can be used, whereby very small volumes of the effluent are directed through the detectors so that the whole effluent does not pass through the detector.

Columns for HPLC. In general two different types of HPLC columns are available. Prepacked columns are those with metal casings with threads at both ends onto which capillary connections are attached. The cartridge HPLC columns are cheaper and are used with cartridge holders. As the cartridge is fitted with a groove for the holding device, no threads are necessary and the connection pieces can be reused.

A large range of HPLC columns (including guard columns, i.e. small pre-columns) are available from Supelco <<http://sigma-aldrich.dirxion.com/WebProject.asp?BookCode=chr09flx#>>, Waters <www.waters.com>, Agilent Technologies <www.chem.agilent.com>, Phenomenex <www.phenomenex.com>, YMC <www.ymc.co.jp/en/>, Merck <www.merck.de>, SGE <www.sge.com>, GE Healthcare <www.gehealthcare.com>, and other leading companies. It is not possible to list the range of columns here that are commercially available because the numbers are too large and include prepared columns for the type of chromatography described below in the *Other Types of Liquid Chromatography* such as Monolithic Chromatography and UPLC (see below). Also, in this range of columns are columns with chiral bonded phases capable of separating enantiomeric mixtures. The number of these, on the other hand, is relatively smaller and some *chiral* columns are listed in Table 15.

Other Types of Liquid Chromatography

New stationary phases for specific purposes in chromatographic separation are being continually developed. *Charge transfer adsorption chromatography* makes use of a stationary phase which contains immobilised aromatic compounds and permits the separation of aromatic compounds by virtue of the ability to form charge transfer complexes (sometimes coloured) with the stationary phase. The separation is caused by the differences in stability of these complexes (Porath & Caldwell *J Chromatogr A* **133** 180 1977, DOI: 10.1016/S0021-9673(00)89219-5).

In *metal chelate adsorption chromatography* a metal is immobilised by partial chelation on a column which contains bi- or tri- dentate ligands. Its application is in the separation of substances which can complex with the bound metals and depends on the stability constants of the various ligands (Porath et al. *Nature* **258** 598 1975, DOI: 10.1038/258598a0; Lönnnerdal et al. *FEBS Lett* **75** 89 1977, DOI: 10.1016/0014-5793(77)80059-8).

An application of chromatography which has found extensive use in biochemistry and has brought a new dimension in the purification of enzymes is *affinity chromatography*. A specific enzyme inhibitor is attached by covalent bonding to a stationary phase (e.g. AH-Sepharose 4B for acidic inhibitors and CH-Sepharose 4B for basic inhibitors, Phenyl-Sepharose for hydrophobic proteins), and will strongly bind only the specific enzyme which is inhibited or preferentially bound, allowing all other proteins to flow through the column. The enzyme is then eluted with a solution of high ionic strength (e.g. 1M sodium chloride) or a solution containing a substrate or reversible inhibitor of the specific enzyme. (The ionic medium can be removed by gel filtration using a mixed-bed gel.) Similarly, an immobilised lectin may interact with the carbohydrate moiety of a glycoprotein. The most frequently used matrixes are cross-linked (4-6%) agarose and polyacrylamide gel. Many adsorbents are commercially available for nucleotides, coenzymes and vitamins, amino acids, peptides, lectins and related macromolecules and immunoglobulins. Considerable purification can be achieved by one passage through the column and the column can be reused several times.

The affinity method may be *biospecific*, for example as an antibody-antigen interaction, or chemical as in the chelation of boronate by *cis*-diols, or of unknown origin as in the binding of certain dyes to albumin and other proteins.

Hydrophobic adsorption chromatography takes advantage of the hydrophobic properties of substances to be separated and has also found use in biochemistry (Hoftsee *Biochem Biophys Res Commun* **50** 751 1973, DOI: 10.1016/0006-291X(73)91308-9; Jennissen & Heilmayer Jr *Biochemistry* **14** 754 1975, DOI: 10.1021/bi00675a017). Specific covalent binding with the stationary phase, a procedure that was called *covalent chromatography*, has been used for the separation of compounds and for immobilising enzymes on a support: the column was then used to carry out specific bioorganic reactions (Mosbach *Method Enzymol* **44** 1976; A.Rosevear et al. *Immobilised Enzymes and Cells: A Laboratory Manual*, Adam Hilger, Bristol, 1987, ISBN 085274515X). See Bibliography for further literature.

More recently *Monolithic Chromatography* has been introduced which is a new type of high-performance liquid chromatography in which the columns are a 'one-piece porous solid', or *monolith*, instead of particles. These columns take a variety of forms for use in adsorption, ion exchange (weak and strong, cation and anion), reverse phase, and are for use in the separation of small and large molecules. The mobile phase in these columns flows

through the whole of the stationary phase. [P. Wang ed., *Monolithic Chromatography and its Modern Applications* ILM Publications, pp 648 2010, ISBN 9781906799038, 1906799032; and for columns see BIA Separations <www.biaseparations.com>].

Ultra performance Liquid chromatography (UPLC) affords a considerable improvement by bringing high performance liquid chromatography to a new level. Great improvements in analysis and purification of amino acids, peptides, proteins, oligonucleotides and glycans can be accomplished. This has been achieved by packing columns with smaller sized particles (1.7-1.8 μ m) and applying pressures of ~15,000psi (~1030 bar) to the mobile phase. NanoACQUITY UPLC trapping and nanoflow columns have been specifically designed for use on Waters nanoACQUITY systems that can be integrated with MS components [see www.waters.com].

Automated column chromatography

Most of the above methods of column chromatography have been, or can be, automated. Devices are available for the automated injection of samples to columns which are useful for analytical evaluation of samples, for repeated analyses, or for repeated separations to obtain larger amounts of material. The specific fractions of the effluent can be collected. Equipment for these purposes can be obtained from several of the supplier listed at the end of the HPLC section above with the corresponding websites. GC systems coupled with mass spectrometers (GC-MS) and HPLC systems coupled to mass spectrometers (LC-MS) are extremely important methods for the separation and identification of substances. These are invariably linked to a computer with internal libraries of mass spectral data which are useful for identifying peaks; and the libraries can be continually updated (see above). With more elaborate equipment LC-MS-MS, where the peaks from the first spectrometer are further analysed by a second mass spectrometer, provide a wealth of information. If not for the costs involved in GC-MS, GC-MS-MS, LC-MS and LC-MS-MS equipment, these systems would be more commonly found in analytical and research laboratories. [For further reading see Bibliography.]

ELECTROPHORESIS

Ionisable substances such as organic and inorganic acids, bases and salts migrate to their respective electrodes (anode or cathode) if a voltage is applied. When they are placed onto a matrix, e.g. paper or gel, then their rate of migration to the electrodes will vary with the charge, nature and structure of the substance. This phenomenon is known as electrophoresis and is very useful for separating and purifying substances. Capillary techniques have been adapted to electrophoresis and ‘capillary electrophoresis’, and ‘capillary zone electrophoresis’ are finding wide use for identification, separation and isolation of ionisable substances (see text in the Bibliography under ‘electrophoresis’ and the ‘Introduction’ in Chapter 6). The method is used extensively for biological substances, e.g. proteins, polypeptides, DNA, RNA, (see Introduction in Chapter 6) but has been used to a limited extent for identifying and purifying small molecules. Elaborate equipment is available commercially which contains essentially an electrolytic cell and a power supply which provides variable voltage for the process. The use of paper (Whatman of various thicknesses) as the matrix on a flat bed or in a vertical descending mode has been completely superseded with polyacrylamide or agarose flat bed gels. These are routinely used mainly for the separation of proteins and nucleic acids. Also capillary electrophoresis (CE) is now widely used for the analysis and detection of biological substances and drugs. It is used for the separation and purification of carbohydrates, nucleic acids, proteins, peptides and for chiral analysis and separation and detection of drugs [see Bibliography].

DRYING

Removal of Solvents

Where substances are sufficiently stable, removal of solvent from recrystallised materials presents no problems. The crystals, after filtering at the pump (and perhaps air-drying by suction), are heated in an oven above the boiling point of the solvent (but below the melting point of the crystals), followed by cooling in a desiccator. Where this treatment is inadvisable, it is still often possible to heat to a lower temperature under reduced pressure, for example in an Abderhalden pistol. This device consists of a small chamber which is heated externally by the vapour of a boiling solvent. Inside this chamber, which can be evacuated (pump) is placed a small boat containing the sample to be dried and also a receptacle with a suitable drying agent. Convenient liquids for use as boiling liquids in an Abderhalden pistol, and their boiling temperatures, are given in Table 16. *Alternatively* an electrically heated drying pistol can also be used. In cases where heating above room temperature cannot be used, drying must be carried out in a vacuum desiccator containing suitable absorbents. For example, hydrocarbons, such as cyclohexane and petroleum ether, can be removed by using shredded paraffin wax, and acetic acid and other acids can be absorbed by pellets of sodium or potassium hydroxide. However, in general, solvent removal is less of a problem than ensuring that the water content of solids and liquids is reduced below an acceptable level.

Removal of Water

Methods for removing water from solids depend on the thermal stability of the solids or the time available. The safest way is to dry in a vacuum desiccator over concentrated sulfuric acid, phosphorus pentoxide, silica gel, calcium chloride, or some other desiccant. Where substances are stable in air and melt above 100°, drying in an air oven may be adequate. In other cases, use of an Abderhalden pistol may be satisfactory.

Often, in drying inorganic salts, the final material that is required is a hydrate. In such cases, the purified substance is left in a desiccator to equilibrate above an aqueous solution having a suitable water-vapour pressure. A convenient range of solutions used in this way is given in Table 17.

Removal of water from gases may be by physical or chemical means, and is commonly by adsorption on to a drying agent in a low-temperature trap. The effectiveness of drying agents depends on the vapour pressure of the hydrated compound - the lower the vapour pressure the less the remaining moisture in the gas.

The most usually applicable of the specific methods for detecting and determining water in organic liquids is due to Karl Fischer. (See J. Mitchell & D.M. Smith, *Aquametry*, 2nd Ed, J Wiley & Sons, New York, 1977-1984, ISBN 0471022640; Fieser & Fieser, *Reagents for Organic Synthesis*, J. Wiley & Sons, NY, Vol 1, 528 1967, ISBN 0271616X), also see Karl Fischer titrant or Hydranal –Titrant Type 5E [64-17-5] and other types in the Aldrich catalogue <<http://www.sigmaaldrich.com/analytical-chromatography/titration/hydranal.html>>. Other techniques include electrical conductivity measurements and observation of the temperature at which the first cloudiness appears as the liquid is cooled (applicable to liquids in which water is only slightly soluble). Addition of anhydrous cobalt (II) iodide (blue) provides a convenient method (colour changing to pink on hydration) for detecting water in alcohols, ketones, nitriles and some esters. Infrared absorption measurements of the broad band for water near 3500 cm⁻¹ can also sometimes be used for detecting water in non-hydroxylic substances.

Cartridges for the removal not only water from solvents or solutions but other specific impurities, e.g. acids, amines, aldehydes, are now commercially available [see supplies listed at the end of the HPLC section together with their respective websites]. For further useful information on mineral and drying agents, go to the SigmaAldrich website <sigmaaldrich.com>, under technical library (Aldrich) for technical bulletin AL-143.

Intensity and Capacity of Common Desiccants

Drying agents are conveniently grouped into three classes, depending on whether they combine with water reversibly, they react chemically (irreversibly) with water, or they are molecular sieves. The first group varies in their drying intensity with the temperature at which they are used, depending on the vapour pressure of the hydrate that is formed. This is why, for example, drying agents such as anhydrous sodium sulfate, magnesium sulfate or calcium chloride should be filtered off from the liquids before the latter are heated. The intensities of drying agents belonging to this group fall in the sequence:

P₂O₅ >> BaO > Mg(ClO₄)₂, CaO, MgO, KOH (fused), conc H₂SO₄, CaSO₄, Al₂O₃ > KOH (pellets), silica gel, Mg(ClO₄)₂·3H₂O > NaOH (fused), 95% H₂SO₄, CaBr₂, CaCl₂ (fused) > NaOH (pellets), Ba(ClO₄)₂, ZnCl₂, ZnBr₂ > CaCl₂ (technical) > CuSO₄ > Na₂SO₄, K₂CO₃.

Where large amounts of water are to be removed, a preliminary drying of liquids is often possible by shaking with concentrated solutions of sodium sulfate or potassium carbonate, or by adding sodium chloride to salt out the organic phase (for example, in the drying of lower alcohols), as long as the drying agent does not react (e.g. CaCl₂ with alcohols and amines, see below).

Drying agents that combine irreversibly with water include the alkali metals, the metal hydrides (discussed in Chapter 2), and calcium carbide.

Suitability of Individual Desiccants

Alumina. (Preheated to 175° for about 7 hours). Mainly as a drying agent in a desiccator or as a column through which liquid is percolated.

Aluminium amalgam. Mainly used for removing traces of water from alcohols *via* refluxing followed by distillation.

Barium oxide. Suitable for drying organic bases.

Barium perchlorate. Expensive. Used in desiccators (*covered with a metal guard*). Unsuitable for drying solvents or organic material where contact is necessary, because of the danger of **EXPLOSION**

Boric anhydride. (Prepared by melting boric acid in an air oven at a high temperature, cooling in a desiccator, and powdering.) Mainly used for drying formic acid.

Calcium chloride (anhydrous). Cheap. Large capacity for absorption of water, giving the hexahydrate below 30°, but is fairly slow in action and not very efficient. Its main use is for preliminary

drying of alkyl and aryl halides, most esters, saturated and aromatic hydrocarbons and ethers. Unsuitable for drying alcohols and amines (which form addition compounds), fatty acids, amides, amino acids, ketones, phenols, or some aldehydes and esters. Calcium chloride is suitable for drying the following gases: hydrogen, hydrogen chloride, carbon monoxide, carbon dioxide, sulfur dioxide, nitrogen, methane, oxygen, also paraffins, ethers, olefins and alkyl chlorides.

Calcium hydride. See Chapter 2.

Calcium oxide. (Preheated to 700-900° before use.) Suitable for alcohols and amines (but does not dry them completely). Need not be removed before distillation, but in that case the head of the distillation column should be packed with glass wool to trap any calcium oxide powder that might be carried over. Unsuitable for acidic compounds and esters. Suitable for drying gaseous amines and ammonia.

Calcium sulfate (anhydrous). (Prepared by heating the dihydrate or the hemihydrate in an oven at 235° for 2-3 hours; it can be regenerated.) Available commercially as *Drierite*. It forms the hemihydrate, $2\text{CaSO}_4 \cdot \text{H}_2\text{O}$, so that its capacity is fairly low (6.6% of its weight of water), and hence is best used on partially dried substances. It is very efficient (being comparable with phosphorus pentoxide and concentrated sulfuric acid). Suitable for most organic compounds. Solvents boiling below 100° can be dried by direct distillation from calcium sulfate.

Copper (II) sulfate (anhydrous). Suitable for esters and alcohols. Preferable to sodium sulfate in cases where solvents are sparingly soluble in water (for example, benzene or toluene). The colourless to fawn coloured powder turns blue as it absorbs water.

Lithium aluminium hydride. See Chapter 2.

Magnesium amalgam. Mainly used for removing traces of water from alcohols by refluxing the alcohol in the presence of the Mg amalgam followed by distillation.

Magnesium perchlorate (anhydrous). (Available commercially as *Dehydrite*. Expensive.) Used in desiccators. Unsuitable for drying solvents or any organic material where contact is necessary, because of the danger of **EXPLOSION**.

Magnesium sulfate (anhydrous). (Prepared from the heptahydrate by drying at 300° under reduced pressure.) More rapid and effective than sodium sulfate but is slightly acidic. It has a large capacity, forming $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ below 48°. Suitable for the preliminary drying of most organic compounds.

Molecular sieves. See below.

Phosphorus pentoxide. Very rapid and efficient, but difficult to handle and should only be used after the organic material has been partially dried, for example with magnesium sulfate. Suitable for anhydrides, alkyl and aryl halides, ethers, esters, hydrocarbons and nitriles, and for use in desiccators. Not suitable with acids, alcohols, amines or ketones, or with organic molecules from which a molecule of water can be eliminated. Suitable for drying the following gases: hydrogen, oxygen, carbon dioxide, carbon monoxide, sulfur dioxide, nitrogen, methane, ethane and paraffins. It is available on a solid support with an indicator under the name *Sicapent* (from Merck). The colour changes in Sicapent depend on the percentage of water present (e.g. in the absence of water, Sicapent is colourless but becomes green with 20% water and blue with 33% w/w water). When the quantity of water in the desiccator is high, a crust of phosphoric acid forms a layer over the phosphorus pentoxide powder and decreases its efficiency. The crust can be removed with a spatula to expose the dry powder and restore the desiccant property.

Potassium (metal). Properties and applications are similar to those for sodium but as the reactivity is greater than that of sodium, the hazards are greater than those of sodium. **Handle with extreme care.**

Potassium carbonate (anhydrous). Has a moderate efficiency and capacity, forming the dihydrate. Suitable for an initial drying of alcohols, bases, esters, ketones and nitriles by shaking with them, then filtering off. Also suitable for salting out water-soluble alcohols, amines and ketones. Unsuitable for acids, phenols, thiols and other acidic substances.

Potassium hydroxide. Solid potassium hydroxide is very rapid and efficient. Its use is limited almost entirely to the initial drying of organic bases. *Alternatively*, sometimes the base is shaken first with a concentrated solution of potassium hydroxide to remove most of the water present. Unsuitable for acids, aldehydes, ketones, phenols, thiols, amides and esters. Also used for drying gaseous amines and ammonia.

Silica gel. Granulated silica gel is a commercially available drying agent for use with gases, in desiccators, and (because of its chemical inertness) in physical instruments (pH meters, spectrometers, balances). Its drying action depends on physical adsorption, so that silica gel must be used at room temperature or below. By incorporating cobalt chloride into the material it can be made self indicating (blue when dry, pink when wet), re-drying in an oven at 110° being necessary when the colour changes from blue to pink.

Sodium (metal). Used as a fine wire or as chips, for more completely drying ethers, saturated hydrocarbons and aromatic hydrocarbons which have been partially dried (for example with calcium chloride or magnesium sulfate). Unsuitable for acids, alcohols, alkyl halides, aldehydes, ketones, amines and esters. Reacts violently if water is present and can cause a fire with highly flammable liquids.

Sodium hydroxide. Properties and applications are similar to those for potassium hydroxide.

Sodium-potassium alloy. Used as lumps. Lower melting than sodium, so that its surface is readily renewed by shaking. Properties and applications are similar to those for sodium.

Sodium sulfate (anhydrous). Has a large capacity for absorption of water, forming the decahydrate below 33°, but drying is slow and inefficient, especially for solvents that are sparingly soluble in water. It is suitable for the preliminary drying of most types of organic compounds.

Sulfuric acid (concentrated). Widely used in desiccators. Suitable for drying bromine, saturated hydrocarbons, alkyl and aryl halides. Also suitable for drying the following gases: hydrogen, nitrogen, carbon dioxide, carbon monoxide, chlorine, methane and paraffins. Unsuitable for alcohols, bases, ketones or phenols. Also available on a solid support with an indicator under the name *Sicacide* (from Merck) for desiccators. The colour changes in *Sicacide* depends on the percentage of water present (e.g. when dry *Sicacide* is red-violet but becomes pale violet with 27% water and pale yellow to colourless with 33% w/w water).

For convenience, many of the above drying agents are listed in Table 18 under the classes of organic compounds for which they are commonly used.

Molecular sieves

Molecular sieves are types of adsorbents composed of crystalline zeolites (sodium and calcium aluminosilicates). By heating them, water of hydration is removed, leaving holes of molecular dimensions in the crystal lattices. These holes are of uniform size and allow the passage into the crystals of small molecules, but not of large ones. This *sieving* action explains their use as very efficient drying agents for gases and liquids. The pore size of these sieves can be modified (within limits) by varying the cations built into the lattices. The four types of molecular sieves currently available are:

Type 3A sieves. A crystalline potassium aluminosilicate with a pore size of about 3 Angstroms. This type of molecular sieves is suitable for drying liquids such as acetone, acetonitrile, methanol, ethanol and 2-propanol, and drying gases such as acetylene, carbon dioxide, ammonia, propylene and butadiene. The material is supplied as beads or pellets.

Type 4A sieves. A crystalline sodium aluminosilicate with a pore size of about 4 Angstroms, so that, besides water, ethane molecules (but not butane) can be adsorbed. This type of molecular sieves is suitable for drying chloroform, dichloromethane, diethyl ether, dimethylformamide, ethyl acetate, cyclohexane, benzene, toluene, xylene, pyridine and diisopropyl ether. It is also useful for low pressure air drying. The material is supplied as beads, pellets or powder.

Type 5A sieves. A crystalline calcium aluminosilicate with a pore size of about 5 Angstroms, these sieves adsorb larger molecules than type 4A. For example, as well as the substances listed above, propane, butane, hexane, butene, higher *n*-olefins, *n*-butyl alcohol and higher *n*-alcohols, and cyclopropane can be adsorbed, but not branched-chain C₆ hydrocarbons, cyclic hydrocarbons such as benzene and cyclohexane, or secondary and tertiary alcohols, carbon tetrachloride or boron trifluoride. This is the type generally used for drying gases, though organic liquids such as THF and dioxane can be dried with this type of molecular sieves.

Type 13X sieves. A crystalline sodium aluminosilicate with a pore size of about 10 Angstroms which enables many branched-chain and cyclic compounds to be adsorbed, in addition to all the substances removed by type 5A sieves.

They are unsuitable for use with strong acids but are stable over the pH range 5-11.

Because of their selectivity, molecular sieves offer advantages over silica gel, alumina or activated charcoal, especially in their very high affinity for water, small polar molecules and unsaturated organic compounds. Their relative efficiency is greatest when the impurity to be removed is present at low concentrations. Thus, at 25° and a relative humidity of 2%, type 5A molecular sieves adsorb 18% by weight of water, whereas for silica gel and alumina the figures are 3.5 and 2.5% respectively. Even at 100° and a relative humidity of 1.3%, molecular sieves adsorb about 15% by weight of water.

The greater preference of molecular sieves for combining with water molecules explains why this material can be used for drying ethanol and why molecular sieves are probably the most universally useful and efficient drying agents. Percolation of ethanol with an initial water content of 0.5% through a 144 cm long column of type 4A molecular sieves reduced the water content to 10ppm. Similar results have been obtained with pyridine.

The main applications of molecular sieves to purification comprise:

1. Drying of gases and liquids containing traces of water.
2. Drying of gases at elevated temperatures.
3. Selective removal of impurities (including water) from gas streams.

(For example, carbon dioxide from air or ethene; nitrogen oxides from nitrogen; methanol from diethyl ether. In general, carbon dioxide, carbon monoxide, ammonia, hydrogen sulfide, mercaptans, ethane, ethene, acetylene (ethyne), propane and propylene are readily removed at 25°. In mixtures of gases, the more polar ones are preferentially adsorbed).

The following applications include the removal of straight-chain from branched-chain or cyclic molecules. For example, type 5A sieves will adsorb *n*-butyl alcohol but not its branched-chain isomers. Similarly, it separates *n*-tetradecane from benzene, or *n*-heptane from methylcyclohexane.

The following liquids have been dried with molecular sieves: acetone, acetonitrile, acrylonitrile, allyl chloride, amyl acetate, benzene, butadiene, *n*-butane, butene, butyl acetate, *n*-butylamine, *n*-butyl chloride, carbon tetrachloride, chloroethane, 1-chloro-2-ethylhexane, cyclohexane, dichloromethane, dichloroethane, 1,2-dichloropropane, 1,1-dimethoxyethane, dimethyl ether, 2-ethylhexanol, 2-ethylhexylamine, *n*-heptane, *n*-hexane, isoprene, isopropyl alcohol, diisopropyl ether, methanol, methyl ethyl ketone, oxygen, *n*-pentane, phenol, propane, *n*-propyl alcohol, propylene, pyridine, styrene, tetrachloroethylene, toluene, trichloroethylene and xylene. In addition, the following gases have been dried: acetylene, air, argon, carbon dioxide, chlorine, ethene, helium, hydrogen, hydrogen chloride, hydrogen sulfide, nitrogen, oxygen and sulfur hexafluoride.

After use, molecular sieves can be regenerated by heating at between 300°–350° for several hours, preferably in a stream of dry inert gas such as nitrogen or preferably under vacuum, then cooling in a desiccator. Special precautions must be taken before regeneration of molecular sieves used in the drying of flammable solvents.

However, care must be exercised in using molecular sieves for drying organic liquids. Appreciable amounts of impurities were *formed* when samples of acetone, 1,1,1-trichloroethane and methyl-*t*-butyl ether were dried in the liquid phase by contact with molecular sieves 4A [Connett *Lab Pract* **21** 545 1972, ISSN: 0023-6853 (Print) 0023-6853 (Linking)]. Other, less reactive types of sieves may be more suitable but, in general, it seems desirable to perform a preliminary test to establish that no unwanted reactions take place. Useful comparative data for Type 4A and 5A sieves are in Table 19. With the advent of nanotechnology, nanoparticles are finding use as porous materials for a variety of purposes [see J.A. Schwartz & C. Contescu (Eds), *Surfaces of Nanoparticles & Porous Materials*, Marcel Dekker Inc, 1999. ISBN 9780824719333].

PROPERTIES USEFUL IN PURIFICATION

Spectroscopic

Spectroscopic instruments of one sort or another are generally available in laboratories and useful for providing some idea of the purity of the specimen in question. Among these are IR, UV-VIS, fluorescence, NMR and mass spectrometers.

Infrared spectra [IR or FT(Fourier Transformed)-IR with frequency range of ν from ~ 600 to 3400 cm^{-1}], generally of the solid ground in a large excess of KBr, or in a mull by grinding into an oil, e.g. Nujol, or in solution, e.g. CHCl_3 , provide a 'fingerprint' of the substance. The KBr spectrum, or the spectrum of a film between NaCl plates if the substance is a liquid, are more useful as they give detailed information without interfering signals from Nujol or solvent which may mask important signals. Since the IR spectra consist of several signals many of which are sharp, impurities show up clearly. However, if the impurities are less than say 10% it may be difficult to say how impure the sample is, or what impurities are present in it. On the other hand, if the sample is very pure then its spectrum will be superimposable on that of the pure authentic sample.

Ultraviolet Spectra (with wavelength range of λ from ~ 200 to $400\text{ m}\mu$) are measured in dilute solution and are generally broad bands. Although the broadness of the bands make it difficult to identify impurities, the values of the molecular absorption extinction coefficients ϵ ($\text{M}^{-1}\text{cm}^{-1}$) at all wavelengths, but usually measured at the peaks

or troughs, are characteristics of the substance in the particular solvent used, and would be different if the sample was impure. Glass cuvettes cannot be used as they are not transparent to UV radiation, and quartz cells should be used. However, quartz cells need only have quartz on the two opposite faces of the four sided cuvette through which the light passes; the other two faces being made of glass. Similarly in the *visible* spectra (wavelength range of λ from ~ 400 to 800 nm) the ϵ values are characteristic of the substance in the solvent used. In this case the cheaper glass cuvettes may be used as they are transparent to visible light.

Fluorescence Spectra are measured in the wavelength range similar to the *visible* range, but from light that is scattered at right angles to the incident excitation (UV) wavelength. Thus at a set excitation wavelength λ_{ex} , the fluorescence spectrum is scanned and the peak maximum λ_{em} and its ϵ_{em} are recorded. In this case the cuvettes must have quartz faces on all four sides, as the UV light has to go through adjacent sides of the cell. This spectroscopy is useful as sometimes impurities in a sample may fluoresce at a particular wavelength. Generally, dyes have fluorescent properties and are identified in this way. Substances with strongly fluorescing properties have found considerable use in biology. Here they have been tagged to biological molecules and their movement into particular tissues and cells has been traced through their fluorescence. Table 20 lists a number of *Fluorochromes* which have found many applications in analytical chemistry (by tagging to non-fluorescent compounds) and in biology. By selecting a mixture of two fluorochromes it is possible to obtain a desired emission wavelength. In this case the emitted fluorescence energy from the excitation of the first fluorochrome is transferred to the second fluorochrome to provide the desired fluorescence.

For other than macromolecules it is important that *at least* the ^1H NMR spectrum and/or the mass spectrum of the substance should be measured routinely. These measurements require no more than one to three milligrams of material and provide a considerable amount of information about the substance. The ^1H NMR and ^{13}C NMR spectra are measured to assess the purity of hydrogen and carbon containing samples. The use of very high magnetic field NMR spectrometers is especially useful for detecting impurities in such samples. The signals and their relative heights can provide valuable information not only about the extent of the impurities, but also some indications about the nature of the impurities. A variety of NMR solvents are available for dissolving the samples, and the hydrogen atoms of the solvents are replaced by deuterium which does not interfere with the ^1H or ^{13}C spectra. However, deuteration is generally just under 100% and signals from residual H in the solvent may appear in the spectrum and need to be identified. Similarly ^{13}C signals from solvents also should be identified. Common solvents and reagents that contain trace impurities will also show minor signals in the NMR spectra. The ^1H NMR signals of trace impurities in some common organic solvents (including water) and some reagents are presented in Tables 21 and 22. Similarly presented in Table 23 are the ^{13}C NMR signals of some common solvents and reagents. In some instances these minor signals have been very useful as internal standards for reporting the chemical shifts of substances, thus avoiding contamination from other added standards, particularly if the samples need to be used for further studies. The NMR spectra of other nuclei such as ^{11}B and ^{31}P are currently also measured routinely for boron and phosphorus containing compounds. Since the compounds invariably have only a small number of these atoms in their molecules, boron or phosphorus containing impurities are readily identified in the ^{11}B or ^{31}P NMR spectra.

References in the bibliography at the end of this chapter to the Aldrich-Sigma catalogues of NMR, IR and mass spectral data for a large number of the compounds are listed. These collections of spectra are extremely useful for identifying compounds and impurities. If the material appears to have several impurities, these spectra should be valuable for identifying the impurities as much as possible. Preliminary chromatographic (e.g. TLC) and spot tests could be devised to monitor the material and its impurities. Purification methods can then be devised to remove these impurities, and a monitoring method will have already been established.

Ionisation Constants — pK

When substances ionise, their neutral species produce positive and negative species. The ionisation constants are those constant values (equilibrium constants) for the equilibria between the charged species and the neutral species, or species with a larger number of charges (e.g. between mono and dications). These ionisation constants are given as **pK** values where **pK = -log K**, and **K** is the dissociation constant for the equilibrium between the species [Albert and Serjeant, *The Determination of Ionisation Constants*, A Laboratory Manual, 3rd Edition, Chapman & Hall, New York, London, 1984, ISBN 0412242907].

The advantage of using pK values (instead of K values) is that theory (and practice) states that the pK values of the ionisable substances are numerically equal to the pH of the solution at which the concentrations of the ionised and

the neutral species are equal. For example acetic acid has a pK^{25} value of 4.76 at 25° in H_2O ; then at pH 4.76 the aqueous solution contains equal amounts of acetic acid $[AcOH]$ and acetate anion $[AcO^-]$, i.e. $[AcOH]/[AcO^-]$ of 50/50. At pH 5.76 ($pK + 1$) the solution contains $[AcOH]/[AcO^-]$ of 10/90, at pH 6.76 ($pK + 2$) the solution contains $[AcOH]/[AcO^-]$ of 1/99 etc; conversely at pH 3.76 ($pK - 1$) the solution contains $[AcOH]/[AcO^-]$ of 90/10, and at pH 2.76 ($pK - 2$) the solution contains $[AcOH]/[AcO^-]$ of 99/1.

One can readily appreciate the usefulness of pK value in purification procedures, e.g. as when purifying acetic acid. If acetic acid is placed in aqueous solution and the pH adjusted to 7.76 $\{[AcOH]/[AcO^-]$ with a ratio of 0.1/99.9 $\}$, and extracted with say diethyl ether, neutral impurities will be extracted into diethyl ether leaving almost all the acetic acid in the form of AcO^- in the aqueous solution. If then the pH of the solution is adjusted to 1.67 where the acid is almost all in the form $AcOH$, almost all of it will be extracted into diethyl ether.

Aniline will be used as a second example. It has a pK^{25} of 4.60 at 25° in H_2O . If it is placed in aqueous solution at pH 1.60 it will exist almost completely (99.9%) as the anilinium cation. This solution can then be extracted with solvents e.g. diethyl ether to remove neutral impurities. The pH of the solution is then adjusted to 7.60 whereby aniline will exist as the free base (99.9%) and can be extracted into diethyl ether in order to give purer aniline.

See Table 24 for the pH values of selected buffers.

A knowledge of the pK allows the adjustment of the pH without the need of large excesses of acids or base. In the case of inorganic compounds, knowledge of the pK is useful for adjusting the ionic species for making metal complexes which could be masked or extracted into organic solvents [Perrin and Dempsey, *Buffers for pH and Metal ion Control*, Chapman & Hall, New York, London, 1974, ISBN 0412117002], or for obtaining specific anionic species in solution e.g. $H_2PO_4^-$, HPO_4^{2-} or PO_4^{3-} .

The pK values that have been entered in Chapters 3, 4 and 6 have been collected directly from the literature or from compilations of literature values for organic bases [Perrin, *Dissociation Constants of Organic Bases in Aqueous Solution*, Butterworths, London, 1965, Supplement 1972, ISBN 040870408X; Albert and Serjeant, *The Determination of Ionisation Constants*, A Laboratory Manual, 3rd Edition, Chapman & Hall, London, New York, 1984, ISBN 0412242907]; organic acids [Kortum, Vogel and Andrussow, *Dissociation Constants of Organic Acids in Aqueous Solution*, Butterworth, London, 1961; Serjeant and Dempsey, *Dissociation Constants of Organic Acids in Aqueous Solution*, Pergamon Press, Oxford, New York, 1979, ISBN 0080223397; and inorganic acids and bases [Perrin, *Ionisation Constants of Inorganic Acids and Bases in Aqueous Solution*, Second Edition, Pergamon Press, Oxford, New York, 1982, ISBN 0080292143; J Russel and R Cohen *Acid dissociation constant* Bookvika Publishing 2015, ISBN 10: 5513329530, ISBN 13: 9785513329534]. Where literature values were not available, values have been predicted and assigned $pK_{Est} \sim$. Most predictions should be so close to true values as to make very small difference for the purposes intended in this book. The success of the predictions, i.e. how close to the true value, depends on the availability of pK values for closely related compounds because the effect of substituents or changes in structures are generally additive [Perrin, Dempsey and Serjeant, *pKa Prediction for Organic Acids and Bases*, Chapman & Hall, London, New York, 1981, ISBN 041222190X].

All the pK values in this book are pK_a values, the acidic pK , i.e. dissociation of H^+ from an acid (AH) or from a conjugate base (BH^+). Occasionally pK_b values are reported in the literature but these can be converted using the equation $pK_a + pK_b = 14$. For strong acids e.g. sulfuric acid, and strong bases, e.g. sodium hydroxide, the pK values lie beyond the 1 to 11 pH scale and have to be measured in strong acidic and basic media. In these cases appropriate scales e.g. the H_o (for acids) and H_- (for bases) have been used [see Katritzky & Waring, *J Chem Soc* 1540 1962, DOI: 10.1039/JR9620001540]. These values will be less than 1 (and negative) for acids and >11 for bases. They are rough guides to the strengths of acids and bases. Errors in the stated pK and $pK_{Est} \sim$ values can be judged from the numerical values given. Thus pK values of 4.55, 4.5 and 4 mean that the respective errors are better than ± 0.05 , ± 0.3 and ± 0.5 . Values taken from the literature are written as pK , and all the values that were estimated because they were not found in the literature are written as pK_{Est} .

pK and Temperature

The temperatures at which the literature measurements were made are given as superscripts, e.g. pK^{25} . Where no temperature is given, it is assumed that the measurements were carried out at room temperature, e.g. 15–25°. No temperature is given for estimated values ($pK_{Est} \sim$), and these have been estimated from data at room temperature.

The variation of pK with temperature is given by the equation:

$$-\delta(\text{pK})/\delta T = (\text{pK} + 0.052\Delta S^\circ)/T$$

where T is in degrees Kelvin and ΔS° is in Joules $\text{deg}^{-1} \text{mol}^{-1}$. The $-\delta(\text{pK})/\delta T$ in the range of temperatures between 5 to 70° is generally small (e.g. between ~ 0.0024 and ~ 0.04), and for chemical purification purposes is not a seriously deterring factor. It does, however, vary with the compound under study because ΔS° varies from compound to compound. The following are examples of the effect of temperature on pK values: for imidazole the pK values are 7.57 (0°), 7.33 (10°), 7.10 (20°), 6.99 (25°), 6.89 (30°), 6.58 (40°) and 6.49 (50°); and for 3,5-dinitrobenzoic acid they are 2.60 (10°), 2.73 (20°), 2.85 (30°), 2.96 (40°) and 3.07 (40°); and for *N*-acetyl- β -alanine they are 4.4788 (5°), 4.4652 (10°), 4.4564 (15°), 4.4488 (20°), 4.4452 (25°), 4.4444 (30°), 4.4434 (35°) and 4.4412 (40°).

pK and solvent

All stated pK values in this book are for data in dilute aqueous solutions unless otherwise stated, although the dielectric constants, ionic strengths of the solutions and the method of measurement, e.g. potentiometric, spectrophotometric etc., are not given. Estimated values are also for dilute aqueous solutions whether or not the material is soluble enough in water. Generally the more dilute the solution the closer is the pK to the real thermodynamic value. The pK in mixed aqueous solvents can vary considerably with the relative concentrations and with the nature of the solvents. For example the pK²⁵ values for *N*-benzylpenicillin are 2.76 and 4.84 in H₂O and H₂O/EtOH (20:80) respectively; the pK²⁵ values for (-)-ephedrine are 9.58 and 8.84 in H₂O and H₂O/MeOCH₂CH₂OH (20:80), respectively; and for cyclopentylamine the pK²⁵ values are 10.65 and 4.05 in H₂O and H₂O/EtOH (50:50) respectively. pK values in acetic acid or aqueous acetic acid are generally lower than in H₂O.

The dielectric constant of the medium affects the equilibria where charges are generated in the dissociations e.g. $\text{AH} \rightleftharpoons \text{A}^- + \text{H}^+$ and therefore affects the pK values. However, its effect on dissociations where there are no changes in total charge such as $\text{BH}^+ \rightleftharpoons \text{B} + \text{H}^+$ is considerably less, with a slight decrease in pK with decreasing dielectric constant.

A selection of pK_a values for inorganic acids and inorganic bases is displayed in Table 24a, and a selection of pK_a values for organic acids and for organic bases is compiled in Tables 24b and 24c respectively.

Solubilities of Gases in Liquids

There are two ways to define the solubilities of gases in water.

The first is the *Bunsen coefficient* (β), which is the ratio of the volume of gas corrected to STP (0°C and 1atm, i.e. 760mmHg) that dissolves in unit volume of solvent at the temperature of the experiment in equilibrium with the gas at 1atm.

The second is the *Ostwald coefficient* (l) which is the ratio of the volume of gas that dissolves in unit volume of solvent at the temperature of the experiment in equilibrium with the gas at 1atm. The latter is a more convenient ratio to use because no correction for volume is required. Note that the volume of an ideal gas occupied by one molecular weight in grams of element or compound is $\sim 22.4\text{L}$ at STP (e.g. 32g of oxygen occupy 22.4L at STP). The discussion will be limited to the solubilities of oxygen, nitrogen and air (which behave almost as ideal gases) in water, water containing salts, and in some organic solvents. Generally the solubility of these three gases in water decreases with increase of temperature and can be 'boiled out' of the liquid. Their solubilities in organic liquids, on the other hand, generally increase with increase of temperature. The presence of salts in water tends to decrease the solubilities of these gases, i.e. a salting out effect, and increase in pressure increases their solubilities. These properties have to be noted in liquid chromatography at atmospheric and at high pressures. They become important when purifying small amounts of compounds by crystallisation or chromatography when large amounts of solvents are used. One must be wary of the presence of oxygen in solution, particularly in the presence of organic matter. Also the formation of reactive oxygen species e.g. 'singlet' oxygen, superoxide and hydroxyl radicals, especially in the presence of trace metals such as iron, and/or of ultraviolet light can result in the formation of impurities.

The composition of air is: 78.08% of N₂, 20.95% of O₂, 0.03% of CO₂, 0.93% of Ar and less than 0.01% of other gases. Although the partial pressure of O₂ in air at 1atm is ~ 0.20 , it has a higher solubility in H₂O than N₂. At STP the solubility of O₂ by volume in H₂O is 34.9% when in equilibrium with excess of air. Thus by successively dissolving air in H₂O, expelling it, and redissolving the expelled air six to seven times it is possible to increase the concentration of oxygen by volume in the expelled air to 90%. The (β) values for O₂ and N₂ in H₂O at STP are

0.028 and 0.014 respectively. There are 55.5 moles of H_2O in 1L of H_2O , so the molar ratios of O_2 to H_2O can be calculated. Note that the concentration of O_2 in liquids is higher when the liquids are in equilibrium with excess O_2 than when they are with excess of air.

The solubility coefficients (β) and/or (l) of some gases in liquids are given in Tables 25–28. Tables of the solubilities of HCl and NH_3 (g/100g of solution) at 760mm (Table 29) and the boiling points of some useful gases at 760mm (Table 30) are included.

MISCELLANEOUS TECHNIQUES

Freeze-pump-thaw and purging

Volatile contaminants, e.g. traces of low boiling solvent residue or oxygen, in liquid samples or solutions can be very deleterious to the samples on storage. These contaminants can be removed by repeated freeze-pump-thaw cycles. This involves freezing the liquid material under high vacuum in an appropriate vessel (which should be large enough to avoid contaminating the vacuum line with liquid that has bumped) connected to the vacuum line *via* efficient liquid nitrogen traps. The frozen sample is then thawed until it liquefies, kept in this form for some time (*ca* 10-15minutes), refreezing the sample and the cycle repeated several times without interrupting the vacuum. This procedure applies equally well to solutions, as well as purified liquids, e.g. as a means of removing oxygen from solutions for NMR and other measurements. If the presence of nitrogen, helium or argon, is not a serious contaminant then solutions can be freed from gases, e.g. oxygen, carbon dioxide, and volatile impurities by purging with N_2 , He or Ar at room, or slightly elevated, temperature. The gases used for purging are then removed by freeze-pump-thaw cycles or simply by keeping in a vacuum for several hours. Special NMR tubes with a screw cap thread and a PTFE valve (Wilmad) are convenient for freeze thawing of NMR samples. *Alternatively*, NMR tubes with 'J Young' valves (Wilmad) can also be used.

Vacuum lines, Schlenk and glovebox techniques

Manipulations involving materials sensitive to air or water vapour can be carried out by these procedures. Vacuum line methods make use of quantitative transfers, and **P**(pressure)-**V**(volume)-**T**(temperature) measurements, of gases, and trap-to-trap separations of volatile substances.

It is usually more convenient to work under an inert-gas atmosphere using **Schlenk** type apparatus. The *principle* of Schlenk methods involve all-glass tubes, flasks or vessels which have standard ground-glass joints with one or more side-arms, one of which may have a tap. The system can be purged by evacuating and flushing with an inert gas (usually dry nitrogen, or in some cases, argon or helium), repeating the process until the contaminants, e.g. O_2 , H_2O or CO_2 in the vapour phases have been diminished to acceptable limits. Many of the reactions using Schlenk equipment require anhydrous conditions and in this case the equipment should be heated in an oven slightly above 100° for 1 to 2 hours (preferably with dry N_2 , He or argon flushing), and allowed to cool to room temperature in the presence of a desiccant. A large range of Schlenk glassware is commercially available. Schlenk equipment in which refluxing of liquids is possible without contact with the atmosphere outside of the apparatus is available commercially. With these, and tailor-made pieces of glassware, inert atmospheres can be maintained during transfer of material, crystallisation, reflux, filtration, and sublimation. Where addition of a solid sample should be made, an L-tube, or a small bulb with a bent neck, with a glass joint is used in which the solid is placed, and can be transferred to the main reaction vessel by simply rotating the tube or bulb. In the case of a liquid, a separating funnel with an equalising tube can be used to allow equilibration of pressure. Also, liquids can be injected, *via* a syringe through 'Sure/Seal' caps which can be stretched over, or insert nicely into, the ground joint necks of the main reaction container.

Syringe techniques have been developed for small volumes, while for large volumes or where much manipulation is required, dryboxes (*glove boxes*) or dry chambers should be used. Disposable glove bags (e.g. Atmosbags see Sigma-Aldrich Labware of various dimensions) with two or four hands which can be sealed, purged and inflated with an inert gas are available and are relatively cheap and disposable. They are useful not only for handling moisture-sensitive substances, but also for toxic materials.

ADVANCES IN PHYSICAL TECHNIQUES USED IN PURIFICATION

The development, and sophistication of instrumentation keep advancing with the avalanching discoveries and applications of ever developing electronics and computer capabilities. These directly and/or indirectly impact on instruments used for the separation and purification of laboratory chemicals. It is beyond the scope of this work to elaborate in any detail on this. Suffice it to say that a large majority of instruments used are in some way or another integrated with computers which not only use programs to configure the instruments but also to process and store the output information. The amount of information produced by some instruments cannot possibly be

stored and processed manually as is evident in instruments that have become available in past few years. UV-VIS, IR, NIR, MS, NMR and fluorescence spectrometers as well as spectropolarimeters and CD-ORD spectrometers, whether stand alone or bench-top type, are now mostly interfaced with computers which have in addition to programs for driving the instruments, files with libraries so that direct comparisons can be made with spectra of known substances. Among the useful new technologies that are continually appearing is one that allows three phases to be studied together in intact living organisms. Developed by André J. Simpson and coworkers [Mobarhan et al. *Chem Sci Advance Article* 17 Apr 2016, DOI: 10.1039/C6SC00329J], the novel NMR technique ‘**Comprehensive Multiphase Nuclear Magnetic Resonance (CM-NMR)**’ makes use of an integrated probe to study solution-, gel-, and solid-state molecular movement together. It can be applied to living animals, it can deliver oxygen effectively for long periods enabling long in-depth study, the various phases (i.e. metabolites in solution, in soft tissue and in solid skeletal tissue) can be fully differentiated using a range of spectral editing methods, and using ^{13}C isotopic labelling and multi-dimensional NMR to assign metabolites and structural components *in vivo*. This technique has great potential for studying insoluble biological substances in membranes, muscle and bone that cannot be studied in solution-phase by NMR. Essential information may be gained by this technique about proteins which crystallise into solid fibres as in Alzheimer and Parkinson brains, and the chemistry traffic across biological interfaces. [See also Richard Massey *Chemistry World* **13**(9) 25 2016.] Great strides have been made recently in the development and production of automated instruments that synthesise and sequence proteins, DNA and RNA; so much so that the sequencing of the human genome which was completed by several laboratories in five years can now be completed on one instrument in one laboratory in only a few months. There has been a continual need in laboratory work to scale down the quantities used, not only because of cost and waste byproducts but also because of speed in providing the required results. Developments in the field of microfluidics are addressing this problem with increasing success (see following).

Laboratory on a chip/microfluidics. Within the past ten years great strides have been made in *miniaturization* in the wake of tremendous developments in *Nanotechnology* (see Chapter 7) and the silicon chip. This has led to the science and technology of microfluidics that manipulate such small volumes as 10^{-9} to 10^{-18} litres which require channels and chambers with dimensions of radii of hundreds to tens of micrometres. The possibilities of carrying out analytical procedures of chromatography in gas and liquid phases, capillary electrophoresis, microseparations combined with sensitive optical laser and/or electronic detection, and more, are slowly coming to fruition. *Micro chips* or microfluidic chemostats of the order of a few centimeters square to which inlet and outlet tubes can be connected and which have the electronics to open and shut intricate plumbing, mixing, and pneumatic-type valves designed for particular purposes are now already in use. The *chips* require fine tubes, special reservoirs, and pumps to move the fluids, and are made for various uses. There are already manufacturers who will custom-make *chips* to satisfy a variety of demands. The development of lithography and associated microelectronics and microelectrochemical systems (MEMS) originally on silicon and glass chips have found general use in microfluidics, but are slowly replaced by plastic or polymer materials for the platform. Some polymers such as poly(dimethylsiloxane) (PDMS) are optically transparent obviating the use of silicon. They are more malleable, more compressible and cheaper to machine to desired specifications. However, in some instances silicon chips are found to have advantages, as for example where organic fluids cause the plastic materials to swell or dissolve slowly.

Microfluidic devices have already found uses for sampling, identifying and assaying in the purification of chemicals and biochemical, the *chips* being connected to instruments used in chromatographic, electrophoretic, spectroscopic and monitoring devices. These in turn transfer their data to computers which process and/or store the data. It should be pointed out that the physics of the movement of fluids under pressure through such fine tubes as are present in the *chips* is quite different from the turbulent type that is found in millimeter and larger diameter tubes. In this case fluids are allowed to move and mix by running the fluid layers together in *laminar flow* movement [see Squires & Quake in ‘Microfluidics: Fluid physics at the nanoliter scale’ *Rev Mod Phys* **77** 977 2005, DOI: 10.1103/RevModPhys.77.977]. *Alternatively*, when the fluids are ionic as with aqueous solutions the microchannels to be used should have fixed charges on the inner walls (e.g. as with silica or surface oxidised PDMS) and an electrical potential is applied, so the fluid moves as a plug which is characteristic of electro-osmotic flow (EOF, see Santiago in ‘Electro-osmotic flows in microchannels with finite inertial and pressure forces’ *Anal Chem* **73** 2353 2001, DOI: 10.1021/ac0101398].

Apart from the applications made above, microfluidic devices have been made for screening conditions under

which proteins crystallise. These devices allow conditions of solvent concentrations and mix, temperature and its gradient to be specifically altered until crystals can be viewed within the wells in the microchannels [see Hansen et al. 'A robust and scalable microfluidic metering method that allows protein crystal growth by free interface diffusion' *Proc Natl Acad Sci USA* **99** 16531 2002, DOI: 10.1073/pnas.262485199; and Zheng et al. 'A droplet-based, composite PDMS/glass capillary microfluidic system for evaluating protein crystallization conditions by microbatch and vapor-diffusion methods with on-chip diffraction' *Angew Chem Int Ed* **43** 2508 2004, DOI: 10.1002/anie.200490056]. Appropriate microchips have been constructed for use in chemical syntheses can be carried out on a chip. For example, Snyder and coworkers [*Angew Chem Int Ed* **88** 1 2006, DOI: 10.1002/hlca.200490304] have described versatile modular microreaction systems for chemical syntheses which could involve homogeneous and heterogeneous catalysis, require moderate reaction times, can rapidly optimise conditions, can be cleaned for re-use, and can be later scalable from milligram to ton production amounts. A complex reaction sequence has been devised by Quake and coworkers [Lee et al. *Science* **310** 1793–1796 2005, DOI: 10.1126/science.1118919] for the synthesis, as an example, of the [^{18}F]-labeled molecular imaging probe 2-deoxy-2-[^{18}F]-fluoro-D-glucose {[^{18}F]FDG} using an integrated microfluidic chip. It involves the following five reaction steps: (i) concentration of a dilute aqueous solution of F^- ions by passage through a miniaturised ion-exchange column in the presence of K_2CO_3 to then form a complex with a cryptant [Kryptofix (K222)], (ii) solvent exchange from H_2O to dry MeCN, in the separate concentration circuit that is connected to the reaction loop, where (iii) fluorination of the sugar precursor (1,3,4,5-tetraacetyl-D-mannose-2-triflate) by the Kryptofix K^+F^- complex to produced 1,3,4,5-tetraacetyl-D-glucose-2-fluoride which is followed by (iv) solvent exchange back to H_2O and finally (v) hydrolysed in 3.0N aqueous HCl to give nanogram amounts of [^{18}F]FDG. The movement of fluids in the various channels is visualised by using different food dyes in the fluids. The actual size of the reaction platform is *ca* 20 x 20 mm. See 'Nanotechnology', Chapter 7, for examples of microfluidics for preparing and purifying nanomaterials for example from CdSe, PbS, Au, Pt, Ag, Au and TiO_2 .

The possibilities are almost infinite. A quick glance at any issue of the periodical *Lab Chip* reveals that large users of this technology are in the field of medical (and veterinary) diagnosis, screening, routine pathology, microbiology, etc, as new diseases and treatments are discovered and developed. The pharmaceutical industry will continuously exploit the need and use for *micro labchips* for drug evaluation, throughput screening as well as in drug synthesis and analysis. Mention should also be made that this technology furthers our knowledge of fluidics in science and engineering at the micro and nano scale.

There are manufacturers who will make *chip-platforms* to order and the cost can drop considerably if large disposable numbers are to be supplied. Some instrument manufacturers such as those that make DNA and RNA sequencers make their own *chips* for their requirements. Some research laboratories also make their own *chips* to meet their new needs. Once used, *chips* become contaminated and are invariably discarded because of difficulties in getting them pristinely clean, although some may be designed for re-use after appropriate treatment.

The bibliography in this field is ever increasing and new journals and periodicals are being published. The Royal Society of Chemistry (London) started a new journal entitled 'Laboratory on a Chip' abbreviated *Lab Chip* in 2001 and has a current Impact Factor of 6.115. A quick glance at the papers published in this journal reveals much about the various applications of this technology (a few titles are listed in 'selected publications' below). The ISSN and date of first issue of relevant periodicals include: *ACS Chem Mater* (ISSN: 0897-4756; 1989), *Biomicrofluidics* (ISSN: 1932-1058; 2007); *Microfluidics and Nanofluidics* (ISSN: 1613-4982; 2004); *BioChip Journal* (ISSN: 1976-0280 print; 2092-7843 online; 2010); *Biomedical Microdevices: bioMEMS and Biomedical Nanotechnology* (ISSN: 1387-2176 print; 1572-8781 online; 1998), *International Journal of Biomedical Nanoscience and Nanotechnology (IJBN)* (ISSN: 1756-0799 print; 1756-0802 online; 2010); *Environmental Nanotechnology, Monitoring and Management* (ISSN: 2215-1532, open access; 2014); *European Journal of Nanomedicine* (ISSN: 1662-596X; 2008). However, some papers in this field will also be found in journals on nanoscience, nanotechnology, e.g. in the journal *Nanoscale* (Impact Factor 7.349; ISSN: 2040-3364, first published in 2009 by the Royal Society of Chemistry, London), as well as in science journals such as *Nature*, *Science*, *PNAS*, the American Chemical Society and Royal Society of Chemistry publications.

Selected publications:

Whitesides 'The origins and the future of microfluidics' *Nature* **442** 368 2006, DOI: 10.1038/nature05058.

deMello 'Control and detection of chemical reactions in microfluidic systems' *Nature* **442** 394 2006, DOI: 10.1038/nature05062.

Chang, H.C., Yeo, Leslie. *Electrokinetically Driven Microfluidics and Nanofluidics*. Cambridge University Press 2009.

Research Highlights — ‘Petra S. Dittrich reviews the current literature in miniaturisation and related technologies’ *Lab Chip* **10** 1507 2010, DOI: 10.1039/C005267C.

Titmarsh et al. ‘Induction of Human iPSC-Derived Cardiomyocyte Proliferation Revealed by Combinatorial Screening in High Density Microbioreactor Arrays’ *Scientific Reports* **6** Article No 24367 2016, DOI:10.1038/srep24637.

Chen et al. ‘Rapid and inexpensive blood typing on thermoplastic chips’ *Lab Chip* **15** 4533 2015, DOI: 10.1039/C5LC01172H.

Volpatti & Yetisen ‘Commercialization of microfluidic devices’ *Trends in Biotechnology* **32**(7) 347 2014, DOI:10.1016/j.tibtech.2014.04.010.

Zhang et al. ‘Fundamentals and applications of inertial microfluidics: a review—a comprehensive review describing the fundamental mechanisms of inertial microfluidics, structure design and applications in biology, medicine and industry’ *Lab Chip* **16** 10 2016, DOI: 10.1039/C5LC01159K.

Zhuang et al. ‘A fully integrated and automated microsystem for rapid pharmacogenetic typing of multiple warfarin-related single-nucleotide polymorphisms — A fully integrated and automated microsystem consisting of disposable plastic chips for DNA extraction and PCR coupled with a reusable glass array-CE chip for rapid pharmacogenetic testing’ *Lab Chip* **16** 86 2016, DOI: 10.1039/C5LC01094B.

Lu et al. ‘Photochemical reactions and on-line UV detection in microfabricated reactors’ *Lab Chip* **1**, 22 2001, DOI: 10.1039/B104037P.

Minagawa et al. ‘Integration of a wet analysis system on a glass chip: determination of Co(II) as 2-nitroso-1-naphthol chelates by solvent extraction and thermal lens microscopy’ *Lab Chip* **1** 72 2001, DOI: 10.1039/B102790P.

Verma et al. ‘Smart material platforms for miniaturized devices: implications in disease models and diagnostics’ *Lab Chip*, 2016, DOI: 10.1039/C6LC00173D.

Herzog et al. ‘Continuous on-chip fluorescence labeling, free-flow isoelectric focusing and marker-free isoelectric point determination of proteins and peptides’ *Lab Chip* **16** 1565 2016, DOI: 10.1039/C6LC00055J.

Shembekar et al. ‘Droplet-based microfluidics in drug discovery, transcriptomics and high-throughput molecular genetics’ *Lab Chip* **16**, 1314 2016, DOI: 10.1039/C6LC00249H.

Hsieh & Kim, ‘A miniature closed-loop gas chromatography system—this work introduces a circulatory chromatography column system that adaptively magnifies the effective column length and the resultant separation capacity’ *Lab Chip*, **16**, 1002 2016, DOI: 10.1039/C5LC01553G.

Kim et al. ‘Pneumatically actuated microvalve circuits for programmable automation of chemical and biochemical analysis— This article reviews programmable microfluidic platforms using pneumatically actuated microvalve array and their applications in biological and chemical analysis’ *Lab Chip* **16** 812 2016, DOI:10.1039/C5LC01397F

CHEMICAL AND BIOCHEMICAL SOURCES

Apart from wishing to obtain a pure substance there are many reasons for wanting to purify a substance. For example the substance may have been in the store for too long and has deteriorated to a smaller or larger extent and needs to be used. Large quantities may be required, so bulk amounts, less pure but of cheaper grade could serve the purpose if they can be purified readily and cheaply. The cost consideration is very important. Substances that are available commercially can be of varying grades of purity and the purer the grade the higher the price. Biological substances may be only available in crude form, e.g. acetone powders for enzymes. There are a large number of suppliers of substances for chemical, biochemical and for biological requirements and they are continually improving quality, increasing their range and introducing recently developed substances. The following is a website list of the more commonly used suppliers from which almost all the substances and equipment described in this book can be purchased. The list also contains suppliers of laboratory ware as well as of scientific instruments. The list is not exhaustive.

USEFUL WEBSITES

http://www.

chemsupply.com.au [organics, inorganics & equipment]
sigmaaldrich.com/ [organics, inorganics, lifescience materials & equipment]
wilmad-labglass.com.aldrich.jsp [glass ware & equipment]
merck-chemicals.com/ [organics, inorganics & equipment]
acros.com/ [organics (Acros organics, inorganics & equipment)]
alfa.com/ [organics, inorganics & equipment]
strem.com/ [general inorganics, metal-organics, catalysts, nanomaterials]
tci-asiapacific.com/ [Tokyo Chemical Industry, chemicals, lab equipment]
thermofisher.com/global/en/home.asp [instruments, chemicals, custom products]
<https://au.vwr.com/app/Home> [VWR International- chemicals/laboratory scientific supplies]
quantum-scientific.com/ [chemical, biochemical & lab equipment]
gelificiences.com [GE Healthcare, chemicals, biochemicals & life science products]
bio-strategy.com [Laboratory technology]
glchina.com [GL biochemical products]
invitrogen.com [Invitrogen, life science products]
lifetechnologies.com [molecular biology products and equipment]
promega.com [Promega, life science products]
www.tocris.com [Tocris Bioscience products, i.e. neurochemicals, biochemical, peptides, DNA]
novachem.com.au
scilabware.com [plastic labware]
waters.com [Waters, chromatography materials]
biaseparations.com [chromatography materials]
[daicel.co.jp/ indexe.html](http://daicel.co.jp/indexe.html) [chromatography materials]
restekcorp.com [chromatography materials]
winlab.com.au/ [chromatography materials]
Fritsch-laser.com [for up to nano particle size and milling]
retsch-technology [for up to nano particle size]
haverstandard.com [for up to nano particle size]
perkinelmer.com [spectral and other instruments]
agilent.com/chem/atomicspec/ap
<http://www.betterworld.com> [books]
<http://www.abebooks.com> [books]
<http://www.booksandcollectibles.com.au/index> [books, collectibles]
<http://www.ebay.com.au/> [books, etc]

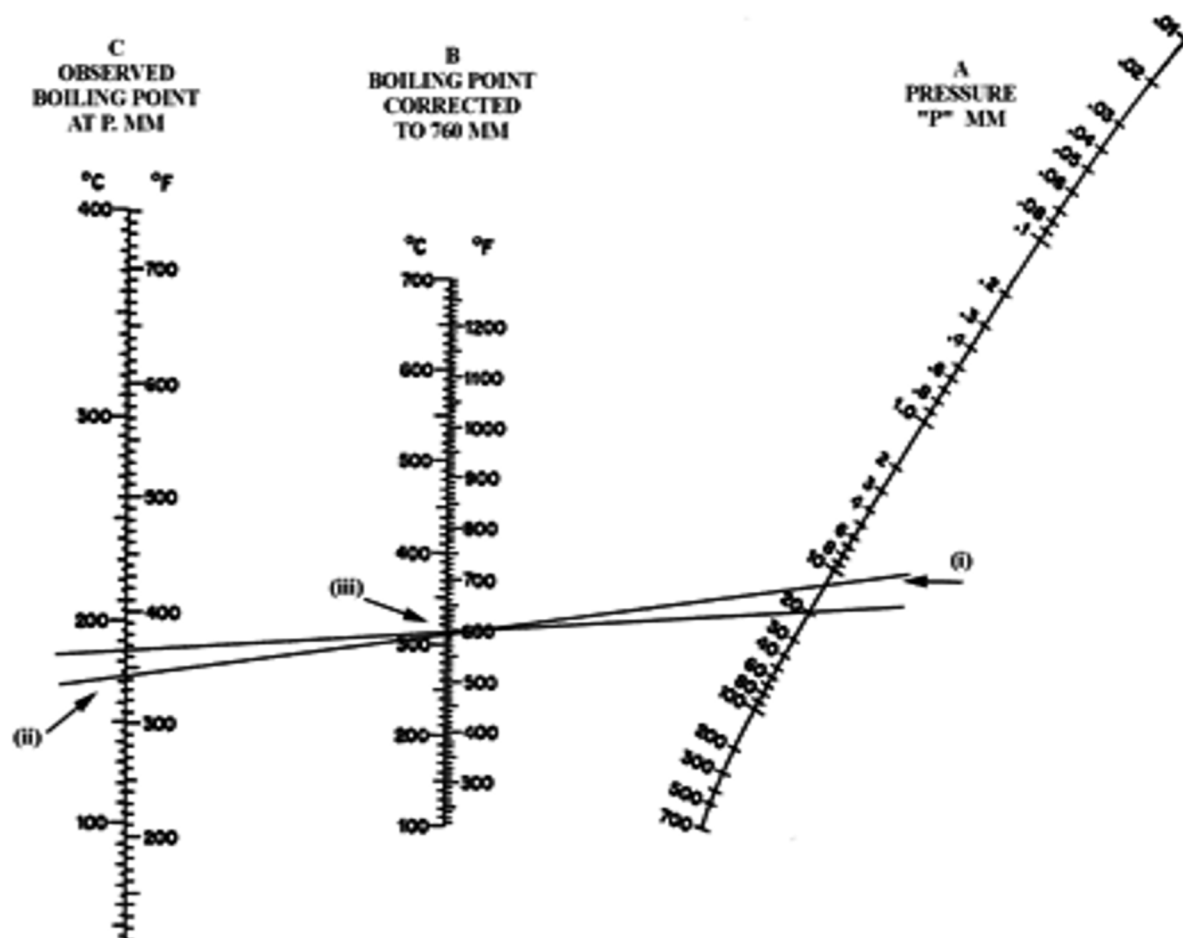
For high purity inorganic compounds, NIST Traceable inorganic reference standards/calibrants and aqueous standard solutions for ICP, ICP-MS, AA, GFAA and IC visit <www.exaxol.com>. Note that all the trace metal analyses in the 'Inorganic Compounds' section of Chapter 4 are by courtesy of Joe Papa (EXAXOL see Preface).

TABLES

TABLE 1. SOME COMMON IMMISCIBLE OR SLIGHTLY MISCIBLE
PAIRS OF SOLVENTS AT AMBIENT TEMPERATURES

Acetonitrile with hexane, heptane, iso-octane, cyclohexane.
Benzene with water, brine and aqueous solutions generally.
Butanol with water, brine and aqueous solutions generally.
Carbon tetrachloride with ethanolamine, ethylene glycol, formamide, water or brine.
Chloroform with glycerol, ethylene glycol, water, aqueous solutions generally
Cyclohexane with alcohols, dimethyl formamide, dimethyl sulfoxide, glycerol, pyridine.
Cyclopropyl methyl ether same as ethyl ether but less so.
Dimethyl formamide or dimethyl acetamide with cyclohexane, pentane, petroleum ether, xylene.
Dimethyl sulfoxide with ethyl ether, pentane, petroleum ethers, cyclohexane, xylene.
Ethyl acetate with aqueous solutions generally or petroleum ethers.
Ethyl ether with ethanolamine, dimethyl sulfoxide, ethylene glycol, glycerol, water or aqueous solutions generally.
Ethanol with carbon disulfide, petroleum ethers, cyclohexanes
Glycerol with benzene, ether, chloroform, carbon tetrachloride, carbon disulfide, petroleum ethers, oils
Iso-octane with acetonitrile, dimethyl formamide, dimethyl sulfoxide, methanol, water.
Methanol with carbon disulfide, hexane, heptane, cyclohexane or petroleum ethers.
N-Methylpyrrolidone with petroleum ethers, cyclohexanes.
Petroleum ether(s) with aniline, benzyl alcohol, dimethyl formamide, dimethyl sulfoxide, formamide, phenol or water and aqueous solutions generally.
Phenol with petroleum ethers, cyclohexanes.
Pyridine with petroleum ethers, hexanes.
Toluene with water, brine, aqueous solutions generally, glycerol but less so than benzene.
Water with aniline, benzene, benzyl alcohol, carbon disulfide, carbon tetrachloride, chloroform, cyclohexane, cyclohexanol, cyclohexanone, diethyl ether, ethyl acetate, isoamyl alcohol, methyl ethyl ketone, nitromethane, tributyl phosphate or toluene.
Xylene with water, brine, aqueous solutions generally, glycerol, dimethyl formamide, dimethyl sulfoxide.

FIGURE 1: NOMOGRAM



How to use Figure 1:

You can use a nomogram to estimate the boiling points of a substance at a particular pressure. For example, the boiling point of 4-methoxybenzenesulfonyl chloride is 173°C/14mm. Thus to find out what the boiling point of this compound will be at 760mm (atmospheric), draw a point on curve A (pressure) at 14mm (this is shown in (i)). Then draw a point on curve C (observed boiling point) corresponding to 173° (or as close as possible). This is shown in (ii). Using a ruler, find the point of intersection on curve B, drawing a line between points (i) and (ii). This is the point (iii) and is the boiling point of 4-methoxybenzenesulfonyl chloride (i.e. approx. 310°C) at atmospheric pressure. If you want to distil 4-methoxybenzenesulfonyl chloride at 20mm, then you will need to draw a point on curve A (at 20mm). Using a ruler, find the point of intersection on curve C drawing through the line intersecting (iii, curve B, i.e. 310°C) and the point in curve A corresponding to 20mm. You should have a value of 185°C; that is, the boiling point of 4-methoxybenzenesulfonyl chloride is estimated to be at 185°C at 20mm.

TABLE 2A. PREDICTED EFFECT OF PRESSURE ON BOILING POINT*

Temperature in degrees Centigrade										
760 mmHg	0	20	40	60	80	100	120	140	160	180
0.1	-111	-99	-87	-75	-63	-51	-39	-27	-15	-4
0.2	-105	-93	-81	-69	-56	-44	-32	-19	-7	5
0.4	-100	-87	-74	-62	-49	-36	-24	-11	2	15
0.6	-96	-83	-70	-57	-44	-32	-19	-6	7	20
0.8	-94	-81	-67	-54	-41	-28	-15	-2	11	24
1.0	-92	-78	-65	-52	-39	-25	-12	1	15	28
2.0	-85	-71	-58	-44	-30	-16	-3	11	25	39
4.0	-78	-64	-49	-35	-21	-7	8	22	36	51
6.0	-74	-59	-44	-30	-15	-1	14	29	43	58
8.0	-70	-56	-41	-26	-11	4	19	34	48	63
10.0	-68	-53	-38	-23	-8	7	22	37	53	68
14.0	-64	-48	-33	-23	-2	13	28	44	59	74
16.0	-61	-45	-29	-14	2	17	33	48	64	79
20.0	-59	-44	-28	-12	3	19	35	50	66	82
30.0	-54	-38	-22	-6	10	26	42	58	74	90
40.0	-50	-34	-17	-1	15	32	48	64	81	97
50.0	-47	-30	-14	3	19	36	52	69	86	102
60.0	-44	-28	-11	6	23	40	56	73	86	107
80.0	-40	-23	-6	11	28	45	62	79	97	114
100.0	-37	-19	-2	15	33	50	67	85	102	119
150.0	-30	-12	6	23	41	59	77	95	112	130
200.0	-25	-7	11	29	47	66	84	102	120	138
300.0	-18	1	19	38	57	75	94	113	131	150
400.0	-13	6	25	44	64	83	102	121	140	159
500.0	-8	11	30	50	69	88	108	127	147	166
600.0	-5	15	34	54	74	93	113	133	152	172
700.0	-2	18	38	58	78	98	118	137	157	177
750.0	0	20	40	60	80	100	120	140	160	180
770.0	0	20	40	60	80	100	120	140	160	180
800.0	1	21	41	61	81	101	122	142	162	182

* *How to use the Table:* Take as an example a liquid with a boiling point of 80°C at 760mm Hg. The Table gives values of the boiling points of this liquid at pressures from 0.1 to 800mm Hg. Thus at 50mm Hg this liquid has a boiling point of 19°C, and at 2mm Hg its boiling point would be -30°C.

TABLE 2B. PREDICTED EFFECT OF PRESSURE ON BOILING POINT*

Temperature in degrees Centigrade											
760mmHg	200	220	240	260	280	300	320	340	360	380	400
0.1	8	20	32	44	56	68	80	92	104	115	127
0.2	17	30	42	54	67	79	91	103	116	128	140
0.4	27	40	53	65	78	91	103	116	129	141	154
0.6	33	40	59	72	85	98	111	124	137	150	163
0.8	38	51	64	77	90	103	116	130	143	156	169
1.0	41	54	68	81	94	108	121	134	147	161	174
2.0	53	66	80	94	108	121	135	149	163	176	190
4.0	65	79	93	108	122	136	151	156	179	193	208
6.0	72	87	102	116	131	146	160	175	189	204	219
8.0	78	93	108	123	137	152	167	182	197	212	227
10.0	83	98	113	128	143	158	173	188	203	218	233
14.0	90	105	120	136	151	166	182	197	212	228	243
18.0	95	111	126	142	157	173	188	204	219	235	251
20.0	97	113	129	144	160	176	191	207	223	238	254
30.0	106	123	139	155	171	187	203	219	235	251	267
40.0	113	130	146	162	179	195	211	228	244	260	277
50.0	119	135	152	168	185	202	218	235	251	268	284
60.0	123	140	157	174	190	207	224	241	257	274	291
80.0	131	148	165	182	199	216	233	250	267	284	301
100.0	137	154	171	189	206	223	241	258	275	293	310
150.0	148	166	184	201	219	237	255	273	290	308	326
200.0	156	174	193	211	229	247	265	283	302	320	338
300.0	169	187	206	225	243	262	281	299	318	337	355
400.0	178	197	216	235	254	273	292	311	330	350	369
500.0	185	205	224	244	263	282	302	321	340	360	379
600.0	192	211	231	251	270	290	310	329	349	368	388
700.0	197	217	237	257	277	296	316	336	356	376	396
750.0	200	220	239	259	279	299	319	339	359	379	399
770.0	200	220	241	261	281	301	321	341	361	381	401
800.0	202	222	242	262	282	302	322	342	362	382	403

* *How to use the Table:* Taking as an example a liquid with a boiling point of 340°C at 760mm Hg, the column headed 340°C gives values of the boiling points of this liquid at each value of pressures from 0.1 to 800mm Hg. Thus, at 100mm Hg its boiling point is 258°C, and at 0.8mm Hg its boiling point will be 130°C.

TABLE 3. HEATING BATHS

Up to 100°	Water baths
-20 to 200°	Glycerol or di- <i>n</i> -butyl phthalate
Up to about 200°	Medicinal paraffin
Up to about 250°	Hard hydrogenated cotton-seed oil (m 40-60°) or a 1:1 mixture of cotton-seed oil and castor oil containing about 1% of hydroquinone.
-40 to 250° (to 400° under N ₂)	D.C. 550 silicone fluid
Up to about 260°	A mixture of 85% orthophosphoric acid (4 parts) and metaphosphoric acid (1 part)
Up to 340°	A mixture of 85% orthophosphoric acid (2 parts) and metaphosphoric acid (1 part)
60 to 500°	Fisher bath wax (highly unsaturated)
73 to 350°	Wood's Metal*
250 to 800°	Solder*
350 to 800°	Lead*

* In using metal baths, the container (usually a metal crucible) should be removed while the metal is still molten.

TABLE 4. WHATMAN FILTER PAPERS

Grade No.	1	2	3	4	5	6	113
Particle size retained (in microns)	11	8	5	12	2.4	2.8	28
Filtration speed*(sec/100ml)	40	55	155	20	<300	125	9

Routine ashless filters

Grade No.	40	41	42	43	44
Particle size retained (in microns)	7.5	12	3	12	4
Filtration speed* (sec/100ml)	68	19	200	38	125

Hardened**Hardened ashless**

Grade No.	50	52	54	540	541	542
Particle size retained(in microns)	3	8	20	9	20	3
Filtration speed* (sec/100ml)	250	55	10	55	12	250

Glass microfilters

Grade No	GF/A	GF/B	GF/C	GF/D	GF/F
Particle size retained (in microns)	1.6	1.0	1.1	2.2	0.8
Filtration speed (sec/100ml)*	8.3	20.0	8.7	5.5	17.2

*Filtration speeds are rough estimates of initial flow rates and should be considered on a relative basis.

TABLE 5. MICRO FILTERS*

Nucleopore (polycarbonate) Filters						
Mean Pore Size (microns)	8.0	2.0	1.0	0.1	0.03	0.015
Av. pores/cm ²	10 ⁵	2x10 ⁶	2x10 ⁷	3x10 ⁸	6x10 ⁸	1-6x10 ⁹
Water flow rate(ml/min/cm ²)	2000	2000	300	8	0.03	0.1-0.5
Millipore Filters						
Type	—Cellulose ester—		—Teflon—		—Microweb [#] —	
	MF/SC	MF/VF	LC	LS	WS	WH
Mean Pore Size (microns)	8	0.01	10	5	3	0.45
Water flow rate (ml/min/cm ²)	850	0.2	170	70	155	55
Gelman Membranes						
Type	—Cellulose ester—		—Copolymer—			
	GA-1	T CM-450	VM-1	DM-800	AN-200	Tuffryn-450
Mean Pore Size (microns)	5	0.45	5	0.8	0.2	0.45
Water flow rate (ml/min/cm ²)	320	50	700	200	17	50
Sartorius Membrane Filters (SM)						
Application	Gravi-metric	Biological clarification	Sterilisation	Particle count in H ₂ O	For acids & bases	
Type No.	11003	11004	11006	11011	12801	
Mean PoreSize (microns)	1.2	0.6	0.45	0.01	8.0	
Water flow rate (ml/min/cm ²)	300	150	65	0.6	1100	

* Only a few representative filters are tabulated (available ranges are more extensive). # Reinforced nylon.

TABLE 6. COMMON SOLVENTS USED IN RECRYSTALLISATION

Acetic acid (118°)	*Cyclohexane (81°)	*Methanol (64.5°)
*Acetone (56°)	Dichloromethane (41°)	*Methyl ethyl ketone (80°)
Acetylacetone (139°)	*Diethyl ether (34.5°)	Methyl isobutyl ketone (116°)
Acetonitrile (82°C)	Dimethyl formamide (76°/39mm)	Nitrobenzene (210°)
*Benzene (80°)	*Dioxane (101°)	Nitromethane (101°)
Benzyl alcohol (93°/10mm)	*Ethanol (78°)	*Petroleum ether (various)
<i>n</i> -Butanol (118°)	2-Ethoxyethanol (cellosolve 135°)	Pyridine (115.5°)
Butyl acetate (126.5°)	*Ethyl acetate (78°)	Pyridine trihydrate (93°)
<i>n</i> -Butyl ether (142°)	Ethyl benzoate (98°/19mm)	*Tetrahydrofuran (64-66°)
γ -Butyrolactone (206°)	Ethylene glycol (68°/4mm)	Toluene (110°)
Carbon tetrachloride (77°)	Formamide (110°/10mm)	Trimethylene glycol (59°/11mm)
Chlorobenzene (132°)	Glycerol (126°/11mm)	Water (100°)
Chloroform (61°)	Isoamyl alcohol (131°)	Xylenes (<i>o</i> 143-145°, <i>m</i> 138-139°, <i>p</i> 138°)

*Highly flammable, should be heated or evaporated on steam or electrically heated water baths only (preferably under nitrogen). None of these solvents should be heated over a naked flame.

TABLE 7. PAIRS OF MISCIBLE SOLVENTS

Acetic acid: with chloroform, ethanol, ethyl acetate, acetonitrile, petroleum ether, or water.
Acetone: with benzene, butyl acetate, butyl alcohol, carbon tetrachloride, chloroform, cyclohexane, ethanol, ethyl acetate, methyl acetate, acetonitrile, petroleum ether or water.
Ammonia: with ethanol, methanol, pyridine.
Aniline: with acetone, benzene, carbon tetrachloride, ethyl ether, <i>n</i> -heptane, methanol, acetonitrile or nitrobenzene.
Benzene: with acetone, butyl alcohol, carbon tetrachloride, chloroform, cyclohexane, ethanol, acetonitrile, petroleum ether or pyridine.
Butyl alcohol: with acetone or ethyl acetate.
Carbon disulfide: with petroleum ether.
Carbon tetrachloride: with cyclohexane.
Chloroform: with acetic acid, acetone, benzene, ethanol, ethyl acetate, hexane, methanol or pyridine.
Cyclohexane: with acetone, benzene, carbon tetrachloride, ethanol or diethyl ether.
Diethyl ether: with acetone, cyclohexane, ethanol, methanol, methylal (dimethoxymethane), acetonitrile, pentane or petroleum ether.
Dimethyl formamide: with benzene, ethanol or ether.
Dimethyl sulfoxide: with acetone, benzene, chloroform, ethanol, diethyl ether or water.
Dioxane: with benzene, carbon tetrachloride, chloroform, ethanol, diethyl ether, petroleum ether, pyridine or water.
Ethanol: with acetic acid, acetone, benzene, chloroform, cyclohexane, dioxane, ethyl ether, pentane, toluene, water or xylene.
Ethyl acetate: with acetic acid, acetone, butyl alcohol, chloroform, or methanol.
Glycerol: with ethanol, methanol or water.
Hexane: with benzene, chloroform or ethanol.
Methanol: with chloroform, diethyl ether, glycerol or water.
Methylal: with diethyl ether.
Methyl ethyl ketone: with acetic acid, benzene, ethanol or methanol.
Nitrobenzene: with aniline, methanol or acetonitrile.
Pentane: with ethanol or diethyl ether.
Petroleum ether: with acetic acid, acetone, benzene, carbon disulfide or diethyl ether.
Phenol: with carbon tetrachloride, ethanol, diethyl ether or xylene.
Pyridine: with acetone, ammonia, benzene, chloroform, dioxane, petroleum ether, toluene or water.
Toluene: with ethanol, diethyl ether or pyridine.
Water: with acetic acid, acetone, ethanol, methanol, or pyridine.
Xylene: with ethanol or phenol.

TABLE 8. MATERIALS FOR COOLING BATHS

Temperature	Composition	Temperature	Composition
0°	Crushed ice	-72°	Solid CO ₂ with ethanol
-5° to -20°	Ice-salt mixtures	-77°	Solid CO ₂ with chloroform or acetone
Up to -20°	Ice-MeOH mixtures	-78°	Solid CO ₂ (powdered; CO ₂ snow)
-33°	Liquid ammonia	100°	Solid CO ₂ with diethyl ether
-40° to -50°	Ice (3.5-4 parts) - CaCl ₂ 6H ₂ O (5 parts)	-196°	liquid nitrogen (see footnote*)

13°	<i>p</i> -Xylene	-55°	Diacetone
12°	Dioxane	-56°	<i>n</i> -Octane
6°	Cyclohexane	-60°	Di-isopropyl ether
5°	Benzene	-73°	Trichloroethylene or isopropyl acetate
2°	Formamide	-74°	<i>o</i> -Cymene or <i>p</i> -cymene
-8.6°	Methyl salicylate	-77°	Butyl acetate
-9°	Hexane-2,5-dione	-79°	Isoamyl acetate
-10.5°	Ethylene glycol	-83°	Propylamine
-11.9°	<i>tert</i> -Amyl alcohol	-83.6°	Ethyl acetate
-12°	Cycloheptane or methyl benzoate	-86°	Methyl ethyl ketone
-15°	Benzyl alcohol	-89°	<i>n</i> -Butanol
-16.3°	<i>n</i> -Octanol	-90°	Nitroethane
-18°	1,2-Dichlorobenzene	-91°	Heptane
-22°	Tetrachloroethylene	-92°	<i>n</i> -Propyl acetate
-22.4°	Butyl benzoate	-93°	2-Nitropropane or cyclopentane
-22.8°	Carbon tetrachloride	-94°	Ethyl benzene or hexane
-24.5°	Diethyl sulfate	-94.6°	Acetone
-25°	1,3-Dichlorobenzene	-95.1°	Toluene
-29°	<i>o</i> -Xylene or pentachloroethane	-97°	Cumene
-30°	Bromobenzene	-98°	Methanol or methyl acetate
-32°	<i>m</i> -Toluidine	-99°	Isobutyl acetate
-32.6°	Dipropyl ketone	-104°	Cyclohexene
-38°	Thiophene	-107°	Isooctane
-41°	Acetonitrile	-108°	1-Nitropropane
-42°	Pyridine or diethyl ketone	-116°	Ethanol or diethyl ether
-44°	Cyclohexyl chloride	-117°	Isoamyl alcohol
-45°	Chlorobenzene	-126°	Methylcyclohexane
-47°	<i>m</i> -Xylene	-131°	<i>n</i> -Pentane
-50°	Ethyl malonate or <i>n</i> -butylamine	-160°	Isopentane
-52°	Benzyl acetate or diethylcarbitol		

For other organic materials used in low temperature slush-baths with liquid nitrogen see R.E.Rondeau [*J Chem Eng Data* **11** 124 1966, DOI: 10.1021/je60028a037]. *NOTE: Use high quality pure nitrogen; do not use liquid air or liquid nitrogen that has been in contact with air for a long period (due to the dissolution of oxygen in it) as this could EXPLODE in contact with organic matter.

TABLE 9. LIQUIDS FOR STATIONARY PHASES IN GAS CHROMATOGRAPHY*

Material	Temp.	Retards
Dimethylsulfolane	0-40°	Olefins and aromatic hydrocarbons
Di- <i>n</i> -butyl phthalate	0-40°	General purposes
Squalane	0-150°	Volatile hydrocarbons and polar molecules
Silicone oil or grease	0-250°	General purposes
Diglycerol	20-120°	Water, alcohols, amines, esters, and aromatics
Dinonyl phthalate	20-130°	General purposes
Polydiethylene glycol succinate	50-200°	Aromatic hydrocarbons, alcohols, ketones, esters
Polyethylene glycol	50-200°	Water, alcohols, amines, esters and aromatics
Apiezon grease	50-200°	Volatile hydrocarbons and polar molecules
Tricresyl phosphate	50-250°	General purposes

* See Suppliers at the end of section on HPLC and E.F. Barry and R.L. Grob, *Columns for Gas Chromatography*, J. Wiley and Sons NY, 2007, ISBN 9780471740438.

TABLE 10. METHODS OF VISUALISATION OF TLC SPOTS

Reagent	Compound	Preparation	Observations
Iodine	General	Iodine crystals in a closed chamber or spray 1% methanol solution of Iodine	Brown spots which may disappear upon standing.
H ₂ SO ₄	General	50% solution, followed by heating to 150°C	Black or coloured spots
Molybdate	General	5% (NH ₄) ₆ Mo ₇ O ₂₄ + 0.2% Ce(SO ₄) ₂ in 5% H ₂ SO ₄ , followed by heating to 150°C.	Deep blue spots
Vanillin	General	0.5g vanillin, 0.5 ml H ₂ SO ₄ , 9 ml ethanol	various coloured spots
Ammonia	phenols	Ammonia vapour in a closed chamber	various coloured spots
FeCl ₃	phenols, enols	1% aqueous FeCl ₃	various coloured spots
2,4-DNP	aldehydes, ketones	0.5% 2,4-dinitrophenylhydrazine/2M HCl	red to yellow spots
HCl	aromatic acids and amines	HCl vapour in a closed chamber	various coloured spots
Ninhydrin	amino acids, and amines	0.3% ninhydrin in <i>n</i> -BuOH with 3% AcOH, followed by heating to 125°C/10 min	blue spots
PdCl ₂	S and Se compds	0.5% aq. PdCl ₂ + few drops of conc. HCl	red and yellow spots
Anisaldehyde	carbohydrates	0.5 ml anisaldehyde in 0.5 ml conc H ₂ SO ₄ + 95% EtOH + a few drops of AcOH Heat at 100-110°C for 20-30 minutes	various blue spots

TABLE 11. GRADED ADSORBENTS AND SOLVENTS FOR CHROMATOGRAPHY

Adsorbents (decreasing effectiveness)	Solvents (increasing eluting ability)
----------------------------------------------	----------------------------------------------

Fuller's earth (hydrated aluminosilicate) Magnesium oxide Charcoal Alumina Magnesium trisilicate Silica gel Calcium hydroxide Magnesium carbonate Calcium phosphate Calcium carbonate Sodium carbonate Talc Inulin Sucrose≈starch	Petroleum ether, b. 40-60°. Petroleum ether, b. 60-80°. Carbon tetrachloride. Cyclohexane. Benzene. Diethyl ether. Chloroform. Ethyl acetate. Acetone. Ethanol. Methanol. Pyridine. Acetic acid. Acetic acid.
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TABLE 12. REPRESENTATIVE ION-EXCHANGE RESINS USED IN CHROMATOGRAPHY

Sulfonated polystyrene Strong-acid cation exchanger AG 50W-x8 Amberlite IR-120 Dowex 50W-x8 Duolite 225 Permutit RS Permutite C50D Carboxylic acid-type Weak acid cation exchangers Amberlite IRC-50 Bio-Rex 70 Chelex 100 Duolite 436 Permutit C Permutits H and H-70	Aliphatic amine-type weak base anion exchangers Amberlites IR-45 and IRA-67 Dowex 3-x4A Permutit E Permutit A 240A Strong Base, anion exchangers AG 2x8 Amberlite IRA-400 Dowex 2-x8 Duolite 113 Permutit ESB Permutite 330D
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TABLE 13. MODIFIED FIBROUS CELLULOSES FOR ION-EXCHANGE CHROMATOGRAPHY

Cation exchange CM cellulose (carboxymethyl) CM 22, 23 cellulose P cellulose (phosphate) SE cellulose (sulfoethyl) SM cellulose (sulfomethyl)	Anion exchange DEAE cellulose (diethylaminoethyl) DE 22, 23 cellulose PAB cellulose (<i>p</i> -aminobenzyl) TEAE cellulose (triethylaminoethyl) ECTEOLA cellulose
---------------------------------------------------------------------------------------------------------------------------------------------------------------------	------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------

SE and SM are much stronger acids than CM, whereas P has two ionisable groups (pK 2-3, 6-7), one of which is stronger, the other weaker, than for CM (3.5-4.5). For basic strengths, the sequence is: TEAE » DEAE (pK 8-9.5) > ECTEOLA (pK 5.5-7) > PAB. Their exchange capacities lie in the range 0.3 to 1.0 mg equiv/g.

TABLE 14. BEAD FORM ION-EXCHANGE PACKAGINGS FOR CHROMATOGRAPHY¹

Cation exchange	Capacity (meq/g)	Anion exchange	Capacity (meq/g)
CM-Sephadex C-25, C-50. ² (weak acid)	4.5±0.5	DEAE-Sephadex A-25, A-50. ⁷ (weak base)	3.5±0.5
SP-Sephadex C-25, C-50. ³ (strong acid)	2.3±0.3	QAE-Sephadex A-25, A-50. ⁸ (strong base)	3.0±0.4
CM-Sepharose CL-6B. ⁴	0.12±0.02	DEAE-Sepharose CL-6B. ⁴	0.13±0.02
		DEAE-Sephacel. ⁹	1.4±0.1
Fractogel EMD, CO ₂ ⁻ (pK ~ 4.5), SO ₃ ²⁻ (pK ~ <1). ⁵		Fractogel EMD, DMAE (pK ~9), DEAE (pK ~10.8), TMAE (pK >13). ⁵	
CM-32 Cellulose.		DE-32 Cellulose.	
CM-52 Cellulose. ⁶		DE-52 Cellulose	

¹May be sterilised by autoclaving at pH 7 and below 120°. ²Carboxymethyl. ³Sulfopropyl. ⁴Crosslinked agarose gel, no pre-cycling required, pH range 3-10. ⁵Hydrophilic methacrylate polymer with very little volume change on change of pH (equivalent to *Toyopearl*, Sigma), available in superfine 650S, and medium 650M particle sizes. ⁶Microgranular, pre-swollen, no pre-cycling required. ⁷Diethylaminoethyl. ⁸Diethyl(2-hydroxy-propyl)aminoethyl. ⁹Bead form cellulose, pH range 2-12, no pre-cycling required. Sephadex and Sepharose are from GE Healthcare, Fractogel is from Merck and Cellulose is from Whatman.

TABLE 15 SELECTED CHIRAL COLUMNS FOR CHROMATOGRAPHY*

Column name	Chiral ligand	Attributes	Manufacture
CHIRA-chrom-1	<i>R</i> - or <i>S</i> - phenylglycine	High capacity	HICHROM
CHIRA-chrom-2	<i>S</i> -leucine	& efficiency	
CHIRAL-AGP	dinitrophenyl tartramide	Used in wide	CHROM TECH
CHIRAL-CBH	α 1-acid glycoprotein	pH range	
CHIRAL-HAS	cellobiohydrolase		
CHIRALCEL OD/OJ	human serum albumin		
CHIRALPAK AD/AS/H	cellulose derivative	Particular separations	DIACEL
CHIRALPAK 1A	amylose derivative	versatile	
CHIRALPAK 1B	immobilised amylose	broad range	
CROWNPAK	immobilised cellulose	broad range	
	18-crown 6 type ether	for amino acids and primary amines	
CHIROFIL	(18-crown 6)(CO ₂ H) ₄	for amino acids and primary amines	RStech

(Continued)

TABLE 15 (Continued) SELECTED CHIRAL COLUMNS FOR CHROMATOGRAPHY*

Column name	Chiral ligand	Attributes	Manufacture
CHIROBIOTIC R	Ristocetin A	Broad selectivity	Astech 1
CHIROBIOTIC T	Teicoplanin	Broad selectivity	
CHIROBIOTIC V	Vancomycin	Broad selectivity	
CYCLOBOND I	β -cyclodextrin	form chiral	Astech 1
CYCLOBOND II	γ -cyclodextrin	inclusion complexes	
KROMASIL DMB	Acetylated <i>N,N'</i> -diallyl	stable, high capacity	Eka Chemicals
KROMASIL TBB	<i>S</i> -tartardiamide	for large prep work	
NUCLEODEX β -OH	β -cyclodextrin	reverse phase work	Macherey— Nagel
NUCLEODEX α -, β - and γ -PM	permethylated α -, β - and γ -cyclodextrins	reverse phase work	
NUCLEOSIL Chiral-1	<i>S</i> -hydroxyproline-Cu ²⁺ complex	reverse phase work	
		ligand exchange, e.g. α -amino acids	
RESOLVOSIL BSA-7PX	bovine serum albumen	various applications	Shinwa Chem Industries
ULTRON ES-OVM	ovomucoid protein	stable phase	
ULTRON ES-Pepsin	pepsin protein	stable phase	REGIS
DACH-DNB	3,5-dinitrobenzoyl derivs	π -electron acceptor/ donor — widely used	
ULMO	3,5-dinitrobenzoyl derivs	π -electron acceptor	
α -Burke 2	3,5-dinitrobenzoyl derivs	π -electron acceptor	
β -GEM 1	3,5-dinitrobenzoyl derivs	π -electron acceptor	
Leucine	3,5-dinitrobenzoyl derivs	π -electron acceptor	
Phenyglycine	β -lactamase chiral selector	π -electron acceptor	
PIRKLE-1J	<i>S</i> -naphthylleucine	π -electron donor	
Naphthylleucine	ligand exchange	good for underivatised	
DAVAKOV	ligand exchange	amino acids	

* These data were generously provided by Gordon Wingate, Operational Director of Winlab Pty Ltd., POBox 5007, Brendale, Queensland 4500, Australia. Further details about these and other chromatographic columns may be obtained from him, and at < <http://www.winlab.com.au/> >.

TABLE 16. LIQUIDS FOR ABDERHALDEN DRYING PISTOLS

Boiling points (760mm)		Boiling points (760mm)	
Ethyl chloride	12.2°	Toluene	110.5°
Dichloromethane	39.8°	Tetrachloroethylene	121.2°
Acetone	56.1°	Chlorobenzene	132.0°
Chloroform	62.0°	<i>m</i> -Xylene	139.3°
Methanol	64.5°	Isoamyl acetate	142.5°
Carbon tetrachloride	76.5°	Tetrachloroethane	146.3°
Ethanol	78.3°	Bromobenzene	155.0°
Benzene	79.8°	<i>p</i> -Cymene	176.0°
Trichloroethylene	86.0°	<i>o</i> -Chlorobenzene	180.5°
Water	100.0°	Tetralin	207.0°

TABLE 17.

**VAPOUR PRESSURES (mm Hg) OF SATURATED AQUEOUS
SOLUTIONS IN EQUILIBRIUM WITH SOLID SALTS**

Salt	Temperature					% Humidity at 20°
	10°	15°	20°	25°	30°	
LiCl.H ₂ O			2.6			15
CaBr ₂ .6H ₂ O	2.1	2.7	3.3	4.0	4.8	19
KOAc			3.5			20
CaCl ₂ .6H ₂ O	3.5	4.5	5.6	6.9	8.3	20
CrO ₃			6.1			32
Zn(NO ₃) ₂ .6H ₂ O			7.4			42
K ₂ CO ₃ .2H ₂ O			7.7	10.7		44
KCNS			8.2			47
Na ₂ Cr ₂ O ₇ .2H ₂ O			9.1			52
Ca(NO ₃) ₂ .4H ₂ O	6.0	7.7	9.6	11.9	14.2	55
Mg(NO ₃) ₂ .6H ₂ O			9.8			56
NaBr.2H ₂ O	5.8	7.8	10.3	13.5	17.5	58
NaNO ₂			11.6			66
NaCl	6.9	9.6	13.2	17.8	21.4	75
NaOAc			13.3			76
NH ₄ Cl			13.8			79
(NH ₄) ₂ SO ₄			14.2			81
KBr			14.7			84
KHSO ₄			15.1			86
KCl			15.1	20.2	27.0	86
K ₂ CrO ₄			15.4			88
ZnSO ₄ .7H ₂ O			15.8			90
NH ₄ .H ₂ PO ₄			16.3			93
KNO ₃			16.7	22.3	29.8	95
Pb(NO ₃) ₂			17.2			98
H ₂ O	9.21	12.79	17.53	23.76	31.82	100

TABLE 18. DRYING AGENTS FOR CLASSES OF COMPOUNDS

Class	Dried with
Acetals	K ₂ CO ₃ .
Acids (organic)	CaSO ₄ , MgSO ₄ , Na ₂ SO ₄ .
Acyl halides	MgSO ₄ , Na ₂ SO ₄ .
Alcohols	CaO, CaSO ₄ , MgSO ₄ , K ₂ CO ₃ , followed by Mg and I ₂ .
Aldehydes	CaSO ₄ , MgSO ₄ , Na ₂ SO ₄ .
Alkanes and alkenes	MgSO ₄ , CaCl ₂ , CaSO ₄ , H ₂ SO ₄ , P ₂ O ₅ .
Alkyl halides	CaCl ₂ , CaSO ₄ , MgSO ₄ , P ₂ O ₅ , Na ₂ SO ₄ , H ₂ SO ₄ .
Amines	BaO, CaO, KOH, NaOH, Na ₂ CO ₃ .
Aryl halides	CaCl ₂ , CaSO ₄ , MgSO ₄ , Na ₂ SO ₄ , P ₂ O ₅ .
Esters	MgSO ₄ , K ₂ CO ₃ , Na ₂ SO ₄ .
Ethers	CaCl ₂ , CaSO ₄ , MgSO ₄ , Na, LiAlH ₄ .
Heterocyclic bases	MgSO ₄ , K ₂ CO ₃ , KOH.
Hydrocarbons	CaCl ₂ , CaSO ₄ , MgSO ₄ , P ₂ O ₅ , Na (not for olefins).
Ketones	CaSO ₄ , MgSO ₄ , K ₂ CO ₃ , Na ₂ SO ₄ .
Mercaptans	MgSO ₄ , Na ₂ SO ₄ .
Nitro compounds and nitriles	CaCl ₂ , MgSO ₄ , Na ₂ SO ₄ .
Sulfides	CaCl ₂ , CaSO ₄ . [See also Molecular Sieves in text above]

TABLE 19.		STATIC DRYING FOR SELECTED LIQUIDS (25°C)			
Liquid	Water	Linde Type 4 A	Linde Type 5 A	Activated Alumina	Silicic Acid Gel
MeOH	Residual H ₂ O %	0.54	0.55	—	0.60
	Wt % absorbed	2.50	1.50	—	—
EtOH	Residual H ₂ O %	0.25	0.25	0.45	0.68
	Wt % absorbed	7.00	6.80	1.50	—
1-Butylamine	Residual H ₂ O %	1.65	1.31	1.93	2.07
	Wt % absorbed	10.40	18.20	3.40	—
2-Ethyl- hexylamine	Residual H ₂ O %	0.25	0.08	0.43	0.53
	Wt % absorbed	15.10	21.10	6.10	1.70
Diethyl ether	Residual H ₂ O %	0.001	0.013	0.16	0.27
	Wt % absorbed	9.50	9.20	6.20	4.30
Amyl acetate	Residual H ₂ O %	0.002	—	0.33	0.38
	Wt % absorbed	9.30	—	7.40	1.80

TABLE 20

FLUOROCHROMES*

	CAS registry No	λ_{ex} (nm)#	λ_{em} (nm)#	Applications
Acriflavin	[8048-52-0]	463	490	Stains many compounds
Allophycocyanin (APC)		604 (650)	650 (660)	Polypeptide with exceptional Fluorescence for cytometry
3-Aminocoumarin	[1635-31-0]	350	445	Forms fluorescent derivatives <i>via</i> reaction with the 3-NH ₂ group
Auramine O (basic yellow 2)	[2465-27-2]	440	530	Fluorescent stain for bacteria and probe for certain enzymes, e.g. alcohol dehydrogenases
4-Bromomethyl-7- methoxycoumarin	[35231-44-8]	360 (340)	410 (421)	Forms fluorescent probes with various compds for TLC/HPLC
Chromomycin A ₃	[7059-24-7]	445	575	Stains DNA
Ethidium bromide	[1239-45-8]	493	620	Stains DNA
Fluorescamine	[38183-12-9]			Reacts with various R-NH ₂ to form fluorescent compds
Fluorescein	[2321-07-5]	490	514	Fluorescent nucleus for staining a Variety of compounds
Fluorescein- Isothiocyanate (FITC)	[3326-32-7]	492	518	Microsequencing of peptides & proteins, makes fluorescent Amino acids
Hoechst H 2495 (benzoxanthene yellow)	[72845-94-4]			Fluorochrome for various compds
Hoechst H33342 (bisBenzimide 3HCl)	[23491-52-3]	346	640	Intercalates DNA for cytometry and fluorescence microscopy
R-Phycoerythrin	[11016-17-4]	488	572	Forms fluorescent probes with proteins
PKH 2 and 72 (green)		490	504	A Protein Kinase polypeptide that links to cells causing fluorescence
PKH 26 (red)		557	567	Used for labeling cells (cf PKH 2 or 72)
Tetramethyl rhodamine Isothiocyanate (TRITC)	[95197-95-8]	529 (492)	596 (518)	Protein fluorochrome, used for immunofluorescence

** Of the many fluorochromes reported in this book only a very small selection is tabulated here. # These wavelengths may vary somewhat depending on the solvent used, and on the pH if measured in aqueous solutions.

TABLE 21. RESIDUAL SOLVENT SIGNALS OF COMMON NMR SOLVENTS(Adapted from Gottlieb et al. *J Org Chem* **62** 7512 1997, DOI: 10.1021/jo971176v; Fulmer et al. *Organometallics* **29**, 2176 2010, DOI: 10.1021/om100106e).

Solvents	¹ H NMR	¹³ C NMR
Deuterated chloroform, CDCl ₃	7.26	77.16
Deuterated dichloromethane CD ₂ Cl ₂	5.32	53.84
Deuterated benzene, C ₆ D ₆	7.16	128.06
Deuterated acetone, (CD ₃) ₂ CO	2.05	29.84; 206.26
Deuterated dimethylsulfoxide, (CD ₃) ₂ SO	2.50	35.52
Deuterated acetonitrile, CD ₃ CN	1.94	1.32; 118.26
Deuterated methanol, CD ₃ OD	3.31	49.00
Deuterated water, D ₂ O	4.79	-

TABLE 22. ¹H NMR CHEMICAL SHIFTS OF TRACE IMPURITIES OF COMMON SOLVENTS AND REAGENTS

	CDCl ₃	CD ₂ Cl ₂	C ₆ D ₆	(CD ₃) ₂ CO	(CD ₃) ₂ SO	CD ₃ CN	CD ₃ OD	D ₂ O
H ₂ O	1.56	1.52	0.40	2.84	3.33	2.13	4.87	4.79
Acetone	2.17	2.12	1.55	2.05	2.09	2.09	2.15	2.22
Acetonitrile	2.10	1.97	0.58	2.05	2.07	1.94	2.03	2.06
Chloroform	7.26	7.32	6.15	8.02	8.32	7.58	7.90	-
Dichloromethane	5.30	5.33	4.27	5.63	5.76	5.44	5.49	-
Diethyl ether	1.21 (t), 3.48 (q)	1.15 (t), 3.43 (q)	1.11 (t), 3.26 (q)	1.11 (t), 3.41 (q)	1.09 (t), 3.38 (q)	1.12 (t), 3.42 (q)	1.18 (t), 3.49 (q)	1.17 (t), 3.56 (q)
DMF	8.02, 2.96, 2.88	7.96, 2.91, 2.82	7.63, 2.36, 1.86	7.96, 2.94, 2.78	7.95, 2.89, 2.73	7.92, 2.89, 2.77	7.97, 2.99, 2.86	7.92, 3.01, 2.85
DMSO	2.62		1.68	2.52	2.50	2.50	2.65	2.71
Dioxane	3.71	3.65	3.35	3.59	3.57	3.60	3.66	3.75
Ethanol*	1.25 (t), 3.72 (q), 1.32 (s, OH)	1.19 (t), 3.66 (q), 1.33 (s, OH)	0.96 (t), 3.34 (q), 0.50 (s, OH)	1.12 (t), 3.57 (q), 3.39 (s, OH)	1.06 (t), 3.44 (q), 4.63 (s, OH)	1.12 (t), 3.54 (q), 2.47 (s, OH)	1.19 (t), 3.60 (q)	1.17 (t), 3.65 (t)
Ethyl acetate	2.05 (s), 4.12 (q), 1.26 (t)	2.00 (s), 4.08 (q), 1.23 (t)	1.65 (s), 3.89 (q), 0.92 (t)	1.97 (s), 4.05 (q), 1.20 (t)	1.99 (s), 4.03 (q), 1.17 (t)	1.97 (s), 4.06 (q), 1.20 (t)	2.01 (s), 4.09 (q), 1.24 (t)	2.07 (s), 4.14 (q), 1.24 (t)
Methanol*	3.49 (s), 1.09 (s, OH)	3.42 (s), 1.09 (s, OH)	3.07 (s)	3.31 (s), 3.12 (s, OH)	3.16 (s), 4.01 (s, OH)	3.28 (s), 2.16 (s, OH)	3.31	3.34
pyridine	8.62 (m), 7.29 (m), 7.68 (m)	8.59 (m), 7.28 (m), 7.68 (m)	8.53 (m), 6.66 (m), 6.98 (m)	8.58 (m), 7.35 (m), 7.76 (m)	8.58 (m), 7.39 (m), 7.79 (m)	8.57 (m), 7.33 (m), 7.73 (m)	8.53 (m), 7.44 (m), 7.85 (m)	8.52 (m), 7.45 (m), 7.87 (m)
Silicon grease	0.07 (s)	0.09 (s)	0.29 (s)	0.13 (s)	-0.06 (s)	0.08	0.10	-
THF	3.76 (m), 1.85 (m)	3.69 (m), 1.82 (m)	3.57 (m), 1.40 (m)	3.63 (m), 1.79 (m)	3.60 (m), 1.76 (m)	3.64 (m), 1.80 (m)	3.71 (m), 1.87 (m)	3.74 (m), 1.88 (m)
Toluene	2.36 (s), 7.17 (m), 7.25 (m)	2.34 (s), 7.15 (m), 7.24 (m)	2.11 (s), 7.02 (m), 7.13 (m)	2.32 (s), 7.10-7.20 (m), 7.10-7.20 (m)	2.30 (s), 7.18 (m), 7.25 (m)	2.33 (s), 7.10-7.30 (m), 7.10-7.30 (m)	2.32 (s), 7.16 (m), 7.16 (m)	-
Triethylamine	1.03 (t), 2.53 (q)	0.99 (t), 2.48 (q)	0.96 (t), 2.40 (q)	0.96 (t), 2.45 (q)	0.93 (t), 2.43 (q)	0.96 (t), 2.45 (q)	1.05 (t), 2.58 (q)	0.99 (t), 2.57 (q)
Silicon grease	0.07	0.09	0.29	0.13	-0.06	0.08	0.10	-

Shaded region denotes residual undeuterated solvent. Multiplicities are singlet unless denoted otherwise (s=singlet, t=triplet, m=multiplet).

*The multiplicities of trace impurities may be different for methanol and ethanol if coupling with the OH proton is observed.

TABLE 23. ^{13}C NMR CHEMICAL SHIFTS OF TRACE IMPURITIES OF COMMON SOLVENTS AND REAGENTS

	CDCl_3	CD_2Cl_2	C_6D_6	$(\text{CD}_3)_2\text{CO}$	$(\text{CD}_3)_2\text{SO}$	CD_3CN	CD_3OD	D_2O
Acetone	30.92 207.07	31.00 206.78	30.14 204.43	30.60 205.87	30.56 206.31	30.91 207.43	30.67 209.67	30.89 215.94
Acetonitrile	1.89 116.43	2.03 116.92	0.20 116.02	1.12 117.60	1.03 117.91	1.79 118.26	0.85 118.06	1.47 119.68
Chloroform	77.36	77.99	77.79	79.19	79.16	79.17	79.44	-
Dichloromethane	53.52	54.24	53.46	54.95	54.84	55.32	54.78	-
Diethyl ether	15.20 65.91	15.44 66.11	15.46 65.94	15.78 66.12	15.12 62.05	15.63 66.32	15.46 66.88	14.77 66.42
DMF	31.45 36.50 162.62	31.39 36.56 162.57	30.72 35.25 162.13	31.03 36.15 162.79	30.73 35.73 162.29	31.32 36.57 163.31	31.61 36.89 164.73	32.03 37.54 165.53
DMSO	40.76		40.03	41.23	40.45	41.31	40.45	39.39
Dioxane	67.14	67.47	67.16	67.60	66.36	67.72	68.11	67.19
Ethanol	18.41 58.28	18.69 58.57	18.72 57.86	18.89 57.72	18.51 56.07	18.80 57.96	18.40 58.26	17.47 58.05
Ethyl acetate	14.19 21.04 60.49 171.36	14.37 21.15 60.63 171.24	14.19 20.56 60.21 170.44	14.50 20.83 60.56 170.96	14.40 20.68 59.74 170.31	14.54 21.16 60.98 171.68	14.49 20.88 61.50 172.89	13.92 21.15 62.32 175.26
Methanol	50.41	50.45	49.97	49.77	48.59	49.90	49.86	49.50
Pyridine	123.75 135.96 149.90	124.06 136.16 150.27	123.58 135.28 150.27	124.57 136.56 150.67	123.84 136.05 149.58	127.76 136.89 150.76	125.53 138.35 150.05	125.12 138.27 149.18
THF	25.62 67.97	25.98 68.16	25.72 67.80	26.15 68.07	25.14 67.03	26.27 68.33	26.48 68.83	25.67 68.68
Toluene	21.46 125.33 128.26 129.07 137.89	21.53 125.62 128.54 129.35 138.36	21.10 125.68 128.56 129.33 137.91	21.46 126.12 129.03 129.76 138.48	20.99 125.29 128.18 128.88 137.35	21.50 126.28 129.23 129.94 138.90	21.50 126.29 129.20 129.91 138.85	
Triethylamine	11.61 46.25	12.12 46.75	12.35 46.77	12.49 47.07	11.74 45.74	12.38 47.10	11.09 46.96	9.07 47.19
Silicon grease	1.19	1.22	1.38	1.40			2.10	

TABLE 24. AQUEOUS BUFFERS

Approx. pH	Composition
0	2N sulfuric acid or N hydrochloric acid
1	0.1N hydrochloric acid or 0.18N sulfuric acid
2	Either 0.01N hydrochloric acid or 0.013N sulfuric acid Or 50 ml of 0.1M glycine (also 0.1M NaCl) + 50 ml of 0.1N hydrochloric acid
3	Either 20 ml of the 0.2M Na_2HPO_4 + 80 ml of 0.1M citric acid Or 50 ml of 0.1M glycine + 22.8 ml of 0.1N hydrochloric acid in 100 ml
4	Either 38.5 ml of 0.2M Na_2HPO_4 + 61.5 ml of 0.1M citric acid Or 18 ml of 0.2M NaOAc + 82 ml of 0.2M acetic acid
5	Either 70 ml of 0.2M NaOAc + 30 ml of 0.2M acetic acid Or 51.5 ml of 0.2M Na_2HPO_4 + 48.5 ml of 0.1M citric acid
6	63 ml of 0.2M Na_2HPO_4 + 37 ml of 0.1M citric acid
7	82 ml of M Na_2HPO_4 + 18 ml of 0.1M citric acid
8	Either 50 ml of 0.1M Tris buffer + 29 ml of 0.1N hydrochloric acid, in 100 ml Or 30 ml of 0.05M borax + 70 ml of 0.2M boric acid
9	80 ml of 0.05M borax + 20 ml of 0.2M boric acid
10	Either 25 ml of 0.05M borax + 43 ml of 0.1N NaOH, in 100 ml Or 50 ml of 0.1M glycine + 32 ml of 0.1N NaOH, in 100 ml
11	50 ml of 0.15M Na_2HPO_4 + 15 ml of 0.1N NaOH
12	50 mL of 0.15M Na_2HPO_4 + 75 ml of 0.1N NaOH
13	0.1N NaOH or KOH
14	N NaOH or KOH

*These buffers are suitable for use in obtaining ultraviolet spectra. Alternatively, for a set of accurate buffers of low, but constant, ionic strength ($I = 0.01$) covering a pH range 2.2 to 11.6 at 20°, see Perrin Aust J Chem **16** 572 1963, DOI: 10.1071/CH9630572. 'In 100 ml' means that the solution is made up to 100 ml with pure water.*

TABLE 24a. pKa OF SELECTED INORGANIC ACIDS AND BASES[§] in H₂O at 25°C.

Acids	pKa	Bases	pKa
Amidophosphoric	2.74, 8.30	Aluminium (Al ³⁺)	5
Aminodisulfonic	8.50	Ammonia	9.25
Arsenic	11.2, 6.98	Ammonia (deuterated) *	9.76
Arsenious	9.18	Barium	13.36
Boric	9.23	Bismuth (Bi ³⁺)	1.58
Carbonic	6.35, 10.33	Cadmium	9.10
Chlorous	1.94	Calcium	12.9
Cyanic (HCNO)	3.46	Chromium (Cr ³⁺)	4
Diamidophosphoric	1.23, 4.94	Cobalt (Co ²⁺)	9.85
Hydrazoic	4.72	Hydrazine * (NH ₂ NH ₂)	-0.88, 8.11
Hydriodic	-9	Methylhydrazine ^{#†}	7.87
Hydrobromic	-8	Tetramethylhydrazine ^{#†}	6.30
Hydrochloric	-6.1	Phenylhydrazine [†]	5.27
Hydrocyanic	9.22	Hydroxylamine (NH ₂ OH)	5.69
Hydrofluoric	3.18	O-Methylhydroxylamine ^{††}	4.60
Hydrogen peroxide	11.65	N,N-DiMe-hydroxylamine ^{††}	5.20
Hydrogen sulfide*	7.05, 14.0	Trimethylhydroxylamine ^{††}	3.65
Hypophosphorous	1.23	Cyanamide (H ₂ NC≡N)*	-0.34, 10.27
Iodic	0.80	Phosphine	-14
Nitric	-1.44	Iron (Fe ²⁺)	6.74
Nitrous *	3.20	Iron (Fe ³⁺)	2.46
Perchloric	< 1.0	Lead (Pb ²⁺)	8
Phosphoric	2.15, 7.20, 12.38	Lithium	13.82
Phosphorous	1.43, 6.67	Magnesium	11.42
Pyrophosphoric	1.53, 2.36, 6.60, 9.25	Mercury (Hg ⁺)	5.0
Silicic	9.77	Mercury (Hg ²⁺)	3.49
Selenic acid	1.99	Nickel	9.86
Selenous acid	2.62, 8.32	Palladium (Pd ²⁺)	1
Sulfamic	0.99	Potassium	16
Sulfuric	-3, 1.96	Silver	>11
Deuterosulfuric	2.33	Sodium	14.77
Sulfurous	1.89, 7.21	Thallium (Tl ⁺)	13.2
Thiocyanic (HCNS)	-1.83	Thallium (Tl ³⁺)	1.16
		Tin (Sn ²⁺)	1.70
		Zinc	8.96

*At 20°, #at 30°, † Not strictly inorganic but a derivative of hydrazine, and †† Not strictly inorganic but a derivative of hydroxylamine.

§Selected from Albert and Serjeant, *The Determination of Ionisation Constants*, A Laboratory Manual, 3rd Edition, Chapman & Hall, New York, London, 1984, ISBN 0412242907.

TABLE 24b. pKa OF SELECTED ORGANIC ACIDS[§] in H₂O at 25°C.

Acids	pKa	Acids	pKa
Methanol	15.5	2-Fluorophenol	8.73
Ethanol	15.5	3-Fluorophenol	9.29
Allyl alcohol	15.5	4-Fluorophenol	9.89
Ethylene glycol	15.1	2-Bromophenol	8.45
Methoxyethanol	14.8	3-Bromophenol	9.03
Formic	3.75	4-Bromophenol	9.37
Acetic	4.76	2-Formylphenol	8.37
Acetic (in D ₂ O)	5.31	3-Formylphenol	8.98
Acetic-2- <i>d</i> ₃	4.77	4-Formylphenol	7.61
Propionic	4.87	2-nitrophenol	7.23
Pentanoic (Valeric) [*]	4.83,	3-nitrophenol	8.36
Octanoic	4.89	4-nitrophenol	7.15
Dimethylacetic [*]	4.84	2,4-dinitrophenol	4.07
Trimethylacetic [*]	5.03	2,4,6-trinitrophenol	0.42
Trifluoroacetic	0.52	Benzoic	4.02
Trichloroacetic [*]	0.66	2-methylbenzoic	3.91
Nitroacetic	1.48, 8.90	3-methylbenzoic [#]	4.25
Cyanoacetic	2.47	4-methylbenzoic	4.37
Fluoroacetic	2.59	2-methoxybenzoic	4.08
Chloroacetic	2.87	3-methoxybenzoic	4.10
Iodoacetic	3.18	4-methoxybenzoic	4.50
Phenoxyacetic	3.17	2-nitrobenzoic	2.17
Ethoxyacetic	3.65	3-nitrobenzoic	3.46
Phenylacetic	4.31	4-nitrobenzoic	3.43
Methylsulfonic	2.36	Pentafluorobenzoic	1.75
Trifluoromethylsulfonic	2.06	2,4-dinitrobenzoic	1.43
Trifluoromethylsulfonic	1.88	2,4,6-dinitrobenzoic	0.65
Acylic acid	4.25	4-Toluenesulfonic	-1.34
Cinnamic	4.41	Benzenesulfonic [*]	1.3
Propiolic	1.84	Benzeneboronic	8.83
Cyclopentane carboxylic	4.99	2-Furan carboxylic	3.16
Oxalic	1.25, 3.81	3-Furan carboxylic	3.9
Malonic	2.85, 5.70	2-Thiophene carboxylic	3.49
Succinic	4.21, 5.46	3-Thiophene carboxylic	4.1
Maleic (<i>cis</i>)	1.42, 6.23	2-Pyrrole carboxylic [*]	4.45
Fumaric (<i>trans</i>)	3.02, 4.38	3-Pyrrole carboxylic [*]	5.00
Phenylacetic	4.39	Thiophenol	6.62
2-phenylpropionic	4.66	Pyrrole	>15
Phenol	9.99	Pyrazole (1 <i>N</i> -2 <i>NH</i>)	~14
2-Methylphenol	10.29	Imidazole (1 <i>N</i> -3 <i>NH</i>)	14.4
3-Methylphenol	10.09	1,2,3-Triazole [*]	9.42
4-Methylphenol	10.26	1,2,4-Triazole	10.26
2- <i>tert</i> -Butylphenol	10.62	Tetrazole	4.89
3- <i>tert</i> -Butylphenol	10.12	Purine [*]	8.93
4- <i>tert</i> -Butylphenol	10.23	8-Azapurine [*]	4.84

^{*}At 20°. [§]Selected from Albert and Serjeant, *The Determination of Ionisation Constants*, A Laboratory Manual, 3rd Edition, Chapman & Hall, New York, London, 1984, ISBN 0412242907.

TABLE 24c. pKa OF SELECTED ORGANIC BASES[§] in H₂O at 25°C

Bases	pKa		Bases	pKa
Methylamine	10.66		2-Methylaniline	4.45
Dimethylamine	10.73		3-Methylaniline	4.71
Trimethylamine	9.80		4-Methylaniline	5.08
Ethylamine	10.65		2-Fluoroaniline	3.20
Diethylamine	10.84		3-Fluoroaniline	3.59
Triethylamine	3.75		4-Fluoroaniline	4.65
Propylamine	10.54		2-Cyanoaniline	0.77
<i>tert</i> -Butylamine	10.68		3-Cyanoaniline	2.75
Octylamine	10.65		4-Cyanoaniline	1.74
Hexadecylamine	10.61		2-Nitroaniline	-0.25
Benzylamine	9.34		3-Nitroaniline	2.46
Glycine methyl ester	7.59		4-Nitroaniline	1.02
Allylamine	9.49		2,6-Dimethylaniline	3.89
Cyanomethylamine	5.34		1-Naphthylamine	4.16
Phenethylamine	9.83		2-Naphthylamine	4.02
Ethanolamine	9.50		Pyridine	5.23
Triethanolamine	7.76		4-Methylpyridine	5.99
Acetamide	12.40		4-Methoxypyridine	6.58
Benzamide [*]	11.6		4-Chloropyridine	3.83
Trimethylamine- <i>N</i> -oxide [*]	4.65		4-Nitropyridine	1.61
Piperidine	11.12		3-Aminopyridine	6.04
<i>N</i> -Methylpiperidine	10.38		4-Aminopyridine	9.11
Pyrrolidine	11.31		Pyridine <i>N</i> -oxide ^{**}	0.79
<i>N</i> -Methylpyrrolidine	10.46		Quinoline [*]	4.90
Azetidine	11.29		2-Aminoquinoline [*]	7.34
Aziridine	8.04		4-Aminoquinoline [*]	9.17
Morpholine	8.50		Isoquinoline [*]	5.40
<i>N</i> -Methylmorpholine	7.38		1-Aminoisoquinoline [*]	7.62
Acetamide	-1.4		Acridine [*]	5.58
Benzamide	-2.16		9-Aminoacridine [*]	9.99
Urea	0.10		3,6-Diaminoacridine [*]	9.65
<i>O</i> -Methylisourea ^{**}	9.72		Pyridazine [*]	2.24
Thiourea	-1		Pyrimidine [*]	1.23, -6.9
<i>S</i> -Methylisothiurea	9.81		4-Aminopyrimidine [*]	5.71
Acetylhydrazine (cf hydrazine)	3.24		Pyrazine [*]	0.65, -6.3
Aniline	4.87		Phthalazine [*]	3.47
<i>N</i> -Methylaniline	4.85		Cinnoline [*]	2.37
<i>N,N</i> -Dimethylaniline	5.07		Quinazoline [*]	1.94, 3.43 [#]
<i>N</i> -Ethylaniline	5.12		Quinoxaline [*]	0.56
<i>N-tert</i> -Butylaniline	7.00		Phenazine [*]	1.20
<i>N</i> -Acetylaniline	~0.5		Purine [*]	2.30
Diphenylamine	0.79		Thiazole	2.52
Triphenylamine	-5.0		Oxazole ^{###}	0.8

^{*}At 20°, ^{**}at 24°, [#]Covalent hydrated cation, ^{##}at 33°. [§]Selected from Albert and Serjeant, *The Determination of Ionisation Constants*, A Laboratory Manual, 3rd Edition, Chapman & Hall, New York, London, 1984, ISBN 0412242907.

TABLE 25. SOLUBILITY COEFFICIENTS OF AIR AT 1atm IN WATER

T °C	(β) _{air}	(l) _{air}	% O ₂ (by vol)	(l) O ₂	Conc of O ₂ in H ₂ O
0	0.0292	0.0292	34.91	0.0102	0.455mM
20	0.0187	0.0199	34.03	0.0068	0.282mM
25	0.0171	0.0187	33.82	0.0063	0.258mM
30	0.0156	0.0173	33.62	0.0060	0.237mM

(β) is the Bunsen coeff. in ml gas/ml H₂O at STP. (l) is the Ostwald coeff. in ml gas/ml H₂O at stated temp.
Adapted from W.F. Linke, *A. Seidell's Solubilities of Inorganic and Metal-organic Compounds*, American Chemical Society, 1965.

TABLE 26. SOLUBILITY COEFFICIENTS OF O₂ AT 1atm IN WATER

T °C	(β)	(l)	Conc of O ₂ in H ₂ O
0	0.047	0.047	2.10mM
20	0.031	0.033	1.37mM
25	0.028	0.0306	1.25mM
30	0.026	0.0289	1.16mM
50	0.021	0.0248	0.94mM

(β) is the Bunsen coeff. in ml gas/ml H₂O at STP. (l) is the Ostwald coeff. in ml gas/ml H₂O at stated temp.
Adapted from W.F. Linke, *A. Seidell's Solubilities of Inorganic and Metal-organic Compounds*, American Chemical Society, 1965.

TABLE 27. BUNSEN COEFFICIENTS (β) OF GASES AT 1atm IN ORGANIC SOLVENTS AT 20°C

Solvent	H ₂	He	N ₂	O ₂	CO	CO ₂
H ₂ O	0.017	0.009	0.015	0.031	0.025	0.88
CS ₂	0.031	—	0.049	—	0.076	0.83
CHCl ₃	—	—	0.120	0.205	0.177	0.345
EtOH	0.080	0.028	0.130	0.143	0.177	3.0
Me ₂ CO	0.065	0.030	0.129	0.207	0.198	6.5
Et ₂ O	0.12	—	0.24	0.415	0.38	5.0
C ₆ H ₆	0.066	0.018	0.104	0.163	0.153	—

(β) is the Bunsen coeff. in ml gas/ml H₂O at STP.

Adapted from S. Glasstone, *Textbook of Physical Chemistry*, Macmillan & Co Ltd, London, 1951.

TABLE 28. OSTWALD COEFFICIENTS (<i>l</i>)/L OF O ₂ AT 1atm IN AQUEOUS SOLUTIONS AT 25°C			
	(<i>l</i>)/L		(<i>l</i>)
0.125N NH ₄ Cl 0.25N NH ₄ Cl 1.0N NH ₄ Cl	2.31 1.6 0.07	0.5M HCl 1.0M HCl 2.0M HCl	0.0296 0.0287 0.0267
0.125N NaCl 0.025N NaCl 0.50N NaCl 1.0N NaCl 2.0N NaCl 4.0N NaCl	5.52 5.30 4.92 4.20 3.05 1.62	0.25M H ₂ SO ₄ 0.5M H ₂ SO ₄ 1.0M H ₂ SO ₄ 1.5M H ₂ SO ₄ 2.0M H ₂ SO ₄ 2.5M H ₂ SO ₄	0.0288 0.0275 0.0251 0.0229 0.0209 0.0194
0.125N NaBr 0.5N NaBr 1.0N NaBr 6.0N NaBr	5.65 5.15 4.47 1.28	0.5M HNO ₃ 1.0M HNO ₃ 2.0M HNO ₃	0.0302 0.0295 0.0284
0.125N KCl 0.25N KCl 1.0N KCl 4.0N KCl	5.52 5.30 4.26 1.17	0.5M NaOH 1.0M NaOH 2.0M NaOH	0.0250 0.0204 0.0133
0.125N K ₂ SO ₄ 0.25N K ₂ SO ₄ 0.05N K ₂ SO ₄	5.11 4.66 3.89	0.5M KOH 1.0M KOH	0.0252 0.0206
0.125N BaCl ₂ 0.25N BaCl ₂ 1.0N BaCl ₂	5.40 5.04 3.10	0.125N Sucrose 0.5N Sucrose 1.0N Sucrose 2.0N Sucrose	0.00540 0.00438 0.00320 0.00184
The Ostwald coefficient (<i>l</i>)/L is for ml of gas in 1L of solution, and (<i>l</i>) is for ml of gas in 1ml of solution. Adapted from W.F. Linke, A. <i>Seidell's Solubilities of Inorganic and Metal-organic Compounds</i> , American Chemical Society, 1965.			

TABLE 29. SOLUBILITIES OF HCl AND NH₃ AT 760mm (g/100g OF SOLUTION)

Gas	Temperature °C	MeOH	EtOH	Et ₂ O
Hydrogen Chloride*	-10	54.6	—	37.5 (-9.2°)
	0	51.3	45.4	35.6
	20	47.0 (18°)	41.0	24.9
	30	43.0	38.1	19.47
Ammonia	15	21.6 (27.6g/100g MeOH)	13.2 (9.2g/100ml soln)	—
	25	16.5 (19.8g/100g MeOH)	10.0 (6.0g/100ml soln)	—

* Saturated EtOH with HCl is ~ 5.7M at 25°C, i.e. 21.5g/100ml of solution.

TABLE 30. BOILING POINTS OF SOME USEFUL GASES AT 760 mm

Argon	-185.6°	Krypton	-152.3°
Carbon dioxide (sublimes)	-78.5°	Methane	-164.0°
Carbon monoxide	-191.3°	Neon	-246.0°
Ethane	-88.6°	Nitrogen	-209.9°
Helium	-268.6°	Nitrous oxide	-88.5°
Hydrogen	-252.6°	Nitric oxide	-195.8°
		Oxygen	-182.96°

TABLE 31. PREFIXES FOR QUANTITIES

Fractional	deci (d) = 10 ⁻¹	centi (c) = 10 ⁻²	milli (m) = 10 ⁻³	micro (μ) = 10 ⁻⁶	nano (n) = 10 ⁻⁹	pico (p) = 10 ⁻¹²	femto (f) = 10 ⁻¹⁵	atto (atto) = 10 ⁻¹⁸
Multiple	deca (d) = 10 ¹	hecto (h) = 10 ²	kilo (k) = 10 ³	mega (M) = 10 ⁶	giga (G) = 10 ⁹	tera (T) = 10 ¹²	penta (P) = 10 ¹⁵	eka (E) = 10 ¹⁸

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CHAPTER 2

CHEMICAL METHODS USED IN PURIFICATION

GENERAL REMARKS

Greater selectivity in purification can often be achieved by making use of differences in chemical properties between the substance to be purified and the contaminants. Unwanted metal ions may be removed by precipitation in the presence of a *collector* (see below). Sodium borohydride and other metal hydrides transform organic peroxides and carbonyl-containing impurities such as aldehydes and ketones in alcohols and ethers. Many classes of organic chemicals can be purified by conversion into suitable salts or derivatives, followed by regeneration. This chapter describes some relevant procedures.

REMOVAL OF TRACES OF METALS FROM REAGENTS

METAL IMPURITIES

The presence of metal contaminants in reagents may sometimes affect the chemical or biochemical outcomes of an experiment. In these cases, it is necessary to purify the reagents used. Metal (and other) impurities can be determined qualitatively and quantitatively by atomic absorption spectroscopy (AAA), x-ray photoelectron spectroscopy (XPS), various mass spectrometric methods and/or inductively coupled plasma mass spectrometry (ICP-MS) (see Chapter 1, Question of Purity) and the required purification procedures can be formulated. Metal impurities in organic compounds are usually in the form of ionic salts or complexes with organic compounds and very rarely in the form of free metal. If they are present in the latter form then they can be removed by crystallising the organic compound (whereby the insoluble metal can be removed by filtration), or by distillation in which case the metal remains behind with the residue in the distilling flask. If the impurities are in the ionic or complex forms, then extraction of the organic compound in a suitable organic solvent with aqueous acidic or alkaline solutions will reduce their concentration to acceptable levels.

When the metal impurities are present in inorganic compounds as in metals or metal salts, then advantage of the differences in chemical properties should be taken. Properties of the impurities like the solubility, the solubility product (product of the metal ion and the counter-ion concentrations), the stability constants of the metal complexes with organic complexing agents and their solubilities in organic solvents should be considered. *Alternatively*, the impurities can be masked by the addition of complexing agents which could lower the concentration of the metal ion impurities to such low levels that they would not interfere with the properties of main compound (see **complexation** below). Specific procedures and examples are provided below.

DISTILLATION

Reagents such as water, ammonia, hydrochloric acid, nitric acid, perchloric acid, and sulfuric acid can be purified *via* distillation (preferably under reduced pressure and particularly with perchloric acid) using an all-glass still. Isothermal distillation is convenient for ammonia: a beaker containing concentrated ammonia is placed alongside a beaker of distilled water for several days in an empty desiccator so that some of the ammonia distils over into the water. The redistilled ammonia should be kept in polyethylene or paraffin-waxed bottles. Hydrochloric acid can be purified in the same way. To ensure the absence of metal contaminants from some salts (e.g. ammonium acetate), it may be more expedient to synthesise the salts using distilled components rather than to attempt to purify the salts themselves.

SCAVENGER RESINS AND OTHER SUPPORTS

There is now available an extensive range of supported reactants for high throughput screening that use specific resins, silica, carbons etc, to clean up reactions prior to final purification in the laboratory. Some are specifically electrophilic resins, i.e. to derivatise specific types of compounds, e.g. aldehyde and isocyanate bound resins for removing unwanted hydrazines, aliphatic and aromatic amines; toluenesulfonyl chloride bound resins for removing OH and NH containing impurities. Nucleophilic scavenger resins such as diethylenetriamine bound resins scavenge acids, acid chlorides and anhydrides; sulfonamide bound resins react with acids, acid chlorides, aldehydes, isocyanates and chloroformate impurities have been used. [Varma et al. *Comb Chem High Throughput Screening* **11** 238 2008, DOI: 10.2174/138620708783877753.] Adsorbents such as Florisil, high purity silica coated with specific reagents, e.g. 2,4-dinitrophenylhydrazine, and various charcoals have been used in cartridges prior to purification through desired FPLC and HPLC columns. (cf Biotage, Supelco and Sigma-Aldrich catalogues).

USE OF ION-EXCHANGE RESINS

Application of ion-exchange columns has greatly facilitated the removal of heavy metal ions such as Cu^{2+} , Zn^{2+} and Pb^{2+} from aqueous solutions of many reagents. Thus, sodium salts and sodium hydroxide can be purified by passage through a column of a cation-exchange resin in its sodium form, prepared by washing the resin with 0.1M aqueous NaOH then washing with water until the pH of the effluent is ~ 7 . Similarly, for acids, a resin in its H^+ form [prepared by washing the column with 0.1M aqueous mineral acid (HCl , H_2SO_4) followed by thorough washing with water until the effluent has pH ~ 7 is used]. In some cases, where metals form anionic complexes, they can be removed by passage through an anion-exchange resin. Iron in hydrochloric acid solution can be removed in this way.

Ion-exchange resins are also useful for demineralising biochemical preparations such as proteins. Removal of metal ions from protein solutions using polystyrene-based resins, however, may lead to protein denaturation. This difficulty may be avoided by using a weakly acidic cation exchanger such as Bio-Rex 70.

Heavy metal contamination of pH buffers can be removed by passage of the solutions through a Chelex X-100 column. For example when a solution of 0.02M HEPES [4-(2-hydroxyethyl)piperazine-1-ethanesulfonic acid] containing 0.2M KCl (1L, pH 7.5) alone or with *calmodulin*, is passed through a column of Chelex X-100 ($\sim 60\text{g}$) in the K^+ form, the level of Ca^{2+} ions falls to less than $2 \times 10^{-7} \text{ M}$ as shown by atomic absorption spectroscopy. Such solutions should be stored in polyethylene containers that have been washed with boiling deionised water (5 minutes) and rinsed several times with deionised water. TES [*N,N,N',N'*-Tetraethylsulfamide] and TRIS [Tris-(hydroxymethyl)aminomethane] have been similarly decontaminated from metal ions.

Water, with very low concentrations of ionic impurities (and approaching conductivity standards), is very readily obtained by percolation through alternate columns of cation- and anion-exchange resins, or through a mixed-bed resin, and many commercial devices are available for this purpose. For some applications, this method is unsatisfactory because the final deionised water may contain traces of organic material after passage through the columns. However, organic matter can be removed by using yet another special column in series for this purpose (see Water Purification Systems-Pure and Ultrapure in <www.millipore.com>).

PRECIPITATION

In removing traces of impurities by precipitation, it is necessary to include a material to act as a *collector* of the precipitated substance so as to facilitate its removal by filtration or decantation. The following are a few examples:

Removal of lead contaminants

Aqueous hydrofluoric acid can be freed from lead by adding 1ml of 10% strontium chloride per 100ml of acid, lead being co-precipitated as lead fluoride with the strontium fluoride. If the hydrofluoric acid is decanted from the precipitate and the process repeated, the final lead content in the acid is less than 0.003ppm. Similarly, lead can be precipitated from a nearly saturated sodium carbonate solution by adding 10% strontium chloride dropwise (1-2ml per 100ml) followed by filtration. (If the sodium carbonate is required as a solid, the solution can be evaporated to dryness in a platinum dish.) Removal of lead from potassium chloride uses precipitation as lead sulfide by bubbling H_2S , followed, after filtration, by evaporation and recrystallisation of the potassium chloride.

Removal of iron contaminants

Iron contaminants have been removed from potassium thiocyanate solutions by adding a slight excess of an

aluminium salt, then precipitating aluminum and iron as their hydroxides by adding a few drops of ammonia. Iron is also carried down on the hydrated manganese dioxide precipitate formed in cadmium chloride or cadmium sulfate solutions by adding 0.5% aqueous potassium permanganate (0.5ml / 100ml of solution), sufficient aqueous ammonia to give a slight precipitate, and 1ml of ethanol. The solution is heated to boiling to coagulate the precipitate, then filtered. Ferrous ion can be removed from copper solutions by adding some hydrogen peroxide to the solution to oxidise the iron, followed by precipitation of ferric hydroxide by adding a small amount of sodium hydroxide.

Removal of other metal contaminants

Traces of calcium can be removed from solutions of sodium salts by precipitation at pH 9.5-10 as the 8-hydroxyquinolate. The excess 8-hydroxyquinoline acts as a *collector* and is extracted out with an organic solvent.

EXTRACTION

In some cases, a simple solvent extraction is sufficient to remove a particular impurity. For example, traces of gallium can be removed from titanous chloride in hydrochloric acid by extraction with diisopropyl ether. Similarly, ferric chloride can be removed from aluminium chloride solutions containing hydrochloric acid by extraction with diethyl ether. Usually, however, it is necessary to extract an undesired metal with an organic solvent in the presence of a suitable complexing agent such as dithizone (diphenylthiocarbazone) or sodium diethyl dithiocarbamate. When the former is used, weakly alkaline solutions of the substance containing the metal impurity are extracted with dithizone into chloroform (at about 25mg/L of chloroform) or carbon tetrachloride until the colour of some fresh dithizone solution remains unchanged after shaking. Dithizone complexes metals more strongly in weakly alkaline solutions. Excess dithizone in the aqueous medium is removed by extracting with the pure solvent (chloroform or carbon tetrachloride), the last traces of which, in turn, are removed by aeration. This method has been used to remove metal impurities from aqueous solutions of ammonium hydrogen citrate, potassium bromide, potassium cyanide, sodium acetate and sodium citrate. The advantage of dithizone for such a purpose lies in the wide range of metals with which it combines under these conditions. 8-Hydroxyquinoline (oxine) can also be used in this way. Sodium diethyl dithiocarbamate has been used to remove metals from aqueous hydroxylamine hydrochloride (made just alkaline to thymol blue by adding ammonia) from copper and other heavy metals by repeated extraction with chloroform until no more diethyl dithiocarbamate remained in the solution (which was then acidified to thymol blue by adding hydrochloric acid).

COMPLEXATION

Although not strictly a removal of an impurity, addition of a suitable complexing agent such as ethylenediaminetetraacetic acid often overcomes the undesirable effects of contaminating metal ions by reducing the concentrations of the free metal species to very low levels, i.e. *sequestering* metal ions by complexation. For a detailed discussion of this *masking*, see *Masking and Demasking of Chemical Reactions*, D.D.Perrin, Wiley-Interscience, New York, 1970, ISBN 10: 0471680710 ISBN 13: 9780471680710.

USE OF METAL HYDRIDES

This group of reagents is commercially available in large quantities; some of its members-notably lithium aluminium hydride (LiAlH_4), calcium hydride (CaH_2), sodium borohydride (NaBH_4) and potassium borohydride (KBH_4) have found widespread use in the purification of chemicals.

LITHIUM ALUMINIUM HYDRIDE

This solid is stable at room temperature and is soluble in ether-type solvents. It reacts violently with water, liberating hydrogen, and is a powerful drying and reducing agent for organic compounds. It reduces aldehydes, ketones, esters, carboxylic acids, peroxides, acid anhydrides and acid chlorides to the corresponding alcohols. Similarly, amides, nitriles, aldimines and aliphatic nitro compounds yield amines, while aromatic nitro compounds are converted to azo compounds. For this reason it finds extensive application in purifying organic chemical substances by the removal of water and carbonyl containing impurities as well as peroxides formed by autoxidation. Reactions can generally be carried out at room temperature, or in refluxing diethyl ether, at

atmospheric pressure. *When drying organic liquids with this reagent it is important that the concentration of water in the liquid is below 0.1% - otherwise a violent reaction or **EXPLOSION** may occur. LiAlH_4 should be added cautiously to a cooled solution of organic liquid in a flask equipped with a reflux condenser.*

CALCIUM HYDRIDE

This powerful drying agent is suitable for use with hydrogen, argon, helium, nitrogen, hydrocarbons, chlorinated hydrocarbons, esters and higher alcohols.

SODIUM BOROHYDRIDE

This solid, which is stable in dry air up to 300° , is a less powerful reducing agent than lithium aluminium hydride, from which it differs also by being soluble in hydroxylic solvents and to a lesser extent in ether-type solvents. Sodium borohydride forms a dihydrate melting at $36-37^\circ$, and its aqueous solutions decompose slowly unless stabilised to above pH 9 by alkali. (For example, a useful sodium borohydride solution is one that is nearly saturated at $30-40^\circ$ and containing 0.2% sodium hydroxide.) Its solubility in water is 25, 55 and 88g per 100ml of water at 0° , 25° and 60° , respectively. Boiling or acidification rapidly decomposes aqueous sodium borohydride solutions. The reagent, available either as a hygroscopic solid or as an aqueous sodium hydroxide solution, is useful as a water soluble reducing agent for aldehydes, ketones and organic peroxides. This explains its use for the removal of carbonyl-containing impurities and peroxides from alcohols, polyols, esters, polyesters, amino alcohols, olefins, chlorinated hydrocarbons, ethers, polyethers, amines (including aniline), polyamines and aliphatic sulfonates.

Purifications using sodium borohydride can be carried out conveniently using alkaline aqueous or methanolic solutions of sodium borohydride, allowing the reaction mixture to stand at room temperature for several hours. Other solvents that can be used with this reagent include isopropyl alcohol (without alkali), amines (including liquid ammonia, in which its solubility is 104g per 100g of ammonia at 25° , and ethylene diamine), diglyme, formamide, dimethylformamide and tetrahydrofurfuryl alcohol. *Alternatively*, the material to be purified can be percolated through a column of the borohydride. In the absence of water, sodium borohydride solutions in organic solvents such as dioxane or amines decompose only very slowly at room temperature. Treatment of ethers with sodium borohydride appears to inhibit peroxide formation.

POTASSIUM BOROHYDRIDE

Potassium borohydride is similar in properties and reactions to sodium borohydride, and can similarly be used as a reducing agent for removing aldehydes, ketones and organic peroxides. It is non-hygroscopic and can be used in water, ethanol, methanol or water-alcohol mixtures, provided some alkali is added to minimise decomposition, but it is somewhat *less soluble* than sodium borohydride in most solvents. For example, the solubility of potassium borohydride in water at 25° is 19g per 100ml of water, as compared to 55g of sodium borohydride.

PURIFICATION *via* DERIVATIVES

Relatively few derivatives of organic substances are suitable for use as aids to purification. This is because of the difficulty in regenerating the starting material. For this reason, we list below the common methods of preparation of derivatives that can be used in this way.

Whether or not any of these derivatives is likely to be satisfactory for use in any particular case will depend on the degree of difference in properties, such as solubility, volatility or melting point, between the starting material, its derivative and likely impurities, as well as on the ease with which the substance can be recovered. Purification *via* a derivative is likely to be of most use when the quantity of pure material that is required is not too large. Where large quantities (for example, more than 50g) are available, it is usually more economical to purify the material directly (for example, in distillations and recrystallisations).

The most generally useful purifications *via* derivatives are as follows:

ALCOHOLS

Aliphatic or aromatic alcohols are converted into solid esters. 4-Nitrobenzoates are examples of convenient esters to form because of their sharp melting points, and the ease with which they can be crystallised as well as the ease

with which the parent alcohol can be recovered. The *p*-nitrobenzoyl chloride used in the esterification is prepared freshly by refluxing dry *p*-nitrobenzoic acid with a 3 molar excess of thionyl chloride for 30 minutes on a steam bath (*in a fume cupboard*). The solution is cooled slightly and excess thionyl chloride is distilled off under vacuum, keeping the temperature below 40°. Dry toluene is added to the residue in the flask, then distilled off in a vacuum, the process being repeated two or three times to ensure complete removal of thionyl chloride, hydrogen chloride and sulfur dioxide. (This freshly prepared *p*-nitrobenzoyl chloride cannot be stored without decomposition; it should be used directly.) A solution of the acid chloride (1mol) in dry toluene or alcohol-free chloroform (distilled from P₂O₅ or by passage through an activated Al₂O₃ column) under a reflux condenser is cooled in an ice bath while the alcohol (1mol), with or without a solvent (preferably miscible with toluene or alcohol-free chloroform), is added dropwise to it. When addition is over and the reaction subsides, the mixture is refluxed for 30 minutes and the solvent is removed under reduced pressure. The solid ester is then recrystallised to constant melting point from toluene, acetone, low boiling point petroleum ether or mixtures of these, but not from alcohols (due to unwanted trans-esterification).

Hydrolysis of the ester is achieved by refluxing in aqueous N or 2N NaOH solution until the insoluble ester dissolves. The solution is then cooled, and the alcohol is extracted into a suitable solvent, e.g. ether, toluene or alcohol-free chloroform. The extract is dried (CaSO₄, MgSO₄) and distilled, then fractionally distilled if liquid or recrystallised if solid. (The *p*-nitrobenzoic acid can be recovered by acidification of the aqueous layer.) In most cases where the alcohol to be purified can be readily extracted from ethanol, the hydrolysis of the ester is best achieved with N or 2N ethanolic NaOH or 85% aqueous ethanolic N NaOH. The former is prepared by dissolving the necessary alkali in a minimum volume of water and diluting with absolute alcohol. The ethanolic solution is refluxed for one to two hours and hydrolysis is complete when an aliquot gives a clear solution on dilution with four or five times its volume of water. The bulk of the ethanol is distilled off and the residue is extracted as above. *Alternatively*, use can be made of ester formation with benzoic acid, toluic acid or 3,5-dinitrobenzoic acid, by the above method.

Other derivatives can be prepared by reaction of the alcohol with an acid anhydride. For example, phthalic or 3-nitrophthalic anhydride (1 mol) and the alcohol (1mol) are refluxed for half to one hour in a non-hydroxylic solvent, e.g. toluene or alcohol-free chloroform, and then cooled. The phthalate ester crystallises out, is precipitated by the addition of low boiling petroleum ether or is isolated by evaporation of the solvent. It is recrystallised from water, 50% aqueous ethanol, toluene or low boiling petroleum ether. Such an ester has a characteristic melting point and the alcohol can be recovered by acid or alkaline hydrolysis.

ALDEHYDES

The best derivative from which an aldehyde can be recovered readily is its *bisulfite addition compound*, the main disadvantage being the lack of a sharp melting point. The aldehyde (sometimes in ethanol) is shaken with a cold saturated solution of sodium bisulfite (evolution of heat) until no more solid adduct separates. The adduct is filtered off, washed with a little water, followed by alcohol. A better reagent to use is a freshly prepared saturated aqueous sodium bisulfite solution to which 75% ethanol is added to near-saturation. (Water may have to be added dropwise to render this solution clear.) With this reagent the aldehyde need not be dissolved separately in alcohol and the adduct is finally washed with alcohol. The aldehyde is recovered by dissolving the adduct in the least volume of water and adding an equivalent quantity of sodium carbonate (not sodium hydroxide) or concentrated hydrochloric acid to react with the bisulfite (SO₂ is liberated), followed by steam distillation or solvent extraction. Other derivatives that can be prepared are the *Schiff bases* and semicarbazones. Condensation of the aldehyde with an equivalent of primary aromatic amine yields the Schiff base, for example aniline at 100° for 10-30 minutes. *Semicarbazones* are prepared by dissolving semicarbazide hydrochloride (*ca* 1g) and sodium acetate (*ca* 1.5g) in water (8-10ml) and adding the aldehyde or ketone (0.5-1g) with stirring. The semicarbazone crystallises out and is recrystallised from ethanol or aqueous ethanol. These are hydrolysed by steam distillation in the presence of oxalic acid or better by exchange with pyruvic acid [Hershberg *J Org Chem* **13** 542 1948, DOI: 10.1021/jo01162a011] [see also entry under Ketones].

AMINES

Salts

Amines can also be purified *via* their salts, e.g. hydrochlorides. A solution of the amine in dry toluene, diethyl ether, dichloromethane or chloroform is saturated with dry hydrogen chloride (generated by addition of concentrated sulfuric acid to dry sodium chloride, or to concentrated HCl followed by drying the gas through sulfuric acid, or the HCl gas is obtained from a hydrogen chloride cylinder) and the insoluble hydrochloride is

filtered off and dissolved in water. The solution is made alkaline and the amine is extracted, as above. Hydrochlorides can also be prepared by dissolving the amine in ethanolic HCl and adding diethyl ether. Where hydrochlorides are too hygroscopic or too soluble for satisfactory isolation, other salts, e.g. nitrate, sulfate, bisulfate or oxalate, can be used.

Double salts

The amine (1mol) is added to a solution of anhydrous zinc chloride (1mol) in concentrated HCl (42ml) in ethanol (200ml, or less depending on the solubility of the double salt). The solution is stirred for 1 hour and the precipitated salt is filtered off and recrystallised from ethanol. The free base is recovered by adding excess of 5-10N NaOH (to dissolve the zinc hydroxide that separates) and is steam distilled. Mercuric chloride (highly poisonous) in hot water can be used instead of zinc chloride and the salt is crystallised from 1% hydrochloric acid. Other double salts have been used, e.g. cuprous salts, but are not as convenient as the above salts.

Bis-(arylsulfonyl)imine salts

Bis-alkyl and aryl sulfonylimines [$\text{RSO}_2\text{NHSO}_2\text{R}'$], which are readily prepared, are acidic and form stable sodium or potassium salts that can be recrystallised. The bis-phenyl and bis-4-chlorophenyl derivatives form salts readily via their Na salts [*m* 314-316° (327-329°) and 278-280° (286-287°) respectively and crystallise as colourless needles from H_2O] with a large variety of amines, including some amino acids, basic drugs and natural bases, which can be readily recrystallised for characterisation and for purification. The *free* bases can be readily released on addition of alkali and isolated. [See 'Miscellaneous As, B, P, ...' section in Chapter 3, and Runge et al. *Chem Ber* **88** 533 1955, DOI: 10.1002/cber.19550880414; Runge et al. *Chem Ber* **86** 1571 1953, DOI: 10.1002/cber.19530861216; Runge & Pfeiffer *Chem Ber* **90** 1757 1957, DOI: 10.1002/cber.19570900910; Helferich & Grünert *Chem Ber* **73** 1131 1940, DOI: 10.1002/cber.19400731019.]

TABLE 1. Bis-(phenylsulfonyl)imine salts[§] of Aliphatic Amines.

Amine	Mp	Solvent [†]	Amine	Mp	Solvent [†]
Methylamine	175-176°	A	Tri- <i>n</i> -butylamine	51-53°	C
Dimethylamine	159-160°	A	<i>i</i> -Hexylamine	172-173°	A
Trimethylamine	163-164°	A	Di- <i>i</i> -hexylamine	55-57°	B
Ethylamine	210-211°	A	<i>n</i> -Octylamine	135-136°	B
Diethylamine	113-114°	A	Di- <i>n</i> -octylamine	65-66°	C
Triethylamine	109-110°	A	<i>n</i> -Dodecylamine	102-103°	C
<i>n</i> -Propylamine	185-186°	A	Di- <i>n</i> -dodecylamine	85-86°	B
Di- <i>n</i> -propylamine	75-77°	B	<i>n</i> -Hexadecylamine	84-86°	D
Tri- <i>n</i> -propylamine	55-57°	B	<i>n</i> -Octadecylamine	76-78°	C
<i>n</i> -Butylamine	165-166°	B	Di- <i>n</i> -octadecylamine	94-95°	D
Di- <i>n</i> -butylamine	79-81°	B			

[§] Prepared mostly in Et_2O , but also in H_2O , $\text{MeOH}/\text{H}_2\text{O}$ or EtOH . [†] A=needles from H_2O , B=needles from $\text{Me}_2\text{CO}/\text{Et}_2\text{O}$, C=needles from Et_2O , D= needles from $\text{EtOH}/\text{Et}_2\text{O}$.

TABLE 2. Bis-(4-chlorophenylsulfonyl)imine salts[§] of Aromatic Amines.

Amine	Mp	Solvent [†]	Amine	Mp	Solvent [†]
Aniline	174°	A	Benzidine [#]	253-254°	D
Ethylaniline	107-108°	B	Tetramethylbenzidine [#]	210-211°	E
<i>o</i> -Toluidine	199-200°	A	α -Naphthylamine	176-177°	B
<i>p</i> -Toluidine	192-193°	A	β -Naphthylamine	209-210°	B
<i>o</i> -Aminophenol	176-177°	B	α -Aminodecalin	198-199°	B
<i>p</i> -Aminophenol	202-203°	A	Phenylhydrazine	178-179°	A
<i>o</i> -Phenylenediamine [#]	217-218°	A	Cyclohexylamine	233-235°	A
<i>m</i> -Phenylenediamine [#]	75-77°	C	Dicyclohexylamine	84-86°	D
<i>p</i> -Phenylenediamine [#]	273° dec	C	Sulfanilic acid	No salt formed	A

[§] Prepared mostly in Et_2O , but also in H_2O , $\text{MeOH}/\text{Et}_2\text{O}$, $\text{Me}_2\text{CO}/\text{Et}_2\text{O}$ or Me_2CO . [†] A=needles or plates from H_2O , B=needles from $\text{Me}_2\text{CO}/\text{Et}_2\text{O}$, C=needles from $\text{MeOH}/\text{H}_2\text{O}$, D= needles from $\text{MeOH}/\text{Et}_2\text{O}$, E=needles from MeOH . [#] With 2 mols of sulfimide.

TABLE 3. Bis-(4-chlorophenylsulfonyl)imine salts[§] of Heterocyclic Compounds.

Amine	Mp	Solvent [†]	Amine	Mp	Solvent [†]
Pyridine	170-171°	A	Isoquinoline	161-170°	A
Piperidine	208-209°	A	Hydroquinoline	155-156°	B
α -Picoline	151-152°	A	Quinaldine	151-152°	B
α,α' -Lutidine	151-152°	B	Acridine	195-196°	A
Pyridine-2,6-Di-CO ₂ H	280-281° dec	A	Pyrrole	No salt formed	A
Nicotine	54° (55°)	B	Pyrrolidine	238-239°	A
Nicotinic Acid	188-189°	C	3-Methylpyrazole	134-135°	A
Nicotinamide	214-216°	D	1,2,4-Triazole	163-164°	A
2-Aminopyridine	171-172°	B	3-Amino-1,2,4-triazole	165-166°	A
Melamine	307-308°	D	Benzotriazole	145-146°	A
Kaymethin	241-242°	A	4,5-Dimethylimidazole	119-120°	B
Quinoline	190-191°	B	Isatin	No salt formed	B

[§] Prepared mostly in Me₂CO, but also in Et₂O, some in Me₂CO/H₂O, MeOH, or MeOH/H₂O. [†] A=needles from H₂O, B=needles from Me₂CO/Et₂O, C=needles from Me₂CO, D= needles from MeOH.

TABLE 4. Bis-(4-chlorophenylsulfonyl)imine salts[§] of Alkaloids.

Amine	Mp	Solvent [†]	Amine	Mp	Solvent [†]
Ephedrine	184-185°	A	Narcotine	~90° dec	B
Hordenine	145-146°	A	Hydrastinine	139-140°	B
Nicotine	45° (55°)	B	Berberine	216-217°	C
Atropine	145-146°	B	Emetine	134-136°	B
Hyoscyamine	123-125°	B	Yohimbine	196-197°	D
Scopolamine	202-203°	B	Eserine	77-79°	D
Cinchonine	~90° dec	B	Strychnine	~142° dec	B
Quinidine	~80° dec	B	Brucine	84-86°	B
Papaverine	118-119°	B	Sparteine	~72° (~92°)dec	A

[§] Prepared mostly in H₂O, but also in Me₂CO, MeOH, or Et₂O/CHCl₃. [†] A=needles or plates from H₂O, B=needles from Me₂CO/Et₂O, C=needles from MeOH/H₂O, D= needles from Me₂CO/H₂O.

TABLE 5. Bis-(4-chlorophenylsulfonyl)imine salts[§] of Amino Acids.

Amine	Mp	Solvent [†]	Amine	Mp	Solvent [†]
DL-Serine	160-161°	A	DL- α -Aminobutyric acid	186-187°	A
L-Glutamic acid	186-187° dec	B	DL-Valine	164-165°	C
DL-Alanine	174-178°	A	L-Histidine	165° dec	A
Glycine	175-177°	A	DL-Leucine	168-170° dec	A

[§] Prepared mostly in Me₂CO/H₂O, but also in H₂O/MeOH or H₂O. [†] A=needles or plates from H₂O, B=needles from MeOH/H₂O, C= needles from Me₂CO/H₂O.

Picrates

The most versatile derivative from which the free base can be readily recovered is the picrate. This is very satisfactory for primary and secondary aliphatic amines and aromatic amines and is particularly so for heterocyclic bases. The amine, dissolved in water or alcohol, is treated with excess of a saturated solution of picric acid in water or alcohol, respectively, until separation of the picrate is complete. If separation does not occur, the solution is stirred vigorously and warmed for a few minutes, or diluted with a solvent in which the picrate is insoluble. Thus, solutions of the amine and picric acid in ethanol can be treated with petroleum ether to crystallise out the picrate salt. *Alternatively*, the amine can be dissolved in alcohol and aqueous picric acid added. The picrate is filtered off, washed with water or ethanol and recrystallised from boiling water, ethanol, methanol, aqueous ethanol, methanol or chloroform. The solubility of picric acid at ~20° in water and ethanol is about 1.4 and 6.23%

respectively.

It is not advisable to store large quantities of picrates for long periods, *particularly when they are dry due to their potential **EXPLOSIVE** nature*. Also this method is not advised when large quantities of amines (e.g. >25g) are to be purified. The free base should be recovered as soon as possible. The picrate is suspended in an excess of 2N aqueous NaOH and warmed a little. Because of the limited solubility of sodium picrate, excess hot water must be added. *Alternatively*, because of the greater solubility of lithium picrate, aqueous 10% lithium hydroxide solution can be used. The solution is cooled, the amine is extracted with a suitable solvent such as diethyl ether or toluene, washed with 5N NaOH until the alkaline solution remains colourless, then with water, and the extract is dried with anhydrous sodium carbonate. The solvent is distilled off and the amine is fractionally distilled (under reduced pressure if necessary) or recrystallised.

If the amines are required as their hydrochlorides, picrates can often be decomposed by suspending them in acetone and adding two equivalents of 10N HCl. The hydrochloride of the base is filtered off, leaving the picric acid in the acetone. Dowex No 1 anion-exchange resin in the chloride form is useful for changing solutions of the more soluble picrates (for example, of adenosine) into solutions of their hydrochlorides, from which sodium hydroxide precipitates the free base.

N-Acetyl derivatives

Purification as their *N*-acetyl derivatives is satisfactory for primary, and to a limited extent, secondary amines. Tertiary amines are not acetylated. The base is refluxed with slightly more than one equivalent of acetic anhydride for half to one hour, cooled and poured into ice-cold water. The insoluble derivative is filtered off, dried, and recrystallised from water, ethanol, aqueous ethanol or benzene (**CAUTION toxic!**). The derivative can be hydrolysed to the parent amine by refluxing with 70% sulfuric acid for a half to one hour. The solution is cooled, poured onto ice, and made alkaline. The amine is steam distilled or extracted as above. Alkaline hydrolysis is very slow.

N-Tosyl derivatives

Primary and secondary amines are converted into their tosyl derivatives by mixing equimolar amounts of amine and *p*-toluenesulfonyl chloride in dry pyridine (*ca* 5-10mols) and allowing to stand at room temperature overnight. The solution is poured into ice-water and the pH adjusted to 2 with HCl. The solid derivative is filtered off, washed with water, dried (vacuum desiccator) and recrystallised from an alcohol or aqueous alcohol solution to a sharp melting point. The derivative is decomposed by dissolving in liquid ammonia (*fume cupboard*) and adding sodium metal (in small pieces with stirring) until the blue colour persists for 10-15minutes. Ammonia is allowed to evaporate (*fume cupboard*), the residue treated with water and the solution checked that the pH is above 10. If the pH is below 10, then the solution has to be basified with 2N NaOH. The mixture is extracted with diethyl ether or toluene, the extract is dried (K₂CO₃), evaporated and the residual amine recrystallised if solid or distilled if liquid.

AROMATIC HYDROCARBONS

Adducts

Aromatic hydrocarbons can be purified as their picrates using the procedures described for amines. Instead of picric acid, 1,3,5-trinitrobenzene or 2,4,7-trinitrofluorenone can also be used. In all these cases, following recrystallisation, the hydrocarbon can be isolated either as described for amines or by passing a solution of the adduct through an activated alumina column and eluting with toluene or petroleum ether. The picric acid and nitro compounds are more strongly adsorbed on the column.

Sulfonation

Naphthalene, xylenes and alkyl benzenes can be purified by sulfonation with concentrated sulfuric acid and crystallisation of the sodium sulfonates. The hydrocarbon is distilled out of the mixture with superheated steam.

CARBOXYLIC ACIDS

4-Bromophenacyl esters

A solution of the sodium salt of the acid is prepared. If the salt is not available, the acid is dissolved in an equivalent of aqueous NaOH and the pH adjusted to 8-9 with this base. A solution of one equivalent of 4-bromophenacyl bromide (for a monobasic acid, two equivalents for a dibasic acid, etc) in ten times its volume of ethanol is then added. The mixture is heated to boiling, and, if necessary, enough ethanol is added to clarify the solution which is then refluxed for half an hour to three hours depending on the number of carboxylic groups that have to be esterified. (One hour is generally sufficient for monocarboxylic acids.) On cooling, the ester should crystallise out. If it does not, then the solution is heated to boiling, and enough water is added to produce a slight

turbidity. The solution is again cooled. The ester is collected, and recrystallised or fractionally distilled.

The ester is hydrolysed by refluxing for 1-2 hours with 1-5% of barium carbonate suspended in water or with aqueous sodium carbonate solution. The solution is cooled and extracted with diethyl ether, toluene or chloroform. It is then acidified and the acid is collected by filtration or extraction, and recrystallised or fractionally distilled. 4-Bromophenylphenacyl esters are used similarly.

p-Nitrobenzyl esters can be prepared in an analogous manner using the sodium salt of the acid and *p*-nitrobenzyl bromide. They are readily hydrolysed.

Alkyl esters

Of the alkyl esters, methyl esters are the most useful because of their rapid hydrolysis. The acid is refluxed with one or two equivalents of methanol in excess alcohol-free chloroform (or dichloromethane) containing about 0.1g of *p*-toluenesulfonic acid (as catalyst), using a Dean-Stark apparatus. (The water formed by the esterification is carried away into the trap.) When the theoretical amount of water is collected in the trap, esterification is complete. The chloroform solution in the flask is washed with 5% aqueous sodium carbonate solution, then water, and dried over anhydrous sodium sulfate or magnesium sulfate. The chloroform is distilled off and the ester is fractionally distilled through an efficient column, or recrystallised if it is a solid. The ester is hydrolysed by refluxing with 5-10% aqueous NaOH solution until the insoluble ester has completely dissolved. The aqueous solution is concentrated a little by distillation to remove almost all of the methanol. It is then cooled and acidified. The acid is either extracted with diethyl ether, toluene or chloroform, or filtered off and isolated as above. Other methods for preparing esters are available, e.g. addition of an ethereal solution of diazomethane (yellow in colour, poisonous, use a fume cupboard. CARE: Use diazomethane with extreme care as the reagent is POISONOUS and HIGHLY explosive; special precautions MUST be used; see *Fieser and Fieser's Reagents for Organic Synthesis* **1** pp191-195 1967, abbreviated to 'Fieser' in the text of following chapters) to the acid which dissolves as the acid is esterified, liberating N₂ and the yellow colour. The methyl ester so produced is obtained by evaporating the ethereal solution.

Salts (S-Arylthiuronium)

The most useful salt derivatives for carboxylic, sulfonic and phosphorus acids are the isothiuronium salts. These are prepared by mixing almost saturated solutions containing the acid (carefully neutralised with N NaOH using phenolphthalein indicator) then adding two drops of N HCl and an equimolar amount of *S*-benzylisothiuronium chloride (Table 6), or its 4-chloro derivative (Table 7 and 8) in ethanol and filtering off the salt that crystallises out, and recrystallising to analytical purity (see Tables below). After recrystallisation from water, alcohol or aqueous alcohol the salt is decomposed by suspending or dissolving in 2N HCl and extracting the carboxylic acid from aqueous solution into diethyl ether, chloroform or toluene. *Alternatively*, the free acids can be obtained by passing a ~50% aqueous alcoholic solution through a Dowex 50W x8 resin previously washed with ~50% aqueous alcohol, evaporating to a small volume and extraction into Et₂O (or similar solvent) from which the acid is isolated. [For preparations see 'Miscellaneous As, B, P, ...' in Chapter 3; Donleavy *J Am Chem Soc* **58** 1004 1936, DOI: 10.1021/ja01297a048; and for 4-chlorobenzylisothiuronium salts see Dewey and Sperry *J Am Chem Soc* **61** 3251 1939, DOI: 10.1021/ja01267a005; Harvey & Jensen *J Org Chem* **28** 470 1963, DOI: 10.1021/jo01037a047].

S-Benzylisothiuronium chloride [538-28-3] **m 146-148° and 172-174°**, and **S-4-chlorobenzylisothiuronium chloride** [544-47-8] **m 177-178° and 197°, 201-203°** crystallises in two forms. After standing in a container for

TABLE 6. S-Benzylisothiuronium Salts of some Carboxylic and Sulfonic Acids.

Acid	Mp of salt	Acid	Mp of salt
Formic	146°	3-Nitrobenzoic	163°
Acetic	134°	4-Nitrobenzoic	182°
Propionic	148°	2-Aminobenzoic	142°
<i>n</i> -Butyric	146°	3-Aminobenzoic	149°
Isobutyric(2-Mepropanoic)	143°	4-Aminobenzoic	166°
Isovaleric (3-Mebutanoic)	153°	2-Toluic	140°
Lauric (dodecanoic)	141°	3-Toluic	164°
Myristic (tetradecanoic)	139°	4-Toluic	190°
Palmitic (hexadecanoic)	141°	2-Hydroxybenzoic	146°

TABLE 6. S-Benzylisothiuronium Salts of some Carboxylic and Sulfonic Acids (continued)

Acid	Mp of salt		Acid	Mp of salt
Stearic	143°		3- Chlorobenzoic	155°
Oxalic	193°		4-Bromobenzoic	168°
Succinic	149°		4-Methoxybenzoic	177°
Malic (acid salt)	163°		Phthalic	151°
Fumaric	178°		Cinnamic (<i>trans</i>)	175°
Crotonic	162°		Mandelic	166°
Glycolic	141°		Diphenylacetic	145°
Lactic	153°		Benzene sulfonic	144°
Diglycolic	154°		3-Nitrobenzene sulfonic	140°
Benzoic	166°		4-Toluenesulfonic	178°
2-Nitrobenzoic	159°			

Taken from Donleavy *J Am Chem Soc* **58** 1004 1936, DOI: 10.1021/ja01297a048.

one week, the lower melting crystals remained unchanged, but the higher melting crystals now melted at the same temperature as the lower melting form. Both forms of crystals give the same derivatives with acids. In the case of phosphorus esters the thiuronium (complexes ?) are prepared differently, i.e. the phosphate ester in EtOH is refluxed for 24hrs with an ethanolic solution of EtSNa (from a mixture of Na and EtSH in EtOH), followed by addition of a saturated solution of an equivalent of S-4-chlorobenzylisothiuronium chloride. The precipitated NaCl is filtered off, the filtrate is evaporated to dryness and the residue is recrystallised from the appropriate solvent (see Table 8).

TABLE 7. S-4-Chlorobenzylisothiuronium Salts of some Carboxylic and Sulfonic Acids.

Acid	Mp of salt		Acid	Mp of salt
Formic	148°		3-Bromobenzoic	161°
Acetic	139°		4-Bromobenzoic	172°
<i>n</i> -Butyric	139°		2-Chlorobenzoic	159°
Caproic (hexanoic)	143°		3-Chlorobenzoic	157°
Monochloroacetic	158°		4-Chlorobenzoic	173°
Oleic(octadec-9Z-enoic)	131°		Cinnamic (<i>trans</i>)	167°
Oxalic	194°		2-Iodobenzoic	162°
Palmitic (hexadecanoic)	146°		3-Iodobenzoic	154°
Propionic	143°		4-Iodobenzoic	177°
Succinic	167°		Phthalic	166°
Trichloroacetic	148°		Salicylic, (sulfosalicylic)	162°, (181°)
Valeric (pentanoic)	142°		2-Toluic	150°
Benzene sulfonic	184°		3-Toluic	151°
Benzoic	155°		4-Toluic	161°
2-Bromobenzoic	165°		4-Toluenesulfonic	193°

Taken from Dewey and Sperry *J Am Chem Soc* **61** 3251 1939, DOI: 10.1021/ja01267a005.

TABLE 8. S-4-Chlorobenzylisothiuronium Derivatives of some Phosphonic Acids and Esters.

Acid	Mp of salt	Solvent	Esters	Mp of derivative	Solvent
C ₈ H ₁₇ PO(OH) ₂	192-193°	EtOH	(EtO) ₃ PO	160-161°	Me ₂ CO
PhPO(OH) ₂	200-202°	Pyridine	EtPO(OEt) ₂	166-167°	Me ₂ CO
PhPHO(OH)	181-182°	EtOH	EtSPO(OEt) ₂	151-152°	H ₂ O
(EtO) ₂ POSH	81-82°	Et ₂ O	PhSPO(OEt) ₂	174-176°	H ₂ O

Taken from Harvey & Jensen *J Org Chem* **28** 470 1963, DOI: 10.1021/jo01037a047.

HYDROPEROXIDES

These can be converted to their sodium salts by precipitation below 30° with aqueous 25% NaOH. The salt is then decomposed by addition of solid (powdered) carbon dioxide and extracted with low-boiling petroleum ether. The solvent should be removed under reduced pressure below 20°. **The manipulation should be adequately shielded at all times to guard against EXPLOSIONS for the safety of the operator.**

KETONES

Bisulfite adduct

The adduct can be prepared and decomposed as described for aldehydes. *Alternatively*, because no Cannizzaro reaction is possible, it can also be decomposed with 0.5N NaOH.

Semicarbazones

A powdered mixture of semicarbazide hydrochloride (1mol) and anhydrous sodium acetate (1.3mol) is dissolved in water by gentle warming. A solution of the ketone (1mol) in the minimum volume of ethanol needed to dissolve it is then added. The mixture is warmed on a water bath until separation of the semicarbazone is complete. The solution is cooled, and the solid is filtered off. After washing with a little ethanol followed by water, it is recrystallised from ethanol or dilute aqueous ethanol. The derivative should have a characteristic melting point. The semicarbazone is decomposed by refluxing with excess of oxalic acid or with aqueous sodium carbonate solution. The ketone (which steam distils) is distilled off. It is extracted or separated from the distillate (after saturating with NaCl), dried with CaSO₄ or MgSO₄ and fractionally distilled using an efficient column (under vacuum if necessary). [See entry under Aldehydes.]

PHENOLS

The most satisfactory derivatives for phenols that are of low molecular weight or monohydric are the benzoate esters. (Their acetate esters are generally liquids or low-melting solids.) Acetates are more useful for high molecular weight and polyhydric phenols.

Benzoates

The phenol (1mol) in 5% aqueous NaOH is treated (while cooling) with benzoyl chloride (1mol) and the mixture is stirred in an ice bath until separation of the solid benzoyl derivative is complete. The derivative is filtered off, washed with alkali, then water, and dried (in a vacuum desiccator over NaOH). It is recrystallised from ethanol or dilute aqueous ethanol. The benzoylation can also be carried out in dry pyridine at low temperature (*ca* 0°) instead of in NaOH solution, finally pouring the mixture into water and collecting the solid as above. The ester is hydrolysed by refluxing in an alcohol (for example, ethanol, *n*-butanol) containing two or three equivalents of the alkoxide of the corresponding alcohol (for example, sodium ethoxide or sodium *n*-butoxide) and a few (*ca* 5-10) millilitres of water, for half an hour to three hours. When hydrolysis is complete, an aliquot will remain clear on dilution with four to five times its volume of water. Most of the solvent is distilled off. The residue is diluted with cold water and acidified, and the phenol is steam distilled. The latter is collected from the distillate, dried and either fractionally distilled or recrystallised. It can also be isolated by extraction from a slightly acidified (pH ~3) aqueous solution with diethyl ether.

Acetates

These can be prepared as for the benzoates using either acetic anhydride with 3N NaOH or acetyl chloride in pyridine. They are hydrolysed as described for the benzoates. This hydrolysis can also be carried out with aqueous 10% NaOH solution, completion of hydrolysis being indicated by the complete dissolution of the acetate in the aqueous alkaline solution. On steam distillation, acetic acid also distils off, but in these cases the phenols (see above) are invariably solids which can be filtered off and recrystallised.

PHOSPHATE AND PHOSPHONATE ESTERS

These can be converted to their uranyl nitrate addition compounds. The crude or partially purified ester is saturated with uranyl nitrate solution and the solid adduct is filtered off. It is recrystallised from *n*-hexane, toluene or ethanol. For the more soluble members crystallisation from hexane using low temperatures (-40°) has been successful. The adduct is decomposed by shaking with sodium carbonate solution and water, the solvent is steam distilled (if hexane or toluene is used) and the ester is collected by filtration. *Alternatively*, after decomposition, the organic layer is separated, dried with CaCl₂ or BaO, filtered, and fractionally distilled under high vacuum. See also characterisation and purification *via* their S-aryliothiuronium salts or complexes, Table 8.

MISCELLANEOUS

Impurities can sometimes be removed by conversion to derivatives under conditions where the major component does not react or reacts much more slowly. For example, normal (straight-chain) paraffins can be freed from unsaturated and branched-chain components by taking advantage of the greater reactivity of the latter with chlorosulfonic acid or bromine. Similarly, the preferential nitration of aromatic hydrocarbons can be used to remove e.g. benzene or toluene from cyclohexane by shaking for several hours with a mixture of concentrated nitric acid (25%), sulfuric acid (58%), and water (17%).

GENERAL METHODS FOR THE PURIFICATION OF CLASSES OF COMPOUNDS

Chapters 3-7 list a large number of individual compounds, with a brief statement of how each one may be purified. For substances that are not included in these chapters the following procedures may prove helpful.

PROCEDURES

If the laboratory worker does not know of a reference to the preparation of a commercially available substance, he/she may be able to make a reasonable guess at the synthetic methods used from published laboratory syntheses. In some cases brief syntheses are provided in this book. This information, in turn, can simplify the necessary purification steps by suggesting probable contaminants.

Physical methods of purification depend largely on the melting and boiling points of the materials. For gases and low-boiling liquids use is commonly made of the *freeze-pump-thaw* procedure. Gas chromatography is also useful, especially for low-boiling point liquids. Liquids are usually purified by refluxing with drying agents, acids or bases, reducing agents, charcoal, etc., followed by fractional distillation under reduced pressure. For solids, general methods include fractional freezing of the melted material, taking the middle fraction. Another procedure is sublimation of the solid under reduced pressure. The other commonly used method for purifying solids is by recrystallisation from a solution in a suitable solvent, by cooling with or without the prior addition of a solvent in which the solute is not very soluble (see Chapter 1).

The nature of the procedure will depend to a large extent on the quantity of purified material that is required. For example, for small quantities (50-250mg) of a pure volatile liquid, preparative gas chromatography is probably the best method. Two passes through a suitable column may well be sufficient. Similarly, for smaller amounts (100-500mg) of an organic solid, column chromatography is likely to be satisfactory, the eluate being collected as a number of separate fractions (*ca* 5-10ml) which are examined by FT-IR, NMR or UV spectroscopy, TLC or by some other appropriate analytical technique. (For information on suitable adsorbents and eluents the texts referred to in the bibliography at the end of Chapters 1 and 2 should be consulted.) Preparative thin layer chromatography or HPLC, FC and HPFC can also be used successfully for purifying up to 500mg of solid. The latter chromatographic techniques (see Chapter 1) are more and more commonly used procedures for the purification of small molecules as well as large molecules such as polypeptides and DNA.

Where larger quantities (upwards of 1g) are required, most of the impurities should be removed by preliminary treatments, such as solvent extraction, liquid-liquid partition, or conversion to a derivative (*vide supra*) which can be purified by crystallisation or fractional distillation before being converted to the starting material. The substance is then crystallised or distilled. If the final amounts must be in excess of 25g, preparation of a derivative is sometimes omitted because of the cost involved. In all of the above cases, purification is likely to be more laborious if the impurity is an isomer or a derivative with closely similar physical properties.

CRITERIA OF PURITY

Purification becomes meaningful only insofar as adequate tests of purity are applied: the higher the degree of purity that is sought, the more stringent these tests must be. For this, the experimenter has to resort, in the first place, to preliminary physical methods such as melting and boiling points, chromatographic and spectroscopic procedures which are described in detail in Chapter 1. If the material is an organic solid, its melting point should first be taken and compared with the recorded value. *Note that the melting points of most salts, organic or inorganic, are generally decomposition points and are not reliable criteria of purity.* As part of the preliminary examination, the sample might be examined by thin layer chromatography in several different solvent systems and in high enough concentrations to facilitate the detection of minor components. On the other hand, if the substance is a liquid, its boiling point should be measured. If, further, the boiling point of the liquid is too high, or it

decomposes on heating, then its purity should be assessed by high pressure liquid chromatography. Those liquids that are especially volatile can be studied very satisfactorily by gas chromatography, preferably using at least two different stationary and/or mobile phases. Spectroscopic methods, if facilities are available, such as atomic absorption spectroscopy (AAA), and inductively coupled plasma mass spectrometry (ICP-MS) are useful and sensitive methods for detecting metal impurities and the concentrations of metals and metal salts or complexes.

Application of these tests at successive steps will give a good indication of whether or not the purification is satisfactory and will also show when adequate purification has been achieved. Finally elemental analyses, e.g. of carbon, hydrogen, nitrogen, sulfur, metals etc., are very sensitive to impurities (other than with isomers), and are good criteria of purity.

There are certain requirements for purity of new compounds in most journals. This is especially so for samples which are shown to have biological activity. See instructions to authors for ACS journals especially in *J. Med. Chem* (see 'Guidelines for Authors' under *Purity Criteria of Tested Compounds* < <http://pubs.acs.org/journal/jmcmar> >.)

GENERAL PROCEDURES FOR THE PURIFICATION OF SOME CLASSES OF ORGANIC COMPOUNDS

In the general methods of purification described below, it is assumed that the impurities belong essentially to a class of compounds different from the one being purified. They are suggested for use in cases where substances are not listed in Chapters 3, 4 and the low-molecular-weight compounds in Chapter 6. In such cases, the experimenter is advised to employ them in conjunction with information given in these chapters for the purification of suitable analogues. Also, for a wider range of drying agents and the use of cartridges (e.g. Na₂SO₄ for removal of H₂O, or Celite for removal of tar), solvents for extraction and solvents for recrystallisation, the reader is referred to Chapter 1. A common method of purification of organic compounds is to convert them to a suitable derivative which is purified (with the assumption that the impurity does not form a similar derivative, or if it does its properties are different), and then regenerate the original compound and purify it further. Various derivatives are described for different classes of compounds below, but many more can be considered, and the reader is referred to texts on 'Protecting Groups' which describe ways of selectively protecting functional groups and facile means of deprotecting them, i.e. regenerating the unprotected group [P.J.Kocienski *Protecting Groups* Thieme International Publisher, 2005, ISBN 9783131356031; J.R.Hanson *Protecting Groups in Organic Synthesis* Sheffield Academic Press, 1999, ISBN 9781850759577, or J.Wiley & Sons, NY, 1999, ISBN 980632045068]. See Chapter 6 for general purification procedures used for macromolecules.

ACETALS

These are generally diethyl or dimethyl acetal derivatives of aldehydes. They are more stable to alkali than to acids. Their common impurities are the corresponding alcohol, aldehyde and water. Drying with sodium wire removes alcohols and water, and polymerises aldehydes so that, after decantation, the acetal can be fractionally distilled. In cases where the use of sodium is too drastic, aldehydes can be removed by shaking with alkaline hydrogen peroxide solution and the acetal is dried with sodium carbonate or potassium carbonate. Residual water and alcohols (up to *n*-propyl) can be removed with Linde type 4A molecular sieves. The acetal is then filtered and fractionally distilled. Solid acetals (i.e. acetals of high-molecular-weight aldehydes) are generally low-melting and can be recrystallised from low-boiling petroleum ether, toluene or a mixture of both.

ACIDS

Carboxylic acids

Liquid carboxylic acids are first freed from neutral and basic impurities by dissolving them in aqueous alkali and extracting with diethyl ether. (The pH of the solution should be at least three units above the pK_a of the acid, see pK in Chapter 1). The aqueous phase is then acidified to a pH at least three units below the pK_a of the acid and again extracted with ether. It is quite unnecessary to add large excesses of mineral acid (e.g. HCl) to liberate the organic acid, as mineral acids dissolve appreciably in organic solvents such as diethyl ether. The extract is dried with magnesium sulfate or sodium sulfate and the ether is distilled off. The acid is fractionally distilled through an efficient column. It can be further purified by conversion to its methyl or ethyl ester (*vide supra*) which is then fractionally distilled. Hydrolysis yields the original acid which is again purified as above.

Acids that are solids can be purified in this way, except that distillation is replaced by repeated crystallisation

(preferable from at least two different solvents such as water, alcohol or aqueous alcohol, toluene, toluene with petroleum ether or acetic acid.) Water-insoluble acids can be partially purified by dissolution in N sodium hydroxide solution and precipitation with dilute mineral acid. If the acid is required to be free from sodium ions, then it is better to dissolve the acid in hot N ammonia, heat to *ca* 80°, adding slightly more than an equal volume of N formic acid and allowing to cool slowly for crystallisation. Any ammonia, formic acid or ammonium formate that adhere to the acid are removed when the acid is dried in a vacuum — these are volatile. Cartridges and columns are available (e.g. the Isolute SCX-2 made of polysulfonic acid bonded to silica as an ion exchange column developed by Biotage Inc <www.biotage.com>), particularly for the purification of acids using flash chromatography.

The separation and purification of naturally occurring fatty acids, based on distillation, salt solubility and low temperature crystallisation, are described by K.S.Markley (Ed.), *Fatty Acids*, 2nd Edn, part 3, Chap. 20, Interscience, New York, 1964, see also N.Reavley *Essential Fatty Acids* Book Media Publ, 2002, ISBN 9780958157643; G.Grati and K.Sato (Eds) *Crystallisation and Polymorphism of Fats and Fatty Acids* Marcel Dekker, 1988, ISBN 9780824778750.

Aromatic carboxylic acids can be purified by conversion to their sodium salts, recrystallisation from hot water, and conversion to the free acids. See also characterisation and purification *via* their S-arylisothiuronium salts, Tables 6 and 7.

Sulfonic acids

The low solubility of sulfonic acids in organic solvents and their high solubility in water makes necessary a treatment different from that for carboxylic acids. Sulfonic acids are strong acids, they have the tendency to hydrate, and many of them contain water of crystallisation. The lower-melting and liquid acids can generally be purified with only slight decomposition by fractional distillation, preferably under reduced pressure. A common impurity is sulfuric acid, but this can be removed by recrystallisation from concentrated aqueous solutions. The wet acid can be dried by azeotropic removal of water with toluene, followed by distillation. The higher-melting acids, or acids that melt with decomposition, can be recrystallised from water or, occasionally, from ethanol. For a typical purification of aromatic sulfonic acids using their barium salts refer to benzenesulfonic acid in the 'Aromatic Compounds' section in Chapter 3. See also characterisation and purification *via* their S-arylisothiuronium salts, Tables 6 and 7.

Sulfinic acids

These acids are less stable, less soluble and less acidic than the corresponding sulfonic acids. The common impurities are the respective sulfonyl chlorides from which they have been prepared, and the thiosulfonates (neutral) and sulfonic acids into which they decompose. The first two of these can be removed by solvent extraction from an alkaline solution of the acid. On acidification of an alkaline solution, the sulfinic acid crystallises out leaving the sulfonic acid behind. The lower molecular weight members are isolated as their metal (e.g. ferric) salts, but the higher members can be crystallised from water (made slightly acidic), or alcohol.

ACID CHLORIDES

The corresponding acid and hydrogen chloride are the most likely impurities. Usually these can be removed by efficient fractional distillation. Where acid chlorides are not readily hydrolysed (e.g. aryl sulfonyl chlorides) the compound can be freed from contaminants by dissolving in a suitable solvent such as alcohol-free chloroform, dry toluene or petroleum ether and shaking with dilute sodium bicarbonate solution. The organic phase is then washed with water, dried with anhydrous sodium sulfate or magnesium sulfate, and distilled or recrystallised. This procedure is *hazardous* with readily hydrolysable *carboxylic* acid chlorides such as acetyl chloride and benzoyl chloride. Solid acid chlorides should be thoroughly dried *in vacuo* over strong drying agents and are satisfactorily recrystallised from toluene, toluene-petroleum ether, petroleum ethers, alcohol-free chloroform/toluene, and, occasionally, from dry diethyl ether. Hydroxylic or basic solvents should be strictly avoided. *All operations should be carried out in a fume cupboard because of the **irritant** nature of these compounds which also attack the skin.*

ALCOHOLS

Monohydric alcohols

The common impurities in alcohols are aldehydes or ketones, and water. [*Ethanol* in Chapter 3 is typical.] Aldehydes and ketones can be removed by adding a small amount of sodium metal and refluxing for 2 hours, followed by distillation. Water can be removed in a similar way but it is preferable to use magnesium metal instead of sodium because it forms a more insoluble hydroxide, thereby shifting the equilibrium more completely from metal alkoxide to metal hydroxide. The magnesium should be activated with iodine (or a small amount of methyl iodide), and the water content should be low, otherwise the magnesium will be deactivated. If the amount of water is large, it should be removed by azeotropic distillation (see below), or by drying over anhydrous MgSO_4 (not CaCl_2 which combines with alcohols). Acidic materials can be removed by treatment with anhydrous Na_2CO_3 , followed by a suitable drying agent, such as calcium hydride, and fractional distillation, using gas chromatography to establish the purity of the product [Ballinger & Long, *J Am Chem Soc* **82** 795 1960, DOI: 10.1021/ja01489a008]. *Alternatively*, the alcohol can be refluxed with freshly ignited CaO for 4 hours and then fractionally distilled [McCurdy & Laidler, *Can J Chem* **41** 1867 1963, DOI: 10.1139/v63-274].

With higher-boiling alcohols it is advantageous to add some freshly prepared magnesium ethoxide solution (only slightly more than required to remove the water), followed by fractional distillation. *Alternatively*, in such cases, water can be removed by azeotropic distillation with toluene. Higher-melting alcohols can be purified by crystallisation from methanol or ethanol, toluene/petroleum ether or petroleum ether. Sublimation in vacuum, molecular distillation and gas and liquid chromatographic methods are also useful means of purification. For purification *via* derivatives, *vide supra*.

Polyhydric alcohols

These alcohols are more soluble in water than are monohydric alcohols. Liquids can be freed from water by shaking with type 4A Linde molecular sieves and can safely be distilled only under high vacuum. Carbohydrate alcohols can be crystallised from concentrated aqueous solutions or, preferably, from mixed solvents such as ethanol/petroleum ether or dimethyl formamide/toluene. Crystallisation usually requires seeding and is extremely slow. Further purification can be effected by conversion to the acetyl or benzoyl derivatives which are much less soluble in water and which can readily be recrystallised, e.g. from ethanol. Hydrolysis of the acetyl derivatives, followed by removal of acetic or benzoic acid and metal ions by ion-exchange chromatography, gives the purified material. On no account should solutions of carbohydrates be concentrated above 40° because of darkening and formation of *caramel*. Ion exchange, charcoal or cellulose column chromatography has been used for the purification and separation of carbohydrates.

ALDEHYDES

Common impurities found in aldehydes are the corresponding alcohols, aldols and water from self-condensation, and the corresponding acids formed by autoxidation. Acids can be removed by shaking with aqueous 10% sodium bicarbonate solution. The organic liquid is then washed with water. It is dried with anhydrous sodium sulfate or magnesium sulfate and then fractionally distilled. Water soluble aldehydes must be dissolved in a suitable solvent such as diethyl ether before being washed in this way. Further purification can be effected *via* the bisulfite derivative (see above) or the Schiff base formed with aniline or benzidine. Solid aldehydes can be dissolved in diethyl ether and purified as above. *Alternatively*, they can be steam distilled, then sublimed and crystallised from toluene or petroleum ether.

AMIDES

Amides are stable compounds. The lower-melting members (such as acetamide) can be readily purified by fractional distillation. Most amides are solids which have low solubilities in water. They can be recrystallised from large quantities of water, ethanol, ethanol/ether, aqueous ethanol, chloroform/toluene, chloroform or acetic acid. The likely impurities are the parent acids or the alkyl esters from which they have been made. The former can be removed by thorough washing with aqueous ammonia followed by recrystallisation, whereas elimination of the latter is by trituration or recrystallisation from an organic solvent. Amides can be freed from solvent or water by drying below their melting points. These purifications can also be used for sulfonamides and acid hydrazides.

AMINES

The common impurities found in amines are nitro compounds (if prepared by reduction), the corresponding halides (if prepared from them) and the corresponding carbamate salts. Amines are dissolved in aqueous acid, the

pH of the solution being at least three units below the pK_a value of the base to ensure almost complete formation of the cation. They are extracted with diethyl ether to remove neutral impurities and to decompose the carbamate salts. The solution is then made strongly alkaline and the amines that separate are extracted into a suitable solvent (ether or toluene) or steam distilled. The latter process removes coloured impurities. Note that chloroform cannot be used as a solvent for primary amines because, in the presence of alkali, poisonous carbylamines (isocyanides) are formed. However, chloroform is a useful solvent for the extraction of heterocyclic bases. In this case it has the added advantage that while the extract is being freed from the chloroform solvent most of the moisture is removed with the solvent.

Alternatively, the amine may be dissolved in a suitable solvent (e.g. toluene), and dry HCl gas is passed through the solution to precipitate the amine hydrochloride. This is purified by recrystallisation from a suitable solvent mixture (e.g. ethanol/diethyl ether). The free amine can be regenerated by adding sodium hydroxide and isolated as above. Cartridges and columns are available (e.g. KP-NH silica column with slightly nitrogenous alkaline chemistry developed by Biotage Inc (<www.biotage.com>)) for the purification of amines using flash chromatography.

Liquid amines can be further purified *via* their acetyl or benzoyl derivatives (*vide supra*). Solid amines can be recrystallised from water, alcohol, toluene or toluene-petroleum ether. *Care should be taken in handling large quantities of amines because their vapours are harmful (possibly carcinogenic) and they are readily absorbed through the skin.*

AMINO ACIDS

Because of their zwitterionic nature, amino acids are generally soluble in water. Their solubility in organic solvents rises as the fat-soluble portion of the molecule increases. The likeliest impurities are traces of salts, heavy metal ions, proteins and other amino acids. Purification of these is usually easy, by recrystallisation from water or ethanol/water mixtures. The amino acid is dissolved in the boiling solvent, decolorised if necessary by boiling with 1g of acid-washed charcoal/100g amino acid, then filtered hot, chilled, and set aside for several hours to crystallise. The crystals are filtered off, washed with ethanol, then ether, and dried.

Amino acids have high melting or decomposition points and are best examined for purity by paper or thin layer chromatography. The spots are developed with ninhydrin. Customary methods for the purification of small quantities of amino acids obtained from natural sources, i.e. 1-5g, are ion-exchange chromatography (see Chapter 1). For general treatment of amino acids see Greenstein and Winitz [*The Amino Acids*, Vols 1-3, J.Wiley & Sons, New York 1961] and individual amino acids in Chapters 3 and 6.

A useful source of details such as likely impurities, stability and tests for homogeneity of amino acids is *Specifications and Criteria for Biochemical Compounds*, 3rd edn, National Academy of Sciences, USA, 1972.

ANHYDRIDES

The corresponding acids, resulting from hydrolysis, are the most likely impurities. Distillation from phosphorus pentoxide, followed by fractional distillation, is usually satisfactory. With high boiling or solid anhydrides, another method involves boiling under reflux for 0.5-1 hours with acetic anhydride, followed by fractional distillation. Acetic acid distils first, then acetic anhydride and finally the desired anhydride. Where the anhydride is a solid, removal of acetic acid and acetic anhydride at atmospheric pressure is followed by heating under vacuum. The solid anhydride is then either crystallised as for acid chlorides or (in some cases) sublimed in a vacuum. A preliminary purification when large quantities of acid are present in a solid anhydride (such as phthalic anhydride) is by preferential solvent extraction of the (usually) more soluble anhydride from the acid (e.g. with $CHCl_3$ in the case of phthalic anhydride). *All operations with liquid anhydrides should be carried out in a fume cupboard because of their LACHRYMATORY properties. Almost all anhydrides attack skin.*

CAROTENOIDS

These usually are decomposed by light, air and solvents, so that degradation products are probable impurities. Chromatography and adsorption spectra permit the ready detection of coloured impurities, and separations are possible using solvent distribution, chromatography or crystallisation. Thus, in partition between immiscible solvents, xanthophyll remains in 90% methanol while carotenes pass into the petroleum ether phase. For small amounts of material, thin-layer or paper chromatography may be used, while column chromatography is suitable for larger amounts. Colourless impurities may be detected by IR, NMR or mass spectrometry. The more common separation procedures are described by P. Karrer and E. Jucker in *Carotenoids*, E.A. Braude (translator), Elsevier, NY, 1950. Purity can be checked by chromatography (on thin-layer plates, Kieselguhr, paper or columns), by UV

or NMR procedures. See 'Carotenoids' in Chapter 6.

ESTERS

The most common impurities are the corresponding acid and hydroxy compound (i.e. alcohol or phenol), and water. A liquid ester from a carboxylic acid is washed with 2N sodium carbonate or sodium hydroxide to remove acid material, then shaken with calcium chloride to remove ethyl or methyl alcohols (if it is a methyl or ethyl ester). It is dried with potassium carbonate or magnesium sulfate, and distilled. Fractional distillation then removes residual traces of hydroxy compounds. This method does not apply to esters of inorganic acids (e.g. dimethyl sulfate) which are more readily hydrolysed in aqueous solution when heat is generated in the neutralisation of the excess acid. In such cases, several fractional distillations, preferably under vacuum, are usually sufficient.

Solid esters are easily crystallisable materials. It is important to note that esters of alcohols must be recrystallised either from non-hydroxylic solvents (e.g. toluene) or from the alcohol from which the ester is derived. Thus methyl esters should be crystallised from methanol or methanol/toluene, but not from ethanol, *n*-butanol or other alcohols, in order to avoid alcohol exchange and contamination of the ester with a second ester. Useful solvents for crystallisation are the corresponding alcohols or aqueous alcohols, toluene, toluene/petroleum ether, and chloroform (ethanol-free)/toluene. Esters of carboxylic acids derived from phenols are more difficult to hydrolyse and exchange, hence any alcoholic solvent can be used freely. Sulfonic acid esters of phenols are even more resistant to hydrolysis: they can safely be crystallised not only from the above solvents but also from acetic acid, aqueous acetic acid or boiling *n*-butanol. Note that sulfonic esters of lower alcohols, e.g. methanol, are good alkylating agents.

Fully esterified phosphoric acid and phosphonic acids differ only in detail from the above mentioned esters. Their major contaminants are alcohols or phenols, phosphoric or phosphonic acids (from hydrolysis), and (occasionally) basic material, such as pyridine, which is used in their preparation. Water-insoluble esters are washed thoroughly and successively with dilute acid (e.g. 0.2N sulfuric acid), water, 0.2N sodium hydroxide and water. After drying with calcium chloride they are fractionally distilled. Water-soluble esters should first be dissolved in a suitable organic solvent and, in the washing process, water should be replaced by saturated aqueous sodium chloride. Some esters (e.g. phosphate and phosphonate esters) can be further purified through their uranyl adducts (*vide supra*). Traces of water or hydroxy compounds can be removed by percolation through, or shaking with, activated alumina (about 100g/L of liquid solution), followed by filtration and fractional distillation in a vacuum. For high molecular weight esters (which cannot be distilled without some decomposition) it is advisable to carry out distillation at as low a pressure as possible. Solid esters can be crystallised from toluene or petroleum ether. Alcohols can be used for recrystallising solid phosphoric or phosphonic esters of phenols.

ETHERS

The purification of diethyl ether (see Chapter 3) is typical of liquid ethers. The most common contaminants are the alcohols or hydroxy compounds from which the ethers are prepared, their oxidation products (e.g. aldehydes), peroxides and water. Dialkyl ethers form peroxides much more readily than other ethers, e.g. ethyl phenyl ethers, on standing in air. Peroxides, aldehydes and alcohols can be removed by shaking with alkaline potassium permanganate solution for several hours, followed by washing with water, concentrated sulfuric acid [CARE], then water. After drying with calcium chloride, the ether is distilled. It is then dried with sodium or with lithium aluminium hydride, redistilled and given a final fractional distillation. The drying process is repeated if necessary. *Alternatively*, methods for removing peroxides include leaving the ether to stand in contact with iron filings or copper powder, shaking with a solution of ferrous sulfate acidified with N sulfuric acid, shaking with a copper-zinc couple, passage through a column of activated alumina, and refluxing with phenothiazine. Cerium(III) hydroxide has also been used.

A simple test for ether peroxides is to add 10ml of the ether to a stoppered cylinder containing 1ml of freshly prepared 10% solution of potassium iodide containing a drop of starch indicator. No colour should develop during one minute if free from peroxides. *Alternatively*, a 1% solution of ferrous ammonium sulfate, 0.1M in sulfuric acid and 0.01M in potassium thiocyanate should not increase appreciably in red colour when shaken with two volumes of the ether. Merck-Chemicals supply peroxide test kits (Perex Test) which use a colorimetric method with test strips which can be used to estimate the amount of hydrogen peroxide, from as low a concentration as 0.2mg/L to as high as 1000mg/L. They are very convenient as they can give an indication of the concentration of peroxide rapidly <see <http://www.merck-chemicals.com>>. As a safety precaution against **EXPLOSION** in case purification from peroxides has been insufficiently thorough, at least a quarter of the total volume of liquid ether should remain in the distilling flask when the distillation is discontinued, as the peroxides are generally

higher boiling than the corresponding ethers. To minimize peroxide formation, ethers should be stored in dark bottles and, if they are liquids, they should be left in contact with type 4A Linde molecular sieves, in a cold place, over sodium amalgam. The rate of formation of peroxides depends on storage conditions and is accelerated by metal impurities, heat, light, air and moisture. Always be vigilant and test for peroxides. The formation of peroxides is inhibited in the presence of diphenylamine, di-*tert*-butylphenol, or other antioxidants which can be used as stabilisers.

Ethers that are solids (e.g. phenyl ethers) can be steam distilled from an alkaline solution which will hold back any phenolic impurity. After the distillate is made alkaline with sodium carbonate, the insoluble ether is collected either by extraction (e.g. with chloroform, diethyl ether or toluene) or by filtration. It is then crystallised from alcohols, alcohol/petroleum ether, petroleum ether, toluene or mixtures of these solvents, sublimed in a vacuum and recrystallised if necessary.

HALIDES

Aliphatic halides are likely to be contaminated with halogen acids and the alcohols from which they have been prepared, whereas in aromatic halides the impurities are usually aromatic hydrocarbons, amines or phenols. In both groups the halogen is less reactive than it is in acid halides. Purification is by shaking with concentrated hydrochloric acid, followed by washing successively with water, 5% sodium carbonate or bicarbonate, and water. After drying with calcium chloride, the halide is distilled and then fractionally distilled using an efficient column. For a solid halide the above purification is carried out by dissolving it in a suitable solvent such as toluene. Solid halides can also be purified by chromatography using an alumina column and eluting with toluene or petroleum ether. They can be crystallised from toluene, petroleum ethers, toluene/petroleum ether or toluene/chloroform/petroleum ether. Care should be taken when handling organic halogen compounds because of their **HIGH TOXICITY**. It should be noted that methyl iodide is a cancer suspect.

Liquid aliphatic halides are obtained alcohol-free by distillation from phosphorus pentoxide. They are stored in dark bottles to prevent oxidation and, in some cases, the formation of phosgene.

A general method for purifying *chlorohydrocarbons* uses repeated shaking with concentrated sulfuric acid [CARE] until no further colour develops in the acid, then washing with water followed by a solution of sodium bicarbonate, then with water again. After drying with calcium chloride, the chlorohydrocarbon is fractionally redistilled to constant boiling point or recrystallised.

HYDROCARBONS

Gaseous hydrocarbons are best freed from water and gaseous impurities by passage through suitable adsorbents and (if olefinic material is to be removed) oxidants such as alkaline potassium permanganate solution, followed by fractional cooling (see Chapter 1 for cooling baths) and fractional distillation at low temperature. To effect these purifications and also to store the gaseous sample, a vacuum line is necessary.

Impurities in hydrocarbons can be characterised and evaluated by gas chromatography and mass spectrometry. The total amount of impurities present can be estimated from the thermometric freezing curve.

Liquid aliphatic hydrocarbons are freed from aromatic impurities by shaking with concentrated sulfuric acid [CARE] whereby the aromatic compounds are sulfonated. Shaking is carried out until the sulfuric acid layer remains colourless for several hours. The hydrocarbon is then freed from the sulfuric acid and the sulfonic acids by separating the two phases and washing the organic layer successively with water, 2N sodium hydroxide, and water. It is dried with CaCl_2 or Na_2SO_4 , and then distilled. The distillate is dried with sodium wire, P_2O_5 , or metallic hydrides, or passage through a dry silica gel column, or preferably, and more safely, with molecular sieves (see Chapter 1) before being finally fractionally distilled through an efficient column. If the hydrocarbon is contaminated with olefinic impurities, shaking with aqueous alkaline permanganate is necessary prior to the above purification. Alicyclic and paraffinic hydrocarbons can be freed from water, non-hydrocarbon and aromatic impurities by passage through a silica gel column before the final fractional distillation. This may also remove isomers. (For the use of chromatographic methods to separate mixtures of aromatic, paraffinic and alicyclic hydrocarbons see references in the bibliography in Chapter 1 under *Liquid and Flash Chromatography*, *Gas Chromatography* and *High Performance Liquid Chromatography*). Another method of removing branched-chain and unsaturated hydrocarbons from straight-chain hydrocarbons depends on the much faster reaction of the former with chlorosulfonic acid.

Isomeric materials which have closely similar physical properties can be serious contaminants in hydrocarbons. With aromatic hydrocarbons, e.g. xylenes and alkyl benzenes, advantage is taken of differences in ease of sulfonation. If the required compound is sulfonated more readily, the sulfonic acid is isolated, crystallised (e.g. from water), and decomposed by passing superheated steam through the flask containing the acid. The sulfonic

acid undergoes hydrolysis, and the liberated hydrocarbon distils with the steam. It is separated from the distillate, dried, distilled and then fractionally distilled. For small quantities (10-100mg), vapour phase chromatography is the most satisfactory method for obtaining a pure sample (for column materials for packings see Chapter 1). Azeotropic distillation with methanol or 2-ethoxyethanol (cellosolve) has been used to obtain highly purified saturated hydrocarbons and aromatic hydrocarbons such as xylenes and isopropylbenzenes. Carbonyl-containing impurities can be removed from hydrocarbons (and other oxygen-lacking solvents such as CHCl_3 and CCl_4) by passage through a column of Celite 545 (100g) mixed with concentrated sulfuric acid (60ml). After first adding some solvent and about 10g of granular Na_2SO_4 , the column is packed with the mixture and a final 7-8cm of Na_2SO_4 is added at the top [Hornstein & Crowe, *Anal Chem* **34** 1037 1962, DOI: 10.1021/ac60188a051]. Alternatively, Celite impregnated with 2,4-dinitrophenylhydrazine can be used.

With solid hydrocarbons such as naphthalene and polycyclic hydrocarbons, preliminary purification by sublimation in vacuum (or high vacuum if the substance is high melting) is followed by zone refining and finally by chromatography (e.g. on alumina) using low-boiling liquid hydrocarbon eluents. These solids can be recrystallised from alcohols, alcohol/petroleum ether or from liquid hydrocarbons (e.g. toluene) and dried below their melting points. Aromatic hydrocarbons that have been purified by zone melting include anthracene, biphenyl, fluoranthrene, naphthalene, perylene, phenanthrene, pyrene and terphenyl, among others. Some polycyclic hydrocarbons, e.g. benzopyrene, are CARCINOGENIC.

Olefinic hydrocarbons have a very strong tendency to polymerise, and commercially available materials are generally stabilised, e.g. with hydroquinone. When distilling compounds such as vinylpyridine or styrene, the stabiliser remains behind and the purified olefinic material is more prone to polymerisation. The most common impurities are higher-boiling dimeric or polymeric compounds. Vacuum distillation in a nitrogen atmosphere not only separates monomeric from polymeric materials but in some cases also depolymerises the impurities. The distillation flask should be charged with a polymerisation inhibitor, and the purified material should be used immediately or stored in the dark and mixed with a small amount of stabiliser (e.g. 0.1% of hydroquinone or di-*tert*-butylcatechol). It is also advisable to add to the flask a small amount (ca 5-10% by volume of liquid in the flask) of a ground mixture of Kieselguhr and NaCl which will provide nuclei for facilitating boiling and finally for cleaning the flask from insoluble polymeric residue (due to the presence of the water soluble NaCl in the residue).

IMIDES

Imides (e.g. phthalimide) can be purified by conversion to their potassium salts by reaction in ethanol with ethanolic potassium hydroxide. The imides are regenerated when the salts are hydrolysed with water or dilute acid. Like amides, imides readily crystallise from alcohols and, in some cases (e.g. quinolinic imide), from glacial acetic acid.

IMINO COMPOUNDS

These substances contain the -C=NH group and, because they are strong, unstable bases, they are kept as their more stable salts, such as the hydrochlorides. (The free base usually hydrolyses to the corresponding oxo compound and ammonia.) Like amine hydrochlorides, the salts are purified by solution in alcohol containing a few drops of hydrochloric acid. After treatment with charcoal, and filtering, dry diethyl ether (or petroleum ether if ethanol is used) is added until crystallisation sets in. The salts are dried and kept in a vacuum desiccator.

KETONES

Ketones are more stable to oxidation than aldehydes and can be purified from oxidisable impurities by refluxing with potassium permanganate until the colour persists, followed by shaking with sodium carbonate (to remove acidic impurities) and distilling. Traces of water can be removed with type 4A Linde molecular sieves. Ketones which are solids can be purified by crystallisation from alcohol, toluene, or petroleum ether, and are usually sufficiently volatile for sublimation in vacuum. Ketones can be further purified *via* their bisulfite, semicarbazone or oxime derivatives (*vide supra*). The bisulfite addition compounds are formed only by aldehydes and methyl ketones but they are readily hydrolysed in dilute acid or alkali.

MACROMOLECULES See Chapter 6.

NITRILES

All purifications should be carried out in an efficient fume cupboard because of the **TOXIC** nature of these compounds. Nitriles are usually prepared either by reacting the corresponding halide or diazonium salts with a cyanide salt or by dehydrating an amide. Hence, possible contaminants are the respective halides or alcohols (from hydrolysis), phenolic compounds, amines or amides. Small quantities of phenols can be removed by chromatography on alumina. More commonly, purification of liquid nitriles or solutions of solid nitriles in a solvent such as diethyl ether is by shaking with dilute aqueous sodium hydroxide, followed by washing successively with water, dilute acid and water. After drying with sodium sulfate, the solvent is distilled off. Liquid nitriles are best distilled from a small amount of P_2O_5 which, besides removing water, dehydrates any amide impurity to the nitrile. About one-fifth of the nitrile should remain in the distilling flask at the end of the distillation (*the residue may contain some inorganic cyanide*). This purification also removes alcohols and phenols. Solid nitriles can be recrystallised from ethanol, toluene or petroleum ether, or a mixture of these solvents. They can also be sublimed under vacuum. Preliminary purification by steam distillation is usually possible. Strong alkali or heating with dilute acids may lead to hydrolysis of the nitrile and should be avoided. They have a characteristic band in the IR spectrum at $\nu_{\max} \sim 2200 \text{ cm}^{-1}$.

NITRO COMPOUNDS

Aliphatic nitro compounds are generally acidic. They are freed from alcohols or alkyl halides by standing for a day with concentrated sulfuric acid, then washed with water, dried with magnesium sulfate followed by calcium sulfate and distilled. The principal impurities are isomeric or homologous nitro compounds. In cases where the nitro compound was originally prepared by vapour phase nitration of the aliphatic hydrocarbon, fractional distillation should separate the nitro compound from the corresponding hydrocarbon. Fractional crystallisation is more effective than fractional distillation if the melting point of the compound is not too low.

The impurities present in aromatic nitro compounds depend on the aromatic portion of the molecule. Thus, benzene, phenols or anilines are probable impurities in nitrobenzene, nitrophenols and nitroanilines, respectively. Purification should be carried out accordingly. Isomeric compounds are likely to remain as impurities after the preliminary purifications to remove basic and acidic contaminants. For example, *o*-nitrophenol may be found in samples of *p*-nitrophenol. Usually, the *o*-nitro compounds are more steam volatile than the *p*-nitro isomers and can be separated in this way. Polynitro impurities in mononitro compounds can be readily removed because of their relatively lower solubilities in solvents. With acidic or basic nitro compounds that cannot be separated in the above manner, advantage may be taken of their differences in pK values (see Chapter 1). The compounds can thus be purified by preliminary extractions with several sets of aqueous buffers of known pH (see for example Table 24, Chapter 1) from a solution of the substance in a suitable solvent such as diethyl ether. This method is more satisfactory and less laborious the larger the difference between the pK value of the impurity and the desired compound. Heterocyclic nitro compounds require similar treatment to the nitroanilines. Neutral nitro compounds can be steam distilled.

NUCLEIC ACIDS See Chapter 6.

PHENOLS

Because phenols are weak acids, they can be freed from neutral impurities by dissolution in aqueous N sodium hydroxide (cf pK) and extraction with a solvent such as diethyl ether, or by steam distillation to remove the non-acidic material. The phenol is recovered by acidification of the aqueous phase with 2N sulfuric acid, and either extracted with ether or steam distilled. In the second case the phenol is extracted from the steam distillate after saturating it with sodium chloride (salting out). A solvent is necessary when large quantities of liquid phenols are purified. The phenol is fractionated by distillation under reduced pressure, preferably in an atmosphere of nitrogen to minimise oxidation. Solid phenols can be crystallised from toluene, petroleum ether or a mixture of these solvents, and can be sublimed under vacuum. Purification can also be effected by fractional crystallisation or zone refining. For further purification of phenols *via* their acetyl or benzoyl derivatives *vide supra*.

POLYPEPTIDES AND PROTEINS See Chapter 6.

QUINONES

These are neutral compounds which are usually coloured. They can be separated from acidic or basic impurities

by extraction of their solutions in organic solvents with aqueous basic or acidic solutions, respectively. Their colour is a useful property in their purification by chromatography through an alumina column with, e.g. toluene, as eluent. They are volatile enough for vacuum sublimation, although with high-melting quinones a very high vacuum is necessary. *p*-Quinones are stable compounds and can be recrystallised from water, ethanol, aqueous ethanol, toluene, petroleum ether or glacial acetic acid. *o*-Quinones, on the other hand, are readily oxidised. They should be handled in an inert atmosphere, preferably in the absence of light.

SALTS

With metal ions

Water-soluble salts are best purified by preparing a concentrated aqueous solution to which, after decolorising with charcoal and filtering, ethanol or acetone is added so that the salts crystallise. They are collected, washed with aqueous ethanol or aqueous acetone, and dried. In some cases, water-soluble salts can be recrystallised satisfactorily from alcohols. With very water-soluble salts, pure crystals are best obtained by dissolving them in water and allowing the solution to evaporate slowly in a desiccator over a suitable desiccant in a cold room. When crystals are formed they are removed, e.g. by centrifugation, washed with a little ice-cold water and dried in a vacuum. Water-insoluble salts are purified by Soxhlet extraction, first with organic solvents and then with water, to remove soluble contaminants. The purified salt is recovered from the thimble.

With organic cations

Organic salts (e.g. trimethylammonium benzoate) are usually purified by recrystallisation from polar solvents (e.g. water, ethanol or dimethyl formamide). If the salt is too soluble in a polar solvent, its concentrated solution should be treated dropwise with a miscible non-polar, or less polar, solvent (see Tables 6 and 7, Chapter 1) until crystallisation begins and cooled.

With sodium alkane sulfonates

These are purified from sulfites by boiling with aqueous HBr. They are purified from sulfates by adding BaBr₂. Sodium alkane disulfonates are finally precipitated by addition of MeOH [Pethybridge & Taba *JCS Faraday Trans 1* **78** 1331 1982, DOI: 10.1039/F19827801331].

SULFUR COMPOUNDS

Disulfides

These can be purified by extracting acidic and basic impurities with dilute aqueous base or acid, respectively. However, they are somewhat sensitive to strong alkali which slowly cleaves the disulfide bond. The lower-melting members can be fractionally distilled under vacuum. The high members can be recrystallised from alcohol, toluene or glacial acetic acid.

Sulfones

Sulfones are neutral and very stable compounds that can be distilled without decomposition. They are freed from acidic and basic impurities in the same way as disulfides. The low-molecular-weight members are quite soluble in water, but the higher members can be recrystallised from water, ethanol, aqueous ethanol or glacial acetic acid.

Sulfonic acids

These are strong acids whose solubilities in water varies. They can be recrystallised from water, sometimes form hydrates and can be characterised and purified *via* their S-aryliothiuronium salts, see Tables 6 and 7, and above.

Sulfoxides

These are odourless, rather unstable compounds because they oxidise to sulfones, and should be distilled under vacuum in an inert atmosphere. They are generally water-soluble but can be extracted from aqueous solution with a solvent such as diethyl ether.

Thioethers (sulfides)

Thioethers are neutral stable compounds that can be freed from acidic and basic impurities as described for disulfides and are more stable than disulfides in alkaline solutions. They can be recrystallised from organic solvents and distilled without decomposition. They have sulfurous odours.

Thiols

Thiols, or mercaptans, are stronger acids than the corresponding aliphatic hydroxy or phenolic compounds, but can be purified in a similar manner. However, care must be exercised in handling thiols to avoid their oxidation to disulfides. For this reason, purification is best carried out in an inert atmosphere in the absence of oxidising agents. Similarly, thiols should be stored out of contact with air. They can be distilled without change, and the higher-melting thiols (which are usually more stable) can be crystallised, e.g. from water or dilute alcohol. They oxidise readily in alkaline solution but can be separated from the disulfide which is insoluble in this medium.

They should be stored in the dark below 0°. *All operations with thiols should be carried out in an efficient fume cupboard because of their very unpleasant odour and their TOXICITY.*

Thiolsulfonates (disulfoxides)

Thiolsulfonates are neutral and are somewhat light-sensitive compounds. Their most common impurities are sulfonyl chlorides (neutral) or the sulfinic acid or disulfide from which they are usually derived. The first can be removed by partial freezing or crystallisation, the second by shaking with dilute alkali, and the third by recrystallisation because of the higher solubility of the disulfide in solvents. Thiolsulfonates decompose slowly in dilute, or rapidly in strong, alkali to form disulfides and sulfonic acids. Thiolsulfonates also decompose on distillation but they can be steam distilled. The solid members can be recrystallised from water, alcohols or glacial acetic acid.

PURIFICATION *via* FLUOROCHROMES

If the purification procedure is proving difficult then by tagging the desired molecule in a mixture with a fluorochrome (see Table 20, Chapter 1) can provide a means of following the substance through the purification process. The fluorochrome should have a group which can react with the desired compound, and it should be possible to remove the fluorochromic group after purification. Such groups are present for example in fluorescein-isothiocyanate (FITC), where the SCN group can react with an RNH₂ compound to form fluorescent thioureas; e.g. 4-bromomethyl-7-methoxycoumarin, which can react with R-OH or R-COONa compounds to form fluorescent ether or ester links with the desired compound; or 3-aminocoumarin, which reacts through its NH₂ group to form fluorescent amides. The fluorescent products can then be readily identified by their fluorescence, separated from impurities, and it should be possible to recover the purified compound after chemically separating it from the fluorescent tag. Such procedures can also be used as analytical tools for detecting specific substances (see *fluorescence spectra* in Chapter 1 and Table 20).

ADVANCES IN CHEMICAL TECHNIQUES USED IN PURIFICATION

Chemical techniques used in purification mainly involve conversion of compounds to be purified into derivatives, e.g. nitrobenzoyl derivatives (where covalent bonds have been formed), into complexes or adducts such as phenanthrene picrate, or into a salt, e.g. aniline bis-4-chlorobenzoylimide salt which can be purified by recrystallisation or some other means. The purified compounds are then recovered by cleaving the covalent bond in the derivatives, by removal of picric acid, e.g. by passage through alumina, or by basifying the salts and subsequently isolating the *free* bases. These procedures can increase the purity of compounds considerably. There are still continuing efforts in devising new ways to derivatise compounds, particularly for selecting desired functional groups in such a manner that selective removal of the derivatising group can be achieved in high yield. This has been very successful for synthesising most elaborate structures. However, with all the previous work and current improvements, [see P. G. M. Wuts, T. W. Greene, *Greene's Protective Groups in Organic Synthesis*, 4th Edition, J.Wiley & Sons Inc, 2006. DOI: 10.1002/0470053488; Print ISBN: 9780471697541, Online ISBN: 9780470053485 and <synarchive.com>], the present armoury of protecting and derivatising groups limit the use of chemical methods for the purification of compounds to small scale work. That is because **atom economy** is not possible on large scale purification due to the necessary wasteful jettisoning or recycling of unwanted products. Medium to large scale purification necessitates the use of physical techniques (see Chapter 1), and a large number have been developed and are continually being improved. The strength of chemical techniques lies in the analytical information that can be derived which not only conveniently provides information of what, and how much, impurities are present but also can help to suggest the best procedures to use for purification.

For advances in *Laboratory on a Chip* and *microfluidics* and their potential use in purification, analyses and chemical syntheses see 'ADVANCES IN PHYSICAL METHODS USED IN PURIFICATION' in Chapter 1.

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CHAPTER 3

PURIFICATION OF ORGANIC CHEMICALS

INTRODUCTION

The general principles, techniques and methods of purification in Chapters 1 and 2 are applicable to this chapter. Most organic liquids and a number of solids can readily be purified by fractional distillation, usually at atmospheric pressure. Sometimes, particularly with high boiling or sensitive liquids, or when in doubt about stability, distillation or fractionation under as low a pressure as possible should be carried out. To save space, the present chapter omits many substances for which the published purification methods involve simple distillation. Where boiling points are given, purification by distillation is another means of removing impurities. Literature references, and in particular *Beilstein* references, are included for most entries which refer the reader directly or indirectly to the original sources. Substances are listed alphabetically in each section, usually with some criteria of purity, giving brief details of how they can be purified. Also noted are the empirical formulae, molecular weights (to the first decimal place), melting points and/or boiling points together with the respective densities and refractive indexes for liquids, and specific optical rotations for chiral compounds. All temperatures are in degrees *centigrade* unless stated otherwise. When temperatures and/or wavelengths are not given for the last three named properties, then they should be assumed to be 20°C and the average of the wavelengths of the sodium D lines respectively; and most densities are relative to water at 4°C. When pressure is stated as **atm** it means that it is not mentioned specifically in the literature, but is assumed to be 760mm of Hg or 101.325 kPa.

Ionisation constants of ionisable compounds are given as **pK** values (published in the literature) and refer to the **pKa** values at room temperature (~ 15°C to 25°C) when not specified. Values at other temperatures are given as superscripts, e.g. **pK³⁰** for 30°C. Estimated values are entered as **pK_{Est}** (see Chapter 1).

Rapid purification procedures are included for commonly used solvents and reagents which make them suitable for general use in synthetic chemistry. Commercially available polymer supported reagents are indicated with § under the appropriate reagent.

Abbreviations of titles of periodicals are defined as in the Chemical Abstracts Service Source Index (CASSI) but without punctuation. References to Fieser & Fieser's *Reagents for Organic Synthesis* are shortened to Fieser throughout, e.g. Fieser **2** 254, **11** 88, etc. Other abbreviations are self evident.

As a good general rule, all low boiling (<100°) organic liquids should be treated as highly flammable and toxic (because they can be inhaled in large quantities) and the necessary precautions should be taken (see Safety precautions associated with the purification of laboratory chemicals in Chapter 1).

Benzene has been used as a solvent successfully and extensively in the past for reactions and purification but is now considered a **very dangerous and CARCINOGENIC substance**. It should be used with extreme care. We emphasise that alternative solvents (e.g. toluene, toluene-petroleum ether, or a petroleum ether to name a few) should be used first. However, if no other solvent system can be found, then all operations involving benzene (and storage) should be performed in an efficiently running fumehood, and precautions taken to avoid inhalation and contact with skin and eyes. An asterisk has been inserted in the text, e.g. ^{*}C₆H₆ or ^{*}benzene, to remind the user that special precautions should be adopted.

This chapter consists of four sections *viz*: **Aliphatic Compounds, Alicyclic Compounds, Aromatic Compounds, Heterocyclic Compounds** and **Miscellaneous As, B, P, Si, S, Se, and Te Compounds**.

Purification of Laboratory Chemicals.

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ALIPHATIC COMPOUNDS

Acetal (acetaldehyde diethylacetal) [105-57-7] $\text{C}_6\text{H}_{14}\text{O}_2$, **M 118.2**, **b 103.7-104°**, **d**₄²⁰ **0.831**, **n**_D²⁵ **1.38054**, **n**_D²⁵ **1.3682**. Dry acetal over Na to remove alcohols and H_2O , and to polymerise aldehydes, then fractionally distil. Or, treat it with alkaline H_2O_2 at 40-45° to remove aldehydes, then saturate with NaCl, separate, dry with K_2CO_3 and distil from Na [Vogel *J Chem Soc* 616 1948, DOI: 10.1039/JR9480000616]. [*Beilstein* 1 IV 3103.]

Acetaldehyde [75-07-0] $\text{C}_2\text{H}_4\text{O}$, **M 44.1**, **b 20.2°/atm**, **d**₄²⁰ **0.788**, **n**_D²⁵ **1.33113**, **pK**²⁵ **13.57** (hydrate). Acetaldehyde is usually purified by fractional distillation in a glass helices-packed column under dry N_2 , discarding the first portion of distillate. Or, it is shaken for 30 minutes with NaHCO_3 , dried with CaSO_4 and fractionally distilled at 760mm through a 70cm Vigreux column. The middle fraction is collected and further purified by standing for 2 hours at 0° with a small amount of hydroquinone (free radical inhibitor), followed by distillation [Longfield & Walters *J Am Chem Soc* 77 810 1955, DOI: 10.1021/ja01608a094]. [*Beilstein* 1 IV 3094.]

Acetaldehyde dimethyl acetal (1,1-dimethoxyethane) [534-15-6] $\text{C}_4\text{H}_{10}\text{O}_2$, **M 90.1**, **b 63-65°/atm**, **d**₄²⁰ **0.852**, **n**_D²⁵ **1.36678**. Distil the dimethyl acetal through a fractionating column and the fraction boiling at 63.8°/751mm is collected. It forms an azeotrope with MeOH. *Alternatively*, purify it as for *acetal* above. It has been purified by GLC. [*Beilstein* 1 IV 3103.]

Acetamide [60-35-5] $\text{C}_2\text{H}_5\text{NO}$, **M 59.1**, **m 81°**, **pK**₁²⁵ **-1.4**, **pK**₂²⁵ **+0.37**. Acetamide is crystallised by dissolving in hot MeOH (0.8ml/g), diluting with Et_2O and allowing to stand [Wagner *J Chem Edu* 7 1135 1930, DOI: 10.1021/ed007p1135]. Alternate crystallisation solvents are acetone, *benzene, chloroform, dioxane, methyl acetate or *benzene/ethyl acetate mixtures (3:1 and 1:1). It has also been recrystallised from hot water after treating with HCl-washed activated charcoal (which had been repeatedly washed with water until free from chloride ions), then crystallised again from hot 50% aqueous EtOH and finally twice from hot 95% EtOH [Christoffers & Kegeles *J Am Chem Soc* 85 2562 1963, DOI: 10.1021/ja00900a005]. Finally it is dried in a vacuum desiccator over P_2O_5 . Acetamide is also purified by distillation (**b 221-223°/atm**) or by sublimation *in vacuo*. It has also been purified by two recrystallisations from cyclohexane containing 5% (v/v) of *benzene. Needle-like crystals separate and are filtered, washed with a small volume of distilled H_2O and dried with a flow of dry N_2 . [Slebocka-Tilk et al. *J Am Chem Soc* 109 4620 1987, DOI: 10.1021/ja00249a027; *Beilstein* 2 H 175, 2 I 80, 2 II 177, 2 III 384, 2 IV 399.]

Acetamidine hydrochloride [124-42-5] $\text{C}_2\text{H}_7\text{ClN}_2$, **M 94.5**, **m 164-166°, 165-170°(dec)**, **174°**, **pK**²⁵ **12.40**. The hydrochloride can be recrystallised from small volumes of EtOH. *Alternatively*, it is dissolved in EtOH, filtered, Et_2O is added; filter the crystalline salt off under N_2 and dry it in a vacuum desiccator over H_2SO_4 . The salt is deliquescent and should be stored in a tightly stoppered container. Its solubility in H_2O is 10% at room temperature and it is soluble in Me_2CO . The *free base* reacts strongly alkaline in H_2O . It has λ_{max} 224nm (ϵ 4000) in H_2O . The *picrate* has **m 252°** (sintering at ~245°). [Dox *Org Synth Coll Vol I* 5 1941, DOI: 10.15227/orgsyn.008.0001; Davies & Parsons *Chem Ind (London)* 628 1958, Barnes et al. *J Am Chem Soc* 62 1281 1940, DOI: 10.1021/ja01862a086, give **m 177-178°**; *Beilstein* 2 H 185, 2 I 85, 2 II 183, 2 III 416, 2 IV 428.]

N-(2-Acetamido)-2-aminoethanesulfonic acid (ACES) [7365-82-4] $\text{C}_4\text{H}_{10}\text{N}_2\text{O}_4\text{S}$, **M 182.2**, **m > 220°(dec)**, **pK**_{Est} **~1.5**, **pK**₂ **6.9**. Recrystallise ACES from hot aqueous EtOH. [Perrin & Dempsey *Buffers for pH and Metal Ion Control* Chapman & Hall, London 1974, ISBN 0412117002; *Beilstein* 4 III 1707.]

N-(2-Acetamido)iminodiacetic acid (ADA) [26239-55-4] $\text{C}_6\text{H}_{10}\text{N}_2\text{O}_5$, **M 190.2**, **m 219°(dec)**, **pK**₁ **~2.3**, **pK**₂ **6.6**. Dissolve ADA in water, add one equivalent of NaOH solution (to final pH of 8-9), then acidify with HCl to precipitate the free acid. This is filtered off, washed with water and dried *in vacuo*. [*Beilstein* 4 IV 2441.]

Acetamidomethanol [625-51-4] $\text{C}_3\text{H}_7\text{NO}_2$, **M 89.1**, **m 47-50°, 54-56°, 55°**. Recrystallise it from freshly

distilled Me_2CO , wash the crystals with dry Et_2O and dry them in a vacuum desiccator over P_2O_5 . R_F 0.4 on paper chromatography with $\text{CHCl}_3/\text{EtOH}$ (2:8) as solvent and developed with ammoniacal AgNO_3 . It also crystallises in needles from EtOAc containing a few drops of Me_2CO . It is *hygroscopic* and should be stored under dry conditions. [Bachmann et al. *J Am Chem Soc* **73** 2775 1951, DOI: 10.1021/ja01150a101; Walter et al. *Chem Ber* **99** 3204 1966, DOI: 10.1002/cber.19660991019; Einhorn & Ladisch *Justus Liebigs Ann Chem* **343** 265 1905, DOI: 10.1002/jlac.19053430207; *Beilstein* **2** IV 405.]

Acetic acid (glacial) [64-19-7] $\text{C}_2\text{H}_4\text{O}_2$, **M 60.1**, **m 16.6°**, **b 118°/atm**, **d₄²⁰ 1.049**, **n_D²⁵ 1.37171**, **n_D²⁵ 1.36995**, **pK_a²⁵ 4.76**. Usual impurities are traces of acetaldehyde and other oxidisable substances and water. Glacial acetic acid is very *hygroscopic*. The presence of 0.1% water lowers its **m** by 0.2°. Purify it by adding some acetic anhydride to react with water present, heat it for 1 hour to just below boiling in the presence of 2g CrO_3 per 100ml and then fractionally distil it [Orton & Bradfield *J Chem Soc* 983 1927, DOI: 10.1039/JR9270000983]. Instead of CrO_3 , use 2-5% (w/w) of KMnO_4 , and boil under reflux for 2-6 hours. Traces of water have been removed by refluxing with tetraacetyl diborate (prepared by warming 1 part of boric acid with 5 parts (w/w) of acetic anhydride at 60°, cooling, and filtering off, followed by distillation [Eichelberger & La Mer *J Am Chem Soc* **55** 3633 1933, DOI: 10.1021/ja01336a023].

Refluxing with acetic anhydride in the presence of 0.2g % of 2-naphthalenesulfonic acid as catalyst has also been used [Orton & Bradfield *J Chem Soc* 983 1927, DOI: 10.1039/JR9270000983]. Other suitable drying agents include anhydrous CuSO_4 and chromium triacetate: P_2O_5 converts some acetic acid to the anhydride. Azeotropic removal of water by distillation with thiophene-free *benzene or with butyl acetate has been used [Birdwhistell & Griswold *J Am Chem Soc* **77** 873 1955, DOI: 10.1021/ja01609a016]. An alternative purification uses fractional freezing. [*Beilstein* **2** H 96, **2** IV 94.] **Rapid procedure:** Add 5% acetic anhydride, and 2% of CrO_3 . Reflux and fractionally distil.

Acetic anhydride [108-24-7] $\text{C}_4\text{H}_6\text{O}_3$, **M 102.1**, **b 138°/atm**, **d₄²⁰ 1.082**, **n_D²⁵ 1.3904**. Adequate purification can usually be achieved by fractional distillation through an efficient column. Acetic acid can be removed by prior refluxing with CaC_2 or with coarse Mg filings at 80-90° for 5 days, or by distillation from a large excess of quinoline (1% AcOH in quinoline) at 75mm pressure. Acetic anhydride can also be dried by standing with Na wire for up to a week, removing the Na and distilling it under vacuum. (Na reacts vigorously with acetic anhydride at 65-70°). Dippy & Evans [*J Org Chem* **15** 451 1950, DOI: 10.1021/jo01149a001] let the anhydride (500g) stand over P_2O_5 (50g) for 3 hours, then decanted it and stood it over ignited K_2CO_3 for a further 3 hours. The supernatant liquid was distilled and the fraction **b** 136-138° was further dried with P_2O_5 for 12 hours, followed by shaking with ignited K_2CO_3 , before two further distillations through a five-section Young and Thomas fractionating column. The final material distilled at 137.8-138.0°/atm. It can also be purified by azeotropic distillation with toluene: the azeotrope boils at 100.6°. After removal of the remaining toluene, the anhydride is distilled [sample had a specific conductivity of $5 \times 10^{-9} \text{ ohm}^{-1}\text{cm}^{-1}$]. [*Beilstein* **2** H 96, **2** I 39, **2** II 91, **2** III 134, **2** IV 94.] **Rapid procedure:** Shake with P_2O_5 , separate, shake with dry K_2CO_3 and fractionally distil. **Acetic hydrazide** [1068-57-1] $\text{C}_2\text{H}_6\text{N}_2\text{O}$, **M 74.1**, **m 67°**, **b 127°/18mm**, crystallises as needles from EtOH . It reduces $\text{NH}_3/\text{AgNO}_3$. [*Beilstein* **2** H 191, **2** IV 435.]

Acetoacetamide [5977-14-0] $\text{C}_4\text{H}_7\text{NO}_2$, **M 101.1**, **m 54-55°, 54-56°**. Recrystallise the amide from CHCl_3 , or Me_2CO /petroleum ether. It also crystallises from pyridine with 4mols of solvent. It is slightly soluble in H_2O , EtOH and AcOH but is insoluble in Et_2O . The *phenylhydrazone* has **m 128°**. [Kato *Chem Pharm Bull Jpn* **15** 921 923 1967, DOI: org/10.1248/cpb.15.921; Claissen & Meyer *Chem Ber* **35** 583 1902, DOI: 10.1002/cber.19020350190; *Beilstein* **3** H 659, **3** I 231, **3** III 1204, **3** IV 1545.]

Acetone [67-64-1] $\text{C}_3\text{H}_6\text{O}$, **M 58.1**, **b 56.2°/atm**, **d₄²⁰ 0.791**, **n_D²⁵ 1.35880**, **pK₁²⁵ -6.1 (basic, mono-protonated)**, **pK₂²⁵ 20.0 (acidic)**. The commercial preparation of acetone by catalytic dehydrogenation of isopropyl alcohol gives relatively pure material. Analytical reagent quality generally contains less than 1% of organic impurities but may have up to about 1% of H_2O . Dry acetone is appreciably *hygroscopic*. The main organic impurity in acetone is mesityl oxide, formed by aldol condensation. It can be dried with anhydrous CaSO_4 , K_2CO_3 or type 4A Linde molecular sieves, and then distilled. Silica gel and alumina, or mildly acidic or basic desiccants cause acetone to undergo the aldol condensation, so that its water content is increased by pass-

age through these reagents. This also occurs to some extent when P_2O_5 or sodium amalgam is used. Anhydrous $MgSO_4$ is an inefficient drying agent, and $CaCl_2$ forms an addition compound. Drierite (anhydrous $CaSO_4$) offers minimum acid and base catalysis for aldol formation and is the recommended drying agent for this solvent [Riddick & Bunger *Organic Solvents* Wiley-Interscience, NY, 3rd edn, 1970]. Acetone can be shaken with Drierite (25g/L) for several hours before it is decanted and distilled from fresh Drierite (10g/L) through an efficient column, maintaining atmospheric contact through a Drierite drying tube.

The equilibrium water content is about $10^{-2}M$. Anhydrous $Mg(ClO_4)_2$ **should not be used as drying agent as there is a high risk of EXPLOSION with acetone vapour.**

Organic impurities have been removed from acetone by adding 4g of $AgNO_3$ in 30ml of water to 1L of acetone, followed by 10ml of M $NaOH$, shaking for 10 minutes, filtering, drying with anhydrous $CaSO_4$ and distilling [Hawley et al. *Analyst (London)* **58** 333 1933, DOI: 10.1039/AN9335800333]. Alternatively, successive small portions of $KMnO_4$ have been added to acetone at reflux, until the violet colour persists, followed by drying and distilling. Refluxing with chromium trioxide (CrO_3) has also been used. Methanol has been removed from acetone by azeotropic distillation (at 35°) with methyl bromide, and treatment with acetyl chloride.

Small amounts of acetone can be purified as the NaI addition compound, by dissolving 100g of finely powdered NaI in 400g of boiling acetone, then cooling in ice and salt to -8° . Crystals of $NaI \cdot 3Me_2CO$ are filtered off and, on warming in a flask, acetone distils off readily. [This method is more convenient than the one using the bisulfite addition compound.] It has also been purified by gas chromatography on a 20% free fatty acid phthalate (on Chromosorb P) column at 100° .

For efficiency of desiccants in drying acetone see Burfield and Smithers [*J Org Chem* **43** 3966 1978, DOI: 10.1021/jo00414a038]. The water content of acetone can be determined by a modified Karl Fischer titration [Koupparis & Malmstadt *Anal Chem* **54** 1914 1982, DOI: 10.1021/ac00248a072]. [*Beilstein* **1** IV 3180.]

Rapid procedure: Dry over anhydrous $CaSO_4$ and distil. *Acetoxime (acetone oxime)* [127-06-0] C_3H_7NO , **M 73.1**, **m** 63° , **b** $135^\circ/760mm$, **d**₄²⁰ **0.901**, **pK**⁴⁰ **0.99**, crystallises from petroleum ether (b $40-60^\circ$) and can be sublimed. [*Beilstein* **1** H 649, **1** IV 3202.] *Acetone semicarbazone* [110-20-3] $C_4H_{10}N_3O$, **M 115.1**, **m** 187° , **pK**²⁵ **1.33**, crystallises from water or from aqueous $EtOH$. [*Beilstein* **3** H 101, **3** I 48, **3** II 81, **3** III 189, **3** IV 179.]

Acetone cyanohydrin [75-86-5] C_4H_7NO , **M 85.1**, **b** $48^\circ/2.5mm$, $68-70^\circ/11mm$, $78-82^\circ/15mm$, **d**₄²⁰ **0.93**. Dry the cyanohydrin with Na_2SO_4 , and distil it as rapidly as possible under vacuum to avoid decomposition. Discard fractions boiling below $78-82^\circ/15mm$. Store it in the dark. **USE AN EFFICIENT FUME HOOD as HCN (POISONOUS) released is always present.** [Cox & Stormont *Org Synth Coll Vol* **2** 7 1940, DOI: 10.15227/orgsyn.015.0001; *Beilstein* **3** H 316, **3** IV 785.]

Acetonedicarboxylic acid (3-oxoglutaric acid) [542-05-2] $C_5H_6O_5$, **M 146.1**, **m** $138^\circ(dec)$, **pK**²⁵ **3.10**. Crystallise it from ethyl acetate and store it over P_2O_5 . It decarboxylates in hot water. [*Beilstein* **3** IV 1816.]

Acetonitrile (methyl cyanide) [75-05-8] C_2H_3N , **M 41.1**, **b** $81.6^\circ/atm$, **d**₄²⁵ **0.77683**, **n**_D²⁰ **1.3441**, **n**_D²⁵ **1.34163**. Commercial acetonitrile is a by-product of the reaction of propylene and ammonia to acrylonitrile. The following procedure that significantly reduces the levels of acrylonitrile, allyl alcohol, acetone, and *benzene was used by Kiesele [*Anal Chem* **52** 2230 1988, DOI: 10.1021/ac50063a062]. Methanol (300ml) is added to 3L of acetonitrile fractionated at high reflux ratio until the boiling temperature rises from 64° to 80° , and the distillate becomes optically clear down to $\lambda = 240nm$. Add sodium hydride (1g) free from paraffin, to the liquid, reflux for 10 minutes, and then distil rapidly until about 100ml of residue remains. Immediately pass the distillate through a column of acidic alumina, discarding the first 150ml of percolate. Add 5g of CaH_2 and distil the first 50ml at a high reflux ratio. Discard this fraction, and collect the following main fraction. The best way of detecting impurities is by gas chromatography.

Usual contaminants in commercial acetonitrile include H_2O , acetamide, NH_4OAc and NH_3 . Anhydrous $CaSO_4$ and $CaCl_2$ are inefficient drying agents. Preliminary treatment of acetonitrile with cold, saturated aqueous KOH is undesirable because of base-catalysed hydrolysis and the introduction of water. Drying by shaking with silica gel or Linde 4A molecular sieves removes most of the water in acetonitrile. Subsequent stirring with CaH_2 until no further hydrogen is evolved leaves only traces of water and removes acetic acid. The acetonitrile is then fractionally distilled at high reflux, taking precaution to exclude moisture by refluxing over CaH_2 [Coetzee *Pure Appl Chem* **13** 427 1966, DOI: org.virtual.anu.edu.au/10.1351/pac196613030427]. Alternatively, 0.5-1% (w/v)

P_2O_5 is often added to the distilling flask to remove most of the remaining water. Excess P_2O_5 should be avoided because it leads to the formation of an orange polymer. Traces of P_2O_5 can be removed by distilling from anhydrous K_2CO_3 .

Kolthoff, Bruckenstein and Chantooni [*J Am Chem Soc* **83** 3927 1961, DOI: 10.1021/ja01480a001] removed acetic acid from 3L of acetonitrile by shaking for 24 hours with 200g of freshly activated alumina (which had been reactivated by heating at 250° for 4 hours). The decanted solvent was again shaken with activated alumina, followed by five batches of 100-150g of anhydrous $CaCl_2$. (Water content of the solvent was then less than 0.2%.) It was shaken for 1 hour with 10g of P_2O_5 , twice; and distilled in a 1m x 2cm column, packed with stainless steel wool and protected from moisture by $CaCl_2$ tubes. The middle fraction had a water content from 0.7 to 2mM. Traces of unsaturated nitriles can be removed by initially refluxing with a small amount of aqueous KOH (1ml of 1% solution per L). Acetonitrile can be dried by azeotropic distillation with dichloromethane, *benzene or trichloroethylene. Isonitrile impurities can be removed by treatment with concentrated HCl until the odour of isonitrile has gone, followed by drying with K_2CO_3 and distilling.

Acetonitrile is refluxed with, and distilled from alkaline $KMnO_4$ and $KHSO_4$, followed by fractional distillation from CaH_2 . (This is better than fractionation from molecular sieves or passage through a type H activated alumina column, or refluxing with KBH_4 for 24 hours and fractional distillation) [Bell et al. *JCS Faraday Trans* **73** 315 1977, DOI: 10.1039/F19777300315; Moore et al. *J Am Chem Soc* **108** 2257 1986, DOI: 10.1021/ja00269a022].

Material suitable for polarography is obtained by refluxing over anhydrous $AlCl_3$ (15g/L) for 1 hour, distilling, refluxing over Li_2CO_3 (10g/L) for 1 hour and redistilling. It is then refluxed over CaH_2 (2g/L) for 1 hour and fractionally distilled, retaining the middle portion. The product is not suitable for UV spectroscopy use. A better purification procedure uses refluxing over anhydrous $AlCl_3$ (15g/L) for 1 hour, distilling, refluxing over alkaline $KMnO_4$ (10g $KMnO_4$, 10g Li_2CO_3 /L) for 15 minutes, and distilling. A further reflux for 1 hour over $KHSO_4$ (15g/L), then distillation, is followed by refluxing over CaH_2 (2g/L) for 1 hour, and fractional distillation. The product is protected from atmospheric moisture and stored under nitrogen [Walter & Ramalay *Anal Chem* **45** 165 1973, DOI: 10.1021/ac60323a041]. Purification of 'General Purity Reagent' for this purpose is not usually satisfactory because very large losses occur at the $KMnO_4/LiCO_3$ step. For electrochemical work involving high oxidation fluorides, further reflux over P_2O_5 (1g/ml for 0.5 hours) and distilling (discarding 3% of first and last fractions) and repeating this step is necessary. The distillate is kept over molecular sieves *in vacuo* after degassing, for 24 hours and distilling in a vacuum onto freshly activated 3A molecular sieves. The MeCN should have absorption at 200nm of <0.05 (H_2O reference) and UV cutoff at *ca* 175nm. Also the working potential range of purified $Et_4N^+ BF_4^-$ (0.1mol.dcm $^{-3}$ in the MeCN) should be +3.0 to -2.7V vs Ag^+/Ag^0 . If these criteria are not realised then further impurities can be removed by treatment with activated neutral alumina (60 mesh) *in vacuo* before final molecular sieves treatment [Winfield *J Fluorine Chem* **25** 91 1984, DOI: 10.1016/S0022-1139(00)81199-9].

Acetonitrile has been distilled from $AgNO_3$, collecting the middle fraction over freshly activated Al_2O_3 . After standing for two days, the liquid is distilled from the activated Al_2O_3 . The specific conductivity should be $0.8-1.0 \times 10^{-8}$ mhos [Harkness & Daggett *Can J Chem* **43** 1215 1965, DOI: 10.1139/v65-158]. Acetonitrile ^{14}C is best purified by gas chromatography, is water free, and distils at $81^\circ/atm$. [Beilstein **2** H 183, **2** IV 419.]

Rapid procedure: Dry over anhydrous K_2CO_3 for 24 hours, followed by further drying for 24 hours over 3A molecular sieves or boric anhydride, followed by distillation. Alternatively, stir over P_2O_5 (5% w/v) for 24 hours then distil. However, this last method is not suitable for reactions with very acid sensitive compounds.

Acetonylacetone (2,5-hexanedione) [110-13-4] $C_6H_{10}O_2$, M 114.2, m -9° , b $76-78^\circ/13mm$, $88^\circ/25mm$, $137^\circ/150mm$, $188^\circ/atm$, d_4^{20} 0.9440, n_D^{20} 1.423, pK^{25} 18.7. Purify it by dissolving in Et_2O , stirring with K_2CO_3 (a quarter of the weight of dione), filtering, drying over anhydrous Na_2SO_4 (not $CaCl_2$), filtering again, evaporating the filtrate and distilling it in a vacuum. Then redistil through a 30cm Vigreux column (oil bath temperature 150°). It is miscible with H_2O and EtOH. The *dioxime* has m 137° (plates from $*C_6H_6$), the *mono-oxime* has b $130^\circ/11mm$, and the *2,4-dinitrophenylhydrazone* has m $210-212^\circ$ (red needles from EtOH). It forms complexes with many metals. [Werner et al. *Chem Ber* **22** 2097 1989, DOI: 10.1002/cber.19891221109; for enol content see Gero *J Org Chem* **19** 1960 1954, DOI: 10.1021/jo01377a013; Beilstein **1** IV 3688.]

Acetoxyacetone (acetonyl acetate, acetol acetone) [592-20-1] $C_5H_8O_3$, M 116.1, b $65^\circ/11mm$, $73-75^\circ/17mm$, $174-176^\circ/atm$, d_4^{20} 1.0757, n_D^{20} 1.4141. Distil it in a vacuum, then redistil it at atmospheric press-

ure. It is miscible with H_2O , but is slowly decomposed by it. Store it in a dry atmosphere. The **2,4-dinitrophenylhydrazones** has **m 115-115.5°** (from CHCl_3 /hexane). [Perkin Jr *J Chem Soc* **59** 786 1891, DOI: 10.1039/CT8915900786; Reich & Samuels *J Org Chem* **21** 68 1956, DOI: 10.1021/jo01377a013; Nef *Justus Liebigs Ann Chem* **335** 247 1904, DOI: 10.1002/jlac.19043350302; *Beilstein* **2** IV 297.]

1-Acetoxy-1,3-butadiene (1,3-butadienyl acetate) cis-trans mixture [1515-76-0] $\text{C}_6\text{H}_8\text{O}_2$, M 112.1, b 42-43°/16mm, 51-52°/20mm, 60-61°/40mm, d_4^{20} 0.9466, n_D^{20} 1.4622. The commercial sample is stabilised with 0.1% of *p*-tert-butylcatechol. If the material contains crotonaldehyde (by IR, used in its synthesis), it should be dissolved in Et_2O , shaken with 40% aqueous sodium bisulfite, then 5% aqueous Na_2CO_3 , water, dried (Na_2SO_4) and distilled several times in a vacuum through a Widmer [*Helv Chim Acta* **7** 59 1924, DOI: 10.1002/hlca.19240070107] or Vigreux column [Wichterle & Hudlicky *Coll Czech Chem Commun* **12** 564 1947, DOI: 10.1135/cccc19470564; Hagemeyer & Hull *Ind Eng Chem* **41** 2920 1949, DOI: 10.1021/ie50480a063]. [*Beilstein* **2** III 295.]

1-Acetoxy-2-butoxyethane (2-butyloxyethyl acetate) [112-07-2] $\text{C}_8\text{H}_{16}\text{O}_3$, M 160.2, b 61-62°/0.2mm, 75-76°/12mm, 185.5°/740mm, 188-192°/atm, d_4^{20} 0.9425, n_D^{20} 1.4121. Shake the ester with anhydrous Na_2CO_3 , filter and distil it in a vacuum. Redistillation can then be carried out at atmospheric pressure. [Dunbar & Bolstad *J Org Chem* **21** 1041 1956, DOI: 10.1021/jo01115a613; *Beilstein* **2** IV 215.]

2-Acetoxyethanol (2-hydroxyethyl acetate) [542-59-6] $\text{C}_4\text{H}_8\text{O}_3$, M 104.1, b 61°/1mm, 79-81°/12mm, 187°/761mm, d_4^{20} 1.108, n_D^{20} 1.42. Dry the ester over K_2CO_3 (not CaCl_2), and distil it. [Davis & Ross *J Chem Soc* 3056 1950, DOI: 10.1039/JR9500003056; rate of hydrolysis: Davis & Ross *J Chem Soc* 2706 1951, DOI: 10.1039/JR9510002706; *Beilstein* **2** H 141, **2** I 66, **2** II 154, **2** III 303, **2** IV 214.]

1-Acetoxy-2-ethoxyethane [111-15-9] $\text{C}_6\text{H}_{12}\text{O}_3$, M 132.2, b 30°/3mm, 49-50°/12mm, 156-159°, d_4^{20} 0.97, n_D^{20} 1.406. Shake the ethoxy-ethane with anhydrous Na_2CO_3 , filter and distil it in a vacuum. Redistillation can then be carried out at atmospheric pressure. [Dunbar & Bolstad *J Org Chem* **21** 1041 1956, DOI: 10.1021/jo01115a613; *Beilstein* **2** IV 214.]

1-Acetoxy-2-methoxyethane [110-49-6] $\text{C}_5\text{H}_{10}\text{O}_3$, M 118.1, b 30°/6mm, 40-41°/12mm, 140-144°/760mm, d_4^{20} 1.009, n_D^{20} 1.4011. Shake the methoxy-ethane with anhydrous Na_2CO_3 , filter and distil it in a vacuum. Redistillation can be then be carried out at atmospheric pressure. [Dunbar & Bolstad *J Org Chem* **21** 1041 1956, DOI: 10.1021/jo015a61311; *Beilstein* **2** IV 214.]

S-(-)-2-Acetoxypropionyl chloride [36394-75-9] $\text{C}_5\text{H}_7\text{ClO}_3$, M 150.6, b 51-53°/11mm, d_4^{20} 1.19, n_D^{20} 1.423, $[\alpha]_D^{27}$ -33, (c 4, CHCl_3), $[\alpha]_{546}^{20}$ -38 (c 4, CHCl_3). It is moisture sensitive and is hydrolysed to the corresponding acid. Check the IR spectrum. If the OH band above 3000cm^{-1} is too large and broad then the mixture should be refluxed with pure acetyl chloride for 1 hour, evaporated and distilled under reduced pressure. [Doolittle & Heath *J Org Chem* **49** 5041 1984, DOI: 10.1021/jo01115a613; *Beilstein* **3** II 189.]

S-Acetoxy succinic anhydride [59025-03-5] $\text{C}_6\text{H}_6\text{O}_5$, M 158.1, m 58° (RS 81.5-82.5°, 86-87°), $[\alpha]_D^{20}$ -26.0 (c 19, Me_2CO), $[\alpha]_D^{20}$ -28.4 (c 13, Ac_2O). Recrystallise it from Ac_2O and dry it in a vacuum over KOH, or by washing it with dry Et_2O due to its deliquescent nature. [Jones *J Chem Soc* 788 1933, DOI: 10.1039/JR9330000788; Henrot et al. *Synth Commun* **16** 183 1986, DOI: 10.1080/00397918608057706; Shiuey et al. *J Org Chem* **53** 1040 1988, DOI: 10.1021/jo00240a021; RS: Cohen et al. *J Am Chem Soc* **88** 5306 1966, DOI: 10.1021/ja00974a051]

Acetylacetone (2,4-pentanedione, acac) [123-54-6] $\text{C}_5\text{H}_8\text{O}_3$, M 100.1, m -23°, 45°/30mm, b 140.4°/atm, $d^{30.2}$ 0.9630, $n^{18.5}_D$ 1.45178, pK_1^{25} -5.0 (enol), -6.6 (keto), pK_2^{25} 8.95. Small amounts of acetic acid are removed by shaking with small portions of 2N NaOH until the aqueous phase remains faintly alkaline. The sample, after washing with water, is dried with anhydrous Na_2SO_4 , and distilled through a modified Vigreux column [Cartledge *J Am Chem Soc* **73** 4416 1951, DOI: 10.1021/ja01153a112]. An additional purification step is fractional crystallisation from the melt. Alternatively, there is less loss of acetylacetone if it is dissolved in four volumes of *benzene and the solution is shaken three times with an equal volume of distilled water (to extr-

act acetic acid): the *benzene is then removed by distillation at 43-53°/20-30mm through a helices-packed column. It is then refluxed over P₂O₅ (10g/L) and fractionally distilled under reduced pressure. The distillate (sp conductivity $4 \times 10^{-8} \text{ ohm}^{-1}\text{cm}^{-1}$) is suitable for polarography [Fujinaga & Lee *Talanta* **24** 395 1977 DOI: 10.1016/0039-9140(77)80028-3]. To recover used acetylacetone, metal ions are stripped from the solution at pH 1 (using 100ml 0.1M H₂SO₄/L of acetylacetone). The acetylacetone is then washed with (1:10) ammonia solution (100ml/L) and with distilled water (100ml/L, twice), then treated as above. It complexes with Al, Be, Ca, Cd, Ce, Cu, Fe²⁺, Fe³⁺, Mn, Mg, Ni, Pb and Zn. [*Beilstein* **1** H 777, **1** I 401, **1** II 831, **1** III 3113, **1** IV 3662.]

Acetyl bromide [506-96-7] **C₂H₃BrO**, **M 123.0**, **b 76-77°/atm**, **d₄²⁰ 1.65**. Boil acetyl bromide with PBr₃/Ac₂O for 1 hour, then distil the latter off and redistil it. Store it dry and in the dark. [Burton & Degering *J Am Chem Soc* **62** 227 1940, DOI: 10.1021/ja01858a502; *Beilstein* **2** IV 398.] **LACHRYMATORY**.

2-Acetylbutyrolactone [517-23-7] **C₆H₈O₃**, **M 128.1**, **b 105°/5mm**, **120-123°/11mm**, **142-143°/30mm**, **d₄²⁰ 1.1846**, **n_D²⁰ 1.459**. Purify the lactone by distillation, which will convert any free acid to the lactone. Alternatively, dissolve it in Et₂O, wash well with 0.5N HCl, dry the organic layer and distil it. Its solubility in H₂O is 20% v/v. The **2,4-dinitrophenylhydrazone** forms orange needles from MeOH, **m 146°**. The lactone hydrolyses in mineral acid to 2-acetyl-4-hydroxybutyric acid which can be converted to the **di-n-propylamine salt** with **m 68-70°**. The lactone is a **SKIN IRRITANT**. [Willman & Schinz *Helv Chim Acta* **35** 2401 1952, DOI: 10.1002/hlca.19520350728; *Beilstein* **17/11** V 16.]

Acetyl chloride [75-36-5] **C₂H₃ClO**, **M 78.5**, **b 52°/atm**, **d₄²⁰ 1.1051**, **n_D²⁰ 1.38976**. Reflux acetyl chloride with PCl₅ for several hours to remove traces of acetic acid, then distil it. Redistil it from one-tenth its volume of dimethylaniline or quinoline to remove free HCl. A.R. quality is freed from HCl by pumping it for 1 hour at -78° and distilling it into a trap at -196°. [*Beilstein* **2** IV 395.] **LACHRYMATORY**.

Acetylene [74-86-2] **C₂H₂**, **M 26.0**, **m -80.8°**, **b -84°/atm**, **pK²⁵ ~25**. If very impure, acetylene should be purified by successive passage through spiral wash bottles containing, in this order, saturated aqueous NaHSO₄, H₂O, 0.2M iodine in aqueous KI (two bottles), sodium thiosulfate solution (two bottles), alkaline sodium hydrosulfite with sodium anthraquinone-2-sulfonate as indicator (two bottles), and 10% aqueous KOH solution (two bottles). The gas is then passed through a Dry-Ice trap and two drying tubes, the first containing CaCl₂, and the second, Dehydrite [Mg(ClO₄)₂] [Conn et al. *J Am Chem Soc* **61** 1868 1939, DOI: 10.1021/ja01876a066]. Acetone vapour can be removed from acetylene by passage through H₂O, then concentrated H₂SO₄, or by passage through two gas traps at -65° and -80°, concentrated H₂SO₄ and a soda lime tower, a tower of 1-mesh Al₂O₃ then through H₂SO₄ [Reichert & Nieuwland *Org Synth Coll Vol I* 229 1941, DOI: 10.15227/orgsyn.004.0023; Wiley *Org Synth Coll Vol 3* 853 1955, DOI: 10.15227/orgsyn.028.0094; Skattebol et al. *Org Synth Coll Vol 4* 792 1963 DOI: 10.15227/orgsyn.039.0056]. Sometimes it contains acetone and air. These can be removed by a series of bulb-to-bulb distillations, e.g. a train consisting of a concentrated H₂SO₄ trap and a cold EtOH trap (-73°), or passage through H₂O and H₂SO₄, then over KOH and CaCl₂. [See Brandsma *Preparative Acetylenic Chemistry*, 1st Edn Elsevier 1971 p15, for pK, ISBN 0444409475, 2nd Edn Elsevier 1988, ISBN 0444429603, and below for sodium acetylide.] It is also available commercially as 10ppm in helium, and several concentrations in N₂ for instrument calibration. [*Beilstein* **1** IV 939.]

Sodium acetylide [1066-26-8] **M 48.0**, is prepared by dissolving Na (23g) in liquid NH₃ (1L) and bubbling acetylene until the blue color is discharged (*ca* 30 minutes) and evaporated to dryness [Saunders *Org Synth Coll Vol 3* 416 1955, DOI: 10.15227/orgsyn.029.0047], and is available commercially as a suspension in xylene/light mineral oil. [See entry in 'Metal-organic Compounds', Chapter 4.]

Acetylenedicarboxamide (Aquamycin, Cellocidin) [543-21-5] **C₄H₄N₂O₂**, **M 112.1**, **m 216-218°(dec)**, **219-221°(dec)**. Acetylenedicarboxamide crystallises from MeOH and H₂O [**m 190-192°(dec)** as *hemihydrate*]. When prepared from the ester + NH₃ it has **m 213°(dec)**. Also a melting point of **290-292°(dec)** has been reported. [Saggimo *J Org Chem* **22** 1171 1957, DOI: 10.1021/jo01361a009; Kharash et al. *J Org Chem* **10** 386 1945, DOI: 10.1021/jo01181a002; Blomquist & Winslow *J Org Chem* **10** 149 1945, DOI: 10.1021/jo01178a010; *Beilstein* **2** I 317, **2** III 1995, **2** IV 2295.]

Acetylenedicarboxylic acid (butynedioic acid) [142-45-0] $C_4H_2O_4$, **M 114.1**, **m 179°(anhydrous)**, **pK₁¹⁹ 1.04**, **pK₂¹⁹ 2.50**. The acid is soluble in Et₂O and crystallises from Et₂O/petroleum ether (**m 183-183.5°**), or H₂O as the *dihydrate* which dehydrates in a desiccator over concentrated H₂SO₄ in a vacuum. The *dipicrate* crystallises from aqueous ether. For the mono K salt see entry in 'Metal-organic Compounds', Chapter 4. [Abbott et al. *Org Synth Coll Vol* **2** 10 1943, DOI: 10.15227/orgsyn.018.0003; Huntress et al. *Org Synth Coll Vol* **4** 329 1963, DOI: 10.15227/orgsyn.032.0055; *Beilstein* **2** H 801, **2** I 317, **2** II 670, **2** III 1991, **2** IV 2290.]

N-Acetylenediamine [1001-53-2] $C_4H_{10}N_2O$, **M 102.1**, **m 50-51°**, **51°**, **b 128°/3mm**, **125-130°/5mm**, **133-139°/27mm**, **pK₂₅ 9.28**. The acetyl-diamine has been fractionated under reduced pressure and fraction **b 125-130°/5mm** was re-fractionated, fraction **b 132-135°/4mm** was collected and it solidified. It is a low melting *hygroscopic* solid which can be recrystallised from dioxane/Et₂O. It is soluble in H₂O, Et₂O and *C₆H₆. The *p-toluenesulfonate salt* can be recrystallised from EtOH/EtOAc (1:8), has **m 125-126°**, but the free base cannot be recovered from it by basifying and extracting with CH₂Cl₂. The *picrate* has **m 175°** (from EtOH) [Aspinall *J Am Chem Soc* **63** 852 1941, DOI: 10.1021/ja01848a060; Hall *J Am Chem Soc* **78** 2570 1956, DOI: 10.1021/ja01592a066]. [*Beilstein* **4** IV 1193.]

Acetyl fluoride [557-99-3] C_2H_3FO , **M 62.0**, **b 20.5°/760mm**, **d₄²⁰ 1.032**. Purify acetyl fluoride by fractional distillation. It attacks glass and is sold in steel cylinders. [*Beilstein* **2** H 172, **2** I 79, **2** II 175, **2** III 385, **2** IV 393.] **TOXIC and LACHRYMATORY.**

Acetyl iodide [507-02-8] C_2H_3IO , **M 170.0**, **b 39.5-40°/64mm**, **108°/760mm**. Purify it by fractional distillation. Store it in the dark. [*Beilstein* **2** II 177, **2** III 393, **2** IV 399.] **TOXIC and LACHRYMATORY.**

3-(S-Acetylmercapto)isobutyric acid [RS 33325-40-5] $C_6H_{10}O_3S$, **M 162.2**, **m 40-40.5°**, **b ca 120°/1.25mm**, **pK_{Est} ~4.0**. Distil the acid under high vacuum and recrystallise it from *C₆H₆. [Fredga & Mastersson *Chem Abstr* **38** 3616 1944.]

Acetyl methanesulfonate [5539-53-7] $C_3H_6O_4S$, **M 138.1**, **b <120°/0.01mm**. The main impurity is methanesulfonic acid. Reflux it with redistilled acetyl chloride for 6-10 hours, i.e. until no further HCl is absorbed in a trap, and exclude moisture. Distil off excess of AcCl and carefully distil it below 0.001mm with the bath temperature below 120° to give the anhydride as a pale yellow oil which solidifies below 0°. Below ~130° it gives the disulfonic anhydride, and above ~130° polymers are formed. It is used for cleaving ethers [Preparation, IR, NMR: Mazur & Karger *J Org Chem* **36** 528, 532 1971, DOI: 10.1021/jo00803a009; DOI: 10.1021/jo00803a010]. [*Beilstein* **2** H 166, **2** III 349.]

3-(Acetylthio)propionic acid [41345-70-4] $C_5H_8O_3S$, **M 148.2**, **m 48-52°**, **52-54°**, **b 127-128°/3mm**, **pK_{Est} ~4.2**. Purify the propionic acid by distillation in a vacuum. It has λ_{max} at 231nm (ϵ 4200). It is a potential enzyme inhibitor [Noda et al. *J Am Chem Soc* **75** 913 1953, DOI: 10.1021/ja01100a041; Clegg et al. *J Am Chem Soc* **121** 5319 1999, DOI: 10.1021/ja9901011; Larsson et al. *Angew Chem Int Ed* **43** 3716 2004, DOI: 10.1002/anie.200454165]. [*Beilstein* **3** III 551, **3** IV 731.]

N-Acetylthiourea [591-08-2] $C_3H_6ON_2S$ **M 118.2**, **m 164-165°**, **166-168°**. Recrystallise the thiourea from AcOH; the solid is washed with Et₂O and dried in air, then at 100°. [Zahradnik *Coll Czech Chem Commun* **24** 3678 1959, DOI:10.1135/cccc19593678; *Beilstein* **3** IV 354.]

cis-Aconitic acid (1,2,3-propenetriscarboxylic acid) [585-84-2] $C_6H_6O_6$, **M 174.1**, **m 126-129°(dec)**. Crystallise the *cis*-acid from water by cooling (solubility is 1g in 2ml of water at 25°). Dry it in a vacuum desiccator. [*Beilstein* **2** IV 2405.] **trans-Aconitic acid** [4023-65-8] $C_6H_6O_6$, **M 174.1**, **m 195°(dec)**, **m 198-199°(dec)**, **204-205°(dec)**, **pK₁²⁵ 2.81**, **pK₂²⁵ 4.46**, is purified dissolving it in AcOH (77g/150ml), filtering and cooling. The acid separates (55g) as colourless needles. A further quantity (10g) can be obtained by reducing the volume of the filtrate. The acid is dried in air then in a vacuum desiccator over NaOH. The acid can be recrystallised from Me₂CO/CHCl₃. The highest melting point is obtained with the very dry acid, and **m 209°** was obtained on a Dennis bar [Dennis & Shalton *J Am Chem Soc* **52** 3128 1930, DOI: 10.1021/ja01371a013; Bruce *Org Synth Coll Vol* **2** 12 1943, DOI: 10.15227/orgsyn.017.0001]. [*Beilstein* **2** IV 2405.]

cis-Aconitic anhydride [6318-55-4] $C_6H_4O_5$, M 156.1, m 75°, 76-78°, 78-78.5°. Reflux it in xylene (7.5 parts) for 1 hour, then evaporate and recrystallise the residue from $*C_6H_6$. Alternatively, reflux it in Ac_2O , evaporate and recrystallise from $*C_6H_6$. It is sensitive to moisture, store dry. [IR: Groth & Dahlén *Acta Chem Scand* **21** 291 1967, DOI: 10.3891/acta.chem.scand.21-0291; Malachowski & Maslowski *Chem Ber* **61** 2521 1928, DOI: 10.1002/cber.19280611119; NMR: Gawron & Mahajan *Biochemistry* **5** 2335 1966, DOI: 10.1021/bi00871a023.] [*Beilstein* **18/8** V 530.]

Acrolein (acraldehyde, 2-propenal) [107-02-8] C_3H_4O , M 56.1, fp -86.95°, b 25°/100mm, 52.69°/760mm (dt/dp 0.0355°/mm, d_4^{20} 0.839, n_D^{25} 1.3992. Purify acrolein by fractional distillation, under nitrogen, drying with anhydrous $CaSO_4$ and then re-distilling under vacuum. Blacet, Young and Roof [*J Am Chem Soc* **59** 608 1937, DOI: 10.1021/ja01283a004] distilled it under nitrogen through a 90cm column packed with glass rings. To avoid formation of diacryl, the vapour is passed through an ice-cooled condenser into a receiver cooled in an ice-salt mixture and containing 0.5g catechol. The acrolein is then distilled twice from anhydrous $CuSO_4$ at low pressure, catechol being placed in the distilling flask and the receiver to avoid polymerisation. [Alternatively, hydroquinone (1% of the final solution) can be used.] **Respiratory irritant, work in an efficient fume cupboard.** [*Beilstein* **1** IV 3435.] **Acrolein semicarbazone** [6055-71-6] $C_4H_8N_3O$, M 114.1, m 171°, crystallises from water in needles. [Auwers & Heineke *Justus Liebigs Ann Chem* **458** 186 1927, DOI: 10.1002/jlac.1927458011; *Beilstein* **1** II 785.]

Acrolein diacetyl acetal (1,1-diacetoxy-2-propene). [869-29-4] $C_7H_{10}O_4$, M 158.2, b 75°/10mm, 184°/atm, d_4^{20} 1.08, n_D^{20} 1.4203. Check the NMR spectrum. If it is not satisfactory, then add Ac_2O and a drop of concentrated H_2SO_4 and heat at 50° for 10 minutes. Then add anhydrous $NaOAc$ (ca 3g/100g of liquid) and fractionate. Note that it forms an azeotrope with H_2O , so do not add H_2O at any time. It is a **highly flammable and TOXIC** liquid; use protective gloves. [Smith et al. *J Am Chem Soc* **73** 5282 1951, DOI: 10.1021/ja01155a080; *Beilstein* **2** H 154, **2** I 72, **2** III 356, **2** IV 291.]

Acrolein diethyl acetal (3,3-diethoxy-1-propene or 1,1-diethoxy-2-propene) [3054-95-3] $C_7H_{14}O_2$, M 130.2, b 120-125°/atm, n_D^{20} 1.398-1.407. Add Na_2CO_3 (ca 3.5%) and distil it using an efficient column, or better use a spinning band column. [Witzemann et al. *Org Synth Coll Vol* **2** 17 1943, DOI: 10.15227/orgsyn.011.0001, *Beilstein* **1** H 727, **1** I 378, **1** III 2960, **1** IV 3437.]

Acrolein dimethyl acetal (1,1-dimethoxy-2-propene) [6044-68-4] $C_5H_{10}O_2$, M 102.1, b 87.5-88°/750mm, 89-90°/760mm, d_4^{20} 0.86, n_D^{20} 1.3962. Fractionally distil it (after adding 0.5g of hydroquinone) under reduced pressure through an all glass column (40cm x 2.5 cm) packed with glass helices and provided with a heated jacket and a total reflux variable take-off head. Stainless steel Lessing rings (1/8 x 1/8 in) or gauze have also been used as packing. It is a **highly flammable and TOXIC** liquid; keep away from the skin. [Hall & Stern *J Chem Soc* 2657 1955, DOI: 10.1039/JR9550002657; *Beilstein* **1** IV 3437.]

Acrylamide [79-06-1] C_3H_5NO , M 71.1, m 84°, b 125°/25mm. Crystallise acrylamide from acetone, chloroform, ethyl acetate, methanol or $*benzene$ /chloroform mixture, then vacuum dry, and store it in the dark under a vacuum. Recrystallise it from $CHCl_3$ by dissolving 200g in 1L, heating to boiling and filtering without suction in a warmed funnel through Whatman 541 filter paper; allowing to cool to room temperature and keeping at -15° overnight. The crystals are collected with suction in a cooled funnel and washed with 300ml of cold MeOH. The crystals are air-dried in a warm oven. [Dawson et al. *Data for Biochemical Research*, Oxford Press 1986 p. 449, *Beilstein* **2** IV 1471.]

CAUTION: Acrylamide is extremely **TOXIC (neurotoxic)**, and precautions must be taken to avoid skin contact or inhalation. Use gloves and handle in a well-ventilated fume cupboard.

Acrylic acid [79-10-7] $C_3H_4O_2$, M 72.1, m 13°, b 30°/3mm, 70°/50mm, d_4^{20} 1.051, pK^{25} 4.25. It can be purified by steam distillation, or vacuum distillation through a column packed with copper gauze to inhibit polymerisation. (This treatment also removes inhibitors, such as methylene blue, that may be present.) Azeotropic distillation of the water with $*benzene$ converts aqueous acrylic acid to the anhydrous material. [*Beilstein* **2** H 397, **2** I 186, **2** II 383, **2** III 1215, **2** IV 1455.]

Acrylonitrile [107-13-1] $\text{C}_3\text{H}_3\text{N}$, **M 53.1**, **b 78°/atm**, **d_4^{20} 0.806**, **n_D^{25} 1.3886**. Wash acrylonitrile with dilute H_2SO_4 or dilute H_3PO_4 , then with dilute Na_2CO_3 and water. Dry it with Na_2SO_4 , CaCl_2 or (better) by shaking with molecular sieves. Fractionally distil it under N_2 . It can be stabilised by adding 10ppm *tert*-butyl catechol. Immediately before use, the stabiliser can be removed by passage through a column of activated alumina (or by washing with 1% NaOH solution if traces of water are permissible in the final material), followed by distillation. Alternatively, shake it with 10% (w/v) NaOH to extract inhibitor, and then wash it in turn with 10% H_2SO_4 , 20% Na_2CO_3 and distilled water. Dry for 24 hours over CaCl_2 and fractionally distil under N_2 collecting fraction boiling at 75.0-75.5°/734mm). Store it with 10ppm *tert*-butyl catechol. Acrylonitrile is distilled off when required. [Burton et al. *JCS Faraday Trans 1* **75** 1050 1979, DOI: 10.1039/F19797501050; *Beilstein 2* IV 1473.]

Acryloyl chloride (acrylyl chloride) [814-68-6] $\text{C}_3\text{H}_3\text{ClO}$, **M 90.5**, **b 72-74°/740mm**, **74°/760mm**, **d_4^{20} 1.1127**, **n_D^{20} 1.4337**. Distil acryloyl chloride rapidly through an efficient 25cm column after adding 0.5g of hydroquinone/200g of chloride, and then re-distil it carefully at atmospheric pressure preferably in a stream of dry N_2 . [Stempel et al. *J Am Chem Soc* **72** 2299 1950, DOI: 10.1021/ja01161a527; *Beilstein 2* IV 1471.] **The liquid is an irritant and is TOXIC.**

Adipic acid [124-04-9] $\text{C}_6\text{H}_{10}\text{O}_4$, **M 146.1**, **m 154-154.5°**, **b 159.5°/0.1mm**, **191°/5mm**, **205.5°/10mm**, **222.5°/20mm**, **337.5°/760mm**, **pK_1^{20} 4.42**, **pK_2^{20} 5.41**; **pK_1^{25} 4.44**, **pK_2^{25} 5.45**; **pK_1^{40} 4.54**, **pK_2^{40} 5.59**. For use as a volumetric standard, adipic acid is crystallised once from hot water with the addition of a little animal charcoal, dried at 120° for 2 hours, then recrystallised from acetone and again dried at 120° for 2 hours. Other purification procedures include crystallisation from ethyl acetate and from acetone/petroleum ether, fusion, followed by filtration and crystallisation from the melt, and preliminary distillation under vacuum. [*Beilstein 2* H 649, **2 I** 277, **2 II** 572, **2 III** 1705, **2 IV** 1956.] **Diethyl adipate** [141-28-6] **M 202.3**, **m -20° to -19°**, **b 111°/5mm**, **125°/10mm**, **133°/15mm**, **251°/atm**, **d_4^{25} 1.0034**, **n_D^{20} 1.42776**, is prepared in the usual way by boiling the acid and excess EtOH in the presence of a catalytic amount of H_2SO_4 and distilled. [*Beilstein 2* H 652, **2 I** 277, **2 II** 574, **2 III** 1721, **2 IV** 1960.]

Adiponitrile (1,4-dicyanobutane) [111-69-3] $\text{C}_6\text{H}_8\text{N}_2$, **M 108.14**, **m 2.4°**, **b 123°/0.5mm**, **153°/6mm**, **175°/26mm**, **184°/30mm**, **295°/atm**, **d_4^{20} 0.9396**, **n_D^{20} 1.4371**. Reflux adiponitrile over P_2O_5 and POCl_3 , and fractionally distil it, then fractionate it through an efficient column. **The liquid is TOXIC and is an IRRITANT.** [Braun & Rudolph *Chem Ber* **67** 1762 1934, DOI: 10.1002/cber.19340671020; Reppe et al. *Justus Liebigs Ann Chem* **596** 127 1955, DOI: 10.1002/jlac.19555960108; Gagnon et al. *Can J Chem* **34** 1662 1956, DOI: 10.1139/v56-214; Copley et al. *J Am Chem Soc* **62** 227 1940, DOI: 10.1021/ja01858a503; *Beilstein 2* IV 1947.]

Agaricic acid [1-(*n*-hexadecyl)citric acid] [666-99-9] $\text{C}_{22}\text{H}_{40}\text{O}_7$, **M 416.6**, **m 142°(dec)**, **$[\alpha]_D$ -9.8 (in NaOH)**, **$\text{pK}_{\text{Est}(1)}$ ~2.7**, **$\text{pK}_{\text{Est}(2)}$ ~4.2**, **$\text{pK}_{\text{Est}(3)}$ ~5.5**. Crystallise the acid from EtOH . The *trihydrazide* has **m 170°(dec)** (from EtOH). [Brandänge et al. *Acta Chem Scand B* **31** 307 1977, DOI: 10.3891/acta.chem.scand.31b-0307; *Beilstein 3* I 186, **3 II** 372, **3 III** 1109, **3 IV** 1284.]

Agmatine sulfate [5-guanidinopent-1-ylamine sulfate] [2482-00-0] $\text{C}_5\text{H}_{16}\text{N}_4\text{O}_4\text{S}$, **M 228.3**, **m 231°**, **$\text{pK}_{\text{Est}(1)}$ ~9.1**, **$\text{pK}_{\text{Est}(2)}$ ~13.0**. Crystallise the salt from aqueous MeOH . The *free base* has [306-60-5] $\text{C}_5\text{H}_{14}\text{N}_4$, **M 130.2**, **m 101.5-103°**, the *gold chloride hydrochloride* crystallises from H_2O with **m 223°(dec)**, and the *picrate* has **m 236-238°**. [Kossel *Z Physiol Chem* **66** 257 1910, DOI: 10.1515/bchm2.1910.66.3.257; Raasch et al. for 'Biological significance of agmatine, an endogenous ligand at imidazoline binding sites' see: *Br J Pharmacol* **133** 755 2001, DOI: 10.1038/sj.bjp.0704153; *Beilstein 4* I 420, **4 II** 703, **4 III** 575, **4 IV** 1291.]

Aldol (3-hydroxybutanal) [107-89-1] $\text{C}_4\text{H}_8\text{O}_2$, **M 88.1**, **b 80-81°/20mm**, **d^{16} 1.109**. An ethereal solution of aldol is washed with a saturated aqueous solution of NaHCO_3 , then with water. The non-aqueous layer is dried with anhydrous CaCl_2 and distilled immediately before use. The fraction, **b 80-81°/20mm**, is collected as a thick liquid which decomposes at 85°/atm. It is a sedative and a hypnotic, but is used in perfumery. [Mason et al. *J Am Chem Soc* **76** 2255 1954, DOI: 10.1021/ja01637a069]. [*Beilstein 1* H 824, **1 I** 419, **1 II** 868, **1 III** 3195, **1 IV** 3984.]

Aleuritic acid [*RS-erythro-9,10,16-trihydroxyhexadecanoic acid*] [533-87-9] $C_{16}H_{32}O_5$, **M 304.4**, **m 100-101°**, **pK_{Est} ~4.9**. Crystallise this *RS*-acid from aqueous EtOH. It is soluble in MeOH, and forms a less soluble crystalline sodium salt. The *methyl ester* **m 72-73°**, **b 235°/0.2mm**, is best prepared by reaction with diazomethane and forms fine feathery needles; it is soluble in MeOH, EtOH, $CHCl_3$, Me_2CO , slightly soluble in $*C_6H_6$ and insoluble in petroleum ether. The *ethyl ester* [6003-09-4] **m 59°**, crystallises in needles from EtOH. The *hydrazide* [6003-10-7] crystallises from EtOH and has **m 139-140°**. The *RS-acid* has been isolated from Shellac although it has two asymmetric carbon atoms, and possibly contains the *RS-erythro* or *RS-cis form* [Gidvani *J Chem Soc* 305 1944, DOI: 10.1039/JR9440000305]. Stereoisomers have been synthesised [Hunsdiecker *Chem Ber* 76 142 1943, DOI: 10.1002/cber.19430760120; Hunsdiecker *Chem Ber* 77 185 1944, DOI: 10.1002/cber.19440770308]. [Beilstein 3 III 901.]

***n*-Alkylammonium chloride** **n=2,4,6**. Recrystallise them from EtOH or an EtOH/Et₂O mixture. [Hashimoto & Thomas *J Am Chem Soc* 107 4655 1985, DOI: 10.1021/ja00302a010; Chu & Thomas *J Am Chem Soc* 108 6270 1986, DOI: 10.1021/ja00280a026.]

***n*-Alkyltrimethylammonium bromide** **n=10,12,16**. Recrystallise them from an EtOH/Et₂O mixture. [Hashimoto & Thomas *J Am Chem Soc* 107 4655 1985, DOI: 10.1021/ja00302a010.]

Allene (propadiene) [463-49-0] C_3H_4 , **M 40.1**, **m -146°**, **b -32°/atm**. Freeze allene in liquid nitrogen, evacuate, then thaw out. This cycle is repeated several times, then the allene is frozen in a methylcyclohexane/liquid nitrogen bath and pumped for some time. It has also been purified by HPLC. [Cripps & Kiefer *Org Synth* 42 12 1962, DOI: 10.15227/orgsyn.042.0012; Beilstein 1 IV 966.]

neo-Allocimene (allocimene B, *tc*-2,6-dimethyl-2,4,6-octatriene) [7216-56-0; *cis/trans mixture* 673-84-7; *trans/trans* 3016-19-1] $C_{10}H_{16}$, **M 136.2**, **b 80°/13mm**, **196-198°/atm**, **d₄²⁰ 0.8161**, **n_D²⁰ 1.5437**. Fractionally distil allocimene through an efficient column and repeatedly distil it at 15mm through a long column of glass helices, with a final distillation from sodium under nitrogen. It should be stabilised with *ca* 0.1% of hydroquinone. Its UV has λ_{max} nm(ϵ M⁻¹cm⁻¹) at 290 (32 500), 279 (41 900) and 278 (42,870). [Alder et al. *Justus Liebigs Ann Chem* 609 1 1957, DOI: 10.1002/jlac.19576090102; O'Connor & Goldblatt *Anal Chem* 26 1726 1954, DOI: 10.1021/ac60095a014; Beilstein 1 IV 1106.]

Allyl acetate [591-87-7] $C_5H_8O_2$, **M 100.1**, **b 103°/atm**, **d₄²⁰ 0.928**, **n_D¹⁷ 1.4004**, **n_D²⁰ 1.4040**. The ester is freed from peroxides by standing with crystalline ferrous ammonium sulfate, then washed with 5% $NaHCO_3$, followed by saturated $CaCl_2$ solution. Dry it with Na_2SO_4 and fractionally distil it in an all-glass apparatus. **FLAMMABLE LIQUID**. [Beilstein 2 H 136, 2 IV 180.]

Allylacetic acid (pent-4-enoic acid) [591-80-0] $C_5H_8O_2$, **M 100.1**, **m -22.5°**, **b 83-84°/12mm**, **90°/15mm**, **187-189°/~760mm**, **d₄²⁰ 0.9877**, **n_D²⁰ 1.4280**, **pK₂₅ 4.68**. Distil the acid through an efficient column (allyl alcohol has **b 95-97°**). It is characterised as the ***S*-benzylisothiuronium salt** **m 155-158°** (from 96% EtOH, or aqueous EtOH) [Friediger & Pedersen *Acta Chem Scand* 9 1425 1955, DOI: 10.3891/acta.chem.scand.09-1425], and the ***4*-bromophenacyl ester** has **m 59.5-60.5°** (from 90% EtOH). Its solubility at 18° in solvents is: pyridine (57%), AcOH (7.3%), MeOH (5.4%), Me_2CO (3.2%), MeOAc (2.8%), EtOH (5.4%), H_2O (1.8%), PrOH (1.6%), isoPrOH (0.27%). [Brown & Berkowski *J Am Chem Soc* 74 1894 1952, DOI: 10.1021/ja01128a007; Beilstein 2 IV 1542.]

Allyl alcohol [107-18-6] C_3H_6O , **M 58.1**, **b 98°/atm**, **d₄²⁰ 0.857**, **n_D²⁰ 1.4134**. It can be dried with K_2CO_3 or $CaSO_4$, or by azeotropic distillation with $*benzene$ followed by distillation under nitrogen. It is difficult to obtain it free of peroxide. It has also been refluxed with magnesium and fractionally distilled [Kamm & Marvel *Org Synth Coll Vol* 1 42 1941, DOI: 10.15227/orgsyn.001.0015]. [Beilstein 1 IV 2079.]

Allylamine [107-11-9] C_3H_7N , **M 57.1**, **b 52.9°/atm**, **d₄²⁰ 0.761**, **n_D²⁰ 1.42051**, **pK₂₅ 9.49**. Purify allylamine by fractional distillation from calcium chloride. It is a strong base and should be stored under N_2 . It causes sneezing and tears. [Beilstein 4 IV 1057.]

Allyl bromide [106-95-6] $\text{C}_3\text{H}_5\text{Br}$, M 121, b 70°/atm, d_4^{20} 1.398, n_D^{20} 1.46924. Wash the bromide with NaHCO_3 solution then distilled water, dry (CaCl_2 or MgSO_4), and fractionally distil. Protect it from strong light. [Beilstein 1 IV 754.] **LACHRYMATORY, HIGHLY TOXIC and FLAMMABLE.**

Allyl butyl ether [3739-64-8] $\text{C}_7\text{H}_{14}\text{O}$, M 114.2, b 64-65°/120mm, 117.8-118°/763mm, d_4^{20} 1.4057, n_D^{20} 0.7829. Check the IR for the presence of OH str vibrations near $\sim 3000\text{cm}^{-1}$; if so then wash it well with H_2O , dry it with CaCl_2 and distil it through a good fractionating column. **The liquid is an irritant.** [Watanabe et al. *J Org Chem* 23 1666 1958, DOI: 10.1021/jo01105a021; Schueler & Hanna *J Am Chem Soc* 73 3528 1951, DOI: 10.1021/ja01151a532; Beilstein 1 IV 2084.]

Allyl chloride [107-05-1] $\text{C}_3\text{H}_5\text{Cl}$, M 76.5, b 45.1°/atm, d_4^{20} 0.939, n_D^{20} 1.4130. Likely impurities include 2-chloropropene, propyl chloride, *iso*-propyl chloride, 3,3-dichloropropane, 1,2-dichloropropane and 1,3-dichloropropane. Purify it by washing with concentrated HCl , then with Na_2CO_3 solution, dry it with CaCl_2 , and distil it through an efficient column [Oae & Vanderwerf *J Am Chem Soc* 75 2724 1953, DOI: 10.1021/ja01107a052]. [Beilstein 1 IV 738.] **LACHRYMATORY, TOXIC.**

Allyl chloroformate [2937-50-0] $\text{C}_4\text{H}_5\text{ClO}_2$, M 120.5, b 56°/97mm, 109-110°/atm, d_4^{20} 1.14, n_D^{20} 1.4223. Wash the chloroformate several times with cold H_2O to remove alcohol and HCl and dry it over CaCl_2 . It is **important** to dry well before distilling *in vacuo*. Note that the receiver should be cooled in ice to avoid loss of distillate into the trap and vacuum pump. The liquid is **highly TOXIC and flammable**. [Fierz-David & Müller *J Chem Soc* 125 26 1924, DOI: 10.1039/JR9330000788; Strain et al. *J Am Chem Soc* 72 1254 1950, DOI: 10.1021/ja01159a052; Beilstein 3 IV 29.]

Allyl cyanide (3-butene nitrile) [109-75-1] $\text{C}_3\text{H}_5\text{N}$, M 67.1, b -19.6°/1.0mm, 2.9°/5mm, 14.1°/5mm, 26.6°/20mm, 48.8°/60mm, 60.2°/100mm, 98°/400mm, 119°/760mm, d_4^{20} 0.8341, n_D^{20} 1.406. The nitrile should be redistilled at atmospheric pressure, then distilled under a vacuum to remove the final traces of HCN from the residue. Note that the residue from the first distillation may be difficult to remove from the flask and should be treated with concentrated HNO_3 then H_2O and finally hot EtOH (**CARE**). Allyl cyanide has an onion-like odour and is stable to heat. It forms a complex with AlCl_3 (2:2) **m** 41°, and (3:2) **m** 120°. **All operations should be done in an efficient fume hood as the liquid is flammable, may contain cyanide and is HIGHLY TOXIC.** [Supniewski et al. *Org Synth Coll Vol I* 46 1941, DOI: 10.15227/orgsyn.008.0004; Beilstein 2 IV 1491.]

Allyl disulfide (diallyl disulfide) [2179-57-9] $\text{C}_6\text{H}_{10}\text{S}_2$, M 146.3, b 58-59°/5mm, 79-81°/20mm, 138-139°/atm, d_4^{20} 1.01, n_D^{20} 1.541. Purify the disulfide by fractional distillation until the molar refractivity is in uniformly good agreement with the calculated value [Small et al. *J Am Chem Soc* 69 1710 1947, DOI: 10.1021/ja01199a040]. It has also been purified by gas chromatography [retention times: Carson & Wong *J Org Chem* 24 175 1959, DOI: 10.1021/jo01084a007; UV: Koch *J Chem Soc* 394 1949, DOI: 10.1039/JR9490000394]. It is present in garlic. [Beilstein 1 IV 2098.]

Allyl iodide (3-iodopropene) [556-56-9] $\text{C}_3\text{H}_5\text{I}$, M 167.98, b 103°/760mm, d^{12} 1.848. Purify allyl iodide in a dark room by washing with aqueous Na_2SO_3 to remove free iodine, then dry with MgSO_4 and distil at 43°/90 mm or at atmospheric pressure to give a very pale yellow liquid. (This material, dissolved in hexane, can be stored in a light-protected tight container at -5° for up to three months before free iodine can be detected, by its colour in the solution.) Store it away from light. [Sibbett & Noyes *J Am Chem Soc* 75 761 1953, DOI: 10.1021/ja01100a001; Beilstein 1 H 202, 1 I 84, 1 II 172, 1 III 714, 1 IV 761.]

Allylisocyanate [1476-23-9] $\text{C}_4\text{H}_5\text{NO}$, M 83.1, b 84°/atm, 87-89°/atm, d_4^{20} 0.94, n_D^{20} 1.417. Purify it as for allylisothiocyanate below and it is **TOXIC**. [Beilstein 4 IV 1081.]

Allylisothiocyanate [57-06-7] $\text{C}_4\text{H}_5\text{NS}$, M 99.2, m -80°, b 84-85°/80mm, 150°/760mm, 151°/atm, d_4^{20} 1.017, n_D^{20} 1.5268. Fractionate the isothiocyanate using an efficient column, preferably in a vacuum. It is a yellow **pungent, irritating and a TOXIC (suspected CARCINOGEN) liquid**. Store it in a sealed tube under N_2 . The *N*-benzylthiourea derivative has **m** 94.5° (from aqueous EtOH) [Weller et al. *J Am Chem Soc* 74 1104 1952,

DOI: 10.1021/ja01124a531]. [*Beilstein* 4 IV 1081.]

N-Allylthiourea (thiosinamine) [109-57-9] $\text{C}_4\text{H}_8\text{N}_2\text{S}$, **M 116.2, m 70-73°, 78°**. Recrystallise it from H_2O . It is soluble in 30 parts of cold H_2O , and is soluble in EtOH but insoluble in C_6H_6 . It has also been recrystallised from acetone, EtOH or ethyl acetate, after decolorising with charcoal. The white crystals have a bitter taste with a slight garlic odour and are **TOXIC**. An unstable crystalline form is obtained by recrystallising from the melt. [McCrone et al. *Anal Chem* 21 421 1949, DOI: 10.1021/ac60027a600; *Beilstein* 4 IV 1072.]

N-Allylurea [557-11-9] $\text{C}_4\text{H}_8\text{N}_2\text{O}$, **M 100.1, m 85°**. It can be recrystallised from EtOH, EtOH/ether, EtOH/chloroform or EtOH/toluene. [*Beilstein* 4 IV 1070.]

Aminoacetaldehyde dimethyl acetal (2,2-dimethoxyethylamine) [22483-09-6] $\text{C}_4\text{H}_{11}\text{NO}_2$, **M 105.1, m <-78°, b 139.5°/768mm, 137-139°/atm, d_4^{20} 0.9676 n_D^{20} 1.4144**. Dry the acetal over KOH pellets and distil it through a 30cm vacuum jacketed Vigreux column. [Lawson *J Am Chem Soc* 75 3398 1953, DOI: 10.1021/ja01110a029; Erickson et al. *J Am Chem Soc* 77 6640 1955, DOI: 10.1021/ja01629a074; *Beilstein* 4 IV 1918.]

Aminoacetonitrile bisulfate [151-63-3] $\text{C}_2\text{H}_4\text{N}_2 \cdot \text{H}_2\text{SO}_4$, **M 154.15, m 125°(dec), pK^{25} 5.34 (NH_2)**. Recrystallise the hydrogen sulfate (1:1) from EtOH/Et₂O (hygroscopic leaflets). The **Sulfate (2:1)** [5466-22-8] crystallises as flat prisms from H_2O /EtOH with **m 166°(dec)**. [Stephen *J Chem Soc* 871 1931, DOI: 10.1039/JR9310000871; Anslow & King *J Chem Soc* 2463 1929, DOI: 10.1039/JR9290002463; *Beilstein* 4 III 1120.]

Aminoacetonitrile hydrochloride [6011-14-9] $\text{C}_2\text{H}_4\text{N}_2 \cdot \text{HCl}$, **M 92.5, m 166-167°, 172-174°, pK^{25} 5.24 (NH_2)**. The salt recrystallises from dilute EtOH as *hygroscopic* leaflets. It is best to crystallise it from absolute EtOH/Et₂O (1:1) and then recrystallise it from absolute EtOH. The melting point recorded ranges from 144° to 174°. The **free base** has **b 58°/15mm** with partial decomposition. [Mange *J Am Chem Soc* 56 2197 1934, DOI: 10.1021/ja01325a063; Goldberg & Kelly *J Chem Soc* 1369 1947, DOI: 10.1039/JR9470001369; *Beilstein* 4 H 344, 4 I 468, 4 II 783, 4 III 1120, 4 IV 2363.]

2-Amino-1-butanol [*RS*(±) 96-20-8, *R*(-) 5856-63-3, *S*(+) 5856-62-2] $\text{C}_4\text{H}_{11}\text{NO}$, **M 89.1, m ~ -2°, b 78-80°/10mm and 179-183°/atm for (±), 172-174°/atm for (+) or (-), pK^0 10.353, pK^{20} 9.672, pK^{60} 8.555**. They are purified by shaking with solid NaOH, filtering and distilling through a short column. The **oxalate salt** of the racemate has **m 176°**. They are strong bases and should be stored under N_2 in the absence of CO_2 . The **enantiomers** have $[\alpha]_D^{20}$ +12.5 and -12.5 (c 2, EtOH). [Johnson & Degering *J Org Chem* 08 7 1943, DOI: 10.1021/jo01189a002; Nagao et al. *J Org Chem* 51 2391 1986, DOI: 10.1021/jo00362a047; Santaniello et al. *JCS Perkin Trans 1* 919 1985, DOI: 10.1039/P19850000919; *Beilstein* 4 H 291, 4 IV 1705.]

2-Aminoethanol (ethanolamine) [141-43-5] $\text{C}_2\text{H}_7\text{NO}$, **M 61.1, f 10.5°, b 72-73°/12mm, 171.1°/760mm, d_4^{20} 1.012, n_D^{20} 1.4539, pK^{25} 9.51**. It decomposes slightly when distilled at atmospheric pressure, with the formation of conducting impurities. Fractional distillation at about 12mm pressure is most satisfactory. After distillation, 2-aminoethanol is further purified by repeated washing with ether and crystallising from EtOH (at low temperature). After fractional distillation in the absence of CO_2 , it is twice crystallised by cooling, followed again by distillation. It is *hygroscopic*, and absorbs CO_2 from the atmosphere. [Reitmeier et al. *J Am Chem Soc* 62 1943 1940, DOI: 10.1021/ja01865a009.] It can be dried by azeotropic distillation with dry C_6H_6 . [*Beilstein* 4 IV 1406.] **2-Aminoethanol hydrochloride** [2002-24-6] $\text{C}_2\text{H}_7\text{NO} \cdot \text{HCl}$, **M 97.6, m 75-77°**, is recrystallised from EtOH. It is deliquescent; store it dry. [*Beilstein* 4 IV 1406.] **2-Aminoethyl hydrogen sulfate (sulfuric acid mono-2-aminoethyl ester)** [926-39-6] $\text{C}_2\text{H}_7\text{NO}_4\text{S}$, **M 141.1, m 285-287° (chars at 275°)**, crystallises from water, or is dissolved in water and EtOH is added. Wash this with Et₂O and dry it *in vacuo*. [*Beilstein* 4 III 1414.]

S-(2-Aminoethyl)isothiuronium bromide hydrobromide [56-10-0] $\text{C}_3\text{H}_9\text{N}_3\text{S} \cdot 2\text{HBr}$, **M 281.0, m 194-195°**. Crystallise the salt from absolute EtOH/ethyl acetate or MeOH. Store dry as it is *hygroscopic* in a humid atmosphere. It is a radioprotective agent. When refluxed in EtOH for 16 hours or H_2O for 30 minutes, it decom-

poses to **2-amino-4(5H)-thiazoline hydrobromide** which on recrystallisation from isoPrOH/EtOAc has **m 175-176°** [Doherty et al. *J Am Chem Soc* **79** 5667 1957, DOI: 10.1021/ja01578a022].

(2-Aminoethyl)trimethylammonium chloride hydrochloride (chloramine chloride hydrochloride) [3399-67-5] **C₅H₁₅N₂Cl·HCl**, **M 175.1**, **m 268°(dec)**. Crystallise the hydrochloride from EtOH. The material is very soluble in H₂O. [Beilstein **4** II 690.]

Aminomalononitrile toluene-4-sulfonate [5098-14-6] **C₃H₄N₂·CH₃C₆H₄SO₃H**, **M 253.3**, **m 168-170°, 172°(dec)**, **pK_{Est} ~ 1.3**. It forms colourless crystals on recrystallisation from MeCN (1.8g in 100mL) using activated charcoal. Wash the crystals with dry Et₂O and dry them at 25°/1mm. Recovery is ~80%. [Ferris et al. *Org Synth Coll Vol* **5** 32 1973, DOI: 10.1522/orgsyn.048.0001.]

(±)-2-Amino-4-methylhexane (Forthane, 1,3-dimethylpentylamine) [105-41-9] **C₇H₁₇N**, **M 115.2**, **b 131-133°/atm, 130-135°/atm, 135-136°/atm**, **d₄²⁰ 0.760**, **n_D¹⁵ 1.4160**, **n_D²⁴ 1.4160**, **pK²⁵ 10.54**. This strong base is obtained by catalytic hydrogenation of 4-methylhexan-2-one oxime (57g) in EtOH (50ml) with Raney Ni (6g) in a bomb at 75-80° and 1000psi of H₂, cool, filter off the catalyst, acidify with HCl, and evaporate to dryness. Basify the residue with aqueous NaOH, extract it with Et₂O, dry the extract (MgSO₄), filter, evaporate, and distil to give the base in 75-80% yield. It forms a **sulfate salt m 215-220°(dec)**, and it readily forms a **carbonate salt**, hence it should be stored in the absence of CO₂. It is physiologically active as an opium *pressor* and is adrenergic. [Rohrmann & Shonle *J Am Chem Soc* **66** 1516 1944, DOI: 10.1021/ja01237a032; Chiang *J Clin Chem Soc* **18** 65 1951, USP to E. Lilly 2350318 (1943) *Chem Abstr* **39** 1510 1945, USP 2386273 (1943) *Chem Abstr* **40** 598 1946, *Beilstein* **4** III 378, **4** IV 747.]

2-Amino-2-methyl-1,3-propanediol [115-69-5] **C₄H₁₁NO₂**, **M 105.1**, **m 111°, b 151-152°/10mm**, **pK²⁵ 8.80**. Crystallise the diol three times from MeOH, dry in a stream of dry N₂ at room temperature, then in a vacuum oven at 55°. Store it over CaCl₂ [Hetzer & Bates *J Phys Chem* **66** 308 1962, DOI: 10.1021/j100808a027]. [*Beilstein* **4** IV 1881.]

2-Amino-2-methyl-1-propanol (β-aminoisobutanol) [124-68-5] **C₄H₁₁NO₂**, **M 89.1**, **m 24°, 31°, b 67°/10mm, 164-166°/760mm**, **d₄²⁰ 0.935**, **n_D²⁰ 1.45**, **pK²⁵ 9.71**. Purify it by distilling and fractional freezing. Store away from CO₂. The **hydrochloride** [3207-12-3] has **m 204°-206°**. [*Beilstein* **4** III 783, **4** IV 1740.]

n-Amyl acetate (n-pentyl acetate) [628-63-7] **C₇H₁₄O₂**, **M 130.2**, **b 149.2°/atm**, **d₄²⁰ 0.876**, **n_D²⁰ 1.40228**. Shake the ester with saturated NaHCO₃ solution until neutral, washed it with water, dry with MgSO₄ and distil it. The ester has also been purified by repeated fractional distillation through an efficient column or spinning band column. [Mumford & Phillips *J Chem Soc* **75** 1950, DOI: 10.1039/JR9500000075; ¹H NMR: Crawford & Foster *Can J Phys* **34** 653 1956, DOI 10.1139/p56-074; *Beilstein* **2** IV 152.]

n-Amyl alcohol (1-pentanol) [71-41-0] **C₅H₁₂O**, **M 88.2**, **b 138.1°/atm**, **d¹⁵ 0.818**, **n_D²⁰ 1.4100**. Dry 1-pentanol with anhydrous K₂CO₃ or CaSO₄, filter and fractionally distil it. It has also been treated with 1-2% of sodium and heated at reflux for 15 hours to remove water and chlorides. Traces of water can be removed from the near-dry alcohol by refluxing it with a small amount of sodium in the presence of 2-3% *n*-amyl phthalate or succinate followed by distillation (see *ethanol*). Small amounts of amyl alcohol have been purified by esterifying with *p*-hydroxybenzoic acid, recrystallising the ester from CS₂, saponifying with ethanolic-KOH, drying with CaSO₄ and fractionally distilling [Olivier *Recl Trav Chim Pays-Bas* **55** 1027 1936, DOI: 10.1002/recl.19360551207]. [*Beilstein* **1** IV 1640.]

tert-Amyl alcohol (2-methyl-2-butanol) [75-85-4] **C₅H₁₂O**, **M 88.2**, **m -12°, b 102.3°/atm**, **d¹⁵ 0.8135**, **n_D²⁰ 1.4058**. Reflux it with K₂CO₃, CaH₂, CaO or sodium, then fractionally distil. The near-dry alcohol is further dried by refluxing with Mg activated with iodine, as described for *ethanol*. Further purification is possible using fractional crystallisation and zone refining at <-10° or preparative gas chromatography. [*Beilstein* **1** IV 1668.]

n-Amylamine [1-aminopentane] [110-58-7] **C₅H₁₃N**, **M 87.2**, **b 105°/atm**, **d₄²⁰ 0.752**, **pK²⁵ 10.63**. Dry it by prolonged shaking with NaOH pellets, then distilling. Store it in a CO₂-free atmosphere. [*Beilstein* **4** IV 674.]

***n*-Amyl bromide (*n*-pentylbromide)** [110-53-2] $C_5H_{11}Br$, M 151.1, b 129.7°/atm, d_4^{20} 1.218, n_D^{20} 1.445. Wash the bromide with conc H_2SO_4 , then water, 10% Na_2CO_3 solution, again with water, dry with $CaCl_2$ or K_2CO_3 , and fractionally distil it just before use. [Beilstein 1 IV 312.]

***n*-Amyl chloride (1-chloropentane)** [543-59-9] $C_5H_{11}Cl$, M 106.6, b 107.8°/atm, d_4^{20} 0.882, n_D^{20} 1.41177. Purify as for *sec*-amyl chloride below. [Beilstein 1 IV 309.]

***sec*-Amyl chloride (1-chloro-2-methylbutane)** [616-13-7] $C_5H_{11}Cl$, M 106.6, b 52.2°/150mm, 96-97° (100°)/760mm, d_4^{20} 0.886, n_D^{20} 1.412. Purify the chloride by stirring vigorously with 95% H_2SO_4 , replacing the acid when it becomes coloured, until the layer remains colourless after 12 hours stirring. The amyl chloride is then washed with saturated Na_2CO_3 solution, then distilled water, and dried with anhydrous $MgSO_4$, followed by filtration, and distillation through a 10-in Vigreux column. Alternatively, a stream of oxygen containing 5% ozone is passed through the amyl chloride for three times as long as it takes to cause the first coloration of starch iodide paper by the exit gas. The liquid is washed with $NaHCO_3$ solution to hydrolyse the ozonides and remove organic acids prior to drying and fractional distillation [Chien & Willard *J Am Chem Soc* 75 6160 1953, DOI: 10.1021/ja01120a016. The *S*(+)-enantiomer [40560-29-0] has b 50-51°/140mm, 100°/760, mm, n_D^{20} 1.404 [α] $_D^{20}$ +1.64 (neat) [Brown et al. *J Am Chem Soc* 62 3435 1940, DOI: 10.1021/ja01869a040]. [Beilstein 1 H 134, 1 I 46, 1 III 356, 1 IV 326.]

***tert*-Amyl chloride (2-chloro-2-methylbutane)** [594-36-5] $C_5H_{11}Cl$, M 106.6, b 86°/atm, d_4^{20} 0.866. Methods of purification commonly used for other alkyl chlorides lead to decomposition. Unsaturated contaminants are removed by chlorination with a small amount of chlorine in bright light, followed by distillation [Chien & Willard *J Am Chem Soc* 75 6160 1953, DOI: 10.1021/ja01120a016]. [Beilstein 1 H 134, 1 I 46, 1 II 100, 1 III 357, 1 IV 324.]

Amylene (β -iso-amylene, 2-methyl-2-butene) [513-35-9] C_5H_{10} , M 70.1, b 37-38°/760mm, d_4^{20} 0.663, n_D^{20} 1.387. Distil amylene and collect the distillate at low temperature. It has also been distilled from sodium. **FLAMMABLE.** It is available in steel cylinders and has a short shelf life. [Beilstein 1 H 211, 1 I 87, 1 II 187, 1 III 788, 1 IV 820.]

Amyl ether (dipentyl ether) [693-65-2] $C_{10}H_{22}O$, M 158.3, b 186.8°/atm, d_4^{20} 0.785, n_D^{20} 1.41195. Repeatedly reflux amyl ether over sodium and distil it. [Beilstein 1 IV 1643.]

Arachidic (eicosanoic C_{20}) acid [506-30-9] $C_{20}H_{40}O_2$, M 312.5, m 77°, pK_{Est} ~5.0. Crystallise the C_{20} acid from absolute EtOH. [Beilstein 2 IV 1276.] **Arachidic alcohol (1-eicosanol C_{20})** [629-96-9] $C_{20}H_{42}O$, M 298.6, m 65.5° (71°), b 200°/3mm. Crystallise the C_{20} alcohol from *benzene or *benzene/petroleum ether. [Beilstein 1 IV 1900.]

Azelaic acid (1,9-nonanedioic acid, heptane-1,7-dicarboxylic acid) [123-99-9] $C_9H_{16}O_4$, M 188.2, m 105-106°, b 225°/10mm, 256°/50mm, pK_1^{25} 4.53, pK_2^{25} 5.33. Recrystallise it from H_2O (charcoal) or thiophene-free *benzene. The acid can be dried by azeotropic distillation with toluene, the residual toluene solution is then cooled and filtered, and the precipitate is dried in a vacuum oven. It has been purified by zone refining or by sublimation onto a cold finger at 10^{-3} torr. It distils above 360° with partial formation of the anhydride. The **dimethyl ester** [1732-10-1] $C_{11}H_{20}O_4$, M 216.3, has m -3.9° and b 140°/8mm, 156°/20mm, d_4^{25} 1.007g/ml, n_D^{20} 1.435. [Hill & McEwen *Org Synth Coll Vol* 2 53 1943, DOI: 10.15227/orgsyn.013.0004; Beilstein 2 IV 2055.]

2,2'-Azobis(isobutyronitrile) [α,α' -bis-azo-(2-methylpropionitrile), AIBN] [78-67-1] $C_8H_{12}N_4$, M 164.2, m 102.1-103.2°, 103°, 103-104°. Crystallise the nitrile from Et_2O , Me_2CO , $CHCl_3$, aqueous EtOH or MeOH. It has also been crystallised from absolute EtOH below 40° in subdued light. Dry it under vacuum at room temperature over P_2O_5 and store it under vacuum in the dark at <-10° until required. Also crystallise it from $CHCl_3$ solution by addition of petroleum ether (b <40°). It is a **radical inhibitor**. [Askham et al. *J Am Chem Soc* 107 7423 1985, DOI: 10.1021/ja00311a034; Ennis et al. *JCS Dalton Trans* 2485 1986, DOI: 10.1039/DT9860002485; Inoue & Anson *J Phys Chem* 91 1519 1987, DOI: 10.1021/j100290a046; Tanner et al. *J Org Chem* 52 2142 1987, DOI: 10.1021/jo00387a005.] It is prepared by the oxidation of *N,N'*-bis(isobutyro-nitrile)

hydrazine by addition of the solid hydrazine (4.15g, 0.025mole) to a solution of HNO_3 (100%, 6.5g, 0.1mol) and Ac_2O (30ml) (0.1mole) which are previously mixed at -30° . As the temperature rises to -20° the solid begins to dissolve and the colour of the solution turns to green, and dissolution is complete at $\sim 5^\circ$. After stirring at 25° for 0.5 hours, the solution is poured onto cracked ice and H_2O (100ml), the product separates out, is filtered off, and washed free from acid with H_2O . The solid (m $95\text{--}100^\circ$) is dried *in vacuo*, recrystallised from 50% EtOH and from Et_2O to give pure *AIBN* (2.5g, 60%), m 105° . [Picard & Boivin *Can J Chem* **29** 223 1951, DOI:10.1139/v51-027.] Alternatively, the *hydrazine* (0.989g, 6mmol) is added to a solution of AcOH (40ml) and conc H_2SO_4 (3ml) at $\sim 0^\circ$, followed by addition of $\text{NaMnO}_2 \cdot 3\text{H}_2\text{O}$ (1.8g) in H_2O (25ml). MnO_2 separates, excess NaMnO_2 and MnO_2 are reduced with aq sodium bisulfite and the mixture is poured into H_2O . *AIBN* separates, is filtered off, washed with H_2O , dried *in vacuo* and recrystallised as above to give pure *azo compound* (0.61g, 63%). [Overberger & Lebovitz *J Am Chem Soc* **76** 2722 1954, DOI: 10.1021/ja01639a039; Overberger et al. *J Am Chem Soc* **71** 2661 1949, DOI: 10.1021/ja01176a018.] [*Beilstein* **4** I 566, **4** III 1750, **4** IV 3377.]

Azomethane (dimethyldiimide) [503-28-6] $\text{C}_2\text{H}_6\text{N}_2$, M **58.1**, m -78° , b $1.5^\circ/\text{atm}$, d_0^{15} **0.981**, n_D^{19} **1.3933**. Purify azomethane by distillation in a vacuum line and store it in the dark at -80° . It is soluble in EtOH, Et_2O and EtOAc. It can be **EXPLOSIVE**. [*Beilstein* **4** H 562, **4** I 566, **4** II 966, **4** III 1747, **4** IV 3366.]

Batyl alcohol (rac-3-[1-octadecyloxy]-1,2-propanediol) [544-62-7] $\text{C}_{21}\text{H}_{44}\text{O}_3$, M **344.6**, m $71\text{--}71.5^\circ$. Batyl alcohol crystallises from aqueous Me_2CO , EtOH or petroleum ether (b $40\text{--}60^\circ$). [Taguchi & Armarego *Med Res Rev* **18** 43 1998, DOI: 10.1002/(SICI)1098-1128(199801); *Beilstein* **1** IV 2758.] Natural **S(+)-alcohol** has m $71\text{--}72^\circ$ (from dry Me_2CO), and $[\alpha]_D^{25}$ **+1.54**, $[\alpha]_{365}^{25}$ **+3.9** (c **6.7** CHCl_3), and *isopropylidene derivative* has m $34\text{--}36^\circ$ and $[\alpha]_D^{40}$ **-11.15** (melt). [Baer & Fischer *J Biol Chem* **140** 397 1941, <http://www.jbc.org/content/140/2/397.citation>; Taguchi et al. *Pteridines* **6** 45 1995, DOI: 10.1515/pteridines.1995.6.2.45.]

Behenoyl chloride (docosanoyl chloride) [21132-76-3] $\text{C}_{22}\text{H}_{43}\text{ClO}$, M **359.0**, m 40° . If the IR shows OH bands, then it should be dissolved in oxalyl chloride in $^*\text{C}_6\text{H}_6$ solution and warmed at 35° for 24 hours in the absence of moisture, evaporated and distilled in a vacuum of 10^{-5}mm . It is soluble in $^*\text{C}_6\text{H}_6$ and Et_2O . It is moisture sensitive. **LACHRYMATORY**. [Francis et al. *J Chem Soc* 999 1937, DOI: 10.1039/JR9370000999; Levene & Taylor *J Biol Chem* **59** 905 1924, <http://www.jbc.org/content/59/3/905>; *Beilstein* **2** III 1076.]

Biacetyl (butan-2,3-dione) [431-03-8] $\text{C}_4\text{H}_6\text{O}_2$, M **86.1**, b $88^\circ/\text{atm}$, d_4^{20} **0.981**, $n_D^{18.5}$ **1.3933**. Dry biacetyl over anhydrous CaSO_4 , CaCl_2 or MgSO_4 , then distil it in a vacuum under nitrogen, taking the middle fraction and storing it at Dry-Ice temperature in the dark (to prevent polymerisation). [*Beilstein* **1** IV 3644.]

Biguanide [56-03-1] $\text{C}_2\text{H}_7\text{O}_5$, M **101.1**, m 130° pK₁²⁵ **3.1**, pK₂²⁵ **12.8** (**11.52**). Crystallise biguanide from EtOH. It gives a red Cu derivative, and it forms salts with many metals. The *monohydrochloride* has m 235° [38664-03-8] and the *dihydrochloride* forms plates with m 248° ($213\text{--}214^\circ$, also reported) [25836-74-2]. [*Beilstein* **3** H 93, **3** I 44, **3** II 76, **3** III 171, **3** IV 162.]

Bis-acrylamide (N,N'-methylene bisacrylamide) [110-26-9] $\text{C}_7\text{H}_{10}\text{N}_2\text{O}_2$, M **154.2**, m $>300^\circ$. Recrystallise the amide from MeOH (100g dissolved in 500ml boiling MeOH) and filter without suction in a warmed funnel. Allow to stand at room temperature and then at -15° overnight. The crystals are collected with suction in a cooled funnel and washed with cold MeOH. The crystals are air-dried in a warm oven. [*Beilstein* **2** IV 1472.] **VERY TOXIC (neurotoxic)**.

Bis(β -chloroethyl)amine hydrochloride [821-48-7] $\text{C}_4\text{H}_9\text{Cl}_2\text{N} \cdot \text{HCl}$, M **178.5**, m $214\text{--}215^\circ$, $216\text{--}217^\circ$, pK_{Est} ~ 5.8 (free base). Crystallise the salt from Me_2CO or MeOH/ Et_2O . The *picrate* has m $112\text{--}113^\circ$ (from EtOH or Me_2CO). [Mann *J Chem Soc* 461 1934, DOI: 10.1039/JR9340000461; Ward *J Am Chem Soc* **57** 914 1935, DOI: 10.1021/ja01308a041; *Beilstein* **4** III 238.]

Bis(β -chloroethyl) ether [111-44-4] $\text{C}_4\text{H}_8\text{Cl}_2\text{O}$, M **143.0**, b $94^\circ/33\text{mm}$, 178.8° , d_4^{20} **1.220**, n_D^{20} **1.45750**. Wash the ether with conc H_2SO_4 , then Na_2CO_3 solution, dry with anhydrous Na_2CO_3 , and finally pass it through a 50cm column of activated alumina before distillation. Alternatively, wash it with 10% ferrous sulfate solution

to remove peroxides, then H₂O, dry with CaSO₄, and distil it in a vacuum. Add 0.2% of catechol to stabilise it. [Beilstein 1 IV 1375, Kamm & Waldo *J Am Chem Soc* 43 2223 1921, DOI: 10.1021/ja01443a013.] **VERY TOXIC.**

2,2'-Bis[di-(carboxymethyl)-amino]diethyl ether, (EEDTA, HOOCCH₂)₂NCH₂CH₂OCH₂CH₂N-(CH₂COOH)₂ [923-73-9] C₁₂H₂₀N₂O₉, M 336.3, pK₁²⁰ 1.8, pK₂²⁰ 2.76, pK₃²⁰ 8.84, K₄²⁰ 9.47. Crystallise it from EtOH or aqueous EtOH. Alternatively, boil in MeOH, filter hot and cool. The tetra-acid is collected, recrystallised from H₂O, and the solid is collected and dried over P₂O₅ at 90°/0.01mm. [Schwarzenbach et al. *Helv Chim Acta* 40 1886 1957, DOI: 10.1002/hlca.19570400640.] It complexes with Ca²⁺, Ba²⁺, Sr²⁺, Mg²⁺, Ag⁺, Hg²⁺, Cd²⁺ and Nd³⁺. [Beilstein 4 II 37, 4 IV 2433.]

Bis(2-ethylhexyl) sebacate ['dioctyl' sebacate, di(2-ethylhexyl) 1,8-octanedicarboxylate] [122-62-3] C₂₆H₅₀O₄, M 426.7, b 212°/1mm, d₄²⁵ 1.914, n_D²⁰ 1.4496. If it is acidic due to hydrolysis (effervesces with NaHCO₃), then purify by dissolving in Et₂O, shaking with aqueous Na₂CO₃, dry (Na₂SO₄), filter, evaporate and distil the residue at a high vacuum. Note that *2-ethylhexan-1-ol* has b 184-185°/760mm with estimated b ~40-50°/1mm (see Figure 1, and Table 2A, Chapter 1) and will distil at a much lower temperature than the ester. Otherwise it should be distilled through a short column under high vacuum and the higher boiling fraction is redistilled. This vacuum pump oil can be prepared by esterifying decanedioic acid with 2-ethylhexanoic acid by Fischer-Speier's method (Fischer & Speier *Chem Ber* 28 1145 1895, DOI: 10.1002/cber.189502801248), whereby dry HCl gas is bubbled through the alcohol until its weight is increased by ~10%, the acid is added, the mixture is heated at 100° until esterification is complete (~2-3 hours), and purified as above. [Bruno USPat 2628249 1953, GBPat 747260 1956, Beilstein 2 IV 2803.]

N,N-Bis(2-hydroxyethyl)-2-aminoethanesulfonic acid (BES) [10191-18-1] M 213.3, m 150-155°, pK²⁵ 7.17. Crystallise BES from aqueous EtOH. [Beilstein 4 IV 3290.]

Bis(2-hydroxyethyl)amino-tris-(hydroxymethyl)methane (BIS-TRIS) [6976-37-0] C₈H₁₉NO₅, M 209.2, m 89°, 104°, pK²⁰ 6.46. Crystallise BIS-TRIS from hot 1-butanol and dry it in a vacuum at 25°.

Bis(2-hydroxypropyl) ether [dipropylene glycol, 4-oxaheptane-2,6-diol, 1,1'-oxybis(propan-2-ol)] [110-98-5, 25265-71-8 non-specific] C₆H₁₄O₃, M 134.2, m 45-46.5°, 46-48°, b 113-114°/10mm, 222.2°/760mm, 229-232°/atm, d₄⁶⁰ 0.9878, n_D⁶⁰ 1.4284. Fractionally distil the diol preferably at low pressure, using a packed column and taking precautions to avoid absorption of water. It has also been purified by crystallisation from ether at Dry Ice temperatures which gives a white solid. There are apparently two forms (*meso* and *d,l* ?) [see Summerbell et al. *J Org Chem* 27 4433 1961, DOI: 10.1021/jo01059a075]. The *bis-p*-nitrobenzoate has m 150-151°, (146-148°) and the *bis*-triphenylmethyl ether has m 145-146° [Sexton & Britton *J Am Chem Soc* 75 4357 1953, DOI: 10.1021/ja01113a525; Beilstein 1 II 537, 1 III 2149, 1 IV 2473.]

Bis(3-hydroxypropyl) ether [4-oxaheptane-1,7-diol, 3,3'-oxybis(propan-1-ol)] [2396-61-4, 25265-71-8 non-specific] C₆H₁₄O₃, M 134.2, b 92-102°/0.01mm, d₄²⁵ 1.023, n_D²⁰ 1.441. Fractionally distil the diol under 15mm pressure, using a packed column and taking precautions to avoid absorption of water. Four isomers are obtained from the condensation of 2 molecules of propylene oxide which have to be separated by fractional distillation [Sexton & Britton *J Am Chem Soc* 75 4357 1953, DOI: 10.1021/ja01113a525, see Summerbell et al. *J Org Chem* 27 4433 1961, DOI: 10.1021/jo01059a075]. However, a clean product was obtained in 23% yield, by reduction of *diethyl 3-oxaheptanoate* with LiAlH₄ in Et₂O under N₂ and distilling under high vacuum. Its ¹H NMR (in CDCl₃/TMS, 60MHz) has δ 1.81 (2 x 2H, quint), 3.55 (2 x 2H, t) and 3.70 (2 x 2H, t); MS *m/z* 134. The *mono*- and *bis*-tosylates are oils. [Samat et al. *JCS Perkin Trans 1* 1717 1985, DOI: 10.1039/JR9550002657.] [Beilstein 1 III 542, 1 IV 2496.] The *bis-phenylurethane* has m 104-105° (from EtOH). The *Di(propylene glycol) dibenzoate* [27138-31-4] C₂₀H₂₂O₄, M 326.4, is purified by redistillation at 232°/5mm, d₄²⁵ 1.12, n_D²⁰ 1.528 [Beilstein 9 II 108].

Bis(2-mercaptoethyl)sulfone (BMS) [145626-87-5] C₄H₁₀O₂S₃, M 186.3, m 57-58°, pK₁²⁵ 7.9, pK₂²⁵ 9.0. BMS recrystallises from hexane as white fluffy crystals. Large amounts are best recrystallised from deoxygenated H₂O (charcoal). It is a good alternative reducing agent to dithiothreitol. Its IR (film) has

ν_{\max} at 2995, 2657, 1306, 1248, 1124 and 729 cm^{-1} . The synthetic intermediate **thioacetate** has **m** 82-83° (white crystals from CCl_4). The **disulfide** is purified by flash chromatography on SiO_2 and elution with 50% EtOAc/hexane, recrystallised from hexane, and has **m** 137-139°. [Lamoureux & Whitesides *J Org Chem* **58** 633 1993, DOI: 10.1021/jo00055a015; *Beilstein* **1** IV 2455.]

Bis(trichloromethyl) carbonate (triphosgene) [32315-10-9] $\text{C}_3\text{Cl}_6\text{O}_3$, **M** 296.8, **m** 79-83°, **81-83°**, **b** 203-206°(slight dec). It is a good solid substitute for phosgene (using a third mol per mol). Crystallise it from petroleum ether (b 60-80°), wash it with anhydrous cold Et_2O , de-gas it at 200mm then dry at 0.1mm (over H_2SO_4). It has IR with ν_{\max} at 900 and 1900 cm^{-1} . It is a **lachrymator**, is **TOXIC**, and should be handled with gloves and in an efficient fume hood. [Hales et al. *J Chem Soc* 618 1957, DOI: 10.1039/JR9570000618; Eckert & Forster, **26** 894 1987, DOI: 10.1002/anie.198708941; *Aldrichimica Acta* **21** 47 1988, *Beilstein* **3** H 17, **3** I 8, **3** II 16, **3** III 36, **3** IV 33.]

Bistrifluoroacetamide (BTFA) [407-24-9] $\text{C}_4\text{HF}_6\text{NO}_2$, **M** 209.1, **m** 85°, **b** 135-136°/744mm, 141°/760mm. A major impurity is trifluoroacetamide. Add trifluoroacetic anhydride to BTFA, reflux for 2 hours and fractionate using a Vigreux column at atmospheric pressure. [Donike *J Chromatogr* **78** 273 1973, Donike *J Chromatogr* **103** 91 1975, DOI 10.1016/S0021-9673(00)83805-4; *Beilstein* **2** IV 471.]

Biuret (allophanic acid amide, carbamoylurea) [108-19-0] $\text{C}_2\text{H}_5\text{N}_3\text{O}_2$, **M** 103.1, **sinters at 218° and chars at 270°**, **pK₁²⁵ -0.88, pK₂²⁵ >4**. Crystallise biuret from EtOH. [*Beilstein* **3** IV 141.]

N-Bromoacetamide [79-15-2] $\text{C}_2\text{H}_4\text{BrNO}$, **M** 138.0, **m** 102-105°, **107-109°**, **108°(anhydrous)**. A possible contaminant is $\text{CH}_3\text{CONBr}_2$. Recrystallise it from CHCl_3 /hexane (1:1, seed if necessary) or water and dry over CaCl_2 . It is a brominating agent. [Oliveto & Gerold *Org Synth Coll Vol* **4** 104 1963, DOI: 10.15227/orgsyn.031.0017.] Alternatively, dissolve it in the minimum volume of warm H_2O (60°), then cool in an ice bath, collect the crystals and dry them in an anhydrous atmosphere, dissolve in Et_2O , chill and evaporate till crystallisation occurs. Dry the crystals *in vacuo* at 25°, then at 45° (**m** 108°). Crystallise from CHCl_3 (**m** 103°). Estimate the available Br iodometrically [Buckles et al. *J Org Chem* **23** 483 1958, DOI: 10.1021/jo01097a605]. [*Beilstein* **2** H 181, **2** I 82, **2** II 180, **2** III 406, **2** IV 417.]

Bromoacetic acid [79-08-3] $\text{C}_2\text{H}_3\text{BrO}_2$, **M** 138.9, **m** 50°, **b** 118°/15mm, **208°/760mm**, **d₄²⁵ 1.93, pK_a²⁵ 2.92**. Crystallise bromoacetic acid from petroleum ether (b 40-60°). A diethyl ether solution of it is passed through an alumina column, and the ether is evaporated at room temperature under vacuum. It is best obtained by distillation from a Claisen flask (immersed in an oil bath) fitted with an insulated Vigreux column and the fraction **b** 108-110°/30mm is collected. It is light and moisture sensitive. [Natelson & Gottfried *Org Synth Coll Vol* **3** 381 1955, DOI: 10.15227/orgsyn.023.0037; *Beilstein* **2** IV 526.] **LACHRYMATORY and is a skin IRRITANT.**

Bromoacetone [598-31-2] $\text{C}_3\text{H}_5\text{BrO}$, **M** 137.0, **b** 31.5°/8mm, **63.5-64°/50mm**, **137°/atm**, **d₄²³ 1.643**. Stand bromoacetone over anhydrous CaCO_3 , filter, distil it under low vacuum, and store it with CaCO_3 in the dark at 0°. [Levene *Org Synth Coll Vol* **2** 88 1943, DOI: 10.15227/orgsyn.010.0012.] **Violently LACHRYMATORY and skin IRRITANT.**

2-Bromobutane [78-76-2] $\text{C}_4\text{H}_9\text{Br}$, **M** 137.0, **b** 91.2°/atm, **d₄²⁰ 1.255, n_D²⁰ 1.4367, n_D²⁵ 1.4341**. Wash 2-bromobutane with concentrated HCl, water, 10% aqueous NaHSO_3 , and then water. Dry it with CaCl_2 , Na_2SO_4 or anhydrous K_2CO_3 , and fractionally distil it through a 1m glass helices packed column. [*Beilstein* **1** IV 261.]

Bromoform [75-25-2] CHBr_3 , **M** 252.7, **m** 8.1°, **55-56°/35mm**, **149.6°/760mm**, **d₁₅ 2.9038, d₃₀ 2.86460, n_D¹⁵ 1.60053, n_D²⁰ 1.5988**. The storage and stability of bromoform and chloroform are similar. Ethanol, added as a stabiliser, is removed by washing with H_2O or with saturated CaCl_2 solution, and the CHBr_3 , after drying with CaCl_2 or K_2CO_3 , is fractionally distilled. Prior to distillation, CHBr_3 has also been washed with concentrated H_2SO_4 until the acid layer is no longer coloured, then dilute NaOH or NaHCO_3 , and H_2O . A further purification step is fractional crystallisation by partial freezing. [*Beilstein* **1** IV 82.]

(±)-2-Bromohexadecanoic acid (2-bromopalmitic acid) [18263-25-7] $C_{16}H_{31}BrO_2$, M 335.3, m 51-53°, 52.3-52.5°, 53°, $pK_{Est} \sim 3.2$. Recrystallise the acid from petroleum ether (b 60-80°, charcoal) and finally from EtOH. The *ethyl ester* has b 177-178°/2mm, d_{28}^{28} 1.0484, n_D^{20} 1.4560. [IR: Sweet & Estes *J Org Chem* 21 1426 1956, DOI: 10.1021/jo01118a026; *Beilstein* 2 IV 1184.]

(±)2-Bromohexanoic acid (α-bromocaproic acid) [616-05-7] $C_6H_{11}BrO_2$, M 195.1, b 118-125°/8mm, 148-153°/30mm, 240°/atm, d_4^{20} 1.37g/ml, n_D^{20} 1.472, $pK_{Est} \sim 3.1$. It can be synthesised as described by Clarke & Taylor [*Org Synth* 4 9 1925, and *Org Synth Coll Vol I* 115 1941, DOI: 10.15227/orgsyn.004.0009] and finally purified by distillation in a vacuum. It is soluble in EtOH and Et₂O. Optical Resolution *via* the strychnine salt, [Levene et al. *J Biol Chem* 75 337 1927, <http://www.jbc.org/content/75/1/337.citation.full.html>; Levene & Mardashev *J Biol Chem* 117 707 1937, <http://www.jbc.org/content/117/2/707.citation.full.html>] and final distillation at 106-107°/4mm or 129-130°/14mm gives the (-)-bromo acid with $[\alpha]_D^{20}$ -27.0 (in Et₂O), and the (+)-bromo acid with b 86-89°/0.35mm and $[\alpha]_D^{25}$ +5.6 (neat). [*Beilstein* 2 II 287, 2 III 736, 2 IV 938.] The (±)-ethyl ester $C_8H_{15}BrO_2$, is a liquid with an aniseed odour and b 95-96°/9mm, 148-153°/30mm, 120-122°/41mm, 206-210°/atm, d_4^{25} 1.2210, n_D^{20} 1.4456.

6-Bromohexanoic acid (6-bromocaproic acid) [4224-70-8] $C_6H_{11}BrO_2$, M 195.1, m 32-33°, 35°, 34-36°, b 129-130°/5mm. It has been prepared by the oxidation of 6-bromohexanol with concentrated HNO₃ (sp. gr. 1.42, 1 hour for addition at 25-30°, stir at ~25° for 4 hours then at 100° for 45 minutes) [Degering & Boatright *J Am Chem Soc* 72 5137 1950, DOI: 10.1021/ja01167a091]. It is made more conveniently by oxidation of cyclohexanone (174 g) with Caro's acid (using a mixture prepared from 919g of potassium persulfate for 10-15 minutes as in Barger et al. *J Chem Soc* 718 1937, DOI: 10.1039/JR9570000618), and the crude lactone (200g) that is formed is treated with a cooled mixture of 48% HBr (1L) and concentrated H₂SO₄ (240ml). After standing at ~25° for 2 hours, then for 4 hours at 100°, the mixture is cooled, poured into H₂O, the organic layer is separated, the aqueous layer is saturated with (NH₄)₂SO₄, and is extracted with Et₂O. The combined organic layers and extracts are washed with saturated (NH₄)₂SO₄, dried (Mg₂SO₄), evaporated and distilled in a vacuum. [See Brown & Partridge *J Am Chem Soc* 66 839 1944, DOI: 10.1021/ja01233a503; Degering & Boatright *J Am Chem Soc* 72 5137 1950, DOI: 10.1021/ja01167a091]. [*Beilstein* 2 II 287, 2 III 737, 2 IV 940.] When the acid is refluxed with SOCl₂ in the absence of moisture until all the HCl and SO₂ have evolved, then evaporated, and the residue is distilled in a vacuum, it provides **6-bromohexanoyl chloride** [22809-37-6] M 213.5, b 130°/20mm, d_4^{25} 1.395, n_D^{20} 1.486 as an almost colourless oil which is typically reactive and corrosive like most acid chlorides. [Osbond *J Chem Soc* 3464 1951, DOI: 10.1039/JR9510003464]. **6-Bromohexanamide** has m 107-108°, and the *anilide* has m 84°. [*Beilstein* 2 IV 940.]

S-(+)-1-Bromo-2-methylbutane [534-00-9] $C_5H_{11}Br$, M 151.1, b 38.2°/39mm, 49°/62mm, 60.8°(57-58°)/100mm, 65-65.6°/140mm, 116-122°/atm, d_4^{20} 1.2232, n_D^{20} 1.4453, $[\alpha]_D^{20}$ +5.1 (c 5, CHCl₃) (neat, +5.8°). Wash the bromobutane with ice-cold H₂O, dry by freezing, shake it twice with an equal volume of H₂SO₄ at 0°, and twice with an equal volume of H₂O at 0°. Freeze-dry and keep over freshly heated (and then cooled) K₂CO₃, and distil it through a vacuum jacketed column of broken glass. *Alternatively*, dissolve it in petroleum ether (b 40-60°), wash it with 5% NaOH, concentrated H₂SO₄ (at 0°), then H₂O, dry (CaCl₂), evaporate it and distil. [Heller *J Am Chem Soc* 74 4858 1952, DOI: 10.1021/ja01139a037; Foley et al. *J Am Chem Soc* 81 2779 1959, DOI: 10.1021/ja01520a042; Easton & Hargreaves *J Chem Soc* 1413 1959, DOI: 10.1039/JR9590001413; Crombie & Harper *J Chem Soc* 2685 1950, DOI: 10.1039/JR9500002685; *Beilstein* 1 IV 327.]

2-Bromo-2-methylpropane [507-19-7] C_4H_9Br , M 137.0, b 71-73°, d_4^{20} 1.218, n_D^{20} 1.429. Neutralise the bromomethylpropane with K₂CO₃, distil, and dry it using molecular sieves (5A), then distil it in a vacuum and degas it by the freeze-pump-thaw technique. Seal it in ampoules under vacuum. [*Beilstein* 1 IV 295.]

1-Bromooctadecane [112-89-0] $C_{18}H_{37}Br$, M 333.4, m 26°, 27.3°, 28-30°, b 178-179°/2mm, 214-218°/15mm, d_4^{20} 0.976, n_D^{20} 1.461. Twice recrystallise bromooctadecane from the melt, then distil it under vacuum three times taking the middle cut. *Alternatively*, wash the oil with aqueous Na₂SO₄, then concentrated H₂SO₄ (cool) and again with aqueous Na₂SO₄, and then fractionally distil it. [Meyer & Reid *J Am Chem Soc* 55 1574 1933, DOI: 10.1021/ja01520a042; Hoffmann & Smyth *J Am Chem Soc* 72 171 1950, DOI: 10.1021/ja01157a048; IR: LeFèvre et al. *Aust J Chem* 12 743 1959, DOI:10.1071/CH9590743; *Beilstein* 1 IV 555.]

(±)-2-Bromopentane [107-81-3] $C_5H_{11}Br$, M 151.1, b 117.2°/753mm, 116-117°/atm, 117.5°/740mm, d_4^{20} 1.2190, n_D^{20} 1.4401. Dry it over K_2CO_3 and distil it through a short Vigreux column. [IR: Pines et al. *J Am Chem Soc* **74** 4063 1952, DOI: 10.1021/ja01136a027; Brown & Wheeler *J Am Chem Soc* **78** 2199 1956, DOI: 10.1021/ja01591a049; *Beilstein* **1** IV 312.]

Bromopicrin (tribromonitromethane) [464-10-8] CBr_3NO_2 , M 297.8, m 10.2-10.3°, b 85-87°/16mm, d_4^{20} 2.788, n_D^{20} 1.579. Steam distil it, dry it with anhydrous Na_2SO_4 and distil it again in a vacuum. **HIGHLY TOXIC**. [*Beilstein* **1** H 77, **1** I 21, **1** II 43, **1** III 115, **1** IV 106.]

R-(+)-2-Bromopropionic acid [10009-70-8] $C_3H_5BrO_2$, M 153.0, b 78°/4mm, d_4^{20} 1.474, $[\alpha]_D^{25}$ +27.2 (neat), pK^{25} 4.07. Dissolve it in Et_2O , dry ($CaCl_2$), evaporate and distil it through a short column. Distillation through a Podbielniak column (see *S*-(-)-2-chloropropionic acid below) led to decomposition. Store it in the dark under N_2 , preferably in sealed ampoules. Even at -10° it slowly decomposes. **LACHRYMATORY**. [Fu et al. *J Am Chem Soc* **76** 6054 1954, DOI: 10.1021/ja01591a049; *Beilstein* **2** IV 761.]

3-Bromopropionic acid [590-92-1] $C_3H_5BrO_2$, M 153.0, m 62.5°, 62.5-63.5°, 63-64°, pK^{25} 4.01. The acid crystallises as plates from CCl_4 . It is soluble in organic solvents and H_2O . Its *methyl ester* has b 65°/18mm and 80°/27mm. The *S*-benzylisothiuronium salt has m 136°. [Kendall & McKenzie *Org Synth Coll Vol I* 131 1941, DOI: 10.15227/orgsyn.003.0025; *Beilstein* **2** IV 764.]

Bromopyruvic acid (3-bromo-2-oxopropionic acid) [1113-59-3] $C_3H_3BrO_3$, M 167.0, m 79-82°, pK_{Est} ~1.6. Dry it by azeotropic distillation (with toluene), and then recrystallise it from dry $CHCl_3$. Dry for 48 hours at 20° (0.5 torr) over P_2O_5 . Store it at 0° away from light. [Labaudiniere et al. *J Org Chem* **52** 157 1987, DOI: 10.1021/jo00377a030; *Beilstein* **3** III 1167.]

N-Bromosuccinimide [128-08-5] $C_4H_4BrNO_2$, M 178.0, m 183-184°(dec). *N*-Bromosuccinimide (30g) is purified by dissolving rapidly in boiling water (300ml) and filtering through a fluted filter paper into a flask immersed in an ice bath, and left for 2 hours. The crystals are filtered off, washed thoroughly with ice-cold water (ca 100ml) and drained on a Büchner funnel before drying under vacuum over P_2O_5 or $CaCl_2$ [Dauben & McCoy *J Am Chem Soc* **81** 4863 1959, DOI: 10.1021/ja01527a027.] This *brominating agent* has also been recrystallised from acetic acid or water (10 parts see above), washed in water and dried extensively (i.e. to constant weight) *in vacuo* [Wilcox et al. *J Am Chem Soc* **108** 7693 1986, DOI: 10.1021/ja00284a038; Skell et al. *J Am Chem Soc* **108** 121 1986, DOI: 10.1021/ja00261a020; *Beilstein* **21/9** V 543.]

Bromotetronic acid (2-bromo-4-hydroxyacetoacetic lactone) [21151-51-9] $C_4H_3BrO_3$, M 179.0, m 182.8°, pK^{25} 2.23. Decolourise with Norit in $EtOAc$, evaporate, and crystallise from $EtOAc$ or $*C_6H_6$. [Schuler et al. *J Phys Chem* **78** 1063 1974, DOI: 10.1021/j100604a004; Gillespie & Price *J Org Chem* **22** 780 1957, DOI: 10.1021/jo01358a016; *Beilstein* **17** III/IV 5819.]

Bromotrichloromethane [75-62-7] $CBrCl_3$, M 198.3, f -5.6°, m 21°, b 104.1°/atm, d_D^{20} 2.01, n_D^{20} 1.5061. Wash it with aqueous $NaOH$ solution or dilute Na_2CO_3 , then with H_2O , and dry with $CaCl_2$, BaO , $MgSO_4$ or P_2O_5 before distilling in diffuse light and storing in the dark. It has also been purified by treatment with charcoal and fractional crystallisation by partial freezing. It is purified also by vigorous stirring with portions of concentrated H_2SO_4 until the acid does not discolour during several hours stirring. Wash with Na_2CO_3 and water, dry with $CaCl_2$ and then illuminate it with a 1000W projection lamp at 15cm for 10 hours, after making it 0.01M in bromine. Pass it through a 30 x 1.5cm column of activated alumina and fractionally redistil it through a 12-in Vigreux column. [Firestone & Willard *J Am Chem Soc* **83** 3551 1961, DOI: 10.1021/ja01478a002; see also Cadogan & Duell *J Chem Soc* 4154 1962, DOI: 10.1039/JR9620004154; *Beilstein* **1** IV 77.]

1-Bromo-2,2,2-trifluoroethane [421-06-7] $C_2H_2BrF_3$, M 163.0, m -94°, b 26-27°/atm, d_4^{20} 1.788, n_D^{20} 1.332. Wash it with water, dry ($CaCl_2$) and distil it. [*Beilstein* **1** III 179, **1** IV 154.]

Bromotrifluoromethane (Freon 13B1) [75-63-8] $CBrF_3$, M 148.9, b -59°/atm, d_4^{20} 1.590. Purify the gas by passing it through a tube containing P_2O_5 on glass wool into a vacuum system where it is frozen out in a quartz

tube and degassed by cycles of freezing, evacuating and thawing. [Beilstein 1 III 83, 1 IV 73.]

2-Bromovaleric (α -bromopentanoic) acid [584-93-0] $\text{C}_5\text{H}_9\text{BrO}_2$, M 181.0, b 132-136°/25mm, d_4^{20} 1.381, $pK_{\text{Est}} \sim 3.2$. Purify it by repeated fractional distillation and use the middle fraction. [Beilstein 2 II 268, 2 III 680, 2 IV 883.] The *amide* has m 83.4-84° [Stevens & Holland *J Org Chem* 18 1112 1953, DOI: 10.1021/jo50015a007].

5-Bromovaleric (γ -bromopentanoic) acid [2067-33-6] $\text{C}_5\text{H}_9\text{BrO}_2$, M 181.0, m 40°, $pK_{\text{Est}} \sim 4.6$. Crystallise the acid from petroleum ether. [Beilstein 2 IV 883.]

1,3-Butadiene [106-99-0] C_4H_6 , M 54.1, b -2.6°/atm. Dry the gas by condensing it into a solution of triethylaluminium in decahydronaphthalene; then it is flash distilled. It has also been dried by passage over anhydrous CaCl_2 or distilled from NaBH_4 . Also purify by passing through a column packed with molecular sieves (4A), followed by cooling in a Dry-ice/MeOH bath overnight, filtering off the ice and drying over CaH_2 at -78°, then distilling in a vacuum line. [Beilstein 1 IV 976.]

n-Butane [106-97-8] C_4H_{10} , M 58.1, m -135°, b -0.5°/atm. Dry by passing over anhydrous $\text{Mg}(\text{ClO}_4)_2$ and molecular sieves type 4A. Air is removed by prolonged and frequent degassing at -107°. [Beilstein 1 IV 236.]

1,4-Butanediol (tetramethylene glycol) [110-63-4] $\text{C}_4\text{H}_{10}\text{O}_2$, M 90.1, f 20.4°, b 107-108°/4mm, 127°/20mm, d_4^{20} 1.02, n_D^{20} 1.4467. Distil the glycol, and store it over Linde type 4A molecular sieves, or crystallise it twice from anhydrous diethyl ether/acetone, and redistil it. It has been recrystallised from the melt and doubly distilled *in vacuo* in the presence of Na_2SO_4 . [Beilstein 1 IV 2515.]

erythro-2,3-Butanediol (meso-2,3-butylene glycol) [5341-95-7] $\text{C}_4\text{H}_{10}\text{O}_2$, M 90.1, m 32-34°, 34.4°, b 89°/16mm, 181.7°/742mm, d_4^{20} 0.9939, n_D^{20} 1.443, n_D^{25} 1.4324. The *meso*-form is prepared from *trans*-2,3-epoxybutane and is recrystallised from isopropyl ether at low temperature. [Wilson & Lucas *J Am Chem Soc* 58 2396 1936, DOI: 10.1021/ja01303a010; Beilstein 1 II 546, 1 III 2178, 1 IV 2524.] **threo-2,3-Butanediol (rac \pm -2,3-butylene glycol)** [513-85-9] $\text{C}_4\text{H}_{10}\text{O}_2$, M 90.1, m 7.6°, b 86°/16mm, 172.7°/742mm, 183-184°/760mm, d_4^{25} 0.995, n_D^{20} 1.443, n_D^{25} 1.4310, is prepared from *cis*-2,3-epoxybutane and is recrystallised from isopropyl ether at low temperature. [Beilstein 1 II 546, 1 III 2180, 1 IV 2524.]

threo-2,3-Butanediol [*R,R*(-) 24347-58-8, *S,S*(+) 19132-06-0] $\text{C}_4\text{H}_{10}\text{O}_2$, M 90.1, m 16-19°, 19.7°, b 77.5-78°/10mm, 179-180°/atm, d_4^{25} 0.987, n_D^{20} 1.443, n_D^{25} 1.4310, $[\alpha]_D^{20}$ (-) or (+) 13.1 (neat). Purify by fractional distillation. The *bis*-(*p*-nitrobenzoate) ester has m 141-142°, and $[\alpha]_D^{20}$ (-) or (+) 52 (c 4, CHCl_3). [Ghirardelli & Lucas *J Am Chem Soc* 79 734 1957, DOI: 10.1021/ja01560a064; Rubin et al. *J Am Chem Soc* 74 425 1952, DOI: 10.1021/ja01122a043; Neish *Can J Res* 27 6 1949, DOI: 10.1139/cjr49b-002; Neish & Ledingham *Can J Res* 27 694 1949, DOI: 10.1139/cjr49b-070; Beilstein 1 IV 2524-2525.] When (-)-2*R,3R*-butane-2,3-diol (22g, 21ml, 244.5mmol) is added slowly to a stirred mixture of toluene-*p*-sulfonyl chloride (100g, 525mmol) in dry pyridine at 0° (ice-water bath), kept thus for 20 minutes, and the semi-solid mixture is set aside at ~25° overnight, then is shaken vigorously with crushed ice-water for 2 hours, poured rapidly with stirring into a mixture of concentrated HCl (70ml) and crushed ice, and the slurry is filtered, it provides solid (+)-2*R,3R*-butane-2,3-diol bis(tosylate) after washing thoroughly with H_2O and drying (91.0g, 93.5%). Alternatively, the crude dry di-ester is dissolved in CH_2Cl_2 , dried (MgSO_4), evaporated *in vacuo*, and the residue is washed with petroleum ether (b 20-60°), and stored over solid KOH in a desiccator. It has m 62-64° (65.1-65.5°), $[\alpha]_D^{20}$ +37.2 (c 2.105, CHCl_3). [Corey & Mitra *J Am Chem Soc* 84 2938 1962, DOI: 10.1021/ja00874a019, Lucas et al. *J Am Chem Soc* 72 2138 1950, DOI: 10.1021/ja01161a076; Fryzuk & Bosnich *J Am Chem Soc* 99 6262 1977, DOI: 10.1021/ja00461a014]. This (+)-2*R,3R*-ditosylate provides the chiral ligand (-)-*S,S*-CHIRAPHOS [cf. 64896-28-2] after reaction with 2 mols of Ph_2PLi , *via* inversion of configuration at the two chiral centres. (+)-2*R,3R*-butane-2,3-diol bis(methanesulfonate) has m 123-125°, $[\alpha]_D^{22}$ +2 (c 1, CHCl_3). [Beilstein 1 I 250, 1 II 547, 1 III 2181-3, 1 IV 2525.]

1-Butanesulfonyl chloride [2386-60-9] $\text{C}_4\text{H}_9\text{ClO}_2\text{S}$, M 156.6, b 75-76°/7mm, 98°/13mm, 100-103°/27-28mm, d_4^{20} 1.2078, n_D^{20} 1.4559. It has a pungent odour and is LACHRYMATORY. If IR shows OH bands,

then dissolve in Et₂O, wash with cold saturated aqueous NaHCO₃ (care since CO₂ will be generated) then H₂O, dry it over solid Na₂SO₄, filter, evaporate and distil the residue twice. Characterise it by shaking a solution in Et₂O or *C₆H₆ with aqueous NH₃, collect the solid **1-butanefulfonamide** with **m 48°** after recrystallisation from CHCl₃, CCl₄ or Et₂O/petroleum ether. [Douglass & Johnson *J Am Chem Soc* **60** 1486 1938, DOI: 10.1021/ja01273a063; Lee & Dougherty *J Org Chem* **5** 81 1940, DOI: 10.1021/jo01208a001; *Beilstein* **4** IV 45.]

1-Butanethiol [109-79-5] C₄H₁₀S, **M 90.2, b 98.4°/atm, d₄²⁵ 0.837, n_D²⁰ 1.443, n_D²⁵ 1.440, pK_{Est} ~11.3**. Dry the thiol with CaSO₄ or Na₂SO₄, then reflux it over magnesium, or dry with, and distil it from CaO, under nitrogen [Roberts & Friend *J Am Chem Soc* **108** 7204 1986, DOI: 10.1021/ja01273a063]. It has been separated from hydrocarbons by extractive distillation with aniline.

Dissolve it also in 20% NaOH, extract with a small amount of *C₆H₆, then steam distil it until the distillate is clear. The distillate is then cooled and acidified slightly with 15% H₂SO₄. The thiol is distilled out, dried with CaSO₄ or CaCl₂, and fractionally distilled under N₂ [Mathias & Filho *J Phys Chem* **62** 1427 1958, DOI: 10.1021/j150569a018]. It has also been purified by precipitation as the lead mercaptide from alcoholic solution, then regeneration by addition of dilute HCl to the residue followed by steam distillation. *All operations should be carried out in a fume cupboard due to the TOXICITY and obnoxious odour of the thiol.* [*Beilstein* **1** IV 1555.]

2-Butanethiol [513-53-1] C₄H₁₀S, **M 90.2, b 37.4°/134mm, d₄²⁵ 0.846, n_D²⁵ 1.4338, pK_{Est} ~11.4**. Purify it as for 1-butanethiol above. [*Beilstein* **1** IV 1584.]

n-Butanol [71-36-3] C₄H₁₀O, **M 74.1, b 117.7°/760mm, d₄²⁵ 0.80572, n_D²⁰ 1.39922, n_D¹⁵ 1.40118**. Dry it with MgSO₄, CaO, K₂CO₃, or solid NaOH, followed by refluxing with, and distilling from, small amounts of calcium, magnesium activated with iodine, or aluminium amalgam. It can also be dried with molecular sieves (4A), or by refluxing with *n*-butyl phthalate or succinate. (For method, see *Ethanol*.) *n*-Butanol can also be dried by efficient fractional distillation, water passing over in the first fraction as a binary azeotrope (contains about 37% water). An ultraviolet-transparent distillate has been obtained by drying with magnesium and distilling from sulfanilic acid. To remove bases, aldehydes and ketones, the alcohol is washed with dilute H₂SO₄, then NaHSO₄ solution; esters are removed by boiling for 1.5 hours with 10% NaOH.

It has also been purified by adding NaBH₄ (2g) to butanol (1.5L), under gentle bubbling of argon, and a reflux condenser for 1 day at 50°. Then add freshly cut sodium (2g, washed with butanol) and refluxed for 1 day. Distil, and collect the middle fraction [Jou & Freeman *J Phys Chem* **81** 909 1977, DOI: 10.1021/j100524a021]. [*Beilstein* **1** IV 1506.]

2-Butanone (methyl ethyl ketone, MEK) [78-93-0] C₄H₈O, **M 72.1, b 79.6°/atm, d₄²⁰ 0.853, n_D²⁰ 1.37850, n_D²⁵ 1.37612, pK₂₅ -7.2 (aqueous H₂SO₄)**. In general, purification methods are the same as for acetone. Aldehydes can be removed by refluxing with KMnO₄ + CaO, until the Schiff aldehyde test is negative, prior to distillation. Shaking with saturated K₂CO₃, or passing through a small column of activated alumina, removes cyclic impurities. The ketone can be dried by careful distillation (an azeotrope containing 11% water boils at 73.4°), or over CaSO₄, P₂O₅, Na₂SO₄, or K₂CO₃, followed by fractional distillation. Purification as the bisulfite addition compound is achieved by shaking with excess saturated Na₂SO₃, cooled to 0°, filtering off the precipitate, washing with a little ethyl ether and drying in air; this is followed by decomposition with a slight excess of Na₂CO₃ solution and steam distillation, the distillate being saturated with K₂CO₃ so that the ketone can be separated, dried with K₂CO₃, filtered, and distilled. Purification as the *NaI addition compound* (**m 73-74°**) is more convenient. (For details, see *Acetone*.) Small quantities of 2-butanone can be purified by conversion to the semicarbazone, recrystallisation to constant melting point, drying it under vacuum over CaCl₂ and paraffin wax, refluxing for 30 minutes with excess oxalic acid, followed by steam distillation, salting out, drying and distilling [Cowan et al. *J Chem Soc* 171 1940, DOI: 10.1039/JR9400000171]. [*Beilstein* **1** IV 3243.]

1-Butene [106-98-9] C₄H₈, **M 56.1, m -185°, b -81°/10mm, -6.3°/atm, d₄^{-6.47} 0.6255**. Present in petroleum oils and coal gas. The gas is dried with CaH₂ and purified by gas chromatography. *Alternatively*, the gas (from Linde Air Products Company, USA) is cooled with liquid nitrogen, evacuated until about one tenth of the sample has evaporated, distilled into a second receiver and the last tenth is discarded. This operation is repeated seven times. During the first three cycles decreasingly small amounts of a white solid are observed when the sample is

cooled, but they disappear by the fourth cycle. This butane was used for the studies of vapour pressure, orthobaric liquid density and critical constants. [Calingaert *J Am Chem Soc* **45** 130 1923, DOI: 10.1021/ja01654a019; Coffin & Maas *J Am Chem Soc* **50** 1427 1928, DOI: 10.1021/ja01392a028; Beattie & Marple *J Am Chem Soc* **72** 1449 1950, DOI: 10.1021/ja01160a006.] [Beilstein **1** H 205, **1** II 176, **1** III 728, **1** IV 765.] **HIGHLY FLAMMABLE GAS.**

cis-2-Butene [590-18-1] C_4H_8 , **M 56.1, b 2.95-3.05°/746mm.** The gas is dried with CaH_2 and purified by gas chromatography. [Beilstein **1** H 205, **1** II 176, **1** III 728, **1** IV 778.] **HIGHLY FLAMMABLE.** **trans-2-Butene** [624-64-6] **b 0.3-0.4°/744mm,** is also a gas which is purified as the *cis*-isomer. [Beilstein **1** H 205, **1** II 176, **1** III 730, **1** IV 781.] **HIGHLY FLAMMABLE.**

2-Butene-1,4-dicarboxylic acid (trans-3-hexenedioic acid, trans-β-hydromuconic acid) [4436-74-2] $C_4H_8O_4$, **M 144.1, m 194-197°, 195-196°, $pK_{Est(1)} \sim 4.2$, $pK_{Est(2)} \sim 5.00$.** Crystallise the acid from boiling water, then dry it at 50-60° in a vacuum oven. [Beilstein **2** IV 2237.]

2-Butoxyethanol (butyl cellosolve) [111-76-2] $C_6H_{14}O_2$, **M 118.2, b 171°/745mm, d_4^{20} 0.903, n_D^{20} 1.4191.** Peroxides can be removed by refluxing with anhydrous $SnCl_2$ or by passage under slight pressure through a column of activated alumina. Dry with anhydrous K_2CO_3 and $CaSO_4$, filter and distil, or reflux with, and distil from NaOH. [Beilstein **1** IV 2380.]

n-Butyl acetate [123-86-4] $C_6H_{12}O_2$, **M 116.2, b 126.1°/atm, d_4^{20} 0.882, n_D^{20} 1.394.** Distil, reflux with successive portions of $KMnO_4$ until the colour persists, dry with anhydrous $CaSO_4$, filter and redistil. [Beilstein **2** IV 143.]

tert-Butyl acetate [540-88-5] $C_6H_{12}O_2$, **M 116.2, b 97-98°/atm, d_4^{20} 0.866, n_D^{20} 1.387.** Wash the ester with 5% Na_2CO_3 solution, then saturated aqueous $CaCl_2$, dry with $CaSO_4$ and distil it. [McCloskey et al. *Org Synth Coll Vol* **4** 261 1963, DOI: 10.15227/orgsyn.034.0026; Mangia et al. *Org Prep Proc Int* **18** 13 1986, DOI:10.1080/00304948609356822; Beilstein **2** IV 151.]

tert-Butyl acetoacetate [1694-31-1] $C_8H_{14}O_3$, **M 158.2, b 71°/10mm, 85°/20mm, d_4^{20} 0.954, n_D^{20} 1.42.** Distil it under reduced pressure through a short column. [Lawesson et al. *Org Synth Coll Vol* **5** 155 1973, DOI: 10.15227/orgsyn.042.0028; Beilstein **3** IV 1536.] **HARMFUL VAPOUR.**

tert-Butylacetyl chloride [7065-46-5] $C_6H_{11}ClO$, **M 134.6, b 68-71°/100mm, 81°/180mm, 128-132°/atm, d_4^{20} 0.964, n_D^{20} 1.423.** Distil it under vacuum. If IR shows OH group, then treat with thionyl chloride or oxalyl chloride at *ca* 50° for 30 minutes, evaporate and fractionate the residue using a short column. Strongly **LACHRYMATORY**, use a good fume hood. [Berliner & Berliner *J Am Chem Soc* **72** 222 1950, DOI: 10.1021/ja01157a063; Traynham & Battiste *J Org Chem* **22** 1551 1957, DOI: 10.1021/jo01363a004; Beilstein **2** IV 956.]

Butyl acrylate [141-32-2] $C_7H_{12}O_2$, **M 128.2, b 59°/25mm, d_4^{20} 0.894, n_D^{25} 1.4254.** Wash it repeatedly with aqueous NaOH to remove inhibitors such as hydroquinone, then with distilled water. Dry with $CaCl_2$. Fractionally distil under reduced pressure in an all-glass apparatus. The middle fraction is sealed under N_2 and stored at 0° in the dark until required or with a stabiliser [Mallik & Das *J Am Chem Soc* **82** 4269 1960, DOI: 10.1021/ja01501a038]. [Beilstein **2** IV 1463.]

tert-Butyl acrylate [1663-39-4] $C_7H_{12}O_2$, **M 128.2, b 30.0-30.8°/26mm, 61-63°/15mm, 117-120°/760mm, d^{25} 0.875, n_D^{20} 1.410.** Purify the ester by fractional distillation. If it contains acid (OH bands in the IR), then dissolve it in Et_2O , wash it with aqueous $NaHCO_3$, dry the organic layer (Na_2SO_4), filter it and distil it under reduced pressure. Stabilise it by adding hydroquinone monomethyl ether (~0.05%). It forms a crystalline *tert*-butyl acrylate polymer which is soluble in organic solvents [Garrett et al. *J Am Chem Soc* **81** 1007 1959, DOI: 10.1021/ja01513a064]. [Beilstein **2** IV 1465.] For other alkyl acrylates see Rehberg *Org Synth Coll Vol* **3** 146 1955, DOI: 10.15227/orgsyn.026.0018.

(±)-*sec*-Butyl alcohol (± 2-butanol) [78-92-2; 15892-23-6] $C_4H_{10}O$, M 74.1, b 99.4°/atm, d_4^{20} 0.808. Purification methods are the same as for *n*-Butanol. These include drying with K_2CO_3 or $CaSO_4$, followed by filtration and fractional distillation, refluxing with CaO , distillation, then refluxing with magnesium and redistillation, and refluxing with, then distilling from CaH_2 . Calcium carbide has also been used as a drying agent. The anhydrous alcohol is obtained by refluxing with *sec*-butyl phthalate or succinate. (For method see *Ethanol*.) Small amounts of alcohol can be purified *via* conversion to the alkyl hydrogen phthalate and recrystallisation [Hargreaves *J Chem Soc* 3674, 3679(note page) 1956, DOI: 10.1039/JR9560003674]. For purification of optical isomers, see Timmermans and Martin [*J Chim Phys. Phys-Chim Biol* 25 411 1928]. [Beilstein 2 III 1566.]

tert-Butyl alcohol [75-65-0] $C_4H_{10}O$, M 74.1, m 23-25°, 25.7°, b 28.3°/60mm, 43.3°/123.8mm, 61.8°/315mm, 72.5°/507mm, 82.45°/760mm, d_4^{20} 0.7858, n_D^{20} 1.3878. It is synthesised commercially by the hydration of 2-methylpropene in dilute H_2SO_4 . Dry it with CaO , K_2CO_3 , $CaSO_4$ or $MgSO_4$, filter and fractionally distil it. Dry further by refluxing with, and distilling from, either magnesium activated with iodine, or small amounts of calcium, sodium or potassium, under nitrogen. Passage through a column of type 4A molecular sieves is another effective method of drying; as well as refluxing with *tert*-butyl phthalate or succinate. (For method see *Ethanol*.) Other methods include refluxing with excess aluminium *tert*-butylate, or standing over CaH_2 , and distilling as needed. Further purification is achieved by fractional crystallisation by partial freezing, taking care to exclude moisture. *tert*-Butyl alcohol samples containing much water can be dried by adding *benzene, so that the water distils off as a tertiary azeotrope, b 67.3°. Traces of isobutylene have been removed from dry *tert*-butyl alcohol by bubbling dry pre-purified nitrogen through for several hours at 40-50° before using. It forms azeotropic mixtures with a large number of compounds. It has also been purified by distillation from CaH_2 onto Linde 4A molecular sieves which had been activated at 350° for 24 hours [Jaeger et al. *J Am Chem Soc* 101 717 1979, DOI: 10.1021/ja00497a039]. [Beilstein 1 IV 1609.]

Rapid purification: Dry *tert*-butanol over CaH_2 (5% w/v), distil and store it over 3A molecular sieves.

n-Butylamine [109-73-9] $C_4H_{11}N$, M 73.1, b 77.8°/atm, d_4^{20} 0.740, n_D^{20} 1.4009, n_D^{25} 1.399, pK^{25} 10.66. Dry *n*-butylamine with solid KOH , K_2CO_3 , $LiAlH_4$, CaH_2 or $MgSO_4$, then reflux it with, and fractionally distil it from P_2O_5 , CaH_2 , CaO or BaO . Further purification is by precipitation as the *hydrochloride*, m 213-213.5°, from ethereal solution by bubbling HCl gas into it. This is re-precipitated three times from $EtOH$ by adding ether, followed by liberation of the free amine using excess strong mineral base. The amine is extracted into ether, which is separated, dried with solid KOH , the ether is removed by evaporation and then the amine is distilled. It is stored in a desiccator over solid $NaOH$ [Bunnett & Davis *J Am Chem Soc* 82 665 1960, DOI: 10.1021/ja01488a043; Lycan et al. *Org Synth Coll Vol* 2 318 1943, DOI: 10.1522/orgsyn.011.0058]. [Beilstein 4 IV 540.] **SKIN IRRITANT.**

R-(*-*)-*sec*-Butylamine [13250-12-9] $C_4H_{11}N$, M 73.1, b 61-63°/atm, 62.5°/atm, d_4^{20} 0.731, n_D^{20} 1.393, $[\alpha]_D^{20}$ -7.5 (neat), pK^{25} 10.56. Dry it over solid $NaOH$ overnight and fractionate it through a short helices packed column. The *L*-hydrogen tartrate salt has m 139-140° (from H_2O), the $1H_2O$ has m 96° $[\alpha]_D^{21}$ +18.1 (c 11, H_2O), the *hydrochloride* has m 152° $[\alpha]_D^{21}$ -1.1 (c 13, H_2O) and the *benzoyl* derivative crystallises from $EtOH$ as needles with m 97°, $[\alpha]_D^{21}$ -34.9 (c 11, H_2O). [Bruck et al. *J Chem Soc* 921 1956, DOI: 10.1039/JR9560000921; Kjaer & Hansen *Acta Chem Scand* 11 898 1957, DOI: 10.3891/acta.chem.scand.11-0898.] [Beilstein 4 H 161, 4 I 372, 4 III 308, 4 IV 617.] The *S*-(+)-*enantiomer* has the same properties except for the optical rotation which has the opposite sign.

tert-Butylamine [75-64-9] $C_4H_{11}N$, M 73.1, b 42°/atm, d_4^{20} 0.696, pK^{25} 10.68. Dry it with KOH or $LiAlH_4$, and/or distil it from CaH_2 or BaO . [Beilstein 4 IV 657.] *tert*-Butylammonium bromide [60469-70-7] $C_4H_{11}N$. HBr , M 154.1, m >250°(dec), is recrystallised several times from absolute $EtOH$ and thoroughly dried at 105° *in vacuo*. **Toxic vapours.** [Beilstein 4 IV 657.]

n-Butyl bromide [109-65-9] C_4H_9Br , M 137.0, b 101-102°/atm, d_4^{25} 1.2678, n_D^{20} 1.4399, n_D^{25} 1.4374. Wash the bromide with concentrated H_2SO_4 , water, 10% Na_2CO_3 and again with water. Dry it over $CaCl_2$, $CaSO_4$ or K_2CO_3 , and distil it. Redistil it after drying with P_2O_5 , or pass it through two columns containing 5:1 silica gel/Celite mixture and store it with freshly activated alumina. [Beilstein 1 IV 258.]

tert-Butyl bromoacetate [5292-43-3] $C_6H_{11}BrO_2$, M 195.1, b 52°/10mm, 74-76°/25mm, d_4^{20} 1.324, n_D^{25} 1.4162. Dissolve the ester in Et_2O , wash it well with ice cold 10% aqueous K_2CO_3 , dry it over $CaCl_2$, filter and evaporate the Et_2O , then fractionate it through a Vigreux column in a vacuum. **LACHRYMATORY**. [Abramovitch et al. *J Am Chem Soc* **64** 2272 1942, DOI: 10.1021/ja01262a01; Abramovitch & Hauser *J Am Chem Soc* **65** 986 1943, DOI: 10.1021/ja01245a501; *Beilstein* **2** III 482.]

tert-Butyl carbazate [870-46-2] $C_5H_{12}NO_2$, M 132.2, m 41-42°, b 64°/0.01mm, 55-57°/0.4mm. Distil it in a Claisen flask with a water or oil bath at *ca* 80°. After a couple of drops have distilled, the carbazate is collected as an oil which solidifies to a snow white solid. It can be crystallised with 90% recovery from a 1:1 mixture of petroleum ether (b 30-60°) and petroleum ether (b 60-70°). [Carpino et al. *Org Synth Coll Vol* **5** 166 1973, DOI: 10.15227/orgsyn.044.0020; *Beilstein* **3** IV 175.]

n-Butyl chloride [109-69-3] C_4H_9Cl , M 92.6, b 78°/atm, d_4^{20} 0.886, n_D^{25} 1.4021. Shake it repeatedly with concentrated H_2SO_4 (until no further colour develops in the acid layer), then wash it with water, aqueous $NaHCO_3$ or Na_2CO_3 , and more water. Dry it with $CaCl_2$, or $MgSO_4$ (then with P_2O_5 if desired), decant and fractionally distil it. *Alternatively*, a stream of oxygen continuing *ca* three times as long as is necessary to obtain the first coloration of starch iodide paper by the exit gas. After washing with $NaHCO_3$ solution to hydrolyse ozonides and to remove the resulting organic acid, the liquid is dried and distilled [Chien & Willard *J Am Chem Soc* **75** 6160 1953, DOI: 10.1021/ja01120a016]. [*Beilstein* **1** IV 246.]

tert-Butyl chloride [507-20-0] C_4H_9Cl , M 92.6, f -24.6°, b 50.4°/atm, d_4^{20} 0.851, n_D^{25} 1.38564. Purification methods commonly used for other alkyl halides lead to decomposition. Some impurities can be removed by photochlorination with a small amount of chlorine prior to use. The liquid is washed with ice water, dried with $CaCl_2$ or $CaCl_2 + CaO$ and fractionally distilled. It has been further purified by repeated fractional crystallisation by partial freezing. [*Beilstein* **1** IV 288.]

tert-Butyl chloroacetate [107-59-5] $C_6H_{11}ClO_2$, M 150.6, b 48-49°/11mm, 60.2°/15mm, 155°/atm (dec), d_4^{25} 1.4204, n_D^{20} 1.4259. Check the NMR spectrum; if satisfactory then distil in a vacuum; if not then dissolve in Et_2O , wash with H_2O , 10% H_2SO_4 until the acid extract does not become cloudy when made alkaline with $NaOH$. Wash the organic layer again with H_2O , then saturated aqueous $NaHCO_3$, dry over Na_2SO_4 , evaporate and fractionate it through a carborundum-packed column or a 6-inch Widmer column (*see tert-butyl ethyl malonate for precautions to avoid decomposition during distillation*). [Johnson et al. *J Am Chem Soc* **75** 4995 1953, DOI: 10.1021/ja01116a033; Baker *Org Synth Coll Vol* **3** 141 1944, DOI: 10.15227/orgsyn.024.0018; *Beilstein* **2** III 444.]

tert-Butyl cyanide (trimethylacetoneitrile, pivalonitrile) [630-18-2] C_5H_9N , M 83.1, m 16-18°, b 104-106°/atm, d_4^{20} 0.765. Purify it by a two-stage vacuum distillation and de-gas by the freeze-pump-thaw technique. Store it under vacuum at 0°. **TOXIC**, use an efficient fume hood. [*Beilstein* **2** IV 875.]

tert-Butyl cyanoacetate [1116-98-9] $C_7H_{11}NO_2$, M 141.2, b 40-42°/0.1mm, 54-56°/0.3mm, 90°/10mm, 107-108°/23mm, d_4^{20} 0.989, n_D^{20} 1.4198. The IR spectrum of a film should have bands at 1742 (ester CO) and 2273 ($C\equiv N$), but no band at *ca* 3500 broad (OH) cm^{-1} . If it does not have the last-named band, then fractionally distil; otherwise dissolve in Et_2O , wash with saturated aqueous $NaHCO_3$, dry over K_2CO_3 , evaporate Et_2O , and distil the residue under a vacuum (*see tert-butyl ethyl malonate for precautions to avoid decomposition during distillation*). [Beech & Piggott *J Chem Soc* 423 1955, DOI: 10.1039/JR9550000423; Dahn & Hauth *Helv Chim Acta* **42** 1214 1959, DOI: 10.1002/hlca.19590420416; *Beilstein* **2** I 255.]

tert-Butyl diazoacetate [3505950-8] $C_6H_{10}N_2O_2$, M 142.2, b 51-53°/12mm, d_4^{25} 1.026, n_D^{20} 1.443. It is a poisonous orange-yellow liquid which is explosive, and the necessary precautions should be strictly adhered to (i.e. efficient fume cupboard, and face and body protection; see reference below). Check the purity by TLC on Merck Kieselgel F₂₅₄ or Eastman Kodak Silica Gel without indicator using $CHCl_3$ as eluent (R_F 0.72). If the ester is suspect, then dissolve it in Et_2O , wash it with brine, H_2O , dry the organic layer (Na_2SO_4), and filter. Remove most of the Et_2O at 30°/water pump vacuum (rotovap), the remaining ether during distillation under lower pressure (<12mm), and the residual oil then distils as a coloured liquid using a water bath at ~50° (no higher) as

heat source. [Regitz et al. *Org Synth Coll Vol* **5** 179 1972, DOI: 10.15227/orgsyn.048.0036.]

***n*-Butyl disulfide** [629-45-8] $C_8H_{18}S_2$, M 178.4, b 110-113°/15mm, d_4^{20} 0.938, n_D^{22} 1.494. Shake it with lead peroxide, filter and distil it in a vacuum under N_2 . [Beilstein **1** IV 1560.]

***n*-Butyl ether (di-*n*-butyl ether)** [142-96-1] $C_8H_{18}O$, M 130.2, b 52-53°/26mm, 142.0°/760mm, d_4^{20} 0.764, n_D^{20} 1.39925, n_D^{25} 1.39685, pK^{25} -5.40 (aqueous H_2SO_4). Peroxides (detected by the liberation of iodine from weakly acid HCl solutions of 2% KI) can be removed by shaking 1L of ether with 5-10ml of a solution comprising of ferrous sulfate (6.0g) in concentrated H_2SO_4 (6ml) and water (110ml), with aqueous Na_2SO_3 , or with acidified NaI, water, then aqueous $Na_2S_2O_3$. After washing with dilute NaOH, KOH, or Na_2CO_3 , then water, the ether is dried with $CaCl_2$ and distilled. It can be further dried by distillation from CaH_2 or Na (after drying with P_2O_5), and stored in the dark with Na or NaH. The ether can also be purified by treating with CS_2 and NaOH, expelling the excess sulfide by heating. The ether is then washed with water, dried with NaOH and distilled [Kusama & Koike *J Chem Soc Jpn, Pure Chem Sect* **72** 229 1951]. Other purification procedures include passage through an activated alumina column to remove peroxides, or through a column of silica gel, and distillation after adding about 3% (v/v) of a 1M solution of MeMgI in *n*-butyl ether. [Beilstein **1** IV 1520.]

***n*-Butyl ethyl ether** [628-81-9] $C_6H_{14}O$, M 102.2, b 92.7°/atm, d_4^{20} 0.751, n_D^{20} 1.38175, n_D^{25} 1.3800. Purify by drying with $CaSO_4$, by passage through a column of activated alumina (to remove peroxides), followed by prolonged refluxing with Na and then fractional distillation. [Beilstein **4** IV 1518.]

***tert*-Butyl ethyl ether** [637-92-3] $C_6H_{14}O$, M 102.2, b 71-72°/atm, d_4^{20} 0.741. Dry the ether with $CaSO_4$, pass it through an alumina column, and fractionally distil it. [Beilstein **1** IV 1618.]

***tert*-Butyl ethyl malonate** [32864-38-3] $C_9H_{16}O_4$, M 188.2, b 83-85°/8mm, 93-95°/17mm, 107-109°/24mm, d_4^{25} 0.994, n_D^{24} 1.4150. A likely impurity is monoethyl malonate; check IR for OH bands at 3330 cm^{-1} . To ca 50g of ester add ice cold NaOH (50g in 200ml of H_2O and 200g of ice). Swirl a few times (filter off ice if necessary), place it in a separating funnel and extract with Et_2O (2 x 75ml). Dry the extract ($MgSO_4$) (since traces of acid decompose the *t*-Bu group of the ester, the distillation flask has to be washed first with aqueous NaOH, rinsed with H_2O and allowed to dry). Addition of some K_2CO_3 or MgO before distilling is recommended to inhibit decomposition. Distil it under reduced pressure through a 10cm Vigreux column. *Decomposition is evidenced by severe foaming due to autocatalytic decomposition and cannot be prevented from accelerating except by stopping the distillation and rewashing the distillation flask with alkali again.* [Breslow et al. *J Am Chem Soc* **66** 1286 1944, DOI: 10.1021/ja01236a022; Hauser et al. *J Am Chem Soc* **64** 2714 1942, DOI: 10.1021/ja01263a054; Strube *Org Synth Coll Vol* **4** 417 1963, DOI: 10.15227/orgsyn.037.0034; Beilstein **2** IV 1884.]

***n*-Butyl formate** [592-84-7] $C_5H_{10}O_2$, M 102.1, b 106.6°/atm, d_4^{20} 0.891, n_D^{20} 1.3890. Wash the formate with saturated $NaHCO_3$ solution in the presence of saturated NaCl, until no further reaction occurs, then with saturated NaCl solution, dry ($MgSO_4$), filter and fractionally distil the filtrate. [Beilstein **2** IV 28.]

Butyl glycolate [7397-62-8] $C_6H_{12}O_3$, M 132.2, b 191-192°/755mm, 187-190°/atm, d_4^{20} 1.019, n_D^{20} 1.4263. Dissolve the ester in $CHCl_3$ ($EtOH$ -free), wash with 5% $KHCO_3$ until effervescence ceases (if free acid is present), dry over $CaCl_2$, filter, evaporate and distil through a short column. [Bøhme & Opfer *Z Anal Chem* **139** 255 1953, Filachione et al. *J Am Chem Soc* **73** 5265 1951, DOI: 10.1021/ja01155a075; Beilstein **3** IV 589.]

***tert*-Butyl hydroperoxide (TBHP)** [75-91-2] $C_4H_{10}O_2$, M 90.1, f 5.4°, m 0.5-2.0°, b 38°/18mm, d_4^{20} 0.900, n_D^{20} 1.4013, pK^{20} 12.8. Care should be taken when handling this peroxide because of the possibility of EXPLOSION. It explodes when heated over an open flame. Work in an efficient fume cupboard, behind a thick plastic transparent shield, and with eye protection. Alcoholic and volatile impurities can be removed by prolonged refluxing at 40° under reduced pressure, or by steam distillation. For example, Bartlett, Benzing and Pincock [*J Am Chem Soc* **82** 1762 1960, DOI: 10.1021/ja01492a055] refluxed at 30mm pressure in an apparatus for azeotropic liquid separation until the two phases no longer separated, and then distilled at 41°/23mm. Pure material is stored under N_2 , in the dark at 0°. Crude commercial material has been added to

25% NaOH below 30°, and the crystals of the sodium salt have been collected, washed twice with *benzene and dissolved in distilled water. After the pH of the solution is adjusted to 7.5 by adding solid CO₂, the peroxide is extracted into petroleum ether, from which, after drying with K₂CO₃, TBHP is recovered by distilling off the solvent under reduced pressure at room temperature [O'Brien et al. *J Am Chem Soc* **79** 6238 1957, DOI: 10.1021/ja01580a034]. **The temperatures should be kept below 75°.** It has also been distilled through a helices packed column (*ca* 15 plates) and the material with **b 34-35°/20mm** is collected. Similarly, a solution in petroleum ether has been extracted with cold aqueous NaOH, and the hydroperoxide has been regenerated by adding at 0°, KHSO₄ to a pH not higher than 4.5, then extracted into diethyl ether, dried with MgSO₄, filtered and the ether evaporated in a rotary evaporator under reduced pressure at as low a temperature as possible [Milas & Djokic *J Am Chem Soc* **84** 3098 1962, DOI: 10.1021/ja00875a012].

A 3M solution of TBHP in CH₂Cl₂ is prepared by swirling of commercial TBHP (85ml, 0.61mol, 70% TBHP-30% H₂O, **d** 0.935 *ca* 7.2mmol/ml) with of CH₂Cl₂ (140ml) in a separating funnel. The milky mixture is allowed to stand until the phases separate (*ca* 30 minutes). The organic (lower) layer (*ca* 200ml) containing 0.60mole of TBHP is separated from the aqueous layer (*ca* 21ml) and used without further drying. TBHP is assayed by iodometric titration. With 90% grade TBHP (w/w, **d** 0.90, *ca* 9.0mmole/ml) no separation of layers occurs, i.e. when TBHP (66.67ml, 0.60mole) is added to CH₂Cl₂ (140ml) the resulting solution (*ca* 200ml) should be clear. [Walling & Buckler *J Am Chem Soc* **77** 6032 1955, DOI: 10.1021/ja01627a069; Rogers & Campbell *J Am Chem Soc* **74** 4742 1952, DOI: 10.1021/ja01139a004; Akashi et al. *J Org Chem* **43** 2063 1978 DOI: 10.1021/jo00404a052 state the quality of available grades, handling and compatibility for reactions, *Beilstein* **1** IV 1616.]

***n*-Butyl iodide (1-iodobutane)** [542-69-8] C₄H₉I, **M 184.0, b 130.4°/atm, d₄²⁰ 1.616, n_D²⁵ 1.44967.** Dry the iodide with MgSO₄ or P₂O₅, fractionally distil it through a column packed with glass helices, taking the middle fraction and storing over calcium or mercury in the dark. *Alternatively*, purify it by prior passage through activated alumina or by shaking with concentrated H₂SO₄ then washing with Na₂SO₃ solution. It has also been treated carefully with sodium to remove free HI and H₂O, before distilling through a column containing copper turnings at the top. Another purification procedure consisted of treatment with bromine, followed by extraction of free halogen with Na₂S₂O₃, washing with H₂O, drying and fractionally distilling. [*Beilstein* **1** IV 271.]

***tert*-Butyl iodide** [558-17-8] C₄H₉I, **M 184.0, m -38°, b 98.1°/760mm, 100°(dec)/atm, d₄²⁰ 1.544, d_D²⁵ 1.622g/cm³, n_D²⁰ 1.502.** Vacuum distillation has been used to obtain a distillate which remained colourless for several weeks at -5°. More extensive treatment has been used by Boggs, Thompson & Crain [*J Phys Chem* **61** 1625 1957, DOI: 10.1021/j150558a013] who washed it with aqueous NaHSO₃ solution to remove free iodine, dried this for 1 hour over Na₂SO₃ at 0°, and purified it by four or five successive partial freezings of the liquid to obtain colourless material, and was stored at -78° with Ag wool. [For resonance Raman spectra see Phillips et al. *J Phys Chem* **95** 9085 1991, DOI: 10.1021/j100176a014; for steric hindrance see Marcellin & Brooks *J Am Chem Soc* **97** 1710 1975, DOI: 10.1021/ja00840a015; *Beilstein* **1** IV 300.]

***tert*-Butyl isocyanate** [1609-86-5] C₅H₉NO, **M 99.1, m 10.5-11.5°, b 30.5-32°/10mm, 64°/52mm, d₂₅²⁵ 0.9079, n_D²⁵ 1.470.** It is **LACHRYMATORY** and **TOXIC**, and should have IR with ν_{\max} at 2251 (C≡N) cm⁻¹ but no OH bands. The NMR should have one band at 1.37 ppm from TMS. Purify it by fractional distillation under reduced pressure. [Stowell et al. *J Org Chem* **36** 3056 1971, DOI: 10.1021/jo00819a039; Curtius *J Prakt Chem* **125** 152 1930, *Beilstein* **4** IV 669.]

***tert*-Butyl isocyanide** [7188-38-7] C₅H₉N, **M 83.1, b 91-92°/730mm, 90°/758mm, d₄²⁰ 0.735.** Dissolve it in petroleum ether (b 40-60°), wash it with H₂O, dry (Na₂SO₄), filter, remove petroleum ether under slight vacuum, and distil it using a vacuum-jacketed Vigreux column at atmospheric pressure, its IR has a band at ν_{\max} 2134 cm⁻¹. [Ugi & Meyr *Chem Ber* **93** 239 1960, DOI: 10.1002/cber.19600930136; *Beilstein* **4** IV 661.] It has **toxic vapours**.

***tert*-butyl isocyanoacetate** [2769-72-4] C₇H₁₁NO, **M 141.2, b 50°/0.1mm, 49-50°/10mm, 63-65°/15mm, d₄²⁰ 0.970, n_D²⁰ 1.420.** If it contains some free acid (OH bands in IR), then dissolve it in Et₂O, shake with 20% Na₂CO₃, dry over anhydrous K₂CO₃, evaporate and distil it. [Ugi et al. *Chem Ber* **94** 2814 1961, DOI: 10.1002/cber.19610941033; Schöllkopf *Angew Chem* **89** 339 1977, DOI: 10.1002/anie.197703393.]

***n*-Butyl methacrylate** [97-88-1] $C_8H_{14}O_2$, M 142.2, b 49-52°/0.1mm, 163-165°/atm, d_4^{20} 0.896, n_D^{20} 1.424. Purify it as for butyl acrylate above, see [141-32-2]. ***tert*-Butyl methacrylate** [585-07-9], f -48°, b 135-136°/760mm, d_4^{20} 0.878, n_D^{20} 1.415, is similarly purified. [Beilstein 2 IV 1582.]

***n*-Butyl methyl ether** [628-28-4] $C_5H_{12}O$, M 88.2, b 70°, d_4^{20} 0.744, pK^{25} -3.50 (aqueous H_2SO_4). Dry it with $CaSO_4$, pass it through an alumina column to remove peroxides, and fractionally distil it. [Beilstein 1 IV 1518.]

***tert*-Butyl methyl ether (methyl *tert*-butyl ether, MTBE)** [1634-04-4] $C_5H_{12}O$, M 88.2, b 56°, n 1.369. Purify it as for *n*-butyl methyl ether. [Beilstein 1 IV 1615.]

***tert*-Butyl methyl ketone (3,3-dimethyl-2-butanone, pinacolone)** [75-97-8] $C_8H_{12}O$, M 100.2, b 105°/746mm, 106°/760mm, d_4^{20} 0.814, n_D^{20} 1.401. Reflux the ketone with a little $KMnO_4$. Dry it with $CaSO_4$ and distil it. [Beilstein 1 IV 3310.]

***tert*-Butyl nitrite** [540-80-7] $C_4H_9NO_2$, M 103.1, b 34°/250mm, 61-63°/atm, d_4^{20} 0.8671, n_D^{25} 1.3660. If it is free from OH bands (IR) then distil it through a 12inch helices packed column under reduced pressure, otherwise wash with aqueous 5% $NaHCO_3$ (effervescence), then H_2O , dry (Na_2SO_4) and fractionate it through a 10 theoretical plates column at ca 10mm pressure. [Allen *J Chem Soc* 1968 1954, DOI: 10.1039/JR9550000423; Coe & Doumani *J Am Chem Soc* 70 1516 1948, DOI: 10.1021/ja01184a065; UV: Ungnade & Smiley *J Org Chem* 21 993 1956, DOI: 10.1021/ja01184a065; IR: Tarte *Bull Soc Chim Belg* 60 240 1951, DOI: 10.1002/bscb.19510600307; Beilstein 1 IV 1622.]

***tert*-Butyl peracetate** [107-71-1] $C_6H_{12}O_3$, M 132.2, b 23-24°/0.5mm, n_D^{25} 1.4030. Wash the ester with $NaHCO_3$ from a *benzene solution, then redistil to remove *benzene [Kochi *J Am Chem Soc* 84 774 1962, DOI: 10.1021/ja00864a020]. Handle with adequate protection due to possible **EXPLOSIVE** nature. [Beilstein 2 IV 391.]

***tert*-Butylperoxy isobutyrate** [109-13-7] $C_8H_{16}O_3$, M 160.2, f -45.6°. After diluting the material (90ml) with petroleum ether (120ml), the mixture is cooled to 5° and shaken twice with 5% $NaOH$ solution (90ml portions, also at 5°). The non-aqueous layer, after washing once with cold water, is dried at 0° with a mixture of anhydrous $MgSO_4$ and $MgCO_3$ containing ca 40% MgO . After filtering, this material is passed, twice, through a column of silica gel at 0° (to remove *tert*-butyl hydroperoxide). The solution is then evaporated at 0°/0.5-1mm to remove the solvent, and the residue is recrystallised several times from petroleum ether at -60°, then subjected to high vacuum to remove traces of solvent [Milas & Golubovic *J Am Chem Soc* 80 5994 1958, DOI: 10.1021/ja01555a027]. Handle with adequate protection due to possible **EXPLOSIVE** nature.

Butyl stearate [123-95-5] $C_{22}H_{44}O_2$, M 340.6, m 26.3°, d_4^{20} 0.861. Acidic impurities are removed by shaking with 0.05M $NaOH$ or a 2% $NaHCO_3$ solution, followed by several water washes, then purified by fractional freezing of the melt and fractional crystallisation from solvents with boiling points below 100°. [Beilstein 2 IV 1219.]

***S*-*tert*-Butyl thioacetate** [999-90-6] M 132.2, b 31-32°/11mm, 38°/14mm, 44-45°/28mm, 67°/54mm, 135.6-135.9°/773mm, d_4^{25} 0.9207, n_D^{20} 1.4532. Dissolve it in $CHCl_3$ (EtOH-free), wash with H_2O , 10% H_2SO_4 , saturated aqueous $NaHCO_3$ (care CO_2 liberated), H_2O again, dry over Drierite and anhydrous K_2CO_3 , and fractionate under reduced pressure. [Rylander & Tarbell *J Am Chem Soc* 72 3021 1950, DOI: 10.1021/ja01163a061; Beilstein 2 IV 546.]

***N*-*tert*-Butyl urea** [1118-12-3] $C_6H_{12}OS$, M 116.2, m 182°, 185°(dec). Possible impurity is *N,N'*-di-*tert*-butyl urea which is quite insoluble in H_2O . Recrystallise it from hot H_2O , filter off insoluble material, and cool from 0° to -5° with stirring. Dry in vacuum at room temperature over KOH or H_2SO_4 . If dried at higher temperatures, it sublimes slowly. It can be recrystallised from EtOH as long white needles or from 95% aqueous EtOH as plates. During melting point determination the bath temperature has to be raised rapidly as the urea sublimes slowly above 100° at 760mm. [Smith & Emerson *Org Synth Coll Vol* 3 148 1955, DOI: 10.15227/

orgsyn.027.0012; *Beilstein* 4 IV 665.]

***n*-Butyl vinyl ether** [111-34-2] $C_6H_{12}O$, M 100.2, b 93.3°/atm, d_4^{20} 0.775. After five washings with equal volumes of water to remove alcohols (made slightly alkaline with KOH), the ether is dried with sodium and distilled under vacuum, taking the middle fraction [Coombes & Eley *J Chem Soc* 3700 1957, DOI: 10.1039/JR9570003700]. Store it over KOH. [*Beilstein* 1 IV 2052.]

2-Butyne (dimethylacetylene, crotonylene) [503-17-3] C_4H_6 , M 54.1, b 0°/253mm, d_4^{20} 0.693. Keep it over Na wire for 24 hours, then fractionally distil it under reduced pressure into a cooled receiver. [*Beilstein* 1 IV 971.]

2-Butyne-1,4-diol [110-65-6] $C_4H_6O_2$, M 86.1, m 54-57°, 56-58°, b 238°/atm. Crystallise the diol from EtOAc. [*Beilstein* 1 IV 2687.]

***n*-Butyric acid** [107-92-6] $C_4H_8O_2$, M 88.1, f -5.3°, b 163.3°/atm, d_4^{20} 0.961, n_D^{25} 1.396, pK^{25} 2.82. Distil the acid, then mix it with $KMnO_4$ (20g/L), and fractionally redistil, discarding the first third of the distillate [Vogel *J Chem Soc* 1814 1948, DOI: 10.1039/JR9480001814]. [*Beilstein* 2 IV 779.] **Butyramide** [514-35-5] C_4H_7NO , M 87.1, m 115°, b 230°/atm, is recrystallised from acetone, *benzene, CCl_4 /petroleum ether, 20% EtOH or water. Dry it under vacuum over P_2O_5 , $CaCl_2$ or 99% H_2SO_4 . [*Beilstein* 2 H 275, 2 I 122, 2 II, 251, 2 III 616, 2 IV 804.] ***n*-Butyric anhydride** [106-31-0] $C_8H_{14}O_3$, M 158.2, b 198°/atm, d_4^{20} 0.968, is dried by shaking with P_2O_5 , then distilling it. [*Beilstein* 2 IV 802.]

γ -Butyrolactone [96-48-0] $C_4H_6O_2$, M 86.1, b 83.8°/12mm, d_4^{20} 1.124. Dry the lactone over anhydrous $CaSO_4$, then fractionally distil it. *Handle it in a fume cupboard due to its TOXICITY.* [*Beilstein* 17 V 7.]

Butyronitrile [109-74-0] C_4H_7N , M 69.1, b 117.9°/atm, d_4^{20} 0.793, n_D^{20} 1.3846, n_D^{30} 1.37954. Treat it with concentrated HCl until the smell of the isonitrile had gone, then dry with K_2CO_3 and fractionally distil [Turner *J Chem Soc* 1681 1956, DOI: 10.1039/JR9560001686]. *Alternatively*, it is twice heated at 75° and stirred for several hours with a mixture of Na_2CO_3 (7.7g) and $KMnO_4$ (11.5g) per L of butyronitrile. The mixture is cooled, then distilled. The middle fraction is dried over activated alumina. [Schoeller & Niemann *J Am Chem Soc* 108 22 1986, DOI: 10.1021/ja00261a005; *Beilstein* 2 IV 806.]

Butyryl chloride (butanoyl chloride) [141-75-3] C_4H_7ClO , M 106.6, f -89°, b 101-102°/atm, d_4^{20} 1.026, n_D^{20} 1.412. Check IR to see if there is a significant peak at 3000-3500 cm^{-1} (br) for OH. If OH is present then reflux it with less than one mole equivalent of $SOCl_2$ for 1 hour and distil directly. The fraction boiling between 85-100° is then refractionated at atmospheric pressure. Keep all apparatus free from moisture and store the product in sealed glass ampoules under N_2 . **LACHRYMATORY**; *handle in a good fume hood.* [Helferich & Schaeffer *Org Synth Coll Vol I* 147 1941, DOI: 10.15227/orgsyn.009.0032; *Beilstein* 2 IV 803.]

Capric acid (decanoic acid) [334-48-5] $C_{10}H_{20}O_2$, M 172.3, m 31.5°, b 148°/11mm, d_4^{20} 0.886, n_D^{25} 1.424, $pK_{Est} \sim 4.9$. The acid is best purified by conversion into its *methyl ester* [110-42-9] $C_{11}H_{22}O_2$, M 186.3, m -14-11°, b 108°/10mm 114.0°/15mm, d_4^{20} 0.871 g/ml, n_D^{20} 1.425 (using excess MeOH, in the presence of H_2SO_4). The H_2SO_4 and MeOH are removed, the ester is distilled *in vacuo* through a 3ft column packed with glass helices. The acid is then obtained from the ester by saponification and vacuum distillation. [Trachtman & Miller *J Am Chem Soc* 84 4828 1962, DOI: 10.1021/ja00883a039; *Beilstein* 2 IV 1041.]

Caproic acid (hexanoic acid) [142-62-1] $C_6H_{12}O_2$, M 116.2, b 205.4°, d_4^{20} 0.925, n_D^{20} 1.417, pK^{25} 4.85. Dry the acid with $MgSO_4$ and fractionally distil it from $CaSO_4$. [*Beilstein* 2 IV 917.] ***n*-Caproamide (*n*-hexanamide)** [628-02-4] $C_6H_{13}NO$, M 115.2, m 100°, 100.5°, is recrystallised from hot water. [*Beilstein* 2 H 324, 2 I 141, 2 II 286, 2 III 732.]

Capronitrile (hexanenitrile) [628-73-9] $C_6H_{11}N$, M 97.2, m -80°, b 163.7°, n_D^{20} 1.4069, n_D^{25} 1.4048. Wash the nitrile twice with half-volumes of conc HCl, then with saturated aqueous $NaHCO_3$, dry over $MgSO_4$, filter,

and distil it. [*Beilstein* 2 H 324, 2 I 141, 2 II 286, 2 III 733, 2 IV 930.]

Caprylonitrile (heptylcyanide) [124-12-9] $C_7H_{13}N$, M 125.2, m -45° , b $198-200^\circ/760\text{mm}$, d_4^{20} 0.812, n_D^{20} 1.420. Wash the nitrile twice with half-volumes of concentrated HCl, then with saturated aqueous $NaHCO_3$, dry over $MgSO_4$, filter and distil it. [*Beilstein* 2 H 349, 2 I 148, 2 II 303, 2 III 798, 2 IV 993.]

Carbon Black [1333-86-4] C, M 12.0, d 1.887/ml, bulk d 0.056g/ml, surface area $>200\text{m}^2/\text{g}$, av pore diameter 64Å. Leach the carbon for 24 hours with 1:1 HCl to remove oil contamination, then wash it repeatedly with distilled water. Dry it in air, and elute for one day each with *benzene and acetone. Again dry it in air at room temperature, then heat it in a vacuum for 24 hours at 600° to remove adsorbed gases. [Tamamushi & Tamaki *Trans Faraday Soc* 55 1007 1959, DOI: 10.1039/TF9595501007.]

Carbon tetrabromide [558-13-4] CBR_4 , M 331.6, m 92.5° . Reactive bromide is removed from CBR_4 by refluxing with dilute aqueous Na_2CO_3 , then steam distilling, crystallising from EtOH, and drying in the dark under vacuum. [Sharpe & Walker *J Chem Soc* 157 1962, DOI: 10.1039/JR9620000157.] It can be sublimed at 70° and low pressure. **It must not be dried with sodium.** [*Beilstein* 1 IV 85.]

Carbon tetrachloride [56-23-5] CCl_4 , M 153.8, b $76.8^\circ/\text{atm}$, d^{25} 1.5842. For many purposes, careful fractional distillation gives adequate purification. Carbon disulfide, if present, can be removed by shaking vigorously for several hours with saturated KOH, separating, and washing with water: this treatment is repeated. The CCl_4 is shaken with concentrated H_2SO_4 until there is no further coloration, then washed with water, dried with $CaCl_2$ or $MgSO_4$ and distilled (from P_2O_5 if desired). **It must not be dried with sodium.** An initial refluxing with mercury for 2 hours removes sulfides. Other purification steps include passage of dry CCl_4 through activated alumina, and distillation from $KMnO_4$. Carbonyl containing impurities can be removed by percolation through a Celite column impregnated with 2,4-dinitrophenylhydrazine (DNPH), H_3PO_4 and water. (Prepared by dissolving 0.5g DNPH in 6ml of 85% H_3PO_4 by grinding together, then mixing with 4ml of distilled water and 10g Celite.) [Schwartz & Parks *Anal Chem* 33 1396 1961, DOI: 10.1021/ac60178a036]. Photochlorination of CCl_4 has also been used: CCl_4 to which a small amount of chlorine has been added is illuminated in a glass bottle (e.g. for 24 hours with a 200W tungsten lamp near it), and, after washing out the excess chlorine with 0.02M Na_2SO_3 , the CCl_4 is washed with distilled water and distilled from P_2O_5 . It can be dried by passing through 4A molecular sieves and distilled. Another purification procedure is to wash CCl_4 with aqueous NaOH, then repeatedly with water and N_2 gas is bubbled through the liquid for several hours. After drying over $CaCl_2$ it is percolated through silica gel and distilled under dry N_2 before use [Klassen & Ross *J Phys Chem* 91 3668 1987, DOI: 10.1021/j100297a041]. [*Beilstein* 1 IV 56.]

Rapid purification: Distil, discarding the first 10% of distillate or until the distillate is clear. The distilled CCl_4 is then stored over 5A molecular sieves.

Carbon tetrafluoride [75-73-0] CF_4 , M 88.0, b $-15^\circ/\text{atm}$. Purify CF_4 by repeated passage over activated charcoal at solid- CO_2 temperatures. Traces of air are removed by evacuating while alternately freezing and melting. *Alternatively*, liquefy CF_4 by cooling in liquid air and then fractionally distil it under vacuum. (The chief impurity originally present is probably CF_3Cl). Use brass equipment. It is non-flammable, but is TOXIC. [Priest *Inorg Synth* 3 178 1950, Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 203 1963, *Beilstein* 1 H 59, 1 I 8, 1 II 11, 1 III 35, 1 IV 26.]

Carbon tetraiodide [507-25-5] CI_4 , M 519.6, m $168^\circ(\text{dec})$, [$171^\circ(\text{dec})$]. Sublime CI_4 *in vacuo*. It hydrolyses in hot EtOH or MeOH, and in H_2O it decomposes slowly into CHI_3 and I_2 , but is soluble and stable in $CHCl_3$ and $*C_6H_6$. [McArthur & Simons *Inorg Synth* 3 37 1950, *Beilstein* 1 H 74, 1 IV 98.]

Cetyl acetate (hexadecyl acetate) [629-70-9] $C_{18}H_{36}O_2$, M 284.5, m 18.3° , b $158^\circ/2\text{mm}$, $204^\circ/18\text{mm}$, d_4^{20} 0.861. Distil the ester in a vacuum twice, then recrystallise it several times from diethyl ether/MeOH. [*Beilstein* 2 H 136, 2 II 146, 2 III 265, 2 IV 171.]

Cetyl alcohol (1-hexadecanol) [36653-82-4] $C_{16}H_{34}O$, M 242.5, m 49.3° . Crystallise the alcohol from aqueous EtOH or from cyclohexane. *Alternatively*, purify it by zone refining. The purity can be checked by gas

chromatography. [Beilstein 1 H 429, 1 I 219, 1 II 466, 1 III 1815, 1 IV 1876.]

Cetylamine [629-54-9] $C_{16}H_{34}NO$, M 255.4, m 104-104.5°, 106-107°, b 235-236°/12mm. Crystallise the amide from thiophene-free *benzene and dry it under vacuum over P_2O_5 . It is slightly soluble in EtOH, Me_2CO , $CHCl_3$ and toluene but insoluble in H_2O . [Beilstein 2 H 374, 2 II 341, 2 III 975, 2 IV 1182.]

Cetylamine (HDA, 1-hexadecylamine) [143-27-1] $C_{16}H_{35}N$, M 241.5, m 48°, b 162-165°/5.2mm, pK^{25} 10.60. Crystallise the base from thiophene-free *benzene and dry under vacuum over P_2O_5 . Store away from CO_2 [Beilstein 4 IV 818.] **Cetylammmonium chloride** [1602-97-7] $C_{16}H_{35}N \cdot HCl$, M 278.0, m 178°, crystallises from MeOH. [Beilstein 4 IV 818.]

Cetyl bromide (1-bromohexadecane) [112-82-3] $C_{16}H_{33}Br$, M 305.4, m 15°, 16-18°, b 144.8°/1mm, 171.5-172°/5mm, 190°/11mm, 193-196°/14mm, d^{25}_D 0.999, n^{20}_D 1.461. It is prepared by boiling hexadecan-1-ol with 48% aqueous HBr and 25% H_2SO_4 for several hours, distilling and purifying by shaking the bromide with H_2SO_4 , washing with H_2O , drying with K_2CO_3 and fractionally distilling it *in vacuo*. [Bezzi & Lanza *Gazzetta* 80 180 1950, Vogel *J Chem Soc* 146 636 1943, DOI: 10.1039/JR9430000636; Heston et al. *J Am Chem Soc* 72 2071 1950, DOI: 10.1021/ja01161a054; Beilstein 1 H 17, 1 I 68, 1 II 138, 1 III 559, 1 IV 542.]

Cetyl ether (dihexadecyl ether) [4113-12-6] $C_{32}H_{66}O$, M 466.9, m 53-56°, 54°, 55°, b 503.8°/760mm. Distil the ether in under high vacuum, then crystallise it several times from MeOH/*benzene. [Gooch *Encyclopedic Dictionary of Polymers* Springer NY, p 209 2011, DOI: 10.1007/978-1-4419-6247-8_3544, Online ISBN: 978-1-4419-6247-8; Beilstein 1 H 430, 1 II 467, 1 III 1820, 1 IV 1878.]

Cetyltrimethylammonium bromide (cetrimonium bromide, CTAB) [57-09-0] $C_{19}H_{42}N \cdot Br$, M 364.5, m 227-235°(dec). Crystallise it from EtOH, EtOH/*benzene or from wet acetone after extracting twice with petroleum ether or dry Et_2O . Shake it with anhydrous Et_2O , filter and dissolve it in a little hot MeOH. After cooling in the refrigerator, the precipitate is filtered off at room temperature and re-dissolved in MeOH. Anhydrous Et_2O is added and, after warming to obtain a clear solution, it is cooled and the crystalline material is collected and dried *in vacuo*. Dry in a vacuum oven at 60° before use. [Dearden & Wooley *J Phys Chem* 91 2404 1987, DOI: 10.1021/j100293a040; Hakemi et al. *J Am Chem Soc* 91 120 1987, DOI: 10.1021/j100285a028; Beilstein 4 IV 819.]

Cetyltrimethylammonium chloride [112-02-7] $C_{19}H_{42}N \cdot Cl$, M 320.0. Crystallise the chloride from acetone/ether mixture, EtOH/ether, or from MeOH. [Moss et al. *J Am Chem Soc* 109 4363 1987, DOI: 10.1021/ja00248a038; Beilstein 4 IV 819.]

Charcoal [7440-44-0] C, M 12.0, m ~3550°. Charcoal (50g) is added to 1L of 6M HCl and boiled for 45 minutes. The supernatant is discarded, and the charcoal is boiled with two more lots of HCl, then with distilled water until the supernatant no longer gives a test for chloride ion. The charcoal (now phosphate-free) is filtered onto a sintered-glass funnel and dried in air at 120° for 24 hours. [Lippin et al. *J Am Chem Soc* 76 2871 1954, DOI: 10.1021/ja01640a004.] The purification can be carried out using a Soxhlet extractor (without cartridge), allowing longer extraction times. Treatment with concentrated H_2SO_4 instead of HCl has been used to remove reducing substances.

Chimyl alcohol (1-O-n-hexadecylglycerol) [(+) 506-03-6, 10550-58-0 (chimyl alcohol)] $C_{19}H_{40}O_3$, M 316.5, m 64°. It is soluble in hexane, $CHCl_3$, Me_2CO and is crystallised from hexane and dried *in vacuo*. The natural **S(+)-alcohol** (from the testes of boars or bulls) crystallises from Et_2O /hexane, and has m 64° (m 62.5-63.5°) and $[\alpha]^{20}_D$ +3.0 (c 1.16, $CHCl_3$); $[\alpha]^{20}_D$ +2.5 \pm 0.5 (c 1.0, $CHCl_3$) was also reported. The **di-phenylurethane** crystallises from petroleum ether as colourless plates with m 97.5-98.5°. The **di-p-nitrobenzoate** has m 52° and $[\alpha]^{20}_D$ -29.2 (c 8.2, 1,1,2,2-tetrachloroethane). [Baer & Fischer *J Biol Chem* 140 397 1941, <http://www.jbc.org/content/140/2/397>; Prelog et al. *Helv Chim Acta* 27 674 1944, DOI: 10.1002/hlca.19440270186; Taguchi et al. *Pteridines* 6 45 1995, DOI: 10.1515/pteridines.1995.6.2.45; Beilstein 1 III 2322.]

Chloral (trichloroacetaldehyde) [75-87-6, 302-17-0 (hydrate)] C_2HCl_3O (anhydr), $C_2H_3Cl_3O_2$ (hydrate),

M 147.4 (anhydrous), 165.4 (hydrate), b 26°/35mm, 98°/760mm. pK²⁵ 10.04. Distil chloral, then dry it by distilling through a heated column of CaSO₄. It readily forms a *hydrate m 57°*, an *alcoholate m 47.5°* [515-83-3] and an *ammonia adduct m 72-74°* [507-47-1]. It is a sedative and a hypnotic. [*Beilstein* 1 H 616, 1 I 328, 1 II 467, 1 III 2663, 1 IV 3142 for anhydr, 1 IV 3143 for hydrate.]

Chloralacetone chloroform (2,2,2-trichloro-1-[2,2,2-trichloro-1,1-dimethylethoxy]ethanol) [512-47-0] M 324.9, m 65°. Crystallise it from *benzene. It sublimes on careful heating and hydrolyses in cold H₂SO₄ to chloral acetone and chloroform. [*Beilstein* 1 H 622.]

Chloroacetaldehyde dimethyl acetal [97-97-2] C₄H₉ClO₂, M 124.6, m -34.4°, b 64°/23mm, 71-72°/35mm, d₄²⁰ 1.0172, n_D²⁰ 1.4175. Purify the acetal by fractional distillation, preferably in a vacuum. [Melhotra *J Indian Chem Soc* 36 4405 1959, *Beilstein* 1 IV 3134.]

Chloroacetic acid [79-11-8] C₂H₃ClO₂, M 94.5, m 62.8°, b 189°, pK²⁵ 2.87. Crystallise the acid from CHCl₃, CCl₄, *benzene or water. Dry it over P₂O₅ or concentrated H₂SO₄ in a vacuum desiccator. Further purification is by distillation from MgSO₄, and by fractional crystallisation from the melt. Store it under vacuum or under dry N₂. [Bernasconi et al. *J Am Chem Soc* 107 3621 1985, DOI: 10.1021/ja00298a035; *Beilstein* 2 IV 474.] **α-Chloroacetamide [79-07-2] C₂H₄ClNO, M 93.5, m 121°, b 224-225°/743mm,** is recrystallised from acetone and dried under vacuum over P₂O₅. [*Beilstein* 2 IV 490.]

Chloroacetic anhydride [541-88-8] C₄H₄Cl₂O₃, M 171.0, m 46°, b 122-123°/20mm, 203°/760mm, d₄²⁰ 1.5494. Crystallise it from *benzene, and distil it preferably under a vacuum. [Eglinton et al. *J Chem Soc* 1860 1954, DOI: 10.1039/JR9540001860; *Beilstein* 2 IV 487.] **IRRITANT, keep away from eyes and skin.**

Chloroacetone [78-95-5] C₃H₅ClO, M 92.5, b 61°/50mm, 119°/763mm, d₄²⁰ 1.15. Dissolve it in water and shake it repeatedly with small amounts of diethyl ether which extracts, preferentially, 1,1-dichloroacetone present as an impurity. The chloroacetone is then extracted from the aqueous phase using a large volume of diethyl ether, and distil at slightly reduced pressure. It is dried with CaCl₂ and stored at Dry-ice temperature. Alternatively, it is kept over CaSO₄ for several hours, distilled and stored over CaSO₄. It is steam volatile. The **2,4-dinitrophenylhydrazones** forms yellow needles from EtOH with **m 120° or 124°**. [*Beilstein* 1 IV 3215.] **LACHRYMATOR with toxic vapour.**

Chloroacetonitrile [107-14-2] C₂H₂ClN, M 75.5, b 125°/atm. Reflux it with P₂O₅ for one day, then distil it through a helices-packed column. Also purified by gas chromatography. [*Beilstein* 2 IV 492.] **LACHRYMATOR AND HIGHLY TOXIC.**

2-Chlorobutane [78-86-4] C₄H₉Br, M 92.6, b 68.5°/atm, d₄²⁰ 0.873, n_D²⁵ 1.3945. Purify it in the same way as *n*-butyl chloride. [*Beilstein* 1 IV 248.]

N-Chlorocarbonyl isocyanate [27738-96-1] C₂ClNO₂, M 105.5, m -68°, b 63.6°/atm, d₄²⁰ 1.310. Fractionally distil it at 760mm using a 40cm column. **TOXIC vapour, use a good fume hood.** Store it dry, its IR (film) has ν_{max} at 2260 (NCO), 1818 (CO) and 1420 (NCO sym) cm⁻¹. [Jäckh & Sundermeyer *Chem Ber* 106 1752 1973, DOI: 10.1002/cber.19731060609.]

2-Chloroethanol (ethylene chlorohydrin) [107-07-3] C₂H₅ClO, M 80.5, b 51.0°/31mm, 128.6°/760mm, d₄²⁰ 1.201, n_D¹⁵ 1.444. Dry it with, then distil it from, CaSO₄ in the presence of a little Na₂CO₃ to remove traces of acid. [*Beilstein* 1 IV 1372.]

2-Chloroethyl bromide (1-bromo-2-chloroethane) [107-04-0] C₂H₄BrCl, M 143.4, b 106-108°/atm, d₄²⁰ 1.723, n_D²⁰ 1.490. Wash it with concentrated H₂SO₄, water, 10% Na₂CO₃ solution, and again with water, then dry with CaCl₂ and fractionally distil before use. [*Beilstein* 1 H 89, 1 I 28, 1 II 61, 1 III 179, 1 IV 155.]

2-Chloroethyl chloroformate [627-11-2] C₃H₄Cl₂O₂, M 143.0, b 52-54°/12mm, 153°/760mm, d₄¹⁸ 1.3760, n_D²⁰ 1.446. Purify it by fractional distillation, preferably in a vacuum, and store it in a dry atmosphere. [Jones *J*

Chem Soc 2735 1957, DOI: 10.1039/JR9570002735; *Beilstein* 3 IV 24.]

2-Chloroethyl vinyl ether [110-75-8] C_4H_7ClO , M 106.6, b 109°/760mm, d_4^{20} 1.048, n_D^{20} 1.437. Wash the ether repeatedly with equal volumes of water made slightly alkaline with KOH, dry with sodium, and distil it under a vacuum. Stabilise it with ~0.01% of triethanolamine. [*Beilstein* 1 IV 2051.] **TOXIC**.

Chlorofluoroacetonitrile (ClFCHCN) [\pm 92484-61-2 and 359-05-7] C_2HClFN , M 93.4, b 66°/atm, d_4^{20} 1.267, n_D^{25} 1.3627. It is prepared from ethyl chlorofluoroacetate (see below) by conversion to the *amide* (0.5mol, b 72-77°/1mm) which is added slowly to P_2O_5 (0.25mol), and the nitrile (3.5g) is distilled off at 50-70°/atm and redistilled (30g, 66°/atm). It is slowly hydrolysed by H_2O but rapidly by aqueous alkali. [Young & Tarrant *J Am Chem Soc* 71 2432 1949, DOI: 10.1021/ja01175a055; *Beilstein* 2 III 454]. Theoretical studies of the molecular dynamics of the enantiomers {(+)- [85196-70-9] and (-)- [85196-71-0]} have been made although they had not been separated [Roselli et al. *J Mol Liq* 29 1 1984, DOI:10.1016/0167-7322(84)80031-8; Evans *Phys Rev A* 30 2062 1984, DOI: org/10.1103/PhysRevA.30.2062; Craig & Elsum *Chem Phys* 73 349 1982, DOI:10.1016/0301-0104(82)85174-4]. Samarium (II) iodide [32248-43-4] promotes the reaction of chlorofluoroacetonitrile with aldehydes to form cyanofluorohydrins in the presence of HMPA [Asano et al. *Synthesis* 1309 2007, DOI: 10.1055/s-2007-966006].

Chloroform [67-66-3] $CHCl_3$, M 119.4, b 61.2°/atm, d^{15} 1.49845, d^{10} 1.47060, n^{15} 1.44858. It reacts slowly with oxygen, or oxidising agents, when exposed to air and light, giving, mainly, phosgene, Cl_2 and HCl . Commercial $CHCl_3$ is usually stabilised with up to 1% of EtOH or of dimethylaminoazobenzene. Simplest purifications involve washing with water to remove the EtOH, drying with K_2CO_3 or $CaCl_2$, refluxing with P_2O_5 , $CaCl_2$, $CaSO_4$ or Na_2SO_4 , and distilling. **It must not be dried with sodium.** The distilled $CHCl_3$ should be stored in the dark to avoid photochemical formation of phosgene. In an alternative purification, $CHCl_3$ (500ml) was shaken (mechanically) with several small portions of 12% H_2SO_4 for 1 hour, washed thoroughly with water, saturated $NaHCO_3$, washed again with water, and dried over $CaCl_2$ or K_2CO_3 (100g) for 1 hour before filtering and distilling. After further drying for a short time over P_2O_5 , the $CHCl_3$ was redistilled and stored over Drierite in the dark [Reynolds & Evans *J Am Chem Soc* 60 2559 1938, DOI: 10.1021/ja01277a080]. EtOH can be removed from $CHCl_3$ by passage through a column of activated alumina, or through a column of silica gel 4-ft long by 1.75-in diameter at a flow rate of 3ml/minute. (The alumina column, which can hold about 8% of its weight of EtOH, is regenerated by air drying and then heating at 600° for 6 hours. It is pre-purified by washing with $CHCl_3$, then EtOH, leaving in concentrated H_2SO_4 for about 8 hours, washing with water until the washings are neutral, then air drying, followed by activation at 600° for 6 hours. Just before use it is reheated for 2 hour at 154°.) [McLaughlin et al. *Anal Chem* 30 1517 1958, DOI: 10.1021/ac60141a021.]

Carbonyl-containing impurities can be removed from $CHCl_3$ by percolation through a Celite column impregnated with 2,4-dinitrophenylhydrazine (DNPH), phosphoric acid and water. (Prepared by dissolving 0.5g DNPH in 6ml of 85% H_3PO_4 by grinding together, then mixing with 4ml of distilled water and 10g of Celite.) [Schwartz & Parks *Anal Chem* 33 1396 1961, DOI: 10.1021/ac60178a036]. Chloroform can be dried by distillation from powdered type 4A Linde molecular sieves. For use as a solvent in IR spectroscopy, chloroform is washed with water (to remove EtOH), then dried for several hours over anhydrous $CaCl_2$ and fractionally distilled. This treatment removes material absorbing near 1600 cm^{-1} . (Percolation through activated alumina increases this absorbing impurity). [Goodspeed & Millson *Chem Ind (London)* 1594 1967, *Beilstein* 1 IV 42.]

Rapid purification: Pass through a column of basic alumina (Grade I, 10g/ml of $CHCl_3$), and either dry by standing over 4A molecular sieves, or *alternatively*, distil from P_2O_5 (3% w/v). Store away from light (to avoid formation of phosgene which is tested by shaking with NH_3 forming solid urea) and use as soon as possible.

Chloromethyl methyl ether (MOMCl) [107-30-2] C_2H_5ClO , M 80.5, b 55-57°/atm, d_4^{20} 1.060, n_D^{20} 1.396. If suspect (check IR), shake it with saturated aqueous $CaCl_2$ solution, dry over $CaCl_2$ and fractionally distil taking the middle fraction. *Alternatively*, it can be prepared afresh. While working in a fume cupboard, a rapid stream of HCl gas ([7647-73-6] from a cylinder, or generated from concentrated H_2SO_4 on $NaCl$) is bubbled through a solution of MeOH (438ml, 10.9moles) and technical formalin (900ml, 252g, 8.4moles, [50-00-0]) which is cooled with running water. After 2 hours two layers are formed, but HCl flow is continued for 2 to 3 hours further until saturation. The MOMCl layer is separated, the aqueous phase is saturated with $CaCl_2$, and the additional MOMCl that is salted out is combined with the first lot, dried over $CaCl_2$, filtered and fractionally

distilled at atmospheric pressure. The fraction that distils at 55-60° is redistilled to give pure MOMCl (586-600g, 86-89%). [Marvel & Porter *Org Synth Coll Vol I* 377 1941, DOI: 10.15227/orgsyn.009.0058; *Beilstein* 1 H 580, 1 I 304, 1 II 645, 1 III 2587, 1 IV 3046.] **VERY TOXIC VAPOUR** and **CARCINOGENIC**.

1-Chloro-2-nitroethane [625-47-8] $C_2H_4ClNO_2$, M 109.5, b 37-38°/20mm, 55°/60mm, 127.5°/atm, n_D^{20} 1.4224, n_D^{25} 1.4235. Dissolve it in alkali, extract with ether (discard), then the aqueous phase is acidified with hydroxylamine hydrochloride, and the nitro compound is collected and fractionally distilled under reduced pressure. [Pearson & Dillon *J Am Chem Soc* 75 2439 1953, DOI: 10.1021/ja01106a048; *Beilstein* 1 H 101, 1 III 202, 1 IV 173.]

Chloropicrin (trichloronitromethane) [76-06-2] CCl_3NO_2 , M 164.5, m -69°, b 112°/atm. Dry it over $MgSO_4$ and fractionally distil. A *fumigant*, soil insecticide and also used as a war gas. [Jackson *Chem Rev* 14 251 1934, Wilhelm US pat 3106588 1963, *Beilstein* 1 IV 106.] **EXTREMELY NEUROTOXIC**, use appropriate precautions.

RS-2-Chloropropionic acid [598-78-7] $C_3H_5ClO_2$, M 108.5, b 98°/3mm, d_4^{20} 1.182, n_D^{20} 1.453 pK^{25} 2.89. Dry it over P_2O_5 and fractionally distil it under vacuum. [*Beilstein* 2 IV 745.] **S-(-)-2-Chloropropionic acid** [29617-66-1] $C_3H_5ClO_2$, M 108.5, b 77°/10mm, 80.7°/10mm, 185-188°/atm, d_4^{25} 1.2485, n_D^{25} 1.436, $[\alpha]_D^{25}$ -14.6 (neat), is purified by fractionating twice through a 115cm Podbielniak column (calculated 50 theoretical plates at atmospheric pressure) using a take-off ratio of 1:5. The *acid chloride* is prepared by dissolving the acid in $SOCl_2$, adding a few drops of PCl_3 , refluxing and then distilling through a 30cm column, b 53°/100mm, $[\alpha]_D^{25}$ -4.6 (neat), d_4^{25} 1.2689, n_D^{25} 1.4368. [Fu et al. *J Am Chem Soc* 76 6054 1954, DOI: 10.1021/ja01652a057; *Beilstein* 2 IV 745.]

3-Chloropropionic acid [107-94-8] $C_3H_5ClO_2$, M 108.5, m 41°, pK^{25} 4.08. Crystallise the acid from petroleum ether or *benzene. [*Beilstein* 2 IV 748.] **3-Chloropropyl bromide (1-bromo-3-chloropropane)** [109-70-6] C_3H_6BrCl , M 157.5, b 142-145°, n_D^{25} 1.4732, is washed with concentrated H_2SO_4 , water, 10% Na_2CO_3 solution, water again and then dried with $CaCl_2$ and fractionally distilled just before use [Akagi et al. *J Am Chem Soc* 78 4034 1956, DOI: 10.1021/ja01597a046]. [*Beilstein* 1 H 109, 1 IV 212.]

2-Chloro-1,1,1-triethoxyethane [51076-95-0] $C_8H_{17}ClO_3$, M 196.7, b 75-80°/5mm, 91°/25mm, 147°/atm, d_4^{25} 1.024, n_D^{20} 1.422. Distil the chloroethane in a vacuum; but if it is discoloured or suspect, then it is better to prepare it anew. To $(EtO)_3CH$ (ethyl orthoformate see [122-51-0]) in CCl_4 add a slight excess of *N*-chlorosuccinimide and heat to 60° while exposing to a sun lamp. An exothermic reaction occurs during 30 minutes; filter off the succinimide formed, evaporate and distil the residue *in vacuo*. [McElvain & Nelson *J Am Chem Soc* 64 1825 1942, DOI: 10.1021/ja01260a024; Mylari et al. *Synth Commun* 19 2921 1989, DOI:10.1080/00397918908052683; Ueno et al. *J Med Chem* 34 2468 1991, DOI: 10.1021/jm00112a023; *Beilstein* 2 III 442, 2 IV 482.]

2-Chlorotriethylamine hydrochloride (2-chloro-*N,N*-diethylethylamine hydrochloride) [869-24-9] $C_6H_{14}ClN$. HCl, M 172.1, m 208-210°, pK_{Est} ~8.6 (free base). Crystallise the salt from absolute MeOH (to remove highly coloured impurities). [*Beilstein* 4 III 240.]

Chlorotrifluoroethylene (CTFE) [79-38-9] C_2F_3Cl , M 116.5, m -158°, b -26 to -24°/atm (-28.4°/atm). Scrub it with 10% KOH solution, then 10% H_2SO_4 solution to remove inhibitors, dry and pass it through silica gel. It is stabilised with ~1% tributylamine. Use brass equipment. [*Beilstein* 1 III 646.] **TOXIC GAS**.

Chlorotrifluoromethane [75-72-9] CF_3Cl , M 104.5, m -180°, b -81.5°/atm. Main impurities are CO_2 , O_2 , and N_2 . The CO_2 is removed by passage through saturated aqueous KOH, followed by concentrated H_2SO_4 . The O_2 is removed using a tower packed with activated copper on Kieselguhr at 200°, and the gas is dried over P_2O_5 . [Miller & Smyth *J Am Chem Soc* 79 20 1957, DOI: 10.1021/ja01558a004; *Beilstein* 1 III 42, 1 IV 34.] **VERY TOXIC GAS**.

Citraconic acid (methylmaleic acid) [498-23-7] $C_5H_6O_4$, M 130.1, m 91°, pK_1^{25} 2.2, pK_2^{25} 5.60 (cis). Steam

distil, and recrystallise it from EtOH/ligroin. [*Beilstein* 2 H 768, 2 I 309, 2 II 652, 2 III 1938, 2 IV 2230.] **Citraconic anhydride** (methylmaleic anhydride) [616-02-4] $C_5H_4O_3$, M 112.1, m 8-9°, b 109°/30mm, 132°/74mm, 213°/760mm, d_4^{20} 1.245, n_D^{20} 1.472, is possibly contaminated with the acid formed by hydrolysis. If the IR has OH bands, then reflux it with Ac_2O for 30 minutes, evaporate, and distil the residue in a vacuum; otherwise distil in a vacuum. Store it in a dry atmosphere. [Vaughan & Andersen *J Am Chem Soc* 77 6702 1955, DOI: 10.1021/ja01629a122; Vaughan & Andersen *J Org Chem* 21 673 1956, DOI: 10.1021/jo01112a023; *Beilstein* 17 H 440, 17 I 234, 17 II 448, 17 III/IV 5912, 17/11 V 65.]

Citric acid (H_2O) [5949-29-1 (monohydrate); 77-92-9 (anhydrous)] $C_6H_8O_7 \cdot H_2O$, $C_6H_8O_7$, M 210.1 (hydrate), 292.1 (anhydrous), m 156-157°, 153° (anhyd), pK_1^{25} 2.96, pK_2^{25} 4.38, pK_3^{25} 5.68. Crystallise it from hot H_2O solution (w/w solubility is 54% at 10°, 71% at 50° and 84% at 100°). The *monohydrate* (softens at ~75° and melts at ~100°) dehydrates in air or when heated gently above 40°. The *triethylester* ([77-93-0] $C_{12}H_{20}O_7$, M 276.3, b 127°/1mm, 294°/atm, d_4^{20} 1.137, n_D^{20} 1.4420) is a bitter tasting oil. [*Beilstein* 3 H 556 and 568, 3 IV 1272.]

Citronellal (3,7-dimethyloctan-6-al) [*R*(+) 2385-77-5, *S*(-) 5949-05-3, *RS*(±) 106-23-3] $C_{10}H_{18}O$, M 154.3, b 67°/4mm, 89°/14mm, 104-105°/21mm, 207°/760mm, $[\alpha]_{546}^{20}$ (+) and (-) 20, $[\alpha]_D^{20}$ (+) and (-) 16.5 (neat). Fractionally distil it. Alternatively, extract it with $NaHSO_3$ solution, wash it with Et_2O , then acidify it to decompose the bisulfite adduct and extract with Et_2O , dry (Na_2SO_4), evaporate and distil. Check for purity by hydroxylamine titration. The ORD in MeOH (c 0.167) has: $[\alpha]_{700} +9$, $[\alpha]_{589} +11$, $[\alpha]_{275} +12$ and $[\alpha]_{260} +12$. The *semicarbazone* has m 85°, and the *2,4-dinitrophenylhydrazone* has m 79-80°. [(+)-compound: Tietze & Beifuss *Org Synth* 71 167 1993, DOI: 10.15227/orgsyn.071.0167; IR: Carroll et al. *J Chem Soc* 3457 1950, DOI: 10.1039/JR9500003457; ORD: Djerassi & Krakower *J Am Chem Soc* 81 237 1959, DOI: 10.1021/ja01510a055; *Beilstein* 1 IV 3515.]

β-Citronellene (2,6-dimethylocta-2,7-diene) [*S*(+) 2436-90-0, *R*(-) 10281-56-8] $C_{10}H_{18}$, M 138.3, b 153-154°/730mm, 155°/atm, d_4^{22} 0.757, n_D^{22} 1.431, $[\alpha]_{546}^{20}$ (+) and (-) 13, $[\alpha]_D^{20}$ (+) and (-) 10 (neat). Purify it by distillation over Na (three times) and fractionation. [(-) Arigoni & Jeger *Helv Chim Acta* 37 881 1954, DOI: 10.1002/hlca.19540370330; for the (+)-isomer see Eschenmoser & Schinz *Helv Chim Acta* 33 171 1950, DOI: 10.1002/hlca.19500330127; *Beilstein* 1 IV 1059-1060.]

β-Citronellol (3,7-dimethyloctan-6-ol) [*R*(+): 11171-61-9, *S*(-): 7540-51-4, *RS*(±): 106-22-9] $C_{10}H_{20}O$, M 156.3, b 47°/1mm, 102-104(110°)/10mm, 112-113°/12mm, 221-224°/atm, 225-226°/atm, d_4^{24} 0.8551, n_D^{24} 1.4562, $[\alpha]_{546}^{20}$ (+) and (-) 6.3, $[\alpha]_D^{20}$ (+) and (-) 5.4 (neat). Purify them by distillation through a cannon packed (Ni) column and the main cut collected at 84°/14mm and redistilled. Also purify *via* the benzoate. [IR: Eschinazi *J Org Chem* 26 3072 1961, DOI: 10.1021/jo01067a007; Naves *Bull Soc Chim Fr* 505 1951, *Beilstein* 1 IV 2188.]

Crotonaldehyde (2-butenal) [123-73-9, *trans*] C_4H_6O , M 70.1, b 104-105°/atm, d_4^{20} 0.851, n_D^{20} 1.437. Fractionally distil it under N_2 , through a short Vigreux column. Stabilise it with 0.01% of 2,6-di-tert-butyl-*p*-cresol and store it in sealed ampoules. [*Beilstein* 1 IV 3447.]

***trans*-Crotonic acid** (*trans*-2-butenic acid) [107-93-7] $C_4H_6O_2$, M 86.1, m 72-72.5°, pK_1^{25} -6.17 (aqueous H_2SO_4), pK_2^{18} 4.71. Distil the acid under reduced pressure and/or recrystallise it from petroleum ether (b 60-80°) or water, or by partial freezing of the melt. [*Beilstein* 2 IV 1498.]

***E*- and *Z*-Crotonitrile** (mixture) [4786-20-3] C_4H_5N , M 67.1, b 120-121°/atm, d_4^{20} 1.091, n_D^{20} 1.4595. Separate the mixture by preparative GLC on a column using 5% FFAP on Chromosorb G. [Lewis et al. *J Am Chem Soc* 108 2818 1986, DOI: 10.1021/ja00271a006; *Beilstein* 2 IV 1507.]

***trans*-Crotonoyl chloride** [625-35-4] C_4H_5ClO , M 104.5, b 42-45°/20mm, 124-125°/760mm, d_4^{20} 1.080, n_D^{20} 1.4570. If the IR of a film has no OH bands, then fractionally distil it, taking the middle fraction and redistilling it. If OH bands are present then add excess of oxalyl chloride, reflux for 3 hours, then distil off the reagent and fractionally distil the crotonoyl chloride as before. Stabilise the distillate with 160ppm of hydroquinone. The

amide forms needles **m 158°** from aqueous ammonia, and the **anilide** also forms needles from H₂O, but with **m 115-118°**. [Beilstein 2 H 411, 2 I 188, 2 II 392, 2 III 1265, 2 IV 1506.]

Crotyl bromide [29576-14-5 *trans*, 591-97-9 *cis/trans*] C₄H₇Br, **M 135.0, b 103-105°/740mm, n_D²⁵ 1.4792**. Dry the bromide with MgSO₄/CaCO₃ mixture and fractionally distil it through an all-glass Todd column. [Beilstein 1 IV 789.]

Cyanoacetic acid [372-09-8] C₃H₃NO₂, **M 85.1, m 70.9-71.1°, pK_a²⁵ 2.47**. Recrystallise the acid to constant melting point from *benzene/acetone (2:3), and dry it over silica gel. [Beilstein 2 H 583, 2 I 253, 2 II 530, 2 III 1626, 2 IV 1888.] **Cyanoacetamide** [107-91-5] C₃H₄N₂O, **M 84.1, m 119.4°**, is crystallised from MeOH/dioxane (6:4), then water and dried over P₂O₅ under vacuum. [Beilstein 2 IV 1891.] **Cyanoacetic acid hydrazide** [140-87-4] C₃H₅N₃O, **M 99.1, m 114.5-115°**, crystallises from EtOH. The **hydrochloride** has **m 178-180°** and the **benzylidene derivative** has **m 178°**. It is converted to 3-oxo-5-iminopyrazolidine in hot 40% aqueous NaOH. [Beilstein 2 H 591, 2 I 256, 2 III 1636.] **IRRITANT**.

Cyanoguanidine (dicyandiamide) [461-58-5] C₂H₄N₄, **M 84.1, m 209.5°, pK_a²⁵ -0.4**. Recrystallise cyanoguanidine from water or EtOH. [Beilstein 3 IV 160.]

***n*-Decane** [124-18-5] C₁₀H₂₂, **M 142.3, m -29.7°, b 57.6°/10mm, 174.1°, d₄²⁰ 0.7300, n_D²⁰ 1.4102, n_D²⁵ 1.40967**. It can be purified by shaking with concentrated H₂SO₄, washing with water, aqueous NaHCO₃, and more water, then drying with MgSO₄, refluxing with Na and distilling. Also purify through a column of silica gel or alumina. It has been purified by azeotropic distillation with 2-butoxyethanol, the alcohol being washed out of the distillate using water, the decane is next dried and redistilled. It can be stored with NaH. Further purification can be achieved by preparative gas chromatography on a column packed with 30% SE-30 (General Electric methyl-silicone rubber) on 42/60 Chromosorb P at 150° and 40psig, using helium carrier gas [Chu *J Chem Phys* 41 226 1964, DOI: org.virtual.anu.edu.au/10.1063/1.1725626]. It is soluble in EtOH and Et₂O. [Beilstein 1 IV 484.]

Decan-1,10-diol [112-47-0] C₁₀H₂₂O, **M 174.3, m 72.5-74°**. Crystallise the diol from dry ethylene dichloride. [Beilstein 1 IV 2613.]

***n*-Decanol (*n*-decyl alcohol)** [112-30-1] C₁₀H₂₂O, **M 158.3, f 6.0°, b 109°/8mm, 231°/atm, d₄²⁰ 0.823, n_D²⁰ 1.434**. Fractionally distil *n*-decanol in an all-glass unit at 10mm pressure (**b 110°**), then crystallise by partial freezing. Also purify by preparative GLC, and passage through alumina before use. [Beilstein 1 IV 1815.]

***n*-Decyl bromide (1-bromodecane)** [112-29-8] C₁₀H₂₁Br, **M 221.2, b 117-118°/15.5mm, d₄²⁰ 1.066**. Shake the it with H₂SO₄, wash with H₂O, dry with K₂CO₃, and fractionally distil it. [Beilstein 1 IV 470.]

Decyltrimethylammonium bromide [2082-84-0] C₁₃H₃₀BrN, **M 280.3, m 239-242°**. Crystallise the salt from 50% (v/v) EtOH/Et₂O, or from acetone and wash with ether. Dry it under vacuum at 60°. Also recrystallise it from EtOH and dry it over silica gel. [McDowell & Kraus *J Am Chem Soc* 73 2170 1952, DOI: 10.1021/ja01149a074; Dearden & Wooley *J Phys Chem* 91 2404 1987, DOI: 10.1021/j100293a040; Beilstein 4 IV 784.]

Diacetamide [625-77-4] C₄H₇NO₂, **M 101.1, m 75.5-76.5°, b 222-223°/atm**. Purify the amide by recrystallisation from MeOH [Arnett & Harrelson *J Am Chem Soc* 109 809 1987, DOI: 10.1021/ja00237a028]. It forms complexes with Mn²⁺, Fe²⁺, Co²⁺, Ni²⁺, Cu²⁺, and Zn²⁺ perchlorates. [Beilstein 2 H 181.]

(+)-Di-*O*-acetyl-L-tartaric anhydride [(*R,R*)-2,3-diacetoxysuccinic anhydride] [6283-74-5] C₈H₈O₇, **M 216.2, m 129-132°, 133-134°, 135°, 137.5°, [α]_D²⁰ +97.2 (c 0.5, dry CHCl₃), [α]_D²⁰ +60 (c 6, Me₂CO)**. If the IR is good, i.e. no OH bands, then keep it in a vacuum desiccator overnight (over P₂O₅/paraffin) before use. If OH bands are present then reflux 4g in Ac₂O (12.6ml) containing a few drops of concentrated H₂SO₄ for 10 minutes (use a relatively large flask), pour onto ice, collect the crystals, wash with dry *C₆H₆ (2 x 2ml), stir with 17ml of

cold Et₂O, filter and dry in it a vacuum desiccator as above, or store it in dark evacuated ampoules under N₂ in small aliquots. It is not very stable in air, the melting point of the crystals drop one degree in the first four days then remains constant (132-134°). If placed in a stoppered bottle, it becomes gummy and the **m** falls 100° in three days. Recrystallisation leads to decomposition. If good quality anhydride is required it, should be prepared freshly from tartaric acid. It sublimes in a CO₂ atmosphere. [Shriner & Furrow *Org Synth Coll Vol* **4** 242 1963, DOI: 10.15227/orgsyn.035.0049; Bell *Aust J Chem* **34** 671 1981, DOI:10.1071/CH9810671; *Beilstein* **18** III/IV 2296.]

Diallyl amine (N-2-propenyl-2-propen-1-amine) [124-02-7] C₆H₁₁N, **M** 97.2, **b** 107-111°/760mm, 112°/760mm, d₄²⁰ 0.789, n_D²⁰ 1.440, pK²⁰ 9.42. Keep the amine over KOH pellets overnight, decant and distil it from a few pellets of KOH at atmospheric pressure (**b** 108-111°), then fractionate through a Vigreux column. [Vliet *J Am Chem Soc* **46** 1307 1924, DOI: 10.1021/ja01670a026; Vliet *Org Synth Coll Vol* **1** 201 1941, DOI: 10.15227/orgsyn.005.0043]. It is a strong base, store under N₂ away from CO₂. The **hydrochloride** has **m** 164-165° (from Me₂CO/EtOH). [Butler & Angelo *J Am Chem Soc* **79** 3128 1957, DOI: 10.1021/ja01569a037.]

(+)-N,N'-Diallyl tartardiamide (DATD) [58477-85-3] C₁₀H₁₆N₂O₄, **M** 228.3, **m** 184° (186-188°), [α]_D²⁰ +108 (c 2.4, H₂O), [α]₅₄₆ +141 (c 3, MeOH). Wash DATD with Et₂O containing 10% EtOH until the washings are clear and colourless, and dry *in vacuo*. [Anker *FEBS Lett* **7** 293 1970, DOI:10.1016/0014-5793(70)80185-5; *Beilstein* **4** H 218.]

1,4-Diaminobutane dihydrochloride (putrescine 2HCl) [333-93-7] C₄H₁₂N₂·2HCl, **M** 161.1, **m** >290°, pK₁²⁵ 9.63, pK₂²⁵ 10.80. Crystallise the salt from EtOH/H₂O. [*Beilstein* **4** IV 1284.]

2,2'-Diaminodiethylamine (diethylenetriamine) [111-40-0] C₄H₁₃N₃, **M** 103.2, **m** -35°, **b** 208°, d₄²⁰ 0.95, n_D²⁰ 1.483, pK₁²⁵ 4.34, pK₂²⁵ 9.13, pK₃²⁵ 9.94. Dry the amine with Na and distil, preferably under reduced pressure, or in a stream of N₂. [*Beilstein* **4** IV 1284.]

§ Polymer-bound diethylenetriamine is commercially available.

3,3'-Diaminodipropylamine [Norspermidine, bis-(3-aminopropyl)amine] [56-18-8] C₆H₁₇N₃, **M** 131.2, **m** -14°, **b** 152°/50mm, d₄²⁰ 0.938, n_D²⁰ 1.481, pK₁²⁵ 7.72, pK₂²⁵ 9.57, pK₃²⁵ 10.65. Dry the amine with Na and distil it under vacuum. It is a strong base, store under N₂ away from CO₂. [*Beilstein* **4** IV 1278.]

1,8-Diaminooctane [373-44-4] C₈H₂₀N₂, **M** 144.3, **m** 50-52°, 51-52°, 52-53°, **b** 121°/18mm, 120°/24mm, pK₁²⁰ 10.1, pK₂²⁰ 11.0. Distil the diamine under vacuum in an inert atmosphere (N₂ or Ar), cool and store the distillate in an inert atmosphere in the dark. The **dihydrochloride** has **m** 273-274°. [Naegeli & Lendorff *Helv Chim Acta* **15** 55 1932, DOI: 10.1002/hlca.19320150109; *Beilstein* **4** III 612.]

1,5-Diaminopentane (Cadaverine) [462-94-2] C₅H₁₄N₂, **M** 102.2, **m** 14-16°, **b** 78-80°/12mm, 101-103°/35mm, 178-180°/750mm, d₄²⁰ 0.869, n_D²⁰ 1.458, pK₁²⁰ 10.02, pK₂²⁰ 10.96. Purify the base by distillation, after standing over KOH pellets (at room temperature, i.e. liquid form). It is a strong base, store under N₂ away from CO₂. Its **dihydrochloride** has **m** 275° (sublimes in a vacuum), and its **tetraphenyl boronate** has **m** 164°. [Schwarzenbach et al. *Helv Chim Acta* **35** 2333 1952, DOI: 10.1002/hlca.19520350719; *Beilstein* **4** IV 1310.]

1,3-Diaminopropane dihydrochloride [10517-44-9] C₃H₁₀N₂·2HCl, **M** 147.1, **m** 243°, pK₁²⁵ 8.29, pK₂²⁵ 10.30. Crystallise the salt from EtOH/H₂O. [*Beilstein* **4** IV 1258 free base.]

1,3-Diaminopropan-2-ol [616-29-5] C₃H₁₀N₂O, **M** 90.1, **m** 38-40°, pK₁²⁵ 7.94, pK₂²⁵ 9.57. Dissolve it in an equal amount of water, shake it with charcoal and distil it at 68°/0.1mm. The distillate solidifies. It is too viscous to be distilled through a packed column. [*Beilstein* **4** IV 1694.]

1,11-Diamino-3,6,9-trioxaundecane [2,2'-oxybis(ethylenoxy)bisethylamine] [929-75-9] C₈H₂₀N₂O₃, **M** 192.3, **b** 115°/0-2mm, 138-153°/1mm, 145-150°/2mm, 135°/3mm, n_D²⁰ 1.440. Distil the diamine *in vacuo*, but if it is suspect then dissolve it in EtOH add 6N HCl boil for 1 hour, evaporate to dryness, and dry the residue in a vacuum. The solid di-hydrochloride residue is treated with 4N NaOH to release the free base which is extra-

cted into CH_2Cl_2 , the extract is dried over K_2CO_3 , filtered, evaporated and then distilled in a vacuum. It is a hydrophilic linker in dendrimer formation or biotinylation. It should be a strong base, store under N_2 away from CO_2 . [Zhang et al. *J Am Chem Soc* **123** 8914 2001, DOI: 10.1021/ja0041369; Yinglin et al. *Synth Commun* **21** 79 1991, DOI:10.1080/00397919108020793; Bogatiskii et al. *J Org Chem USSR* (English Transl) **16** 1124 1980, McReynolds et al. *Bioorg Med Chem* **10** 625 2002, DOI:10.1016/S0968-0896(01)00325-X.]

Diazomethane [H_2CN_2] [334-88-3] $\text{C}_2\text{H}_2\text{N}_2$, **M 42.0, m -145° , b $-23^\circ/\text{atm}$** . Diazomethane is produced when *N*-methyl-*N*-nitroso-amides, e.g. *N*-methyl-*N*-nitroso-urea, *N,N'*-dimethyl-*N,N'*-dinitroso-oxamide, *bis*-(*N*-methyl-*N*-nitroso)terephthalamide or *p*-toluenesulfonyl-*N*-methyl-*N*-nitrosoamide (Diazald), are treated with strong solutions of NaOH or KOH. [Fieser **1** 191, **2** 102.] It has been commonly prepared from Diazald as it is commercially available, although *N*-methyl-*N*-nitroso-urea, which is readily prepared and stored below 5° [Arndt *Org Synth Coll Vol* **2** 461 1943, DOI: 10.15227/orgsyn.015.0048] is also frequently used. *Diazomethane is a sweet smelling, highly irritating yellow gas which is TOXIC when inhaled, causing severe irritation, pulmonary oedema, asthma, chest pains, headaches, weakness as well as producing hypersensitivity reactions and skin irritation. It is a carcinogen. Diazomethane and its precursors should be handled in an efficient fume cupboard; and because of its potentially explosive nature experiments should be carried out behind a safety screen with face and eye protection.*

Preparation: A distilling flask (100ml) containing KOH (5g) in H_2O (8ml) and diluted with 95% EtOH (25ml) is heated to $\sim 65^\circ$ using a water bath, and a solution of Diazald (21.4g, see [80-11-5]) in Et_2O (130ml) is added through a dropping funnel in ~ 25 minutes while the ether which is yellow in colour due to the diazomethane distills off, and the distillate is collected in a receiver immersed in ice. When addition of Diazald is complete, a further volume of Et_2O (20ml) is added dropwise and distillation is continued until the Et_2O distillate is colourless. The yield of diazomethane in the distillate is $\sim 3\text{g}$. [Note that if 95% EtOH is not added to the KOH solution no diazomethane is formed since the base does not dissolve into the Et_2O .] Do **NOT** redistil this ethereal solution as an explosion may result, particularly from the smaller volume left in the flask. [For alcohol-free diazomethane distillate, the distilling flask should contain diethyleneglycol monomethyl ether (35ml), Et_2O (10ml) and KOH (6g) in H_2O (10ml) to which is added the ethereal Diazald as before]. [deBoer & Backer *Org Synth Coll Vol* **6** 943 1963, DOI: 10.15227/orgsyn.034.0096; *Recl Trav Chim Pays-Bas* **73** 229 1954, DOI: 10.1002/recl.19540730308.] By using ^{13}C , ^{14}C and/or ^2H or ^3H methyl labeled Diazald (some of these are also commercially available) appropriately labeled diazomethane can be prepared. The amount of diazomethane is determined by adding a measured volume of 0.2N pure benzoic acid in ether to an aliquot of the ethereal diazomethane solution which will discharge the yellow colour, meaning that all the diazomethane has been consumed. Water is added to this mixture and the excess of benzoic acid is then titrated with 0.1N aqueous NaOH. (See the safer $\text{Me}_3\text{SiCHN}_2$ [18107-18-1] in 'Miscellaneous As, B, P...' section in this chapter.)

For most purposes it is not necessary to distil the diazomethane in ether. In such cases it is better to use *N*-methyl-*N*-nitroso-urea because the unwanted products of the reaction are soluble in water. To a mixture of 50% aqueous KOH (60ml) and pure Et_2O (200ml) at 0° to 5° is added solid nitrosomethylurea (20g) with shaking or stirring. The ethereal layer becomes deep yellow and is decanted or siphoned off; or separated if in a separating funnel. The aqueous layer can be extracted with Et_2O to get a further amount of diazomethane. The ethereal solutions can be dried over KOH pellets and can be used directly or kept at $\sim 0^\circ$ for 2-3 days. [Arndt *Org Synth Coll Vol* **2** 165 1943, DOI: 10.15227/orgsyn.015.003] Diazomethane is a very powerful methylating agent and, in so doing the yellow colour disappears and N_2 bubbles off. It is a versatile reagent in organic synthesis [Black *Aldrichim Acta* **16** 2 1983.]

Di-*O*-benzoyl-(*R* and *S*)-tartaric acid (H_2O) [*R*-(+) 17026-42-5, *S*-(-) 2743-38-6 (anhydrous), 62708-56-9 (hydrate)] $\text{C}_{18}\text{H}_{14}\text{O}_8 \cdot \text{H}_2\text{O}$, **M 376.3 (hydrate), 358.3 (hydrate), m 88-89° (hydrate), 173° (anhydrous), $[\alpha]_{\text{D}}^{20}$ (+) and (-) 136 (c 2, EtOH), $[\alpha]_{\text{D}}^{20}$ (+) and (-) 117 (c 5, EtOH), $\text{pK}_{\text{Est}(1)} \sim 2.9$, $\text{pK}_{\text{Est}(2)} \sim 4.2$** . Crystallise the acid from water (18g from 400 ml boiling H_2O) and stir vigorously while cooling in order to obtain crystals; otherwise an oil will separate which solidifies on cooling. Dry it in a vacuum desiccator over KOH/ H_2SO_4 (yield 16.4g) as **monohydrate, m 88-89°**. It crystallises from xylene as the **anhydrous acid, m 173° (150-153°)**. It **does not** crystallise from $^*\text{C}_6\text{H}_6$, toluene, $^*\text{C}_6\text{H}_6$ /petroleum ether (forms oil), or CHCl_3 /petroleum ether. [Acs et al. *Tetrahedron* **41** 2465 1985, DOI: 10.1016/S0040-4020(01)96641-4; *R*(+) *Beilstein* **9** IV 557, *S*(-) *Beilstein* **9** III 870.] Zetsche & Hubacher [*Helv Chim Acta* **9** 291 1926, DOI: 10.1002/hlca.19260090134] obtained the 2*R*,3*R*(+)-tartaric (L) acid by hydrolysis of the anhydride with dilute NH_3 followed by acidification with HCl.

trans-1,4-Dibromobut-2-ene [821-06-7] $C_4H_6Br_2$, M 213.9, m 54°, b 85°/10mm, 205°/atm,. Crystallise the dibromide from ligroin and/or distil it in a vacuum. [Beilstein 1 IV 79.]

Dibromodichloromethane [594-18-3] CBr_2Cl_2 , M 242.7, m 21.8°, b 66°/81mm, d_4^{20} 2.433, n_D^{25} 1.5499. Crystallise CBr_2Cl_2 , repeatedly from its melt, after washing with aqueous $Na_2S_2O_3$ and drying with BaO. Alternatively, distil it in a vacuum. Store away from light. [Beilstein 1 H 68, 1 III 88, 1 IV 82.]

1,2-Dibromoethane (ethylene dibromide, EDB) [106-93-4] $C_2H_4Br_2$, M 187.9, f 10.0°, b 29.1°/10mm, 131.7°/760mm, d 2.179, n^{15} 1.54160. Wash the dibromide with concentrated HCl or H_2SO_4 , then water, aqueous $NaHCO_3$ or Na_2CO_3 , more water, and dry it with $CaCl_2$. Fractionally distil it. Alternatively, keep in daylight with excess bromine for 2 hours, then extract with aqueous Na_2SO_3 , wash with water, dry with $CaCl_2$, filter and distil. It can also be purified by fractional crystallisation by partial freezing. Store it in the dark. [Beilstein 1 H 90, 1 I 28, 1 II 61, 1 III 182, 1 IV 158.]

Dibromomaleic acid [608-37-7] $C_4H_2Br_2O_4$, M 273.9, m 123.5°, 125°(dec), pK_1^{25} 1.45, pK_2^{25} 4.62. It has been recrystallised from Et_2O or $Et_2O/CHCl_3$. It is soluble in AcOH, also slightly soluble in water, but insoluble in C_6H_6 and $CHCl_3$. [Salmony & Simonis *Chem Ber* 38 2583 1905, DOI: 10.1002/cber.19050380329; Ruggli *Helv Chim Acta* 3 559 1920, DOI: 10.1002/hlca.19200030152; Beilstein 3 IV 2224.]

1,3-Dibromopropane [109-64-8] $C_3H_6Br_2$, M 201.9, f -34.4°, b 63-63.5°/26mm, 76-77°/40mm, 90°/80mm, 165°/atm, d_4^{20} 1.977, n_D^{20} 1.522. Wash the dibromide with dilute aqueous Na_2CO_3 , then water. Dry and fractionally distil it under reduced pressure. [Beilstein 1 IV 216.]

meso-2,3-Dibromosuccinic acid [608-36-6] $C_4H_4Br_2O_4$, M 275.9, m 288-290°(sealed tube, dec), pK_1^{20} 1.56, pK_2^{20} 2.71. Crystallise the acid from distilled water, keeping the temperature below 70°. [Beilstein 2 IV 1930.]

1,2-Dibromotetrafluoroethane [124-73-2] $C_2Br_2F_4$, M 259.8, b 47.3°/760mm. Wash it with water, then with weak alkali. Dry with $CaCl_2$ or H_2SO_4 and distil it. [Locke et al. *J Am Chem Soc* 56 1726 1934, DOI: 10.1021/ja01323a023.] Also purify it by gas chromatography on a silicone DC-200 column. [Hazardine *J Chem Soc* 4259 1952, DOI: 10.1039/JR9520004259; Beilstein 1 III 189.]

Di-n-butylamine [111-92-2] $C_8H_{19}N$, M 129.3, m -62°, b 159°, d_4^{20} 0.761, n_D^{20} 1.41766, pK^{25} 11.25. Dry this strong base with $LiAlH_4$, CaH_2 or KOH pellets, filter and distil it from BaO or CaH_2 . [Beilstein 4 IV 550.]

Di-tert-butylazodicarboxylate [870-50-8] $C_{10}H_{18}N_2O_4$, M 230.3, m 89-92°, 90-92°. The tert-butyl ester has the advantage over the ethyl ester (below) in being a solid and more acid labile. It crystallises from ligroin and is best purified by covering the dry solid (22g) with petroleum ether (b 30-60°, 35-40 ml) heating to boiling and adding ligroin (b 60-90°) until the solid dissolves. On cooling, large lemon yellow crystals of the ester separate (~20g), m 90.7-92°. Evaporation of the filtrate gives a further crop of crystals [Carpino & Crowley *Org Synth* 44 18 1964]. This reagent is useful in the Mitsunobu reaction [Mitsunobu *Synthesis* 1 1981, DOI: 10.1055/s-1981-29317; Gennari et al. *J Am Chem Soc* 108 6394 1986, DOI: 10.1021/ja00280a049; Hughes *Org React* 42 335 1992, Dodge et al. *Org Synth* 73 110 1996, DOI: 10.1522/orgsyn.073.0110; Hughes *Org Prep Proc Int* 28 127 1996, DOI:10.1080/00304949609356516; Ferguson et al. *J Am Chem Soc* 128 4576 2006, DOI: 10.1021/ja058746q; see also DEAD and DIAD below].

Dibutylcarbitol (di[ethyleneglycol]-dibutyl ether, bis[2-butoxyethyl]-ether) [112-73-2] $C_{12}H_{26}O_2$, M 218.3, b 125-130°/0.1mm, 256°/atm, d_4^{20} 0.883, n_D^{20} 1.424. Dibutylcarbitol is freed from peroxides by slow passage through a column of activated alumina. The eluate is then shaken with Na_2CO_3 (to remove any remaining acidic impurities), washed with water, and stored with $CaCl_2$ in a dark bottle [Tuck *J Chem Soc* 3202 1957, DOI: 10.1039/JR9570003202]. [Beilstein 1 IV 2395.]

Di-tert-butyl dicarbonate (di-tert-butyl pyrocarbonate, Boc anhydride) [24424-99-5] $C_{10}H_{18}O_5$, M 218.3, m 23° (21-22°), b 55-56°/0.15mm, 62-65°/0.4mm, d_4^{20} 0.950, n_D^{20} 1.409. Melt the ester by heating at ~35°, and distil it in a vacuum. If the IR and NMR (ν_{max} 1810m and 1765 cm^{-1} , δ in CCl_4 1.50 singlet) suggest that it

is very impure, then wash it with an equal volume of H₂O containing citric acid to make the aqueous layer slightly acidic, collect the organic layer, dry it over anhydrous MgSO₄ and distil it in a vacuum. Store it away from moisture at ~ 4°. If some moisture enters the container pressure may develop since it hydrolyses to *tert*-BuOH and CO₂. It is also available commercially as a 1.0 M solution in THF. It is **FLAMMABLE**. [Pope et al. *Org Synth* **57** 45 1977, DOI: 10.15227/orgsyn.057.0045; Keller et al. *Org Synth* **63** 160 1985, DOI: 10.15227/orgsyn.063.0160.] It is a useful reagent for easy introduction of the *N*-Boc protecting group in amines [Iwanowicz et al. *Synth Commun* **23** 1443 1993, DOI:10.1080/00397919308011234], amino acids, peptides and proteins [Keller et al. *Org Synth* **63** 160 1985, DOI: 10.15227/orgsyn.063.0160], amides [Flynn et al. *J Org Chem* **48** 2424 1983, DOI: 10.1021/jo00162a028; Grehn et al. *Angew Chem Int Ed* **24** 510 1985, DOI: 10.1002/anie.198505101], and *N*-Boc-ylation of sensitive compounds in non-aqueous media [Kemp & Carey *J Org Chem* **54** 3640 1989, DOI: 10.1021/jo00276a026]. Boc protection of alcohols has been achieved via Lewis acid catalysis [Bartoli et al. *Synlett* 2104 2006, DOI: 10.1055/s-2006-949609], and Boc reacted with 4-carboxyphenylhydrazine to form the *N,N,N'*-tris-Boc protected acid which was converted to 4-Fmoc(9-fluorenylmethoxycarbonyl ester)phenyl-hydrazine which is used as a chromophoric reagent to quantify aldehydes attached to a solid-phase since the hydrazine NH₂ group forms a coloured Schiff's base with the aldehyde [Shannon & Barany *J Org Chem* **69** 4586 2004, DOI: 10.1021/jo049830b]. [See the many volumes of *Fieser Reagents for Organic Synthesis*, J Wiley & Sons, for more applications.]

***N,N*-Dibutyl formamide** [761-65-9] C₉H₁₉NO, M 157.3, b 63°/0.1mm, 118-120°/15mm, 244-246°/760mm, d₄²⁰ 0.878, n_D²⁰ 1.445. Purify the amide by fractional distillation, preferably under reduced pressure [Mandel & Hill *J Am Chem Soc* **76** 3978 1954, DOI: 10.1021/ja01644a034]. [*Beilstein* **4** IV 565.]

Di-*tert*-butyl peroxide (tert-butyl peroxide) [110-05-4] C₈H₁₈O₂, M 146.2, b 109-110°/atm (CARE), d 0.794, n_D²⁰ 1.389. Wash the peroxide with aqueous AgNO₃ to remove olefinic impurities, water and dry (MgSO₄). Free it from *tert*-butyl hydroperoxide by passage through an alumina column [Jackson et al. *J Am Chem Soc* **107** 208 1985, DOI: 10.1021/ja00287a038], and if necessary two high vacuum distillations from room temperature to a liquid-air trap [Offenbach & Tobolsky *J Am Chem Soc* **79** 278 1957, DOI: 10.1021/ja01559a009]. [*Beilstein* **1** IV 1619.] *The necessary protection from EXPLOSION should be used.*

Di-*n*-butyl sulfide [544-40-1] C₈H₁₈S, M 146.3, m -76°, α-form b 182°, β-form b 190-230°(dec). Wash the sulfide with aqueous 5% NaOH, then water. Dry with CaCl₂ and distil it from sodium. [*Beilstein* **1** IV 1559.] **Di-*n*-butyl sulfone** [598-04-9] C₈H₁₈SO₂, M 178.3, m 43.5°, b 287-295°/atm, is purified by zone melting. It crystallises from petroleum ether (m 44-44.5°), CHCl₃ (m 44°), and EtOH (m 45°). [*Beilstein* **1** H 371, **1** II 400, **1** III 1524, **1** IV 1561.]

***N,N'*-Di-*tert*-butylthiourea** [4041-95-6] C₉H₂₀N₂S, M 188.3, m 174-175°(evacuated capillary). Recrystallise it from H₂O [Schmidt et al. *Justus Liebigs Ann Chem* **560** 222 1948, DOI: 10.1002/jlac.19485600206; Bortnick et al. *J Am Chem Soc* **78** 4358 1956, DOI: 10.1021/ja01598a043]. [*Beilstein* **4** III 325, **4** IV 668.]

Dichloroacetic acid [79-43-6] C₂H₂Cl₂O₂, M 128.9, m 13.5°, b 95.0-95.5°/17-18mm, b 294°/atm, d₄²⁰ 1.563, n_D²⁰ 1.466, pK²⁵ 1.35. Crystallise this strong acid from *benzene or petroleum ether. Dry it with MgSO₄ and fractionally distil it. [Bernasconi et al. *J Am Chem Soc* **107** 3612 1985, DOI: 10.1021/ja00298a035; *Beilstein* **2** IV 498.]

sym-Dichloroacetone (1,3-dichloropropan-2-one) [534-07-6] C₃H₄Cl₂O, M 127.0, m 41-43°, 45°, b 86-88°/12mm, 75-77°/22mm, 172-172.5°/760mm, 170-175°/760mm, d₄²⁰ 1.383. Crystallise it from CCl₄, CHCl₃ or *benzene and/or distil it under vacuum [Conant & Quayle *Org Synth Coll Vol I* 211 1941, DOI: 10.15227/orgsyn.002.0013; Hall & Sirel *J Am Chem Soc* **74** 836 1952, DOI: 10.1021/ja01123a511]. It is dimorphic [Daasch & Kagarise *J Am Chem Soc* **77** 6156 1955, DOI: 10.1021/ja01628a018]. The *oxime* has m 130-131°, b 106°/25mm [*Arzneimittel-Forsch* **8** 638 1958]. [*Beilstein* **1** IV 3219.]

Dichloroacetonitrile [3018-12-0] C₂HCl₂N, M 110.0, b 110-112°/atm, d₄²⁰ 1.369, n_D²⁰ 1.440. Purify the nitrile by distillation or by gas chromatography. It may liberate HCN. [*Beilstein* **2** IV 506.] **FLAMMABLE.**

2,5-Dichlorobenzoic acid [50-79-3] $\text{C}_7\text{H}_4\text{Cl}_2\text{O}_2$, M 191.0, m 154° , b $301^\circ/760\text{mm}$, pK^{25} 2.47. Recrystallise the acid from water. [Beilstein 9 IV 1005.]

2,3-Dichloro-1,3-butadiene [1653-19-6] $\text{C}_4\text{H}_4\text{Cl}_2$, M 123.0, b $41-43^\circ/85\text{mm}$, $98^\circ/760\text{mm}$, d_4^{20} 1.1872, n_D^{20} 1.4872. Crystallise it from pentane to constant melting point of -40° . A mixture of *meso* and *d,l* forms is separated by gas chromatography on an 8m stainless steel column (8mm i.d.) with 20% DEGS (diethyleneglycolsilyl chloride) on Chromosorb W (60-80 mesh) at 60° and 80ml He/min. Distil it under vacuum. [Su & Ache *J Phys Chem* 80 659 1976, DOI: 10.1021/j100548a002.] [Beilstein 1 III 954, 1 IV 986.]

1,2-Dichloro-1,2-difluoroethane [431-06-1] $\text{C}_2\text{H}_2\text{Cl}_2\text{F}_2$, M 134.9, b $59^\circ/\text{atm}$, n_D^{20} 1.376. Purify it by fractional distillation [Hazeldine *J Chem Soc* 4259 1952, DOI: 10.1039/JR9520004259]. For purification of a diastereoisomeric mixture, with resolution into *meso* and *rac* forms, see Machulla and Stoecklin [*J Phys Chem* 78 658 1974, DOI: 10.1021/j100600a002].

Dichlorodifluoromethane (Freon 12) [75-71-8] CCl_2F_2 , M 120.9, m -158° , b $-29.8^\circ/\text{atm}$, $42.5^\circ/10\text{atm}$. Pass the gas through saturated aqueous KOH then concentrated H_2SO_4 , and a tower packed with activated copper on Kiesselguhr at 200° to remove CO_2 and O_2 . A trap cooled to -29° removes a trace of high boiling material. It is a non-flammable propellant and refrigerant.

1,1-Dichloroethane (ethylidene dichloride) [75-34-3] $\text{C}_2\text{H}_4\text{Cl}_2$, M 99.0, b $57.3^\circ/\text{atm}$, d_4^{15} 1.18350, d_4^{20} 1.177, n^{15} 1.41975. Shake it with concentrated H_2SO_4 or aqueous KMnO_4 , then wash it with water, saturated aqueous NaHCO_3 , again with water, dry with K_2CO_3 and distil it from CaH_2 or CaSO_4 . Store it over silica gel. [Beilstein 1 IV 130.]

1,2-Dichloroethane [107-06-2] $\text{C}_2\text{H}_4\text{Cl}_2$, M 99.0, m -35° , b $83.4^\circ/\text{atm}$, d_4^{20} 1.256, n^{15} 1.44759. It is usually prepared by chlorinating ethylene, so that likely impurities include higher chloro derivatives and other chloro compounds depending on the impurities originally present in the ethylene. It forms azeotropes with water, MeOH, EtOH, trichloroethylene, CCl_4 and isopropanol. Its azeotrope with water (containing 8.9% water, has b 77°) can be used to remove gross amounts of water prior to final drying. As a preliminary purification step, it can be steam distilled, and the lower layer is treated as below.

Shake it with concentrated H_2SO_4 (to remove alcohol added as an oxidation inhibitor), wash with water, then dilute KOH or aqueous Na_2CO_3 and again with water. After an initial drying with CaCl_2 , MgSO_4 or by distillation, it is refluxed with P_2O_5 , CaSO_4 or CaH_2 and fractionally redistilled. Carbonyl-containing impurities can be removed as described for chloroform. [Beilstein 1 IV 131.]

1,2-Dichloroethylene [*cis* + *trans* 540-59-0] $\text{C}_2\text{H}_2\text{Cl}_2$, M 96.9, b 60° (*cis*), d_4^{20} 1.284, b $48^\circ/\text{atm}$ (*trans*), d_4^{20} 1.257. Shake it successively with concentrated H_2SO_4 , water, aqueous NaHCO_3 and water. Dry it with MgSO_4 and fractionally distil it through an efficient column to separate the *cis*- and *trans*-isomers. [Beilstein 1 IV 707-709.] ***cis*-1,2-Dichloroethylene (cis-acetylene dichloride)** [156-59-2] $\text{C}_2\text{H}_2\text{Cl}_2$, M 96.9, m -80° , b $60.4^\circ/\text{atm}$, d_4^{20} 1.2830, n^{15} 1.44903, n_D^{20} 1.4495, is purified by careful fractional distillation, followed by passage through neutral activated alumina. Also by shaking with mercury, drying with K_2CO_3 and distilling from CaSO_4 . Stabilise with 0.02% of 2,6-di-*tert*-butyl-*p*-cresol. [Beilstein 1 IV 707.] ***trans*-1,2-Dichloroethylene (trans-acetylene dichloride)** [156-60-5] $\text{C}_2\text{H}_2\text{Cl}_2$, M 96.9, m -50° , b 47.7° , n^{15} 1.45189, d_4^{20} 1.2551, n_D^{20} 1.4462, is dried with MgSO_4 , and fractionally distilled under CO_2 . Fractional crystallisation at low temperatures has also been used. [Beilstein 1 IV 709.]

2,3-Dichloromaleic anhydride [1122-17-4] $\text{C}_4\text{Cl}_2\text{O}_3$, M 167.0, m $105-115^\circ$, 120° , $121-121.5^\circ$. Purify the anhydride by sublimation *in vacuo* [Katakis et al. *JCS Dalton Trans* 1491 1986, DOI: 10.1039/DT9860001491]. It has also been purified by Soxhlet extraction with hexane, recrystallisation from CHCl_3 and by sublimation. [MS, Relles *J Org Chem* 37 3630 1972, DOI: 10.1021/jo00796a015]. [Beilstein 17/11 V 63.]

Dichloromethane (methylene dichloride) [75-09-2] CH_2Cl_2 , M 84.9, m -97° , b $40.0^\circ/\text{atm}$, d_4^{20} 1.325, n_D^{20} 1.42456, n^{25} 1.4201. Shake it with portions of concentrated H_2SO_4 until the acid layer is colourless, then wash with water, aqueous 5% Na_2CO_3 , NaHCO_3 or NaOH , then water again. Pre-dry it with CaCl_2 , and distil it from

CaSO₄, CaH₂ or P₂O₅. Store it away from bright light in a brown bottle with Linde type 4A molecular sieves, in an atmosphere of dry N₂. Other purification steps include washing with aqueous Na₂S₂O₃, passage through a column of silica gel, and removal of carbonyl-containing impurities as described under **Chloroform**. It has also been purified by treatment with basic alumina, distillation, and is stored over molecular sieves under nitrogen [Puchot et al. *J Am Chem Soc* **108** 2353 1986, DOI: 10.1021/ja00269a036]. Dichloromethane from Japanese sources contained MeOH as stabiliser which is not removed by distillation. It can, however, be removed by standing over activated 3A Molecular Sieves (note that 4A Sieves cause the development of pressure in bottles), passed through activated Al₂O₃ and distilled [Gao et al. *J Am Chem Soc* **109** 5765 1987, DOI: 10.1021/ja00253a032]. It has been fractionated through a platinum spinning band column, degassed, and distilled onto degassed molecular sieves Linde 4A (heated under high vacuum at over 450° until the pressure readings reached the low values of 10⁻⁶ mm, ~1-2 hours). Stabilise it with 0.02% of 2,6-di-*tert*-butyl-*p*-cresol [Kosower & Mohammad *J Am Chem Soc* **93** 2713 1971, DOI: 10.1021/ja00740a022]. [*Beilstein* **1** IV 35.] **Rapid purification:** Reflux over CaH₂ (5% w/v) and distil it. Store it over 3A molecular sieves.

1,2-Dichloropropane (propylene dichloride) [78-87-5] C₃H₆Cl₂, M 113°, m -110°, b 95.9-96.2°/atm, d₄²⁰ 1.158, n_D²⁰ 1.439. Distil the propane from CaH₂. It has a limited shelf life. [*Beilstein* **1** IV 195.]

2,2-Dichloropropane (isopropylidene dichloride) [594-20-7] C₃H₆Cl₂, M 113.0, m -35°, b 69.3°/atm, d₄²⁰ 1.090, n_D²⁰ 1.415. Wash it with aqueous Na₂CO₃ solution, then distilled water, dry it over CaCl₂ and fractionally distil it. [*Beilstein* **1** IV 196.]

Di-*n*-decylamine [1120-49-6] C₂₀H₄₃N, M 297.6, m 34°, b 153°/1mm, 359°/760mm, pK_{Est} ~11.0. Dissolve the amine in *benzene and precipitate it as its bisulfate salt by shaking with 4M H₂SO₄. Filter, wash with *benzene, separate by centrifugation. The free base is obtained by treating with NaOH [McDowell & Allen *J Phys Chem* **65** 1358 1961, DOI: 10.1021/j100826a020], and is a strong base; store away from CO₂. [*Beilstein* **4** IV 780.]

Didodecylamine [3007-31-6] C₂₄H₅₁N, M 353.7, m 51.8°, b 263-265°/27mm, d₄²⁵ 0.806, pK²⁵ 11.00. Crystallise the amine from EtOH/*C₆H₆ under N₂, and store away from CO₂. It provides two crystalline forms: an α-form with m 44.4° and a β-form with m 51.8°. The *hydrochloride* has m 207-208° dec (from isoProOH), the *hydroiodide* has m 23.8-234° dec (sealed capillary), and the *nitrate* has m 125.4-125.2° dec (sealed capillary) when crystallised from MeOH/Me₂CO. [Hoerr et al. *J Am Chem Soc* **65** 328 1943, DOI: 10.1021/ja01243a008; Hoerr & Harwood *J Org Chem* **16** 779 1951, DOI: 10.1021/jo01145a020; *Beilstein* **4** IV 801.]

Didodecyldimethylammonium bromide [3282-73-3] C₂₆H₅₆BrN, M 462.6, m 157-162°. Recrystallise the salt from acetone, acetone/ether mixture, then from ethyl acetate, wash with ether and dry it in a vacuum oven at 60° [Chen et al. *J Phys Chem* **88** 1631 1984, DOI: 10.1021/j150652a038; Rupert et al. *J Am Chem Soc* **107** 2628 1985, DOI: 10.1021/ja00295a012; Rupert et al. *J Am Chem Soc* **108** 3920 1986, DOI: 10.1021/ja00274a011; Allen et al. *J Phys Chem* **91** 2320 1987, DOI: 10.1021/j100293a022]. [*Beilstein* **4** IV 801.]

Diethanolamine (2,2'-iminodiethanol) [111-42-2] C₄H₁₁NO₂, M 105.1, m 28°, b 154-155°/10mm, 270°/760mm, pK²⁵ 8.88. Fractionally distil the amine twice, then fractionally crystallise it from its melt. Its solubility in H₂O is 10% at 20°. It absorbs CO₂ and H₂O from the atmosphere. [Perrin & Dempsey *Buffers for pH and Metal Ion Control* Chapman & Hall, London 1974, *Beilstein* **4** H 283, **4** II 729, **4** III 689, **4** IV 1514.]

***N,N*-Diethylacetamide** [685-91-6] C₆H₁₃NO, M 115.2, b 86-88°/atm, d₄²⁰ 0.994, n_D²⁰ 1.474. Dissolve the amide in cyclohexane, shake with anhydrous BaO and then filter. The procedure is repeated three times, and the cyclohexane is distilled off at atmospheric pressure. The crude amide is also fractionally distilled three times from anhydrous BaO. [*Beilstein* **4** III 349.]

Diethyl acetamidomalonate [1068-90-2] C₉H₁₅NO₅, M 217.2, m 96°, b 185°/10mm. Crystallise the ester from *benzene/petroleum ether. [*Beilstein* **4** III 2993.]

Diethyl acetylenedicarboxylate [762-21-0] C₈H₁₀O₄, M 170.2, b 60-62°/0.3mm, 107-110°/11mm, 118-120°/1mm.

20mm, d_4^{20} 1.0735, n_D^{20} 1.4428. Dissolve the ester in *C_6H_6 , wash it with $NaHCO_3$, H_2O , dry over Na_2SO_4 , filter, evaporate and distil it in a vacuum [IR: Walton & Hughes *J Am Chem Soc* **79** 3985 1957, DOI: 10.1021/ja01572a008; Truce & Kruse *J Am Chem Soc* **81** 5372 1959, DOI: 10.1021/ja01529a030]. [*Beilstein* **2** H 803.]

Diethylamine [109-89-7] $C_4H_{11}N$, **M 73.1, m -50° , b $55.5^\circ/atm$, d_4^{20} 0.707, n_D^{20} 1.38637, pK^{15} 11.38.** Dry diethylamine with $LiAlH_4$ or KOH pellets. Reflux it with, and distil it from, BaO or KOH. Convert it to the *p*-toluenesulfonamide and crystallise it to constant melting point from dry petroleum ether (b $90-120^\circ$), then hydrolyse it with HCl; excess NaOH is added, and the amine is passed through a column of activated alumina. Redistil the amine and dry it with activated alumina before use [Swift *J Am Chem Soc* **64** 115 1942, DOI: 10.1021/ja01253a030]. [*Beilstein* **4** III 313.] § A polystyrene diethylaminomethyl supported version is commercially available. **Diethylamine hydrochloride** [660-68-4] $C_4H_{11}N \cdot HCl$, **M 109.6, m 223.5° , $226-229^\circ$** , is recrystallised from absolute EtOH. Also recrystallise it from dichloroethane/MeOH. *Hygroscopic*. [*Beilstein* **4** III 113.]

Diethyl azodicarboxylate (DEAD) [1972-28-7] $C_6H_{10}N_2O_4$, **M 174.2, b $104.5^\circ/12mm$, $211-213^\circ/atm$, d_4^{20} 1.110, n_D^{20} 1.420.** Dissolve DEAD in toluene, wash it with 10% $NaHCO_3$ till neutral (may require several washes if too much hydrolysis had occurred: check IR for OH bands), then wash it with H_2O (2x), dry it over Na_2SO_4 , filter, evaporate the toluene and distil it through a short Vigreux column at as high a vacuum as possible. The main portion boils at $107-111^\circ/15mm$. *Since it is likely to explode, use an oil bath for heating the still and all operations should be carried out behind an adequate shield with head protection.* [Rabjohn *Org Synth Coll Vol* **3** 375 1955, DOI: 10.15227/orgsyn.028.0058; see Kauer *Org Synth Coll Vol* **4** 411 1963, DOI: 10.15227/orgsyn.000.0007]. [*Beilstein* **3** III 233.] It is commercially available as a 40% solution in toluene. This reagent is useful in the Mitsunobu reaction [Mitsunobu DOI: 10.1055/s-1981-29317; 1 1981, Fürstner & Thiel *J Org Chem* **65** 1738 2000, DOI: 10.1021/jo991611g; Gennari et al. *J Am Chem Soc* **108** 6394 1986, DOI: 10.1021/ja00280a049; Hughes *Org React* **42** 335 1992, Dodge et al. *Org Synth* **73** 110 1996, DOI: 10.15227/orgsyn.073.0110; Hughes *Org Prep Proc Int* **28** 127 1996, DOI: 10.1080/00304949609356516; Ferguson et al. *J Am Chem Soc* **128** 4576 2006, DOI: 10.1021/ja058746q, see also di-*tert*-butyl azodicarboxylate above and **DIAD** below].

§ A polystyrene supported DEAD version is commercially available with a loading of $\sim 1.2mmol/g$.

Diethyl bromomalonate [685-87-0] $C_7H_{11}BrO_2$, **M 239.1, b $116-118^\circ/10mm$, $122-123^\circ/20mm$, d_4^{20} 1.420, n_D^{20} 1.4507.** Purify the ester by fractional distillation in a vacuum. Its IR (film) has ν_{max} 1800 and $1700cm^{-1}$ [Abramovitch *Can J Chem* **37** 1146 1959, DOI: 10.1139/v59-169; Bretschneider & Karpitschka *Monatsh Chem* **84** 1091 1953, DOI: 10.1007/BF00904132]. [*Beilstein* **2** IV 1904.]

Diethyl *tert*-butylmalonate [759-24-0] $C_{11}H_{20}O_4$, **M 216.3, b $40-42^\circ/0.03$, $102-104^\circ/11mm$, $109.5-110.5^\circ/17mm$, $205-210^\circ/760mm$, d_4^{20} 0.980, n_D^{20} 1.425.** Dissolve it in Et_2O , wash with aqueous $NaHCO_3$, H_2O , dry ($MgSO_4$), filter, evaporate and distil the residue. Identify by hydrolysis to the acid and determine the neutralisation equivalent (theor: 80.0). The *acid* has **m $155-157^\circ$** effervescence [Hauser et al. *J Am Chem Soc* **64** 2714 1942, DOI: 10.1021/ja01263a054; Bush & Beauchamp *J Am Chem Soc* **75** 2949 1953, DOI: 10.1021/ja01108a048]. [*Beilstein* **2** IV 2027.]

Diethyl carbonate [105-58-8] $C_5H_{10}O_3$, **M 118.1, m -43° , b $124-125^\circ/atm$, $126.8^\circ/atm$, d_4^{20} 0.975, n_D^{25} 1.38287.** Wash the ester (100ml) with an aqueous 10% Na_2CO_3 (20ml) solution, saturated $CaCl_2$ (20ml), then water (30ml). After drying by standing over solid $CaCl_2$ for 1 hour (note that prolonged contact should be avoided because slow combination with $CaCl_2$ occurs), it should be fractionally distilled. Also dry it over $MgSO_4$ and distil it. [*Beilstein* **3** H 5, **3** I 4, **3** II 4, **3** III 5, **3** IV 5.]

Diethyl disulfide [110-81-6] $C_4H_{10}S_2$, **M 122.3, b $86.2^\circ/87mm$, $154-155^\circ/atm$, d_4^{20} 0.993, n_D^{20} 1.506.** Dry the disulfide over silica gel or $MgSO_4$ and distil it under reduced pressure (optionally from $CaCl_2$). [*Beilstein* **1** H 347, **1** I 173, **1** II 345, **1** III 1377, **1** IV 1379.]

Diethylene glycol [111-46-6] $C_4H_{10}O_3$, **M 106.1, f -10.5° , b $244.3^\circ/atm$, d_4^{20} 1.118, n_D^{15} 1.4490, n_D^{20} 1.4475.**

Fractionally distil it in a vacuum (**b** 133°/14mm, 2.5cm x 1.3m heli-grid column), then recrystallise it by partial freezing. [Feldman et al. *J Am Chem Soc* **73** 4341 1951, DOI: 10.1021/ja01153a091; *Beilstein* **1** III 2090, **1** IV 2390.] **Diethylene glycol ditosylate** [7460-82-4] $C_{18}H_{22}O_7S_2$, **M** 414.5, **m** 86-87°, 87-88°, 88-89°, is purified by recrystallisation from Me_2CO and dried in a vacuum. [*Beilstein* **11** III 225.]

Diethylene glycol diethyl ether [112-36-7] $C_8H_{18}O_3$, **M** 162.2, **b** 76°/32mm, 85-86°/10mm, 188.2-188.3°/751mm, d_4^{20} 0.910, n_D^{20} 1.412. Dry the ether with $MgSO_4$, then CaH_2 or $LiAlH_4$, under N_2 . If sodium is used, the ether should be redistilled alone to remove any products which may be formed by the action of sodium on the ether. As a preliminary purification, the crude ether (2L) can be refluxed for 12 hours with 25ml of concentrated HCl in 200ml of water, under reduced pressure, with slow passage of N_2 to remove aldehydes and other volatile substances. After cooling, add sufficient solid KOH pellets (slowly and with shaking until no more dissolves) to give two liquid phases. The upper of these is decanted, dried with fresh KOH pellets, decanted, then refluxed over, and distilled from sodium. It can be passed through (alkaline) alumina prior to purification. [*Beilstein* **1** IV 2394.]

Diethylene glycol mono-*n*-butyl ether (butyl carbitol) [112-34-5] $C_8H_{18}O_3$, **M** 162.2, **b** 69-70°/0.3mm, 230.5°/760mm, d_4^{20} 0.967, n_D^{20} 1.4286. Dry the ether with anhydrous K_2CO_3 or $CaSO_4$, filter and fractionally distil it. Peroxides can be removed by refluxing with stannous chloride or a mixture of $FeSO_4$ and $KHSO_4$ (or, less completely, by filtration under slight pressure through a column of activated alumina). [*Beilstein* **1** IV 2394.]

Diethylene glycol monoethyl ether (carbitol) [111-90-0] $C_6H_{14}O_3$, **M** 134.2, **b** 201.9°/atm, d_4^{20} 0.999, n_D^{20} 1.4273, n_D^{25} 1.4254. Ethylene glycol can be removed by extracting 250g in 750ml of *benzene with 5ml portions of water, allowing for phase separation, until successive aqueous portions show the same volume increase. Dry, and free from peroxides, as described for diethylene glycol mono-*n*-butyl ether. [*Beilstein* **1** IV 2393.]

Diethylene glycol monomethyl ether [111-77-3] $C_5H_{13}O_3$, **M** 120.2, **m** -70°, **b** 194°/atm, d_4^{20} 1.010, n_D^{20} 1.423. Purify it as for diethylene glycol mono-*n*-butyl ether. [*Beilstein* **1** IV 2392.]

Diethylenetriaminepenta-acetic acid (DTPA, DEPTAPAC) [67-43-6] $C_{14}H_{23}N_3O_{10}$, **M** 393.4, **m** 219-220°, pK_1^{25} 1.79, pK_2^{25} 2.56, pK_3^{25} 4.42, pK_4^{25} 8.76, pK_5^{25} 10.42. Crystallise DTPA from water. Dry it under vacuum or at 110°. [Bielski & Thomas *J Am Chem Soc* **109** 7761 1987, DOI: 10.1021/ja00259a026; NMR: Wenzel et al. *Anal Chem* **54** 615 1982, DOI: 10.1021/ac00241a004; *Beilstein* **4** IV 2454.]

Diethyl ether (ethyl ether) [60-29-7] $C_4H_{10}O$, **M** 74.1, **m** -116°, **b** 34.6°/760mm, d_4^{20} 0.714, n_D^{15} 1.3555, n_D^{20} 1.35272. Usual impurities are water, EtOH, diethyl peroxide (which is explosive when concentrated), and aldehydes. Peroxides [detected by liberation of iodine from weakly acid (HCl) solutions of KI, or by the blue colour in the ether layer when 1mg of $Na_2Cr_2O_7$ and 1 drop of dilute H_2SO_4 in 1ml of water is shaken with 10ml of ether] can be removed in several different ways. The simplest method is to pass dry ether through a column of activated alumina (80g Al_2O_3 /700ml of ether). More commonly, 1L of ether is shaken repeatedly with 5-10ml of a solution comprising 6.0g of ferrous sulfate and 6ml of concentrated H_2SO_4 in 110ml of water. Aqueous 10% Na_2SO_3 or stannous chloride can also be used. The ether is then washed with water, dried for 24 hours with $CaCl_2$, filtered and dried further by adding sodium wire until it remains bright. The ether is stored in a dark cool place, until distilled from sodium before use. Peroxides can also be removed by wetting the ether with a little water, then adding excess $LiAlH_4$ or CaH_2 and leaving to stand for several hours. (This also dried the ether.) Werner [*Analyst* **58** 333 1933, DOI: 10.1039/AN9335800333] removed peroxides and aldehydes by adding 8g $AgNO_3$ in 60ml of water to 1L of ether, then 100ml of 4% NaOH and shaking for 6 minutes. Fierz-David [*Chimia* **1** 246 1947] shook 1L of ether with 10g of a zinc-copper couple. (This reagent is prepared by suspending zinc dust in 50ml of hot water, adding 5ml of 2M HCl and decanting after 20 seconds, washing twice with water, covering with 50ml of water and 5ml of 5% cuprous sulfate with swirling. The liquid is decanted and discarded, and the residue is washed three times with 20ml of ethanol and twice with 20ml of diethyl ether). Aldehydes can be removed from diethyl ether by distillation from hydrazine hydrogen sulfate, phenylhydrazine or thiosemicarbazide. Peroxides and oxidisable impurities have also been removed by shaking with strongly alkaline saturated- $KMnO_4$ (with which the ether was left to stand in contact for 24 hours), followed by washing with water, concentrated H_2SO_4 , water again, drying ($CaCl_2$), and distillation from sodium, or Na containing

benzophenone to form the ketyl. Other purification procedures include distillation from sodium triphenylmethide or butyl magnesium bromide, and drying with solid NaOH or P₂O₅. [*Beilstein* 1 IV 1314.]

Rapid purification: Same as for 1,4-dioxane.

Diethyl ethoxymethylene malonate [87-13-8] C₁₀H₁₆O₅, M 216.2, b 014°/0.2mm, 109°/0.5mm, 279-283°/atm, d₄²⁰ 1.079, n_D²⁰ 1.4623. Likely impurity is diethyl diethoxymethylene malonate which is difficult to separate from diethyl ethoxymethylene malonate by distillation, and it is necessary to follow the course of the distillation by the change in refractive index instead of boiling point. After a low boiling fraction is collected, there is obtained an intermediate fraction (n_D²⁰ 1.414–1.458), the size of which depends on the amount of the diethoxymethylene compound. This fraction is fractionated through a 5inch Vigreux column at low pressure, and avoiding interruption in heating. Fraction **b 108-110°/0.25mm** is ca 10° higher than the submitters' fraction (b 97.2°/0.25mm, n_D²⁰ 1.4612–1.4623) [Parham & Reed *Org Synth Coll Vol* 3 395 1955, DOI: 10.15227/orgsyn.028.0060; Fuson et al. *J Org Chem* 11 194 1946, DOI: 10.1021/jo01172a014; Duffin & Kendal *J Chem Soc* 893 1948 DOI: 10.1039/JR9480000890]. [*Beilstein* 3 IV 1192.]

N,N'-Dimethylethylenediamine [1,2-bis(methylamino)ethane] [110-70-3] C₄H₁₂N₂, M 88.2, b 110-112°/750mm, 119°/760mm, d₄²⁰ 0.819, n_D²⁰ 1.431, (pK²⁵ 7.01 and 9.88). This strong base has been prepared in various ways including hydrolysis of *N,N'*-di(benzylsulfonyl)-*N,N'*-dimethylethylenediamine (**m 217-219°**, from AcOH) and concentrated HCl (120-130°) [Johnson & Bailey *J Am Chem Soc* 38 2135 1916, DOI: 10.1021/ja02267a024], the reaction of 1,2-dibromoethane with 21-33% aqueous MeNH₂ in EtOH (reflux for 2 hours, 50% yield) [Kermack & Wight *J Chem Soc* 1421 1935, DOI: 10.1039/JR9350001421; Woodburn & O'Gee *J Org Chem* 17 1235 1952, DOI: 10.1021/jo50009a008], hydrolysis of *N,N'*-di(*p*-toluenesulfonyl)-*N,N'*-dimethylethylenediamine (1mol, **m 164°**, from AcOH) with H₂SO₄ (8.2mol) and H₂O (9mol) at 140-145°/7 hours [Boon *J Chem Soc* 307 1947, DOI: 10.1039/JR9470000307], LAH reduction in THF of 3-benzenesulfonyloxy-5,6-dihydrouracil (**m 175-176°**, from iso-PrOH) and isolated as the *dibenzoyl derivative* (41%, **m 177-178°**, from *C₆H₆) [Bauer *J Am Chem Soc* 78 1945 1956, DOI: 10.1021/ja01590a049], and by catalytic hydrogenation of *N*-benzyl-*N,N'*-dimethylethylenediamine (**b 73-74°/0.1mm**) with 10% Pd/C [Jucker & Rissi *Helv Chim Acta* 45 2383 1962, DOI: 10.1002/hlca.19620450710]. General isolation and purification procedures involve steam distillation of the product diamine, acidifying the distillate with HCl, evaporating to dryness, treating the residual salt with 10% excess of cold 32% aqueous NaOH, collecting the organic layer (**care**: very caustic solution), drying it over solid KOH, and distilling from it. Final distillation of the dry base over Na has been reported (see also [107-15-3]). The diamine is a strong base and readily absorbs CO₂ and H₂O from the atmosphere—store it in a dark stoppered bottle, wax the stopper if it should be stored for long periods. The *dihydrochloride* [5752-40-9] has **m 235-236°(dec)**, and the *picrate* has **m 160°** (from EtOH and Me₂CO in rectangular plates). It forms complexes with Cu, Ni, and Pt among other metals. Its FT-IR (neat) has ν_{max} at 3285.5, 2788.2, 1447.2, 1346.5, 1251.0, 1106.5, 1040.5, 876.9 and 763.7 cm⁻¹. [For pK and metal complexes see Gustafson & Martell *J Am Chem Soc* 81 525 1959, DOI: 10.1021/ja01512a005; *Beilstein* 4 H 250, 4 I 415, 4 II 689, 4 III 512, 4 IV 1171.]

N,N-Diethylformamide [617-84-5] C₅H₁₁NO, M 101.2, b 29°/0.5mm, 61-63°/10mm, 178.3-178.5°/760mm, d₄²⁰ 0.906, n_D²⁵ 1.4313. Distil it under reduced pressure, then at atmospheric pressure [Winteler et al. *Helv Chim Acta* 37 2370 1954, DOI: 10.1002/hlca.19540370752; NMR: Hoffmann *Z Anal Chem* 170 177 1959]. [*Beilstein* 4 IV 346.]

Diethyl fumarate [623-91-6] C₁₀H₁₆O₅, M 172.2, m 1-2°, b 218°/atm, d 1.052, n 1.441. Wash the fumarate with aqueous 5% Na₂CO₃, then with saturated CaCl₂ solution, dry with CaCl₂ and distil it. [*Beilstein* 2 IV 2207.] Note that **dimethyl fumarate** [624-49-7] M 144.1, m 102°, b 192-193°/atm is a higher melting solid. [*Beilstein* 2 IV 2205.]

N,N-Diethyl-1,1,2,3,3,3-hexafluoropropylamine (Ishikawa's Reagent, perfluoropropyldiethylamine, PPDa) [309-88-6] C₇H₁₁F₆N, M 223.2, b 56-57°/58mm d₄²⁵ 1.230, n_D²⁰ 1.3460. When this reagent is prepared from diethylamine (11g, 0.15mol) in dry Et₂O (30ml) in a glass pressure bomb cooled to -70° (sealed bomb) is allowed to rise to ~25°, and stirred at this temperature overnight, the reaction is complete. Crystalline Et₂NH.HF is removed by filtration, the solvent (Et₂O) is evaporated, and the residual oil is distilled in a vacuum

to give a liquid (23.7g, ~72%) which boils at **56-57°/58mm**. This liquid is shown, by its ^{19}F NMR signal intensities, to be a 3:1 mixture of PPDA and exclusively (*E*)-*N,N*-diethyl-pentafluoro-propenylamine (*trans*-dppa) with respectively δ_{F} [PPDA] at -3.0 (1- CF_3), 6.0 (3- CF_A), 11.0 (3- CF_B), 131.0 (2-CF), and δ_{F} [*trans*-dppa] at -12.0 (1- CF_3 , $^1J = 14\text{Hz}$, $^4J = 23.5\text{Hz}$), 41.0 (3-CF, $^4J = 23.5\text{Hz}$, $^3J_{\text{trans}} = 117\text{Hz}$), 122 (2-CF, $^1J = 14\text{Hz}$, $^3J_{\text{trans}} = 117\text{Hz}$) upfield from external $\text{CF}_3\text{CO}_2\text{H}$ with 3:1 relative intensities. The mixture is easy to handle, and is stable on storage at room temperature without discolouration or loss of activity for at least 6 months. It reacts with a variety of CHOH groups to form the respective CHF derivatives, with carboxylic acids to form the acid fluorides [Takaoka, Iwakiri and Ishikawa *Bull Chem Soc Jpn* **52** 3377 1979, DOI: org/10.1246/bcsj.52.3377; Watanabe, Fujita, Ushi and Kitazume *J Fluorine Chem* **31** 247 1986, DOI:10.1016/S0022-1139(00)81428-1; Watanabe, Fujita, Sakamoto, Kuramochi and Kitazume *J Fluorine Chem* **36** 361 1987 DOI:10.1016/S0022-1139(00)82078-3], and allylic alcohols to form α -fluoro- α -(trifluoromethyl)- γ,δ -unsaturated amide precursors [Ogu, Akazome and Ogura *Tetrahedron Lett* **39** 305 1998, DOI:10.1016/S0040-4039(97)10548-2]. The crude 3:1 reaction product has been used satisfactorily without distillation. The presence of 33% of *trans*-dppa is not a drawback because HF is liberated in the reaction of PPDA with OH, and the HF that is liberated adds on to the olefinic *trans*-dppa to give PPDA; so that as long as there is at least equal amounts of these two compounds the mixture behaves as if it were completely made up of DDPA.

When the above synthesis is carried out at $\sim 5\text{-}10^\circ$ while *F*-propene is bubbled through (completely absorbed within 2 hours), then stirred at $\sim 25^\circ$ overnight, an oil is obtained (89% yield) with **51-53°/58mm** which is a 1:1 mixture (by ^{19}F NMR) of PPDA and *trans*-dppa. If $n\text{-Bu}_2\text{NH}$, or piperidine, replace Et_2NH in the reaction at -70° in sealed bombs, ($n\text{-Bu}$) $_2\text{N-CF}_2\text{CHFCF}_3$ and (*E*)-($n\text{-Bu}$) $_2\text{N-CF=CFCF}_3$ (**55-57°/5mm**, 2:1, 78% yield) or $\text{C}_5\text{H}_{10}\text{N-CF}_2\text{CHFCF}_3$ only (**49-50°/7mm**, 80% yield) respectively are obtained [Takaoka, Iwakiri and Ishikawa *Bull Chem Soc Jpn* **11** 3377 1979, DOI: org/10.1246/bcsj.52.3377]. For use in fluorination see Koch et al. [*Synlett* 693 2004, DOI:10.1055/s-2004-817771] and Ogu et al. [*Tetrahedron Lett* **39** 305 1998, DOI: 10.1016/S0040-4039(97)10548-2]. [*Beilstein* **4** IV 353.]

Diethyl ketone (3-pentanone) [96-22-0] $\text{C}_5\text{H}_{10}\text{O}$, M 86.1, m -42° , b $102^\circ/751\text{mm}$, $102.1^\circ/\text{atm}$, d_4^{20} 0.8099, n_D^{20} 1.392. The ketone is dried with anhydrous CaSO_4 or CuSO_4 , and distil from P_2O_5 under N_2 or under reduced pressure. Further purification is by conversion to the semicarbazone (recrystallise to constant m **139°**, from EtOH) which, after drying *in vacuo* over CaCl_2 and paraffin wax, is refluxed for 30 minutes with excess oxalic acid, then steam distilled and salted out with K_2CO_3 . Dry it with Na_2SO_4 and distil it [Cowan et al. *J Chem Soc* 171 1940, DOI: 10.1039/JR9400000171]. [*Beilstein* **1** IV 3279.]

Diethyl malonate [105-53-3] $\text{C}_7\text{H}_{12}\text{O}_4$, M 160.2, m $-51\text{-}52^\circ$, b $92^\circ/22\text{mm}$, $198\text{-}199^\circ/760\text{mm}$, d_4^{20} 1.056, d^{25} 1.0507, n_D^{20} 1.413. If too impure (IR, NMR) the ester (250g) is heated on a steam bath for 36 hours with absolute EtOH (125ml) and conc H_2SO_4 (75ml), then fractionally distilled under reduced pressure. Otherwise fractionally distil it under reduced pressure and collect the steady boiling middle fraction. [*Beilstein* **2** IV 1881.]

Diethylnitrosamine (N-nitrosodiethylamine) [55-18-5] $\text{C}_4\text{H}_{10}\text{N}_2\text{O}$, M 102.1, b $172\text{-}173.5^\circ/\text{atm}$, $177^\circ/\text{atm}$, d_4^{20} 0.95, n_D^{20} 1.437. Dry it over anhydrous K_2CO_3 and distil it at atmospheric pressure which removes any Et_2NH . See purification of dimethylnitrosamine below, and preparation in A.I. Vogel *Practical Organic Chemistry* Textbooks. [*Beilstein* **4** IV 3386.] **Carcinogen.**

2,2-Diethyl-1,3-propanediol [115-76-4] $\text{C}_7\text{H}_{16}\text{O}_2$, M 132.2, m $61.4\text{-}61.8^\circ$, b $130\text{-}133^\circ/16\text{mm}$, n_D^{20} 1.4574. Crystallise the diol from petroleum ether (b $65\text{-}70^\circ$). **IRRITANT.** [McKusick *J Am Chem. Soc* **70** 1982 1948, DOI: 10.1021/ja01185a602; *Beilstein* **1** III 2217, **1** IV 2589.]

Diethyl pyrocarbonate (DEP) [1609-47-8] $\text{C}_6\text{H}_{10}\text{O}_5$, M 162.1, b $38\text{-}40^\circ/12\text{mm}$, $160\text{-}163^\circ/\text{atm}$, d_4^{20} 1.119, n_D^{20} 1.398. Dissolve the ester in Et_2O , wash it with dilute HCl, H_2O , dry over Na_2SO_4 , filter, evaporate and distil the residue first *in vacuo* then at atmospheric pressure. It is soluble in alcohols, esters, ketones and hydrocarbon solvents. A 50% w/w solution is usually prepared for general use. **Treat with great CAUTION as DEP irritates the eyes, mucous membranes and skin.** [Boehm & Mehta *Chem Ber* **71** 1797 1938, DOI: 10.1002/cber.19380710904; Thoma & Rinke *Justus Liebigs Ann Chem* **624** 30 1959, DOI: 10.1002/jlac.19596240104; *Beilstein* **3** IV 18.]

Diethyl succinate [123-25-1] $\text{C}_8\text{H}_{14}\text{O}_4$, M 174.2, m -20° , b $105^\circ/15\text{mm}$, $218^\circ/\text{atm}$, d_4^{20} 1.047, n_D^{20} 1.4199. Dry the succinate with MgSO_4 , and distil it at 15mm pressure. [Beilstein 2 IV 1914.]

Diethyl sulfate [64-67-5] $\text{C}_4\text{H}_{10}\text{O}_4\text{S}$, M 154.2, m -24° , b $96^\circ/15\text{mm}$, $118^\circ/40\text{mm}$, d_4^{20} 1.177, n_D^{20} 1.399. Wash the ester with aqueous 3% Na_2CO_3 (to remove acidic material), then distilled water, dry (CaCl_2), filter and distil it in a vacuum. *It is an ethylating agent and blisters the skin.* [Beilstein 1 IV 1236.]

Diethyl sulfide [352-93-2] $\text{C}_4\text{H}_{10}\text{S}$, M 90.2, m -100° , b $0^\circ/15\text{mm}$, $90.1^\circ/760\text{mm}$, d_4^{20} 0.837, n_D^{20} 1.443. Wash the sulfide with aqueous 5% NaOH , then water, dry with CaCl_2 and distil it from sodium. It can also be dried with MgSO_4 or silica gel. *Alternative purification is via the Hg(II) chloride complex $[(\text{Et})_2\text{S} \cdot 2\text{HgCl}_2]$ (see dimethyl sulfide).* [Beilstein 1 IV 1394.]

Diethyl (-)-D- (from the non-natural) [13811-71-7] and (+)-L- (from the natural acid) [89-91-2] tartrate $\text{C}_8\text{H}_{14}\text{O}_6$, M 206.2, m 17° , b $80^\circ/0.5\text{mm}$, $162^\circ/19\text{mm}$, $278-282^\circ/\text{atm}$, d_4^{20} 1.204, n_D^{20} 1.4476, $[\alpha]_D^{20}$ (+) and (+) 26.5 (c 1, H_2O) and (-) and (+) 8.5 (neat), $[\alpha]_{546}^{20}$ (+) and (+) 30 (c 1, H_2O). Distil the esters under high vacuum and store them under vacuum or in an inert atmosphere in a desiccator in round bottomed flasks equipped with a vacuum stopcock. They have also been distilled by Kugelrohr distillation and/or by ‘wiped-film’ molecular distillation. They are slightly soluble in H_2O but miscible with EtOH and Et_2O . [Gao et al. *J Am Chem Soc* 109 5765 1987, DOI: 10.1021/ja00253a032; IR: Pristera *Anal Chem* 25 844 1953, DOI: 10.1021/ac60078a002; Beilstein 3 III 1025 for D(-), 3 IV 1232 for L(+).]

sym-Diethylthiourea [105-55-5] $\text{C}_5\text{H}_{12}\text{N}_2\text{S}$, M 132.2, m $76-77^\circ$, $77-79^\circ$. Crystallise it from *benzene. [Beilstein 4 H 118, 4 I 355, 4 II 610, 4 III 220, 4 IV 375.]

Difluoroacetic acid [381-73-7] $\text{C}_2\text{H}_2\text{F}_2\text{O}_2$, M 96.0, m -0.35° , b $67-70^\circ/20\text{mm}$, $134^\circ/760\text{mm}$, d_4^{20} 1.530, n_D^{20} 1.3428, pK^{25}_1 1.28. Purify the acid by distilling over P_2O_5 . The *acid chloride* is a fuming liquid b $25^\circ/\text{atm}$, the *amide* has b $108.6^\circ/35\text{mm}$, m 52° (from * C_6H_6), and the *anilide* has b $90^\circ/1\text{mm}$, $114^\circ/5\text{mm}$, and m 58° [Henne & Pelley *J Am Chem Soc* 74 1426 1952, DOI: 10.1021/ja01126a019; Coffman et al. *J Org Chem* 14 747 1949, DOI: 10.1021/jo01157a005; NMR: Meyer et al. *J Am Chem Soc* 75 4567 1953, DOI: 10.1021/ja01114a053; pK: Wegscheider *Z Phys Chem* 69 614 1909]. [Beilstein 2 IV 455.] It is a strong acid see pKa.

Diglycolic acid (2-oxapentane-1,5-dioic acid) [110-99-6] $\text{C}_6\text{H}_6\text{O}_5$, M 134.1, m 148° (monohydrate), pK^{25}_1 2.97, pK^{25}_2 4.37. Crystallise diglycolic acid from water. [Beilstein 3 IV 577.]

Diglyme [bis(2-methoxyethyl) ether, diethylene glycol dimethyl ether] [111-96-6] $\text{C}_6\text{H}_{14}\text{O}_3$, M 134.2, m -64° , b $62^\circ/17\text{mm}$, $75^\circ/35\text{mm}$, $160^\circ/760\text{mm}$, d_4^{20} 0.917, n_D^{20} 1.4087. Dry diglyme with NaOH pellets or CaH_2 , then reflux with, and distil it (under reduced pressure) from Na , CaH_2 , LiAlH_4 , NaBH_4 or NaH . These operations are carried out under N_2 . The amine-like odour of diglyme has been removed by shaking with a weakly acidic ion-exchange resin (Amberlite IR-120) before drying and distilling. Addition of 0.01% NaBH_4 to the distillate inhibits peroxidation. Purify it also as for dioxane. It has been passed through a 12-in column of molecular sieves to remove water and peroxides. [Beilstein 1 IV 2393.]

Dihydroxyfumaric (1,2-dihydroxybut-1-ene-1,2-dioic acid dihydrate, dihydroxymaleic acid) [133-38-0, 20688-70-4 dihydrate] $\text{C}_4\text{H}_4\text{O}_6$, 2 H_2O , M 184.1, m $155^\circ(\text{dec})$, pK^{25}_1 1.57, pK^{25}_2 3.36. The acid has been prepared from tartaric acid by oxidation with $\text{H}_2\text{O}_2/\text{FeSO}_4$ at low temperatures ($<0^\circ$). Traces of metals cause the acid to deteriorate rapidly particularly if it is in the hydrated form. Recrystallisation is a way to reduce traces of heavy metals. The acid (100 g) is dissolved in absolute MeOH (400 mL) at room temperature and filtered (using a further 50 mL of MeOH for transferring and washing). Water (450 mL) is added with stirring whereby white hydrated dihydroxyfumaric acid separates. After 15 minutes the acid is filtered off, washed with several 25 mL portions of 50% aqueous MeOH and allowed to dry in air. It is dehydrated over P_2O_5 in a vacuum desiccator at 0.1mm pressure to constant weight (70-78 g). The dihydrate undergoes a slow change in air becoming yellow and sticky, and develops an odour resembling that of commercial sodium pyruvate. The anhydrous acid is stable in a desiccator at room temperature for at least 6 months even with repeated opening of the desiccator. It appears to keep indefinitely if sealed up with P_2O_5 . [The acid has UV with λ_{max} at 292nm (ϵ 9100) in H_2O , and λ_{max} at

308nm (ϵ 7900) in Et₂O)]. Almost complete removal of heavy metals reduces spontaneous decarboxylation and oxidation in aqueous solution.

The acid forms a labile insoluble dimercury salt which can be used for its estimation [Schmalfuss & Barthmeyer *Hoppe-Seyler's Z physiol Chem* **160** 106 1926]. The *anhydrous acid* melts at ~141°. [Hartree *Biochemical Preparations* **3** 56 1953, Powers et al. *Biochemical Preparations* **4** 56 1955, Fox *J Org Chem* **12** 535 1947, DOI: 10.1021/jo01168a007; *Beilstein* **3** I 186, **3** II 346, **3** III 1045, **3** IV 1975.]

1,2-Diiodoethane [624-73-7] C₂H₄I₂, M 281.9, m 81-84°, d₄²⁵ 2.134g/cm³. Dissolve it in ether, wash it with saturated aqueous Na₂S₂O₃, dry it over MgSO₄, and evaporate the ether *in vacuo* then distil it. Store it in the dark. [Molander et al. *J Am Chem Soc* **109** 453 1987, DOI: 10.1021/ja00236a025]. [*Beilstein* **1** IV 169.]

Diiodomethane (methylene diiodide) [75-11-6] CH₂I₂, M 267.8, m 6.1°, b 66-70°/11-12mm, d₄²⁰ 3.325. Fractionally distil it under reduced pressure, then fractionally crystallise it by partial freezing, and stabilise it with silver wool if necessary. It has also been purified by drying over CaCl₂ and fractionally distilling from Cu powder. Store it in the dark. [*Beilstein* **1** IV 97.]

Diisopropanolamine [110-97-4] C₆H₁₅NO₂, M 133.2, m 41-44°, d₄²⁰ 1.004, pK_{Est} ~10.7. Crystallise the amine repeatedly from dry diethyl ether. It is a strong base, store away from CO₂ and H₂O. [*Beilstein* **4** III 761.]

Diisopropylamine [108-18-9] C₆H₁₅N, M 101.2, m -61°, b 83.5°/760mm, d₄²⁰ 0.720, n_D²⁰ 1.39236, pK₂₅ 11.20. Distil the amine from NaOH, or reflux it three minutes over Na wire or NaH, and distil it into a dry receiver under N₂. It is a strong base, store away from CO₂ and H₂O. [*Beilstein* **4** H 154, **4** I 369, **4** II 630, **4** III 274, **4** IV 510.] § A polystyrene supported version of diisopropylamine is commercially available.

Diisopropylazodicarboxylate (DIAD) [2446-83-5] C₈H₁₄N₂O₄, M 202.2, b 75°/0.2mm, d₂₅ 1.420, n_D²⁰ 1.420. Purify the azo compound by distillation at as high a vacuum as possible. Since it is likely to explode, use an oil bath for heating the still, and all operations should be carried out behind an adequate shield. [Kauer *Org Synth Coll Vol* **4** 411 1963, DOI: 10.15227/orgsyn.000.0007; *Beilstein* **3** III 233]. This reagent is useful in the Mitsunobu reaction [Mitsunobu *Synthesis* **1** 1981, DOI: 10.1055/s-1981-29317; Gennari et al. *J Am Chem Soc* **108** 6394 1986, DOI: 10.1021/ja00280a049; Hughes *Org React* **42** 335 1992, Dodge et al. *Org Synth* **73** 110 1996, DOI: 10.15227/orgsyn.073.0110; Hughes *Org Prep Proc Int* **28** 127 1996, DOI: 10.1080/00304949609356516; Ferguson et al. *J Am Chem Soc* **128** 4576 2006, DOI: 10.1021/ja058746q; see also di-*tert*-butyl azodicarboxylate and **DEAD** above].

Diisopropylethylamine (Hünig's base) [7087-68-5] C₈H₁₉N, M 129.3, m <-51-52°, b 119°/731mm, 127°/760mm, d₄²⁷ 1.440, n_D²⁵ 1.4376, pK_{Est} ~10.9. Distil the amine from ninhydrin, then from KOH [Dryland & Sheppard, *JCS Perkin Trans 1* 125 1986, DOI: 10.1039/P19860000125]. It is a strong sterically hindered *strong* base and should be stored in the absence of carbon dioxide. [Hünig & Kiessel *Chem Ber* **91** 380, 387 1958, DOI: 10.1002/cber.19580910223; Wotiz et al. *J Org Chem* **24** 1202 1959, DOI: 10.1021/jo01091a009; *Beilstein* **4** IV 551.]

Diisopropyl ketone (2,4-dimethyl-3-pentanone) [565-80-0] C₇H₁₄O, M 114.2, m -80°, b 124°/atm, d₂₅ 0.801g/cm³, n_D²⁰ 1.400. Dry the ketone with CaSO₄, shake it with chromatographic alumina and fractionally distil it from P₂O₅ under N₂. [*Beilstein* **1** IV 3334.]

2,3-Dimercapto-1-propanol (BAL, British Anti-Lewisite) [59-52-9] C₃H₈OS₂, M 124.2, b 82-84°/0.8mm, 120°/15mm, d₄²⁰ 1.239, n_D²⁰ 1.5732, pK₁²⁵ 8.62, pK₂²⁵ 10.75. Precipitate BAL as the Hg mercaptide [see Sjöberg *Chem Ber* **75** 13 1942, DOI: 10.1002/cber.19420750103], regenerate with H₂S, and distil it under a vacuum [Rosenblatt & Jean *Anal Chem* **27** 951 1955, DOI: 10.1021/ac60102a023]. *It is an antidote for heavy metal (As, Hg, Au etc) poisoning.* [*Beilstein* **1** IV 2770.]

1,3-Dimercapto-2-propanol [584-04-3] C₃H₈OS₂, M 124.2, b 68-69°/0.8mm, 82°/1.5mm, 94°/12mm, n_D²³ 1.5696. Purify the dithiol as for 2,3-dimercapto-1-propanol above. [Johary & Owen *J Chem Soc* 1302 1955, DOI: 10.1039/JR9550001302; *Beilstein* **1** IV 2773 or **2** IV 1102.]

meso-2,3-Dimercaptosuccinic acid (DMSA) [304-55-2] $C_4H_6O_4S_2$, M 182.2, m 191-192°(dec), 210°(dec), 210-211° (dec), pK_1^{25} 2.71, pK_2^{25} 3.48, pK_3^{25} 8.89, pK_4^{25} 10.75. Purify the acid by dissolving it in NaOH and precipitating with dilute HCl, drying and recrystallising from MeOH. IR has ν_{max} at 2544 (SH) and 1689 (CO₂H) cm^{-1} . The **bis-S-acetyl** derivative has m 183-185° (from EtOAc or Me₂CO), and its **Me ester** has m 119-120° (from petroleum ether) [Gerecke et al. *Helv Chim Acta* **44** 955 1961, DOI: 10.1002/hlca.19610440410; Owen & Sultanbawa *J Chem Soc* 3105 1949, DOI: 10.1039/JR9490003109]. [*Beilstein* 3 III 1033.]

1,2-Dimethoxyethane (DME, glycol dimethyl ether, glyme) [110-71-4] $C_4H_{10}O_2$, M 90.1, m -58°, b 84°/atm, d_4^{20} 0.867, n_D^{20} 1.380. Traces of water and acidic materials have been removed from it by refluxing with Na, K or CaH₂, decanting and distilling from Na, K, CaH₂ or LiAlH₄. The reaction has been speeded up by using vigorous high-speed stirring with molten potassium. For virtually complete elimination of water, 1,2-dimethoxyethane has been dried with Na-K alloy until a characteristic blue colour is formed in the solvent at Dry-ice/cellosolve temperatures: the solvent is kept with the alloy until distilled for use [Ward *J Am Chem Soc* **83** 1296 1961, DOI: 10.1021/ja01467a010]. Alternatively, glyme, refluxed with benzophenone and Na-K, is dry enough if, on distillation, it gives a blue colour of the ketyl immediately on addition to benzophenone and sodium [Ayscough & Wilson *J Chem Soc* 5412 1963, DOI: 10.1039/JR9630005412]. It has also been purified by distillation under N₂ from sodium benzophenone ketyl (see above). [*Beilstein* 1 IV 2376.]

N,N-Dimethylacetamide (DMAc) [127-19-5] C_4H_9NO , M 87.1, m -20°, b 58.0-58.5°/11.4mm, 66-67°/15mm, 85-87°/33mm, 96°/80mm, 163-165°/760mm, d_4^{20} 0.940, n_D^{20} 1.4373. Shake the amide with BaO for several days, reflux it over BaO for 1 hour, then fractionally distil it under reduced pressure. Store it over molecular sieves (4A). It is a useful organic solvent for reactions, as it is soluble in many organic solvents and in H₂O. Use it in a fume hood because over-exposure causes liver damage, jaundice, lethargy and can irritate skin. [*Beilstein* 4 IV 180.]

β,β -Dimethylacrylic acid (senecioic acid, 3-methyl-2-butenic acid) [541-47-9] $C_5H_8O_2$, M 100.1, m 68°, pK^{25} -5.4 (aqueous H₂SO₄). Crystallise the acid from hot water or petroleum ether (b 60-80°). [*Beilstein* 2 IV 1555.] It is a strong acid.

Dimethyl adipate [627-93-0] $C_8H_{14}O_4$, M 174.2, m 9-11°, b 109°/10mm, 121-123°/20mm, 235°/760mm, d_4^{20} 1.0642, n_D^{20} 1.4292. Dissolve it in Et₂O, wash with NaHCO₃, H₂O, dry over MgSO₄, filter, evaporate and distil it several times until the IR and NMR are consistent with its structure [Lorette & Brown *J Org Chem* **24** 261 1959, DOI: 10.1021/jo01084a633; Hoffmann & Weiss *J Am Chem Soc* **79** 4759 1957, DOI: 10.1021/ja01574a045]. [*Beilstein* 2 IV 1959.]

Dimethyl adipimidate dihydrochloride [14620-72-5] $C_8H_{16}N_2O_2 \cdot HCl$, M 245.1, m 218-220°, 222-224°. If the salt smells of HCl, then wash it with MeOH and dry Et₂O (1:3) under N₂ until the free HCl is completely removed. Recrystallise it from MeOH/Et₂O (it is very important that the solvents are super dry) [Hartman & Wold *Biochemistry* **6** 2439 1967, DOI: 10.1021/bi00860a021; McElvain & Shroeder *J Am Chem Soc* **71** 40 1949, DOI: 10.1021/ja01169a013].

Dimethylamine [124-40-3] C_2H_7N , M 45.1, m -93°, b 0°/563mm, 6.9°/760mm, pK^{25} 10.73. Dry dimethylamine by passage through a KOH-filled tower, or by standing with sodium pellets at 0° during 18 hours. It is a strong base — *do not inhale its vapours*. [*Beilstein* 4 IV 128.] § A dimethylaminomethyl polystyrene supported version is commercially available. **Dimethylamine hydrochloride** [506-59-2] $C_2H_7N \cdot HCl$, M 81.5, m 171°, crystallises from hot CHCl₃ or absolute EtOH. It also recrystallises from MeOH/ether solution. Dry it in a vacuum desiccator over H₂SO₄, then P₂O₅. *Hygroscopic*. [*Beilstein* 4 IV 132.]

2-Dimethylaminoethanol [108-01-0] $C_4H_{11}NO$, M 89.1, b 134.5-135.5°/atm, d_4^{20} 1.430, n_D^{20} 1.4362, pK^{25} 9.23. Dry the amine with anhydrous K₂CO₃ or KOH, and fractionally distil it. [*Beilstein* 4 IV 1424.]

1-(3-Dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (EDCI, DEC, 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride) [25952-53-8] $C_8H_{17}N_3 \cdot HCl$, M 191.7, m 113.5-114.5°, 114-116°, $pK_{Est} \sim 10.3$. It is an excellent H₂O-soluble peptide coupling reagent. It is purified by dissol-

ving (ca 1g) in CH_2Cl_2 (10ml) at room temperature and then adding dry Et_2O (~110ml) dropwise and the crystals that separate are collected, washed with dry Et_2O , recrystallised from $\text{CH}_2\text{Cl}_2/\text{Et}_2\text{O}$ and dried in a vacuum over P_2O_5 . It is important to work in a dry atmosphere or work rapidly, and then dry the solid as soon as possible. The material is moderately *hygroscopic*, but once it becomes wet it reacts slowly with H_2O . Store it away from moisture at -20° to slow down the hydrolysis process. The *free base* has **b 47-48°/0.27mm, 53-54°/0.6mm**, n_D^{25} 1.4582. The *methiodide* is recrystallised from $\text{CHCl}_3/\text{EtOAc}$, the crystals are filtered off, washed with dry Et_2O , recrystallised from $\text{CHCl}_3/\text{Et}_2\text{O}$, and dried *in vacuo* over P_2O_5 , **m 93-95°, 94-95°**. [Sheehan et al. *J Am Chem Soc* **87** 2492 1965, DOI: 10.1021/ja01089a034; Sheehan & Cruickshank *Org Synth Coll Vol* **5** 555 1973, DOI: 10.15227/orgsyn.048.0083.] § A polymer bound version is commercially available.

N,N-Dimethylbiuret [7710-35-2] $\text{C}_4\text{H}_9\text{N}_3\text{O}_2$, **M 131.1, m 178°**. Purify it by repeated crystallisation from the melt, or from H_2O . [Bredereck & Richter *Chem Ber* **99** 2461 1966, DOI: 10.1002/cber.19660990811; Dunnigan & Close *J Am Chem Soc* **75** 3615 1953, DOI: 10.1021/ja01111a001] [*Beilstein* **4** III 574, **4** IV 225.]

2,3-Dimethyl-1,3-butadiene [513-81-5] C_6H_{10} , **M 82.2, m -76°, b 68-69°/760mm, d_4^{20} 0.727, n_D^{20} 1.4385**. Distil it from NaBH_4 , and store with 100ppm of BHT as radical inhibitor. [*Beilstein* **1** IV 1023; Fieser **1** 276.]

1,3-Dimethylbutadiene sulfone (1,3-dimethylsulfolene, **2,4-dimethyl-2,5-dihydrothiophen-1,1-dioxide**) [10033-92-8] $\text{C}_6\text{H}_{10}\text{O}_2\text{S}$, **M 145.2, m 40.4-41.0°**. Crystallise the sulfone from diethyl ether (three times, **m 40.4-41.0°**), or from CCl_4 (**m 39.5-40°**). [For thermal dissociation see Grummitt et al. *J Am Chem Soc* **72** 5167 1950, DOI: 10.1021/ja01167a103; *Beilstein* **17** III/IV 161, **17/1** V 197.]

2,2-Dimethylbutane [75-83-2] C_6H_{14} , **M 86.2, m -100°, b 49.7°/atm, d_4^{20} 0.649, n_D^{25} 1.36595**. Distil it azeotropically with MeOH, then wash it with water, dry (Na_2SO_4) it, and distil it. [*Beilstein* **1** IV 367.]

2,3-Dimethylbutane [79-29-8] C_6H_{14} , **M 86.2, m -129°, b 58.0°/atm, d_4^{20} 1.375, n_D^{25} 1.37231**. Distil it from sodium, pass it through a column of silica gel (activated by heating in nitrogen to 350° before use) to remove unsaturated impurities, and again distil it from sodium. Also distil it azeotropically with MeOH, then wash with water, dry (Na_2SO_4), filter and redistil it. [*Beilstein* **1** IV 371.]

2,3-Dimethylbut-2-ene [563-79-1] C_6H_{12} , **M 84.2, m -75°, b 72-73°/760mm, d_4^{20} 0.708, n_D^{20} 1.41153**. Purify it by GLC on a column of 20% squalene on chromosorb P at 50° [Flowers & Rabinovitch *J Phys Chem* **89** 563 1985, DOI: 10.1021/j100250a003]. Also wash it with 1M NaOH solution followed by H_2O . Dry it over Na_2SO_4 , distil it over powdered KOH under nitrogen and pass it through activated alumina before use. [Woon et al. *J Am Chem Soc* **108** 7990 1986, DOI: 10.1021/ja00285a018; Wong et al. *J Am Chem Soc* **109** 3428 1987, DOI: 10.1021/ja00245a039; *Beilstein* **1** IV 853.]

Dimethylcarbamoyl chloride [79-44-7] $\text{C}_3\text{H}_6\text{ClNO}$, **M 107.5, m -33°, b 34°/0.1mm, d_4^{20} 1.172, n_D^{20} 1.4511**. It must be distilled under high vacuum to avoid decomposition. It is moisture sensitive. [*Beilstein* **4** IV 224.]

Dimethyl carbonate [616-38-6] $\text{C}_3\text{H}_6\text{O}_3$, **M 90.1, m 2-4°, 4.65°, b 89.5°/755mm, 90.2°/atm, 90-91°/atm, d_4^{20} 1.079, n_D^{20} 1.3687**. If the reagent has broad intense bands at 3300cm^{-1} and above (i.e. OH stretching), then it should be purified further. It will contain small amounts of water and/or alcohol which form azeotropes with it. Wash it successively with 10% Na_2CO_3 solution, saturated CaCl_2 , H_2O , and dry it by shaking mechanically for 1 hour with anhydrous CaCl_2 , and fractionate. *Alternatively*, stand it for several days in contact with Linde type 4A molecular sieves, then fractionally distil it. The middle fraction is frozen slowly at 2° , several times, retaining 80% of the liquid at each cycle. [Bowden & Butler *J Chem Soc* **78** 1939, DOI: 10.1039/JR9390000075; Vogel *J Chem Soc* 1833 1948, DOI: 10.1039/JR9480001833; *Beilstein* **3** IV 3.]

Dimethyl dicarbonate (dimethyl pyrocarbonate, DMPC) [4525-33-1] $\text{C}_4\text{H}_6\text{O}_5$, **M 134.1, m 15.2°, b 45-46°/5mm, d_4^{20} 1.2585, n_D^{20} 1.3950**. If the reagent has broad intense bands at 3300cm^{-1} and above (i.e. OH stretching), then it should be purified further. Dissolve it in Et_2O , shake this with a small volume of 0.1N HCl, dry the Et_2O solution (Na_2SO_4), and distil it *in vacuo* at as low pressure as possible, but below 100° , to give a clear liquid. It decomposes to CO_2 and dimethyl carbonate on heating at $123-149^\circ$. It is readily hydrolysed by

H₂O and is a yeast inhibitor. It is an **IRRITANT**. [Brysov et al. *J Org Chem USSR* **10** 2551 1974, Boehm & Mehta *Chem Ber* **71** 1797 1938, DOI: 10.1002/cber.19380710904; *Beilstein* **3** IV 17.]

1,5-Dimethyl-1,5-diazaundecamethylene polymethobromide (Hexadimethrene, Polybrene) [28728-55-4] **M 5000–10,000 polymer**. Purify it by chromatography on Dowex 50 and/or by filtration through alumina before use [Frank *Hoppe-Seyler's Z Physiol Chemie* **360** 997 1979]. It is hygroscopic, and its solubility in H₂O is 10%. The pH of a 1% solution in brine is 5-9. **Toxic**.

Dimethyldioctadecylammonium bromide (dioctadecyldimethylammonium bromide, distearyldimethylammonium bromide) [3700-67-2] **C₃₈H₈₀BrN, M 630.9, m 161-163°**. Crystallise the bromide from acetone, then MeOH [Luka *J Am Chem Soc* **106** 4386 1984, DOI: 10.1021/ja00328a016]. Also purify it by chromatography on alumina by washing with *C₆H₆ and eluting with Me₂CO, evaporating and crystallising from MeCN [Swain & Kreevoy *J Am Chem Soc* **77** 1122 1955, DOI: 10.1021/ja01610a012]. [*Beilstein* **4** IV 829.]

N,N'-Dimethylethylenediamine [1,2-bis(methylamino)ethane] [110-70-3] **C₄H₁₂N₂, M 88.2, b 110-112°/750mm, 119°/760mm, d₄²⁰ 0.819, n_D²⁰ 1.431, (pK²⁵ 7.01 and 9.88)**. This strong base has been prepared in various ways including hydrolysis of *N,N'*-di(benzylsulfonyl)-*N,N'*-dimethylethylenediamine (**m 217-219°**, from AcOH) and concentrated HCl (120-130°) [Johnson & Bailey *J Am Chem Soc* **38** 2135 1916, DOI: 10.1021/ja02267a024], the reaction of 1,2-dibromoethane with 21-33% aqueous MeNH₂ in EtOH (reflux for 2 hours, 50% yield) [Kermack & Wight *J Chem Soc* 1421 1935, DOI: 10.1039/JR9350001421; Woodburn & O'Gee *J Org Chem* **17** 1235 1952, DOI: 10.1021/jo50009a008], hydrolysis of *N,N'*-di(*p*-toluenesulfonyl)-*N,N'*-dimethylethylenediamine (1mol, **m 164°**, from AcOH) with H₂SO₄ (8.2mol) and H₂O (9mol) at 140-145°/7 hours [Boon *J Chem Soc* 307 1947, DOI: 10.1039/JR9470000307], LAH reduction in THF of 3-benzenesulfonyloxy-5,6-dihydrouracil (**m 175-176°**, from iso-PrOH) and isolated as the *dibenzoyl derivative* (41%, **m 177-178°**, from *C₆H₆) [Bauer *J Am Chem Soc* **78** 1945 1956, DOI: 10.1021/ja01590a049]; and by catalytic hydrogenation of *N*-benzyl-*N,N'*-dimethylethylenediamine (**b 73-74°/0.1mm**) with 10% Pd/C [Jucker & Rissi *Helv Chim Acta* **45** 2383 1962, DOI: 10.1002/hlca.19620450710]. General isolation and purification procedures involve steam distillation of the product diamine, acidifying the distillate with HCl, evaporating to dryness, treating the residual salt with 10% excess of cold 32% aqueous NaOH, collecting the organic layer (**care**: very caustic solution), drying it over solid KOH, and distilling from it. Final distillation of the dry base over Na has been reported (see also [107-15-3]). The diamine is a strong base and readily absorbs CO₂ and H₂O from the atmosphere—store it in a dark stoppered bottle, wax the stopper if it should be stored for long periods. The *dihydrochloride* [5752-40-9] has **m 235-236° (dec)**, and the *picrate* has **m 160°** (from EtOH and Me₂CO in rectangular plates). It forms complexes with Cu, Ni, and Pt among other metals. Its FT-IR (neat) has ν_{max} at 3285.5, 2788.2, 1447.2, 1346.5, 1251.0, 1106.5, 1040.5, 876.9 and 763.7 cm⁻¹. [For pK and metal complexes see Gustafson & Martell *J Am Chem Soc* **81** 525 1959, DOI: 10.1021/ja01512a005; *Beilstein* **4** H 250, **4** I 415, **4** II 689, **4** III 512, **4** IV 1171.]

Dimethyl disulfide (DMDS) [624-92-0] **C₂H₆S₂, M 94.2, f -98°, b 40°/12mm, 110°/760mm, d₄²⁰ 1.0605, n 1.5260**. Pass it through neutral alumina before use. [Trost *Chem Rev* **78** 363 1978, DOI: 10.1021/cr60314a002; *Beilstein* **1** IV 1281.]

N,N-Dimethylformamide (DMF) [68-12-2] **C₃H₇NO, M 73.1, m -61°, b 40°/10mm, 61°/30mm, 88°/100mm, 153°/760mm, d₄²⁰ 0.948, n_D²⁵ 1.4269, pK²⁵ -0.3**. DMF decomposes slightly at its normal boiling point to give small amounts of dimethylamine and carbon monoxide. The decomposition is catalysed by acidic or basic materials, so that even at room temperature DMF is appreciably decomposed if allowed to stand for several hours with solid KOH, NaOH or CaH₂. If these reagents are used as dehydrating agents, therefore, they should not be refluxed with the DMF. Use of CaSO₄, MgSO₄, silica gel or Linde type 4A molecular sieves is preferable, followed by distillation under reduced pressure. This procedure is adequate for most laboratory purposes. Larger amounts of water can be removed by azeotropic distillation with *benzene (10% v/v, previously dried over CaH₂), at atmospheric pressure: water and *benzene distil below 80°. The liquid remaining in the distillation flask is further dried by adding MgSO₄ (previously ignited overnight at 300-400°) to give 25g/L. After shaking for one day, a further quantity of MgSO₄ is added, and the DMF is distilled at 15-20mm pressure through a 3-ft vacuum-jacketed column packed with steel helices. However, MgSO₄ is an

inefficient drying agent, leaving about 0.01M water in the final DMF. More efficient drying (to around 0.001-0.007M water) is achieved by standing with powdered BaO, followed by decanting before distillation, then with alumina powder (50g/L, previously heated overnight to 500-600°), and distilling from more of the alumina, or by refluxing at 120-140° for 24 hours with triphenylchlorosilane (5-10g/L), then distilling at *ca* 5mm pressure [Thomas & Rochow *J Am Chem Soc* **79** 1843 1957, DOI: 10.1021/ja01565a021]. Free amine in DMF can be detected by the colour reaction with 1-fluoro-2,4-dinitrobenzene. It has also been purified by drying overnight over KOH pellets and then distilling from BaO through a 10 cm Vigreux column [Jasiewicz et al. *Exp Cell Res* **100** 213 1976, DOI:10.1016/0014-4827(76)90344-X]. [For efficiency of desiccants in drying dimethylformamide see Burfield & Smithers *J Org Chem* **43** 3966 1978, DOI: 10.1021/jo00414a038; and for a review on purification, tests of purity and physical properties, see Juillard *Pure Appl Chem* **49** 885 1977, doi.org.virtual.anu.edu.au/10.1351/pac197749060885.]

It has been purified by distilling from K₂CO₃ under high vacuum and fractionated in an all-glass apparatus. The middle fraction is collected, degassed (seven or eight freeze-thaw cycles) and redistilled under as high a vacuum as possible [Kosower & Mohammad *J Am Chem Soc* **93** 2713 1971, DOI: 10.1021/ja00740a022]. [Beilstein **4** IV 171.]

Rapid purification: Stir over CaH₂ (5% w/v) overnight, filter, then distil at 20mmHg. Store the distilled DMF over 3A or 4A molecular sieves. For solid phase synthesis, the DMF used must be of high quality and free from amines.

2,2-Dimethylglutaric acid (2,2-dimethylpentane-1,5-dioic acid) [681-57-2] C₇H₁₂O₄, M 160.2, m 82-83.5°, 82-86°, b 110-145°/0.1mm, pK₁²⁵ 4.28, pK₂²⁵ 5.42 (H₂O). Crystallise the acid from water, *benzene or ether/petroleum ether. Dry it in a vacuum. The *mono-S-benzylisothiuronium salt* has m 177-178° (from aqueous Me₂CO). [Karrer et al. *Helv Chim Acta* **13** 1084 1930, DOI: 10.1002/hlca.19300130532; Young & Linden *J Am Chem Soc* **69** 2042 1947, DOI: 10.1021/ja01200a066; Oroshnik et al. *J Am Chem Soc* **74** 295, 304 1952, DOI: 10.1021/ja01122a003; Beilstein **2** I 283, **2** II 590, **2** III 1750, **2** IV 2018.]

Trans-d,l-2,4-Dimethylglutaric acid (d,l-2,4-dimethylpentane-1,5-dioic acid) [2121-67-7] C₇H₁₂O₄, M 160.2, m 139-141°, 140-141°, 142.5-143.5°, 144-145°, pK₁ 4.24, pK₂ 5.82. Distil the acid in steam and recrystallise it from ether/petroleum ether. Alternatively, form the Ca salt obtained by adding an equivalent of CaCO₃ until evolution of CO₂ is complete, recrystallise the salt from H₂O (any *cis(meso)* salt which is more soluble remains in solution). The salt is heated in excess of concentrated HCl, and the *trans-(dl)* acid separates as colourless crystals. Filter it off and dry it over KOH *in vacuo*, or recrystallise the dried acid in *C₆H₆/petroleum ether. [Beilstein **2** H 681, **2** I 284, **2** II 590, **2** III 1755, **2** IV 2022.]

Also prepared by the reaction of ethyl 2-bromoisobutyrate with sodio diethyl methylmalonate [von Auwers & Thorpe *Justus Liebigs Ann Chem* **285** 310, 315 1895, DOI: 10.1002/jlac.18952850304], ethyl 2-bromoisobutyrate with sodio ethyl 2-cyanopropionate [Auwers *Chem Ber* **28** 263 1895, DOI: 10.1002/cber.18950280166] or two mols of sodio ethyl 2-cyanopropionate with one mol of CH₂I₂ [Zelinsky *Chem Ber* **22** 2823 1889, DOI: 10.1002/cber.188902202187] which gave, after hydrolysis and decarboxylation, a mixture of approximately equal amounts of *trans-(dl)-* and *cis-(meso)-2,4-dimethylglutaric acids*, m ~105-107° [for discussion see: Bone & Perkin Jr *J Chem Soc* **67** 416 1895, DOI: 10.1039/CT8956700416]. These can be separated by (1) the higher solubility of the Ca salt or (2) the more facile formation of the anhydride of the latter (*meso*) acid. For example, when the mixture is heated with half its weight of acetyl chloride in the cold or warmed gently for 10 min until evolution of HCl is complete, a clear solution is obtained. The *meso-* isomer readily forms the *anhydride* (see below) whereas the *trans* acid remains virtually unchanged. Evaporation of excess AcCl and drying over KOH provides a solid residue which is extracted with *C₆H₆ whereby the *cis-anhydride* is completely removed and leaves behind the solid *trans(d,l)-acid*. After crystallising the latter from hot hydrochloric acid the pure *d,l-acid*, m 140-141°, is obtained. Evaporation of the former *C₆H₆ extract leaves a semi solid (*meso*)-2,4-dimethylglutaric anhydride m 93-94°, 96°, after crystallisation from Ac₂O [Beilstein **17** H 419, **17** II 436, **17/11** V 23.] Upon hydrolysis with concentrated HCl and cooling, pure white crystals of *meso(cis)-2,4-dimethylglutaric acid* [2RS,4SR 3891-70-1] C₇H₁₂O₄, M 160.2, m 125-127°, 127-128°, 128.5°, b 137-138 °/4mm, n_D²⁰ 1.4438, pK₁ 4.28, pK₂ 5.80, separate. [Bone & Perkin jr *J Chem Soc* **67** 416 1895, DOI: 10.1039/CT8956700416; **68** 253 1896, DOI: 10.1039/CT8966900253.] Note that attempted equilibration of the *cis-anhydride* by boiling for 2 hrs with Na in EtOH followed by addition of KOH in H₂O and boiling further for 1 hr to obtain large amounts of *trans-acid* was unfavourable. Thus 153g of *cis-anhydride* gave only 18g of *trans-(d,l)-acid*. [see Allinger *J Am Chem Soc* **81** 232 1959, DOI: 10.1021/ja01510a053].

The (\pm)-*diethyl ester* [34022-15-6] has **b** 231°/atm.

Diethyl *d,l*-2,4-dimethylglutarate [34022-15-6] $C_{11}H_{20}O_4$, **M** 216.2, has **b** 231°/atm, and the ***d,l*-2,4-dimethylglutaric anhydride** [3891-69-8] $C_7H_{10}O_3$, **M** 142.2, has **m** 33-34°. **2*R*,4*R*-(-)-2,4-Dimethylglutaric acid** [24018-75-5] has **m** 79.5-80° (needles from H_2O), $[\alpha]_D^{20}$ -39.8 (H_2O), **2*S*,4*S*-(+)-2,4-dimethylglutaric acid** has **m** 79-80° (needles from H_2O), $[\alpha]_D^{20}$ +39.8 (H_2O), and **2*R*,4*R*-(-)-2,4-dimethylglutaric anhydride** $C_7H_{10}O_3$, **M** 142.2, has **m** 41.5-42.5° (petroleum ether), $[\alpha]_D^{20}$ +69.6 ($*C_6H_6$).

3,3-Dimethylglutaric acid [4839-46-7] $C_7H_{12}O_4$, **M** 160.2, **m** 103-104°, **b** 89-90°/2mm, 126-127°/4.5mm, **pK**₁²⁵ 3.85, **pK**₂²⁵ 6.45. Crystallise the acid from water, EtOH (needles), *benzene or ether/petroleum ether (prisms). Dry it in a vacuum. On a large scale it is better to boil it under reflux in Ac_2O for 7 hrs, evaporating, and distilling the residual liquid to give the ***anhydride*** (**m** 125°, **b** 156-156°/20mm, 181°/25mm, see below) which solidified on cooling. This is hydrolysed by boiling with 3 moles of aqueous KOH solution (64 w/w%) for 2.5 hrs, whereby a clear solution (containing the di potassium salt) is formed. Acidification with dilute H_2SO_4 and salting out with ammonium sulphate, extracting into Et_2O , drying (Na_2SO_4), evaporating, and a residual solid which is recrystallised from concentrated HCl. The desired glutaric acid that crystallises out is collected, and dried at 70°. It can be further recrystallised as above to give pure acid **m** 103-104°, or distilled in a vacuum; or sublimed in a vacuum. The esters are prepared in the usual way by boiling the acid in a mixture of the required pure alcohol and *benzene with some concentrated H_2SO_4 for a few hours. After workup, ***methyl 3,3-dimethylglutarate*** has **b** 111°/20mm, d_4^{20} 1.0270, n_D^{20} 1.42897; and ***ethyl 3,3-dimethylglutarate*** has **b** 126°/17mm, d_4^{20} 0.9893, n_D^{20} 1.4287. [Thole & Thorpe *J Chem Soc* 99 422 1911, DOI: 10.1039/CT9119900422; Vogel *J Chem Soc* 1758 1934, DOI: 10.1039/JR9340001758.] [Beilstein 21 H 391, 21 I 334, 21 II 309, 21 III/IV 4601, 21/9 V 592.]

2,2-Dimethylglutaric anhydride [2938-48-9] $C_7H_{10}O_3$, **M** 142.2, **m** 34-38°, 38.5°, 38-40°, **b** 150°/15mm, 158-160°/25mm, 175-180°/60mm, 270°/atm. First check the IR to see if OH frequencies at *ca* 3000 cm^{-1} are present. If present then reflux with $AcCl$ until evolution of HCl ceases, distil and collect the fraction at **b** 126°/10mm which should not have bands for OH and should solidify with **m** 37-38°. [Rothstein & Schofield *J Chem Soc* 4566 1965, DOI: 10.1039/JR9650004566; Beilstein 17/11 V 23.] Ac_2O has also been used for dehydration by refluxing for 6 hrs [Perkin, jun. *J Chem Soc* 81 246 (252) 1902, DOI: 10.1039/CT9028100246; Wheeler & Almeida *J Org Chem* 27 4448 1962, DOI: 10.1021/jo01059a079.]

3,3-Dimethylglutaric anhydride [4160-82-1] $C_7H_{10}O_3$, **M** 142.2, **m** 125°, 124-126°, **b** 156-156°/20mm, 181°/25mm. Some hydrolysis may have occurred (check IR), so purify best by boiling with redistilled Ac_2O for a few hours (e.g. 3–7 hrs, depending on how much acid is present), distilling and recrystallising the solid distillate from petroleum ether (**b** 100-120°), $*C_6H_6$ /petroleum ether or Ac_2O . [Perkin *J Chem Soc* 69 1457 1896, DOI: 10.1039/CT8966901457; Perkin & Thorpe *J Chem Soc* 75 48 1899, DOI: 10.1039/CT8997500048; Thorpe & Wood *J Chem Soc* 103 276 1913, DOI: 10.1039/CT9130300276, Thorpe & Wood *J Chem Soc* 103 1592 1913, DOI: 10.1039/CT9130301586; Bruice & Bradbury *J Am Chem Soc* 87 4838 1965, DOI: 10.1021/ja00949a030.] [Beilstein 17 H 419, 17 I 230, 17 III/IV 5853, 17/11 V 24.]

Dimethylglyoxime [95-45-4] $C_4H_8N_2O_2$, **M** 116.1, **m** 240°, **pK**₁²⁵ 10.60, **pK**₂²⁵ 11.85. Crystallise it from EtOH (10ml/g) or aqueous EtOH. [Beilstein 1 III 3105.] **TOXIC.**

2,5-Dimethyl-1,5-hexadiene [627-58-7] C_8H_{14} , **M** 110.2, **m** -75.6°, **b** 114.3°/atm, d_4^{20} 0.7423, n_D^{20} 1.4293. This olefin was prepared by condensation of allyl chloride and methallyl chloride over Mg in Et_2O which provided a mixture of 1,5-hexadiene (**b** 59.4°/atm, 12%), 2-methyl-1,5-hexadiene (**b** 88.1°/atm, 47%), and 2,5-dimethyl-1,5-hexadiene (**b** 114.3°/atm, 30%) which were separated by fractional distillation [Henne et al. *J Am Chem Soc* 63 3474 1941, DOI: 10.1021/ja01857a062]. [Beilstein 1 H 259, 1 I 122, 1 II 237, 1 III 1010, 1 IV 1042.]

2,5-Dimethyl-2,4-hexadiene [764-13-6] C_8H_{14} , **M** 110.2, **m** 14.5°, **b** 132-134°/atm, d_4^{20} 0.773, n_D^{20} 1.4796. Distil it, then repeatedly fractionally crystallise it by partial freezing. Immediately before use, the diene is passed through a column containing Woelm silica gel (activity I) and Woelm alumina (neutral) in separate layers. [Beilstein 1 IV 1043.]

2,2-Dimethylhexane [590-73-8] C_8H_{18} , M 114.2, m -121.2°, b 107°/atm, d_4^{20} 0.695. Dry the hexane over type 4A molecular sieves and distil it. [Beilstein 1 IV 432.] **2,5-Dimethylhexane** [592-13-2] C_8H_{18} , M 114.2, m -91.2°, b 109°/atm, d_4^{20} 0.694, is similarly purified. [Beilstein 1 IV 434.]

2,5-Dimethylhexane-2,5-diol [110-03-2] $C_6H_{18}O_2$, M 146.2, m 88-90°, b 214-215°/atm. Purify the diol by fractional crystallisation. Then the diol is dissolved in hot acetone, treated with activated charcoal, and filtered while hot. The solution is cooled and the diol is filtered off and washed well with cold acetone. The crystallisation process is repeated several times, and the crystals are dried under a vacuum in a freeze-drying apparatus [Goates et al. *JCS Faraday Trans 1* **78** 3045 1982, DOI: 10.1039/F19827803045]. [Beilstein 1 IV 2600.]

1,1-Dimethylhydrazine [57-14-7] $C_2H_8N_2$, M 60.1, b 60.1°/702mm, d_4^{20} 0.790, n_D^{20} 1.408 pK^{30} 7.21. Fractionally distil the hydrazine through a 4-ft column packed with glass helices. Precipitate it as its oxalate salt from diethyl ether solution. After crystallising from 95% EtOH, the salt is decomposed with aqueous saturated NaOH, and the free base is distilled, dried over BaO and redistilled [McBride & Kruse *J Am Chem Soc* **79** 572 1957, DOI: 10.1021/ja01560a020]. Distillation and storage should be under nitrogen as it is a moderately strong organic base. [Beilstein 4 IV 3322.]

Dimethyl itaconate [617-52-7] $C_7H_{10}O_4$, M 158.2, m 38°, 37-41°, b 208°/atm, d_4^{20} 1.124. Crystallise the ester from MeOH by cooling to -78°. Alternatively, distil it at atmospheric pressure. [Beilstein 2 IV 2229.]

Dimethylmalonic acid [595-46-0] $C_5H_8O_4$, M 132.1, m 192-193° pK_1^{25} 3.03, pK_2^{25} 5.73. Recrystallise the acid from *benzene/petroleum ether. It sublimes in a vacuum with slight decomposition. [Beilstein 2 IV 1955.]

Dimethylmaleic anhydride [766-39-2] $C_6H_6O_3$, M 126.1, m 96°, b 225°/760mm. It can be purified by distillation, by recrystallisation from *benzene/ligroin and/or sublimation in a vacuum. [Beilstein 17/11 V 69.]

Dimethylnitrosamine (N-nitrosodimethylamine) [62-75-9] $C_2H_6N_2O$, M 74.0, m -28°, b 149-150°/atm, 151°/767mm, 153°/774mm, d_4^{20} 1.006, n_D^{20} 1.4370. Dry the nitrosamine over anhydrous K_2CO_3 or dissolve it in Et_2O , dry it over solid KOH, filter, evaporate Et_2O and distil the yellow oily residue through a 30cm fractionating column discarding the first fraction which may contain Me_2N . Also dry over $CaCl_2$ and distil it at atmospheric pressure. All operations should be done in an efficient fume cupboard as the vapors are **TOXIC** and **CARCINOGENIC**. [Fischer *Chem Ber* **8** 1587 1875, DOI: 10.1002/cber.187500802203; Romburg *Recl Trav Chim Pays-Bas* **5** 246 1886, DOI: 10.1002/recl.18860050803; Vogel *J Chem Soc* 1825 1948, DOI: 10.1039/JR9480001825; Hatt *Org Synth Coll Vol* **2** 211 1961, DOI: 10.15227/orgsyn.016.0022; Krebs & Mandt *Chem Ber* **108** 1130 1975, DOI: 10.1002/cber.19751080419; Beilstein 4 H 84, 4 I 241, 4 II 585, 4 III 166, 4 IV 3384.] Trace amounts are found in tobacco smoke, cured meats, e.g. bacon, smoked and salted fish.

Dimethylolurea [di-(hydroxymethyl)urea] [140-95-4] $C_3H_8N_2O_3$, M 120.1, m 137-139°. Crystallise it from aqueous 75% EtOH. As it is very soluble in cold H_2O , hot EtOH and MeOH, dissolve it in the minimum of cold H_2O then add hot EtOH and cool. It has antiseptic and pesticidal activity. [Einhorn & Hamburger *Chem Ber* **41** 24 1908, DOI: 10.1002/cber.19080410107; Beilstein 3 IV 107.]

Dimethyl oxalate [553-90-2] $C_4H_6O_4$, M 118.1, m 54°, b 163-165°/atm, d_4^{20} 1.148. Crystallise the ester repeatedly from EtOH. De-gas it under nitrogen at high vacuum and distil it. [Beilstein 2 IV 1847.]

2,4-Dimethylpentane [108-08-7] C_7H_{16} , M 100.2, m -123.4°, b 80.5°/am, d_4^{20} 0.763, n_D^{20} 1.3814, n_D^{25} 1.37882. Extract it repeatedly with concentrated H_2SO_4 , wash with water (CARE), dry and distil it. Alternatively, percolate through silica gel (previously heated in N_2 to 350°). Purify it by azeotropic distillation with EtOH, followed by washing out the EtOH with water, drying and distilling. [Beilstein 1 IV 406.]

4,4-Dimethyl-1-pentene [762-62-9] C_7H_{14} , M 98.2, b 72.5°/760mm, d_4^{20} 0.6827, n_D^{20} 1.3918. Purify it by passing through alumina before use [Traylor et al. *J Am Chem Soc* **109** 3625 1987, DOI: 10.1021/ja00246a019]. [Beilstein 1 IV 869.]

Dimethyl peroxide [690-02-8] $\text{C}_2\text{H}_6\text{O}_2$, M 62.1, b 13.5°/760mm, d_4^{20} 0.8677, n_D^{20} 1.3503. Purify dimethyl peroxide by repeated trap-to-trap fractionation until no impurities could be detected by gas IR spectroscopy [Haas & Oberhammer *J Am Chem Soc* 106 6146 1984, DOI: 10.1021/ja00333a004]. *All necessary precautions should be taken in case of EXPLOSION.* [Beilstein 1 II 271, 1 III 1194, 1 IV 1249.]

2,2-Dimethyl-1,3-propanediol (neopentyl glycol) [126-30-7] $\text{C}_5\text{H}_{12}\text{O}_2$, M 104.2, m 128.4-129.4°, b 208°/760mm. Crystallise the diol from *benzene or acetone/water (1:1). [Beilstein 1 IV 2551.]

2,2-Dimethyl-1-propanol (neo-pentyl alcohol) [75-84-3] $\text{C}_5\text{H}_{12}\text{O}$, M 88.2, m 52°, b 113.1°/760mm. It is difficult to distil because it is a solid at ambient temperatures. Purify it by fractional crystallisation and by sublimation. [Beilstein 1 IV 1690.]

***N,N*-Dimethylpropionamide** [758-96-3] $\text{C}_5\text{H}_{11}\text{NO}$, M 101.2, m -45°, b 175-178°/atm, d_4^{20} 0.920, n_D^{20} 1.440. Shake the amide with BaO for 1-2 days, then distil at atmospheric or reduced pressure. [Beilstein 4 III 126.]

2,2-Dimethylsuccinic acid [597-43-3] $\text{C}_6\text{H}_{10}\text{O}_4$, M 146.1, m 141°, pK_1^{20} 4.15, pK_2^{20} 6.40. Crystallise the acid from EtOH/ether or EtOH/chloroform. [Beilstein 2 IV 1996.]

***meso*-2,3-Dimethylsuccinic acid** [608-40-2] $\text{C}_6\text{H}_{10}\text{O}_4$, M 146.1, m 200° (dec), 211°, pK_1^{25} 3.77, pK_2^{25} 5.36. Crystallise the *meso*-acid from EtOH/ether or EtOH/chloroform.

(±)-2,3-Dimethylsuccinic acid [13545-04-5] $\text{C}_6\text{H}_{10}\text{O}_4$, M 146.1, m 129°, pK_1^{25} 3.82, pK_2^{25} 5.98. Crystallise the *racemic* acid from water. [Beilstein 2 IV 1998.]

Dimethyl sulfide (DMS) [75-18-3] $\text{C}_2\text{H}_6\text{S}$, M 62.1, f -98.27°, b 0°/172mm, 37.5-38°/760mm, d^{21} 0.8458, n^{25} 1.4319. Purify dimethyl sulfide *via* the Hg(II) chloride complex by dissolving 1 mole of Hg(II)Cl₂ in 1250ml of EtOH and slowly adding the boiling alcoholic solution of Me₂S to give the right ratio for 2(CH₃)₂S.3HgCl₂. After recrystallisation of the complex to constant melting point, 500g of complex are heated with 250ml conc HCl in 750ml of water. The sulfide is separated, washed with ice-water, and dried (CaCl₂ and CaSO₄). Finally, it is distilled under reduced pressure from sodium. Precautions should be taken (*efficient fume hood*) because of its very UNPLEASANT ODOUR and TOXICITY. [Beilstein 1 IV 1275.] **Dimethyl sulfone** [67-71-0] $\text{C}_2\text{H}_6\text{O}_2\text{S}$, M 94.1, m 109°, b 238°/atm, is recrystallised from water. Dry it over P₂O₅. [Beilstein 1 IV 1279.]

2,4-Dimethylsulfolane (2,4-dimethyltetrahydrothiophene 1,1-dioxide) [1003-78-7] $\text{C}_6\text{H}_{12}\text{O}_2\text{S}$, M 148.2, b 123°/70mm, 128°/77mm, 280-281°/atm (with some dec), d^{25} 1.1314, n_D^{20} 1.474. Distil the yellow liquid sulfolane in a vacuum. It is soluble in hydrocarbon solvents and slightly soluble in H₂O. [Beilstein 17/1 IV 97.]

Dimethyl sulfoxide (DMSO) [67-68-5] $\text{C}_2\text{H}_6\text{OS}$, M 78.1, m 18.0-18.5°, b 75.6-75.8°/12mm, 190°/760mm, d_4^{20} 1.100, n_D^{20} 1.479. This colourless, odourless, very *hygroscopic* liquid, is synthesised from dimethyl sulfide. The main impurity is water, with a trace of dimethyl sulfone. The Karl-Fischer test is applicable. It is dried with Linde types 4A or 13X molecular sieves, by prolonged contact and passage through a column of the material, then distilled under reduced pressure. Other drying agents include CaH₂, CaO, BaO and CaSO₄. It can also be fractionally crystallised by partial freezing. More extensive purification is achieved by standing overnight with freshly heated and cooled chromatographic grade alumina. It is then refluxed for 4 hours over CaO, dried over CaH₂, and fractionally distilled at low pressure. For efficiency of desiccants in drying dimethyl sulfoxide see Burfield and Smithers [*J Org Chem* 43 3966 1978, DOI: 10.1021/jo00414a038; Martin et al. *Angew Chem Int Ed* 6 318 1967, DOI: 10.1002/anie.196703181;]. [Reddy *Pure Appl Chem* 25 457 1971, DOI: 10.1351/pac197125020457; Beilstein 1 IV 1277.]

Rapid purification: Stand over freshly activated alumina, BaO or CaSO₄ overnight. Filter and distil it over CaH₂ under reduced pressure (~ 12 mm Hg). Store it over 4A molecular sieves.

***N,N*-Dimethylthiocarbamoyl chloride (DMTCC)** [16420-13-6] $\text{C}_3\text{H}_6\text{CINS}$, M 123.6, m 42-43°, b 64-65°/0.1mm. Crystallise it twice from pentane and/or distil it at low pressure. [Ponaras & Zaim *J Heterocycl Chem* 44 487 2007, DOI: 10.1002/jhet.5570440236; Beilstein 4 III 147.] **TOXIC VAPOURS.**

sym-Dimethylurea [96-31-1] $\text{C}_6\text{H}_8\text{N}_2\text{O}$, **M 88.1, m 106°, b 260-270°/atm.** Crystallise the urea from acetone/diethyl ether by cooling in an ice bath. Also crystallise it from EtOH and dry it at 50°/5mm for 24 hours [Bloemendahl & Somsen *J Am Chem Soc* **107** 3426 1985, DOI: 10.1021/ja00298a005]. [*Beilstein* **4** IV 207.]

2,2-Dinitropropane [595-49-3] $\text{C}_3\text{H}_6\text{N}_2\text{O}_4$, **M 162.1, m 53.5°, 53.7-54.3°.** Crystallise it from EtOH or MeOH. Dry it over CaCl_2 or *in vacuo* for 1 hour just above the melting point. Note that it has been purified by slow vacuum sublimation in the presence of P_2O_5 [Whittaker *J Phys Chem* **62** 267 1958, DOI: 10.1021/j150561a004]. [*Beilstein* **1** H 117, **1** II 79, **1** III 261, **1** IV 234.]

N,N-Dioctadecyl methylamine (proton ionophore III) [4088-22-6] $\text{C}_{37}\text{H}_{77}\text{N}$, **M 536.0, m 40°, 44-46°, 48-49°, b 252-259°, $\text{pK}_{\text{Est}} \sim 10$.** It can be distilled at high vacuum, but dissolving in $^*\text{C}_6\text{H}_6$, filtering and evaporating gives a waxy solid suitable for electrode use. It recrystallises from Me_2CO or MeCN. [Hoerr et al. *J Org Chem* **09** 201 1944, DOI: 10.1021/jo01184a011; Wu & Yu *Talanta* **34** 577 1987, DOI:10.1016/0039-9140(87)80193-5; *Beilstein* **4** III 435.]

Di-n-propyl ketone (4-heptanone) [123-19-3] $\text{C}_7\text{H}_{14}\text{O}$, **M 114.2, m -33°, b 143.5°, d_4^{20} 0.8143, n_D^{20} 1.40732.** Dry 4-pentanone with CaSO_4 , then distil it from P_2O_5 under nitrogen. [*Beilstein* **1** IV 3323.]

Di-n-propyl sulfide [111-47-7] $\text{C}_7\text{H}_{14}\text{S}$, **M 118.2, m -103°, b 141-142°/atm, d_4^{20} 0.870, n_D^{20} 1.449.** Wash the sulfide with aqueous 5% NaOH, then water. Dry it with CaCl_2 and distil it from Na [Dunstan & Griffiths *J Chem Soc* 1344 1962, DOI: 10.1039/JR9620001344]. [*Beilstein* **1** IV 1452.]

S-1,2-Distearin (S-glycerol-1,2-distearate) [*S*- 1429-59-0, *RS*- 51063-97-9, *R*- 1188-58-5] $\text{C}_{39}\text{H}_{76}\text{O}_5$, **M 625.0, m 76-77°, $[\alpha]_D^{20}$ -2.8 (c 6.3, CHCl_3), $[\alpha]_{546}^{20}$ +1.4 (c 10, $\text{CHCl}_3/\text{MeOH}$, 9:1).** Crystallise the glyceride from chloroform/petroleum ether. [cf. p 1014, in 'Physiologically Active ...' in Chapter 6, *Beilstein* **2** IV 1231.]

1,4-Dithioerythritol (DTE, erythro-2,3-dihydroxy-1,4-dithiobutane) [6892-68-8] $\text{C}_4\text{H}_{10}\text{O}_2\text{S}_2$, **M 154.3, m 82-84°, pK_1 9.0, pK_2 9.9.** Crystallise DTE from ether/hexane and store it in the dark at 0°. [*Beilstein* **1** III 2360.]

Dithiooxamide (rubeanic acid) [79-40-3] $\text{C}_2\text{H}_4\text{O}_2\text{S}_2$, **M 120.2, m >300°.** Crystallise dithiooxamide from EtOH and sublime it at high vacuum. [*Beilstein* **2** IV 1871.]

RS-1,4-Dithiothreitol (DTT, Cleland's reagent) [3483-12-3, 27565-41-9] $\text{C}_4\text{H}_{10}\text{O}_2\text{S}_2$, **M 154.3, m 42-43°, pK_1 8.3, pK_2 9.5.** Crystallise DTT from ether and sublime it at 37°/0.005mm. It should be stored at 0°. [*Beilstein* **1** III 2360.] It is also available as a 0.1M solution in H_2O .

All-cis-4,7,10,13,16,19-Docosahexaenoic acid (DHA, Cervonic acid) [6217-54-5] $\text{C}_{22}\text{H}_{32}\text{O}_2$, **M 328.5, m -44.1°, -44.7° to -44.5°, n_D^{20} 1.5017, $\text{pK}_{\text{Est}} \sim 4.6$.** Its solubility in CHCl_3 is 5%. It has been purified from fish oil by GLC using Argon as mobile phase and EGA as stationary phase with an ionisation detector [UV: Stoffel & Ahrens *J Lipid Res* **1** 139 1959], and *via* the ester by evaporative molecular distillation using a 'continuous molecular still' at 10^{-4} mm with the highest temperature being 110° and a total contact time with the hot surface being 60 seconds [Farmer & van den Heuvel *J Chem Soc* 427 1938, DOI: 10.1039/JR9380000427]. It was isolated in quantity from cod liver oil by Wright, Kuo & Corey [*J Org Chem* **52** 4399 1987, DOI: 10.1021/jo00228a050] who separated it from eicosapentaenoic acid [10417-94-4] *via* its ability to form an iodo- γ -lactone. The **methyl ester** [2566-90-7] has **b 208-211°/2mm, d_4^{20} 0.9398, n_D^{20} 1.5035.** With Br_2 it forms a **dodecabromide m ca 240°(dec).** Also, the acid was converted to the methyl ester and purified through a three-stage molecular still [as described by Sutton *Chem Ind (London)* 11383 1953] at 96°, and the rate was adjusted so that one-third of the material was removed each cycle of three distillations. The distillate (numbered 4) (13g) was dissolved in EtOH (100ml containing 8g of KOH) at -70° and set aside for 4 hours at 30° with occasional shaking under a vacuum. Water (100ml) was added and the solution was extracted with pentane, washed with HCl, dried (MgSO_4), filtered and evaporated to give a clear oil (11.5g) **m -44.5° to -44.1°.** In the catalytic hydrogenation of the oil six mols of H_2 are absorbed and **docosanoic acid (behenic acid)** is produced with **m 79.0-79.3°** undepressed with an authentic sample (see docosanoic acid below) [Whitcutt *Biochem J* **67** 60 1957, DOI: 10.1042/bj0670060]. [*Beilstein* **2** IV 1812.]

Docosane (C22) [629-97-0] $C_{22}H_{46}$, **M 310.6, m 47°, b 224°/15mm**. Crystallise docosane from EtOH or ether. Disdistil it in a vacuum. [*Beilstein* 1 IV 572.]

Docosanoic acid (behenic acid) [112-85-6] $C_{22}H_{44}O_2$, **M 340.6, m 81-82°, pK_{Est} ~4.9**. Crystallise the acid from ligroin. [Francis & Piper *J Am Chem Soc* **61** 577 1939, DOI: 10.1021/ja01872a011; *Beilstein* 2 IV 1290.]

1-Docosanol (behenyl alcohol) [661-19-8] $C_{22}H_{46}O$, **M 182.3, m 70.8°**. Crystallise docosanol from ether or chloroform/ether. [*Beilstein* 1 IV 1906.]

n-Dodecane [112-40-3] $C_{12}H_{26}$, **M 170.3, m -9.6°, b 97.5-99.5°/5mm, 216°/760mm, d₄²⁰ 0.748, n_D²⁰ 1.42156**. Pass it through a column of Linde type 13X molecular sieves. Store it in contact with, and distil it from sodium. Pass it through a column of activated silica gel. It has been crystallised from diethyl ether at -60°. Unsaturated dry material which remained after passage through silica gel has been removed by catalytic hydrogenation (Pt₂O) at 45lb/in² (3.06 atmospheres), followed by fractional distillation under reduced pressure [Zook & Goldey *J Am Chem Soc* **75** 3975 1953, DOI: 10.1021/ja01112a029]. It has also purified by partial crystallisation from the melt. [*Beilstein* 1 IV 498.]

Dodecane-1,10-dioic acid (decane-1,10-dicarboxylic acid) [693-23-2] $C_{12}H_{22}O_4$, **M 230.3, m 129°, b 245°/10mm, pK_{Est} ~4.8**. Crystallise the dioic acid from water, 75% or 95% EtOH (solubility is ~10%), or glacial acetic acid. Dry it *in vacuo*. [*Beilstein* 2 IV 2126.]

1-Dodecanol (dodecyl alcohol, lauryl alcohol) [112-53-8] $C_{12}H_{26}O$, **M 186.3, m 24°, b 91°/1mm, 135°/10mm, 167°/40mm, 213°/200mm, 259°/760mm, d₄²⁴ 0.8309 (liquid)**. Crystallise dodecanol from aqueous EtOH, and distil it through a spinning-band column under vacuum. [Ford & Marvel *Org Synth* **10** 62 1930, DOI: 10.15227/orgsyn.010.0062; *Beilstein* 1 IV 1844.]

1-Dodecanthiol (lauryl mercaptan) [112-55-0] $C_{12}H_{26}S$, **M 202.4, b 111-112°/3mm, 153-155°/24mm, d₄²⁰ 0.844, n_D²⁰ 1.458, pK_{Est} ~10.8**. Dry it with CaO for several days, then distil it from CaO. [*Beilstein* 1 IV 1851.]

Dodecylamine (lauryl amine) [124-22-1] $C_{12}H_{27}N$, **M 185.4, m 28°, 27-29°, 120-121°/2mm, 134°/15mm, 156°/33mm, pK₂₅ 10.63**. Fractionally distil the amine, preferably under N₂ and in a vacuum. Store it in the absence of CO₂. It can be recrystallised from *n*-hexane at low temperature. The **hydrochloride** crystallises from Me₂CO (**m 182-183°**) or CHCl₃/petroleum ether (**m 185-187°**). [Magnien & Baltzly *J Org Chem* **23** 2029 1958, DOI: 10.1021/jo01106a630; *Beilstein* 4 H 200, III 406, 4 IV 794.] **Dodecylammonium butyrate** [17615-97-3] **M 273.4, m 39-40°, 39-41°, pK₂₅ 10.63 (for free base)**, is recrystallised from *n*-hexane. [Kitahara *Bull Chem Soc Jpn* **28** 234 1955, DOI: org/10.1246/bcsj.28.234; *Beilstein* 4 III 409, 4 IV 797.] **Dodecylammonium propionate** [17448-65-6] **M 259.4, m 55-56°**, is recrystallised from hexanol/petroleum ether (b 60-80°). [Kitahara *Bull Chem Soc Jpn* **28** 234 1955, DOI: org/10.1246/bcsj.28.234; *Beilstein* 4 III 409, 4 IV 797.]

Dodecyldimethylamine N-oxide (DDAO, DDMAO) [1643-20-5] $C_{14}H_{31}NO$, **M 229.4, m 102°, 130-131°**. Crystallise the oxide from acetone, ethyl acetate or dry toluene which forms very hygroscopic needles. It is available as a ~30% solution in H₂O, and has a detergent action. **IT IS AN EYE AND SKIN IRRITANT**. [Bunton et al. *J Org Chem* **52** 3832 1987, DOI: 10.1021/jo00226a020; *Beilstein* 4 III 410, 4 IV 798.]

Dodecyl ether (didodecyl ether, lauryl ether) [4542-57-8] $C_{24}H_{50}O$, **M 354.6, m 32.5-33°, 33°, b 175°/0.15mm, d₃₆ 0.8127, n₃₉ 1.4393**. Distil the ether in a vacuum, then crystallise it from MeOH or MeOH/*benzene. [Mannich & Nadelmann *Chem Ber* **63** 796 1930, DOI: 10.1002/cber.19300630410; Butterworth & Hey *J Chem Soc* 388 1940, DOI: 10.1039/JR9400000388; *Beilstein* 1 III 1785, 1 IV 1846.]

Dodecyl methacrylate (lauryl methacrylate) [142-90-5] $C_{16}H_{30}O_2$, **M 254.4, m -7°, b 142°/4mm, d₂₅ 0.8717, n_D²⁵ 1.4330**. Purify the ester by fractional distillation in a high vacuum. Add 0.05% of hydroquinone monomethyl ether as stabiliser. [Rehberg & Fischer *Ind Eng Chem* **40** 1429 1948, DOI: 10.1021/ie50464a019; *Beilstein* 2 III 1290, 2 IV 1528.]

Dodecyltrimethylammonium bromide [1119-94-4] $C_{15}H_{34}BrN$, **M 308.4, m 246°(dec)**. Purify the salt by repeated crystallisation from acetone. Wash it with diethyl ether and dry it in a vacuum oven at 60° [Dearden & Wooley *J Phys Chem* **91** 2404 1987, DOI: 10.1021/j100293a040]. [*Beilstein* **4** IV 798.] **Dodecyltrimethylammonium chloride** [112-00-5] $C_{15}H_{34}ClN$, **M 263.9, m 246°(dec)**, is purified by dissolving in MeOH, treating with active charcoal, filtering and drying *in vacuo* [Walderhaug *J Phys Chem* **88** 1655 1984, DOI: 10.1021/j150652a043], or recrystallising several times from 10% EtOH in acetone. It has also been repeatedly crystallised from EtOH/ether or MeOH. [Cella et al. *J Am Chem Soc* **74** 2061 1952, DOI: 10.1021/ja01128a061; *Beilstein* **4** IV 79.] A ~50% solution in isopropanol/H₂O (4:1) is commercially available.

Eicosane (C20) [112-95-8] $C_{20}H_{42}$, **M 282.6, m 36-37°, b 205°/15mm, d^{36.7} 0.7779, n⁴⁰ 1.43453**. Crystallise eicosane from EtOH, and/or distil it in a vacuum. [*Beilstein* **1** IV 563.]

Cis-5,8,11,14,17-Eicosapentaenoic acid (EPA, Omega-3, Timnodonic acid) [10417-94-4] $C_{20}H_{30}O_2$, **M 302.5, m -54° to -54°, d²⁵ 0.943, n²⁰ 1.49865**. It is present in cod liver oil together with *docosahexanoic acid* which can be separated from it because it does not form an iodo- γ -lactone (see docosahexanoic acid above) [Wright et al. *J Org Chem* **52** 4399 1987, DOI: 10.1021/jo00228a050; Klenk & Eberhagen *Z Physiol Chem* **307** 42 1957]. Purification is as for docosahexanoic acid above using GLC and/or molecular distillation. [*Beilstein* **2** IV 1808.]

Elaidic (trans-oleic) acid [112-79-8] $C_{18}H_{34}O_2$, **M 282.5, m 44.5°, b 288°/100mm, pK²⁵ 4.9**. Crystallise the acid from acetic acid, then EtOH. [*Beilstein* **2** IV 1647.]

Erucic acid (cis-13-docosenoic acid) [112-86-7] $C_{22}H_{42}O_2$, **M 338.6, m 33.8°, b 358°/400mm, pK_{Est} ~4.9**. Crystallise erucic acid from MeOH. [*Beilstein* **2** IV 1676.]

Ethane [74-84-0] C_2H_6 , **M 30.1, f -172°, b -88°/atm, d₄⁰ 1.0493 (air = 1)**. Ethylene can be removed by passing the gas through a sintered-glass disc into fuming H₂SO₄ then slowly through a column of charcoal saturated with bromine. Bromine and HBr are removed by passage through firebrick coated with *N,N*-dimethyl-*p*-toluidine. The ethane is also passed over KOH pellets (to remove CO₂) and dried with Mg(ClO₄)₂. Further purification is by several distillations of liquified ethane, using a condensing temperature of -195°. Yang and Gant [*J Phys Chem* **65** 1861 1961, DOI: 10.1021/j100827a042] treated ethane by standing it for 24 hours at room temperature in a steel bomb with activated charcoal treated with bromine. They then immersed the bomb in a Dry-ice/acetone bath and transferred the ethane to an activated charcoal trap cooled in liquid nitrogen. (The charcoal had previously been degassed by pumping for 24 hours at 450°.) By allowing the trap to warm slowly, the ethane distils, and only the middle third fraction is kept. Removal of methane is achieved using Linde type 13X molecular sieves (previously degassed by pumping for 24 hours at 450°) in a trap which, after cooling in Dry-ice/acetone, is saturated with ethane. After pumping for 10 minutes, the ethane is recovered by warming the trap to 25°. (The final gas contains less than 10⁻⁴ mole % of either ethylene or methane). [*Beilstein* **1** IV 108.]

Ethanesulfonyl chloride [594-44-5] $C_2H_5ClO_2S$, **M 128.6, b 55°/9mm, 62°/12mm, 74°/19mm, 76-79°/22mm, 95-98°/50mm, 177°/760mm, d₄²⁰ 1.357, n_D²⁰ 1.4539**. Purify the sulfonyl chloride by repeated distillation to remove HCl formed from hydrolysis. **It is a fuming, corrosive liquid, handle in a good fumehood.** It is hydrolysed by aqueous N NaOH at room temperature and is best stored in aliquots in sealed ampoules under N₂. [Davies & Dick *J Chem Soc* 484 1932, DOI: 10.1039/JR9320000483; Klamann & Drahowzal *Monatsh Chem* **83** 463 1952, DOI: 10.1007/BF00938572; Saunders et al. *Biochem J* **36** 368 1942, DOI: 10.1042/bj0360368; *Beilstein* **4** IV 34.]

Ethanethiol (ethyl mercaptan) [75-08-1] C_2H_6S , **M 62.1, b 32.9°/704mm, d⁵² 0.83147, pK²⁵ 10.61**. Dissolve the thiol in aqueous 20% NaOH, extract it with a small amount of *benzene and then steam distil until clear. After cooling, the alkaline solution is acidified slightly with 15% H₂SO₄ and the thiol is distilled off, dried with CaSO₄, CaCl₂ or 4A molecular sieves, and fractionally distilled under nitrogen [Ellis & Reid *J Am Chem Soc* **54** 1674 1932, DOI: 10.1021/ja01343a067]. It has a foul odour, work in an efficient fume cupboard. Any exhausted vapours should be absorbed into an aqueous NaOH solution, and this should be oxidised with dilute H₂O₂.

before disposal. [Beilstein 1 IV 1390.]

Ethanol [64-17-5] $\text{C}_2\text{H}_6\text{O}$, M 46.1, m -114° , b $78.3^\circ/\text{atm}$, d^{15}_4 0.79360, d^{25}_4 0.78506, n^{20}_D 1.36139, pK^{25} 15.93. Usual impurities of fermentation alcohol are fusel oils (mainly higher alcohols, especially pentanols), aldehydes, esters, ketones and water. With synthetic alcohol, likely impurities are water, aldehydes, aliphatic esters, acetone and diethyl ether. Traces of *benzene are present in ethanol that has been dehydrated by azeotropic distillation with *benzene. Anhydrous ethanol is very *hygroscopic*. Water (down to 0.05%) can be detected by formation of a voluminous precipitate when aluminium ethoxide in *benzene is added to a test portion. Rectified spirit (95% ethanol) is converted to *absolute* (99.5%) ethanol by refluxing with freshly ignited CaO (250g/L) for 6 hours, standing overnight and distilling with precautions to exclude moisture.

Numerous methods are available for further drying of *absolute* ethanol for making ‘Super dry ethanol’. Lund and Bjerrum [Chem Ber 64 210 1931, DOI: 10.1002/cber.19310640204] used reaction with magnesium ethoxide, prepared by placing 5g of clean dry magnesium turnings and 0.5g of iodine (or a few drops of CCl_4), to activate the Mg, in a 2L flask, followed by 50-75 ml of *absolute* ethanol, and warming the mixture until a vigorous reaction occurs. When this subsides, heating is continued until all the magnesium is converted to magnesium ethoxide. Up to 1L of ethanol is then added and, after refluxing for an hour, it is distilled off. The water content should be below 0.05%. Walden, Ulich and Laun [Z Phys Chem 114 275 1925] used *amalgamated aluminium* chips, prepared by degreasing aluminium chips (by washing with Et_2O and drying in a vacuum to remove grease from the machining of Al), treating with alkali until hydrogen evolved vigorously, washing with H_2O until the washings were weakly alkaline and then stirring with 1% HgCl_2 solution. After 2 minutes, the chips were washed quickly with H_2O , then alcohol, then ether, and dried with filter paper. (The amalgam became warm.) These chips were added to the ethanol, which was then gently warmed for several hours until evolution of hydrogen ceased. The alcohol was distilled and aspirated for some time with pure dry air. Smith [J Chem Soc 1288 1927, DOI: 10.1039/JR9270001288] reacted 1L of *absolute* ethanol in a 2L flask with 7g of clean dry sodium, and added 25g of pure ethyl succinate (27g of pure ethyl phthalate was an alternative), and refluxed the mixture for 2 hours in a system protected from moisture, and then distilled the ethanol. A modification used 40g of ethyl formate instead, so that sodium formate separated out and, during reflux the excess of ethyl formate decomposed to CO and ethanol.

Drying agents suitable for use with ethanol include Linde type 4A molecular sieves, calcium metal, and CaH_2 . The calcium hydride (2g) is crushed to a powder and dissolved in 100ml of *absolute* ethanol by gently boiling. About 70ml of the ethanol are distilled off to remove any dissolved gases before the remainder is poured into 1L of *ca* 99.9% ethanol in a still, where it is boiled under reflux for 20 hours, while a slow stream of pure, dry hydrogen (better use nitrogen or Ar) is passed through. It is then distilled [Rüber Z Elektrochem 29 334 1923]. If calcium is used for drying, about ten times the theoretical amount should be used, and traces of ammonia (from some calcium nitride in the Ca metal) would be removed by passing dry air into the vapour during reflux. Ethanol can be freed from traces of basic materials by distillation from a little 2,4,6-trinitrobenzoic acid or sulfanilic acid. *Benzene can be removed by fractional distillation after adding a little water (the *benzene/water/ethanol azeotrope distils at 64.9°), the alcohol is then re-dried using one of the methods described above. Alternatively, careful fractional distillation can separate *benzene as the *benzene/ethanol azeotrope (b 68.2°). Aldehydes can be removed from ethanol by digesting with 8-10g of dissolved KOH and 5-10g of aluminium or zinc per L, followed by distillation. Another method is to heat under reflux with KOH (20g/L) and AgNO_3 (10g/L) or to add 2.5-3g of lead acetate in 5ml of water to 1L of ethanol, followed (slowly and without stirring) by 5g of KOH in 25ml of ethanol: after 1 hour the flask is shaken thoroughly, then set aside overnight before filtering and distilling. The residual water can be removed by standing the distillate over activated aluminium amalgam for 1 week, then filtering and distilling. Distillation of ethanol from Raney nickel eliminates catalytic poisons.

Other purification procedures include pre-treatment with concentrated H_2SO_4 (3ml/L) to eliminate amines, and with KMnO_4 to oxidise aldehydes, followed by refluxing with KOH to resinify aldehydes, and distilling to remove traces of H_3PO_4 and other acidic impurities after passage through silica gel, and drying over CaSO_4 . Water can be removed by azeotropic distillation with dichloromethane (azeotrope boils at 38.1° and contains 1.8% water) or 2,2,4-trimethylpentane. [Beilstein 1 IV 1289.]

Rapid purification: Place degreased Mg turnings (grease from machining the turnings is removed by washing with dry EtOH then Et_2O , and drying in a vacuum) (5g) in a dry 2L round bottomed flask fitted with a reflux condenser (protect from air with a drying tube filled with CaCl_2 or KOH pellets) and flushed with dry N_2 . Then

add iodine crystals (~0.5g) and gently warm the flask until iodine vapour is formed and coats the turnings. Cool, then add EtOH (50ml) and carefully heat to reflux until the iodine disappears. Cool again, then add more EtOH (to 1L) and reflux under N₂ for several hours. Distil and store over 3A molecular sieves (pre-heated at 300°–350° for several hours and cooled under dry N₂ or argon).

Ethoxycarbonyl isocyanate (ethyl isocyanatoformate) [19617-43-7] C₄H₅NO₃, M 115.1, b 25°/10mm, 51-55°/13mm, 56°/18mm, 115-116°/781mm, d₄²⁰ 1.15. Distil it twice from P₂O₅ (1-2g) through a small Vigreux column and then through a 20-plate column. All fractional distillations should be under a vacuum. [Lamon *J Heterocycl Chem* 5 837 1968, DOI: 10.1002/jhet.5570050616; *Beilstein* 3 H 36, 3 I 17.]

Ethoxycarbonyl isothiocyanate [16182-04-0] C₁₈H₂₂NO₂S, M 131.2, b 43°/14mm, 51-55°/13mm, 56°/18mm, d₄²⁰ 1.12. Fractionally distil it through a short column. It also distils at 83°/30mm with some decomposition liberating CO₂ and sulfurous gases; best distil below 20mm vacuum. [Capp et al. *J Chem Soc* 1340, 1948, DOI: 10.1039/JR9480001340; Lamon *J Heterocycl Chem* 5 837 1968, DOI: 10.1002/jhet.5570050616; *Beilstein* 3 H 174, 3 I 71, 3 III 279, 3 IV 323.]

2-Ethoxyethanol (Cellosolve, ethylene glycol monoethyl ether) [110-80-5] C₄H₁₀O₂, M 90.1, m -90°, b 134.8°, d₄²⁰ 0.931, n_D²⁰ 1.40751. Dry it with CaSO₄ or K₂CO₃, filter and fractionally distil it. Peroxides can be removed by refluxing with anhydrous SnCl₂ or by filtration under slight pressure through a column of activated alumina. [*Beilstein* 1 IV 2377.]

2-Ethoxyethyl methacrylate [2370-63-0] C₈H₁₄O₃, M 158.2, b 91-93°/35mm, d₄²⁰ 0.965, n_D²⁰ 1.429. Purify the ester as described under methyl methacrylate below. If it has to be stored for some time add hydroxyquinone monomethyl ether [150-76-5] (to 50 ppm as stabiliser). [*Beilstein* 2 III 1291.]

Ethyl acetate [141-78-6] C₄H₈O₂, M 88.1, m -84°, b 77.1°/atm, d₄²⁰ 0.9003, n_D²⁰ 1.37239, n_D²⁵ 1.36979, pK_a²⁵ -6.93 (aqueous H₂SO₄). The most common impurities in EtOAc are water, EtOH and acetic acid. These can be removed by washing with aqueous 5% Na₂CO₃, then with saturated aqueous CaCl₂ or NaCl, and drying with K₂CO₃, CaSO₄ or MgSO₄. More efficient drying is achieved if the solvent is further dried with P₂O₅, CaH₂ or molecular sieves before distillation. CaO has also been used. *Alternatively*, ethanol can be converted to ethyl acetate by refluxing with acetic anhydride (*ca* 1ml per 10ml of ester), the liquid is then fractionally distilled, dried with K₂CO₃ and redistilled. [*Beilstein* 2 III 127.]

Rapid purification: Distil, dry over K₂CO₃, re-distil and store over 4A molecular sieves.

Ethyl acetimidate [1000-84-6] C₄H₉NO, M 87.1, b 92-95°/atm, 89.7-90°/765mm, d₄²⁰ 0.8671, n_D²⁰ 1.4025, pK_{Est} ~5.5. It is best to prepare it freshly from the *hydrochloride* (see below). Dissolve the hydrochloride (123.5g) by adding it slowly to an ice-cold mixture of H₂O (500ml), K₂CO₃ (276g) and Et₂O (200ml) and stirring rapidly. The Et₂O layer is separated, the aqueous layer is extracted with Et₂O (100ml), the combined Et₂O layers are dried (MgSO₄), evaporated and the residual oil is distilled through a glass helices packed column (70x1.2cm). The yield is 19g (22%). It is best to keep it as the hydrochloride (see below), and to liberate the *free base* as described when required. [Glickman & Cope *J Am Chem Soc* 67 1012 1945, DOI: 10.1021/ja01222a035; Chaplin & Hunter *J Chem Soc* 1114 1937, DOI: 10.1039/JR9370001114; Hunter & Ludwig *Methods Enzymol* 25 585 1972, DOI: 10.1016/S0076-6879(72)25058-3.]

Ethyl acetimidate hydrochloride [2208-07-3] C₄H₉NO·HCl, M 123.6, m 98-100°(dec), 110-115°(dec), 112-113°(dec), m 112-114°(dec), pK_{Est} ~5.5. Recrystallise the hydrochloride by dissolving it in the minimum volume of super dry EtOH and adding dry Et₂O, or from dry Et₂O. Dry it in a vacuum and store it in a vacuum desiccator with P₂O₅. *Alternatively*, it could be crystallised from EtOH (containing a few of drops of ethanolic HCl) and adding dry Et₂O. Filter and dry (and store) it in a vacuum desiccator over H₂SO₄ and NaOH. [Pinner *Chem Ber* 16 1643 1883, DOI: 10.1002/cber.18830160235; Glickman & Cope *J Am Chem Soc* 67 1017 1945, DOI: 10.1021/ja01222a034; Chaplin & Hunter *J Chem Soc* 1114 1937, DOI: 10.1039/JR9370001114; McElvain & Schroeder *J Am Chem Soc* 71 40 1949, DOI: 10.1021/ja01169a013; McElvain & Tate *J Am Chem Soc* 73 2233 1951, DOI: 10.1021/ja01149a090; *Methods Enzymol* 25 585 1972, DOI: 10.1016/S0076-6879(72)25058-3; *Beilstein* 2 III 418.]

Ethyl acetoacetate [141-97-9] $C_6H_{10}O_3$, M 130.1, m -43° , b $71^\circ/12\text{mm}$, $100^\circ/80\text{mm}$, d_4^{20} 1.026, n_D^{20} 1.419, pK^{25} 10.68. Shake the ester with small amounts of saturated aqueous $NaHCO_3$ (until no further effervescence), then with water. Dry it with $MgSO_4$ or $CaCl_2$ and distil it under reduced pressure. [Beilstein 3 632, 3 I 224, 3 IV 1528.]

Ethyl acrylate [140-88-5] $C_4H_6O_2$, M 100.1, m -71° , b $20^\circ/40\text{mm}$, $99.5^\circ/\text{atm}$, d_4^{20} 0.922, n_D^{20} 1.406. Wash the ester repeatedly with aqueous $NaOH$ until free from inhibitors such as hydroquinone, then wash it with saturated aqueous $CaCl_2$ and distil it under reduced pressure. Hydroquinone should be added if the ethyl acrylate is to be stored for extended periods. [Beilstein 2 IV 1460.] **LACHRYMATORY.**

Ethylamine [75-04-7] C_2H_7N , M 45.1, m -81° , b $16.6^\circ/760\text{mm}$, d_4^{20} 1.3663, pK^{20} 10.79. Condense it in an all-glass apparatus cooled by circulating ice-water, and store it with KOH pellets below 0° . [Beilstein 4 IV 307.] **Ethylamine hydrochloride** [557-66-4] $C_2H_7N \cdot HCl$, M 81.5, m $109-110^\circ$, is crystallised from absolute $EtOH$ or $MeOH/CHCl_3$, washed with dry ether and dried in a vacuum. [Beilstein 4 IV 310.]

Ethyl bromide [74-96-4] C_2H_5Br , M 109.0, m -119° , b $0^\circ/165\text{mm}$, $38^\circ/745\text{mm}$, d_4^{20} 1.460, n_D^{20} 1.4241. The main impurities are usually $EtOH$ and water, both of which form azeotropes with it. Ethanol and unsaturated compounds can be removed by washing with concentrated H_2SO_4 until no further coloration is produced. The ethyl bromide is then washed with water, aqueous Na_2CO_3 , and water again, then dried with $CaCl_2$, $MgSO_4$ or CaH_2 , and distilled from P_2O_5 . Olefinic impurities can also be removed by storing the ethyl bromide in daylight with elemental bromine, later removing the free bromine by extraction with dilute aqueous Na_2SO_3 , drying the ethyl bromide with $CaCl_2$ and fractionally distilling it. Alternatively, unsaturated compounds can be removed by bubbling oxygen containing ca 5% ozone through the liquid for an hour, then washing with aqueous Na_2SO_3 to hydrolyse ozonides and remove hydrolysis products, followed by drying and distillation. [Beilstein 1 IV 150.]

Ethyl bromoacetate [105-36-2] $C_4H_7BrO_2$, M 167.0, b $158-158.5^\circ/758\text{mm}$, d_4^{20} 1.50, n_D^{20} 1.450. Wash the ester with saturated aqueous Na_2CO_3 (three times), 50% aqueous $CaCl_2$ (three times) and saturated aqueous $NaCl$ (twice). Dry with $MgSO_4$, $CaCl_2$ or $CaCO_3$, and distil it. [Beilstein 2 IV 527.] **LACHRYMATORY.**

Ethyl 6-bromohexanoate (ethyl 6-bromocaproate) [25542-62-5] $C_8H_{15}BrO_2$, M 223.1, b $120-125^\circ/14\text{mm}$, $126-127^\circ/20-21\text{mm}$, $128-130^\circ/16\text{mm}$, $127-130^\circ/19\text{mm}$, d_4^{20} 1.241, d^{25} 1.254, n_D^{20} 1.458, n_D^{25} 1.4566. 6-Bromohexanoic acid (250g) is esterified by refluxing with $EtOH$ (600ml) containing H_2SO_4 (15ml) for 8 hours, evaporating *in vacuo*, the residue is taken up in Et_2O , washed with H_2O , then 5% of aqueous Na_2CO_3 (effervescence), the Et_2O layer is dried ($CaCl_2$), filtered, evaporated, and the residual oil is distilled to give the ester (178-218g). [Brown & Partridge *J Am Chem Soc* 66 839 1844, DOI: 10.1021/ja01233a503.] It has also been prepared from 6-hydroxyhexanoic acid by reaction with PBr_3 in pyridine, or HBr (d 1.5) and H_2SO_4 and then esterifying in the same way [Barger et al. *J Chem Soc* 718 1937, DOI: 10.1039/JR9370000718]. [Beilstein 2 IV 940.]

Ethyl 2-(bromomethyl)acrylate [17435-72-2] $C_6H_9BrO_2$, M 193.1, b $38^\circ/0.8\text{mm}$, d_4^{20} 1.398, n_D^{20} 1.479. If it contains some free acid, add H_2O , cool, and neutralise with $NaHCO_3$ until evolution of CO_2 ceases. Extract the mixture with Et_2O (3x) and dry the combined extracts (Na_2SO_4 , 3 hours). Evaporate Et_2O and distil the ester collecting fraction b $39-40^\circ/0.9\text{mm}$, and check spectra. [Preparation and NMR: Ramarajan et al. *Org Synth Coll Vol* 7 210 1990, DOI: 10.15227/orgsyn.061.0056; Beilstein 2 IV 1541.]

Ethyl 2-bromopropionate [535-11-5] $C_5H_9BrO_2$, M 181.0, b $69-70^\circ/25\text{mm}$, $159-160^\circ/\text{atm}$, d_4^{20} 1.39, n_D^{20} 1.447. Wash the ester with saturated aqueous Na_2CO_3 (three times), 50% aqueous $CaCl_2$ (three times) and saturated aqueous $NaCl$ (twice). Dry with $MgSO_4$, $CaCl_2$ or $CaCO_3$, and distil it. [Beilstein 2 IV 762.] **LACHRYMATORY.**

Ethyl 3-bromopropionate [539-74-2] $C_5H_9BrO_2$, M 181.0, b $64-65^\circ/5\text{mm}$, $62.5^\circ/10\text{mm}$, $67-67.5^\circ/12\text{mm}$, $112^\circ/44\text{mm}$ (?), $98^\circ/49\text{mm}$, d_4^{20} 1.452, n_D^{20} 1.452. Purify by carefully fractionating through a short column under reduced pressure. Store away from light as it becomes yellow on standing. [Oae *J Am Chem Soc* 78 4030 1956, DOI: 10.1021/ja01597a044; Mozingo & Patterson *Org Synth Coll Vol* 3 576 1955, DOI: 10.15227/

orgsyn.020.0064; *Beilstein* 2 H 765, 2 I 112, 2 II 231, 2 III 570, 2 IV 765.]

Ethyl bromopyruvate [70-23-5] $\text{C}_5\text{H}_7\text{BrO}_3$, **M 195.0**, **b 47°/0.5mm, 71-73°/5mm, 87°/9mm, 89-104°/14mm**, **d_4^{20} 1.561**, **n_D^{20} 1.464**. The most likely impurity is the free acid [bromopyruvic or bromoacetic (from decarboxylation) acids]. Dissolve the ester in dry Et_2O or dry CHCl_3 , stir with CaCO_3 until effervescence ceases, filter, (may wash rapidly with a little H_2O , or brine), dry (MgSO_4) and distil it at least twice. The **2,4-dinitrophenylhydrazone** has **m 144-145°**. [Borrows & Holland *J Chem Soc* 672 1947, DOI: 10.1039/JR9470000672; Letsinger & Lasco *J Org Chem* 21 764 1956, DOI: 10.1021/jo01113a012; Kruse et al. *J Am Chem Soc* 76 5796 1954, DOI: 10.1021/ja01651a061; *Beilstein* 3 IV 1519.] **LACHRYMATORY**.

2-Ethyl-1-butanol [97-95-0] $\text{C}_6\text{H}_{14}\text{O}$, **M 102.2**, **m -15°**, **b 146.3°/atm**, **n^{15} 1.4243**, **n^{25} 1.4205**. Dry it with CaSO_4 for several days, filter and fractionally distil it. [*Beilstein* 1 IV 1725.]

2-Ethylbut-1-ene [760-21-4] C_6H_{12} , **M 84.1**, **m -131.5° to -131°**, **b 66.6°/atm**, **d_4^{20} 0.833**, **n_D^{20} 1.423**. Wash it with 10N aqueous NaOH , then water. Dry the organic layer with CaCl_2 , filter and fractionally distil it. [*Beilstein* 1 IV 850.]

Ethyl n-butyrate [105-54-4] $\text{C}_6\text{H}_{12}\text{O}_2$, **M 116.2**, **m -93°**, **b 49°/50mm, 119-120°/760mm**, **d_4^{20} 0.880**, **n_D^{20} 1.393**. Dry the ester with anhydrous CuSO_4 and distil it under dry nitrogen. [*Beilstein* 2 IV 787.]

Ethyl carbazate (N-ethoxycarbonyl hydrazine) [4114-31-2] $\text{C}_3\text{H}_8\text{N}_2\text{O}_2$, **M 104.1**, **m 44-48°**, **51-52°**, **b 95.5°/10mm, 100-102°/11mm**. Fractionate the carbazate using a Vigreux column until the distillate crystallises [Allen & Bell *Org Synth Coll Vol* 3 404 1955, DOI: 10.15227/orgsyn.024.0058; *Beilstein* 3 IV 174].

Ethyl chloride [75-00-3] $\text{C}_2\text{H}_5\text{Cl}$, **M 64.5**, **m -139°**, **b 12.4°/atm**, **d_4^{20} 0.8978**, **n_D^{20} 1.3676**. Pass ethyl chloride through absorption towers containing, successively, concentrated H_2SO_4 , NaOH pellets, P_2O_5 on glass wool, or soda-lime, CaCl_2 , or P_2O_5 . Condense it into a flask containing CaH_2 and fractionally distil it. It has also been purified by illumination in the presence of bromine at 0° using a 1000W lamp, followed by washing, drying and distilling. [*Beilstein* 1 IV 124.]

Ethyl chloroacetate [105-39-5] $\text{C}_4\text{H}_7\text{ClO}_2$, **M 122.6**, **m -26°**, **b 143-143.2°/atm**, **d_4^{20} 1.150**, **n^{25} 1.4192**. Shake the ester with saturated aqueous Na_2CO_3 (three times), aqueous 50% CaCl_2 (three times) and saturated aqueous NaCl (twice). Dry it with Na_2SO_4 or MgSO_4 and distil it. [*Beilstein* 2 IV 481.] **LACHRYMATORY**.

Ethyl (ClFCHCCO₂Et) [± 401-56-9] $\text{C}_4\text{H}_6\text{ClFO}_2$, **M 140.5**, **b 128°/atm, 133°/atm**, **d_4^{25} 1.225**, **n_D^{25} 1.3926**. The ester is prepared from 2-chloro-1,1,2-trifluoroethyl ether (340g, 2.09mol at < 5°) while 96% H_2SO_4 (228ml, 420g, 4.1mol) is added dropwise to it with stirring, and allowing the liberated HF (HIGHLY TOXIC vapour which etches glass) to vacate into an efficient fume cupboard, at a rate such that the temperature is maintained at 5-15° (ca 30-45 minutes). The mixture is stirred further at 10° for 2 hours when all the HF is released, then it is poured carefully onto crushed ice (1 Kg) and H_2O (500ml). The lower white oily layer is allowed to settle, separated, washed until free of acid with saturated aqueous NaHCO_3 (3 x 25ml, **CARE**: CO_2 may be liberated), H_2O (4 x 25ml) and dried over Drierite (10g) [Note that thorough washing, to remove any acid, is essential to avoid decomposition on distillation]. The crude dry ester is filtered, and fractionated through an efficient helix packed column (2 x 12cm) to give pure *ethyl chlorofluoroacetate* (190-100g, 65-68%) boiling at 129-130° and atmospheric pressure. The **free acid** [471-44-3] has **b 162°/atm** (160.5-161°/atm also been reported and is neurotoxic), **d_4^{25} 1.532**, **n_D^{25} 1.4085**, is obtained by hydrolysis of the **methyl ester**, **b 116°/atm**, **d_4^{25} 1.323**, **n_D^{25} 1.3903**, with 10% aqueous NaOH . The ethyl ester provides **chlorofluoroacetyl chloride** **b 69.5°/atm**, **d_4^{25} 1.468**, **n_D^{25} 1.3992**, by Middleton's procedure (*J Org Chem* 44 2291 1979, DOI: 10.1021/jo01327a058), and **chlorofluoroacetamide**, **b 72°/1mm**, **d_4^{25} 1.510**, **n_D^{25} 1.4535**, by reaction with ammonia [Young & Tarrant *J Am Chem Soc* 71 2432 1949, DOI: 10.1021/ja01175a055; Hazeldine *J Chem Soc* 4259 1952, DOI: 10.1039/JR9520004259]. [*Beilstein* 2 III 453, 2 IV 493, Englund *Org Synth Coll Vol* 4 423 1963, DOI: 10.15227/orgsyn.034.0049; Young & Tarrant *J Am Chem Soc* 72 1860 1950, DOI: 10.1021/ja01160a531.] This ester, its acid and acid chloride are chiral, are obtained in (+) and (-) forms from optical resolutions of the acid [Bellucci et al. *Tetrahedron* 25 2979 1969, DOI: 10.1016/s0040-4020(01)82931-3; Molines & Wakselman *Synthesis* 838 1984, DOI: 10.1055/s-30987], and are sterically better alternatives than Mosher's acid (p 387) for chiral discrimination of esters and amides [Ruzicka et al. *J Chem Res (S)* 830 1998, DOI: 10.1039/A806698A].

Ethyl chloroformate [541-41-3] $\text{C}_3\text{H}_5\text{ClO}_2$, M 108.5, m -81° , b $94\text{--}95^\circ/\text{atm}$, d_4^{20} 1.135, n_D^{20} 1.3974. Wash the ester several times with water, redistil it using an efficient fractionating column at atmospheric pressure and a CaCl_2 guard tube to keep free from moisture [Hamilton & Sly *J Am Chem Soc* **47** 435 1925, DOI: 10.1021/ja01679a024; Saunders et al. *J Am Chem Soc* **73** 3796 1951, DOI: 10.1021/ja01152a069]. [Beilstein 3 IV 23.] **LACHRYMATORY AND TOXIC.**

Ethyl trans-crotonate [623-70-1] $\text{C}_6\text{H}_{10}\text{O}_2$, M 114.2, b $137^\circ/\text{atm}$, d_4^{20} 0.917, n_D^{20} 1.425. Wash it with aqueous 5% Na_2CO_3 , then with saturated aqueous CaCl_2 , dry it with CaCl_2 and distil it. [Beilstein 2 IV 1500.]

Ethyl cyanoacetate [105-56-6] $\text{C}_5\text{H}_7\text{NO}_2$, M 113.1, m -22° , b $206.0^\circ/\text{atm}$, d_4^{20} 1.061, n_D^{20} 1.41751. Shake the ester several times with aqueous 10% Na_2CO_3 , wash it well with water, dry with Na_2SO_4 and fractionally distil it. [Beilstein 2 IV 1889.]

Ethyl cyanoformate [623-49-4] $\text{C}_4\text{H}_5\text{NO}_2$, M 99.1, b $113\text{--}114^\circ/740\text{mm}$, $116.5\text{--}116.8^\circ/765.5\text{mm}$, d_4^{20} 1.0112, n_D^{20} 1.3818. Dissolve the cyanoformate in Et_2O , dry it over Na_2SO_4 , filter, evaporate and distil it [Malachowski et al. *Chem Ber* **70** 1012 1937, DOI: 10.1002/cber.19370700527; Adickes et al. *J Prakt Chem* [2] **133** 313 1932, DOI: 10.1002/prac.19321331101; Grundmann et al. *Justus Liebigs Ann Chem* **577** 77 1952, DOI: 10.1002/jlac.19525770110]. [Beilstein 2 IV 1862.]

Ethyl diazoacetate [623-73-4] $\text{C}_4\text{H}_6\text{N}_2\text{O}_2$, M 114.1, m -22° , b $42^\circ/5\text{mm}$, $45^\circ/12\text{mm}$, $85\text{--}86^\circ/88\text{mm}$, $140\text{--}141^\circ/720\text{mm}$, $140\text{--}143^\circ/\text{atm}$, $d_4^{17.6}$ 1.0852, $n_D^{17.6}$ 1.4588. It is a very volatile yellow oil with a strong pungent odour. **EXPLOSIVE [distillation even under reduced pressure is dangerous and may result in an explosion — TAKE ALL THE NECESSARY PRECAUTIONS IF DISTILLATION IS TO BE CARRIED OUT]**. It explodes in contact with concentrated H_2SO_4 -trace acid causes rapid decomposition. It is slightly soluble in H_2O , but is miscible with EtOH , C_6H_6 , petroleum ether and Et_2O . To purify, dissolve it in Et_2O [using CH_2Cl_2 instead of Et_2O , protects the ester from acid], wash it with 10% aqueous Na_2CO_3 , dry (MgSO_4), filter and repeat as many times as possible until the Et_2O layer loses its yellow colour, then remove the solvent below 20° (vacuum). Note that prolonged heating may lead to rapid decomposition and low yields. It can also be purified by steam distillation under reduced pressure but with considerable loss in yield. Place the residual oil in a brown bottle, keep below 10° , and use as soon as possible without distilling. For preparing esters usually the ethereal solution is used directly without purification. [Womack & Nelson *Org Synth Coll Vol* **3** 392 1955, DOI: 10.15227/orgsyn.024.0056; UV: Miller & White *J Am Chem Soc* **79** 5974 1957, DOI: 10.1021/ja01579a035; Fieser **1** 367 1967, Beilstein 3 IV 1495.]

Ethyl dibromoacetate [617-33-4] $\text{C}_4\text{H}_6\text{Br}_2\text{O}_2$, M 245.9, b $81\text{--}82^\circ/14.5\text{mm}$, $194^\circ/\text{atm}$, d_4^{22} 1.9081, n_D^{22} 1.4973. Wash the ester briefly with concentrated aqueous NaHCO_3 , then with aqueous CaCl_2 . Dry it with CaSO_4 and distil it under reduced pressure. [Hornyak & Amis *J Am Chem Soc* **79** 2079 1957, DOI: 10.1021/ja01566a015; Beilstein 2 H 219, 2 I 97, 2 III 484, 2 IV 533.]

Ethyl dichloroacetate [535-15-9] $\text{C}_4\text{H}_6\text{Cl}_2\text{O}_2$, M 157.0, b $54\text{--}55^\circ/11\text{mm}$, $131.0\text{--}131.5^\circ/40\text{mm}$, d_4^{20} 1.28, n_D^{20} 1.438. Shake the ester with aqueous 3% NaHCO_3 to remove free acid, wash with distilled water, dry for 3 days with CaSO_4 and distil it under reduced pressure. [Beilstein 2 IV 501.]

Ethyl 3,3-diethoxypropionate [10601-80-6] $\text{C}_9\text{H}_{18}\text{O}_4$, M 190.2, b $58.5^\circ/1.5\text{mm}$, $65^\circ/2\text{mm}$, $95\text{--}96^\circ/12\text{mm}$, d_4^{20} 0.78, n_D^{25} 1.4101. Dissolve it in dry Et_2O , and dry with solid NaHCO_3 , filter, distil and carefully fractionate it [Dyer & Johnson *J Am Chem Soc* **56** 222 193, DOI: 10.1021/ja01316a070]. [Beilstein 3 II 411.]

Ethylene (ethene) [74-85-1] C_2H_4 , M 28.0, m -169.4° , b $-102^\circ/700\text{mm}$. Purify ethylene by passage through a series of towers containing molecular sieves, or anhydrous CaSO_4 , or cuprous ammonium solution, then concentrated H_2SO_4 , followed by KOH pellets. Alternatively, it has been condensed in liquid nitrogen, with melting, freezing and pumping to remove air before passage through an activated charcoal trap, followed by a further condensation in liquid air. A sputtered sodium trap was used to remove oxygen. [Beilstein 1 IV 677.]

Ethylenediamine (1,2-diaminoethane) [107-15-3] $C_2H_8N_2$, **M 60.1**, **f 11.0°**, **m -8.5°**, **b 117.0°/atm**, **d**₄²⁰ **0.897**, **n**_D²⁰ **1.45677**, **n**_D³⁰ **1.4513**, **pK**₁²⁵ **6.86**, **pK**₂²⁵ **9.92**. It forms a constant-boiling (**b 118.5°**, *monohydrate*, **m 10°**) mixture with water (23w/w%). [It is *hygroscopic* and miscible with water.] Recommended purification procedure [Asthana & Mukherjee in J.F. Coetzee (ed), *Purification of Solvents*, Pergamon Press, Oxford, 1982 cf. p 53]: to 1L of ethylenediamine is added 70g of type 5A Linde molecular sieves and shaken for 12 hours. The liquid is decanted and shaken for a further 12 hours with a mixture of CaO (50g) and KOH (15g). The supernatant is fractionally distilled (at 20:1 reflux ratio) in contact with freshly activated molecular sieves. The fraction distilling at 117.2°/760mm is collected. Finally it is fractionally distilled from sodium metal. All distillations and storage of ethylenediamine should be carried out under nitrogen to prevent reaction with CO₂ and water. The material containing 30% water is dried with solid NaOH (600g/L) and heated on a water bath for 10 hours. Above 60°, separation into two phases takes place. The hot ethylenediamine layer is decanted off, refluxed with 40g of sodium for 2 hours and distilled [Putnam & Kobe *Trans Electrochem Soc* **74** 609 1938, DOI:10.1149/1.3494023]. Ethylenediamine is usually distilled under nitrogen. *Alternatively*, it is dried over type 5A Linde molecular sieves (70g/L), then a mixture of 50g of CaO and 15g of KOH/L, with further dehydration of the supernatant with molecular sieves followed by distillation *from* molecular sieves and, finally, from sodium metal. A spectroscopically improved material is obtained by shaking with freshly baked alumina (20g/L) before distillation. [Beilstein **4** IV 1166.] **Ethylenediamine dihydrochloride** [333-18-6] $C_2H_8N_2 \cdot 2HCl$, **M 133.0**, **m >300°**, **pK**₁²⁵ **6.86**, **pK**₂²⁵ **9.92**, is recrystallised from H₂O or H₂O/EtOH. Wash the crystals with EtOH and dry them *in vacuo*. It sublimes on heating. [Beilstein **4** IV 1168.]

N,N'-Ethylenediaminediacetic acid (EDDA) [5657-17-0] $C_6H_{12}N_2O_4$, **M 176.2**, **m 222-224°(dec)**, **pK**₁²⁵ **6.48**, **pK**₂²⁵ **9.57** (for NH groups). Crystallise EDDA from H₂O. [Beilstein **4** IV 2446.]

Ethylenediaminetetraacetic acid (EDTA) [60-00-4] $C_{10}H_{18}N_2O_8$, **M 292.2**, **m 253°(dec)**, **pK**₁²⁵ **0.26**, **pK**₂²⁵ **0.96**, **pK**₃²⁵ **2.60**, **pK**₄²⁵ **2.67**, **pK**₅²⁵ **6.16**, **pK**₆²⁵ **10.26**. Dissolve EDTA in aqueous KOH or ammonium hydroxide, and precipitate it twice with dilute HCl or HNO₃. Boil it twice with distilled water to remove mineral acid, then recrystallise it from water or dimethylformamide. Dry it at 110°. It also recrystallises from boiling 1N HCl; wash the crystals with distilled H₂O and dry them *in vacuo*. [Ma & Ray *Biochemistry* **19** 751 1980, DOI: 10.1021/bi00545a022s; Beilstein **4** IV 2449.] It is a sequestering agent and a useful buffer

Ethylene dimethacrylate (ethylene glycol dimethacrylate) [97-90-5] $C_{10}H_{14}O_4$, **M 198.2**, **b 98-100°/5mm**, **d**₄²⁰ **1.053**, **n**_D²⁰ **1.456**. Distil it through a short Vigreux column (p 11) at about 1mm pressure, in the presence of 3% (w/w) of phenyl-β-naphthylamine. [Beilstein **2** IV 1532.]

Ethylene dimyristate [1,2-bis(myristoyloxy)ethane] [627-84-9] $C_{30}H_{58}O_4$, **M 482.8**, **m 61.7°**. Crystallise the ester from *benzene/MeOH or diethyl ether/MeOH, and dry it in a vacuum desiccator. It forms an inclusion compound with 25.9 mols of urea. [McGreer et al. *J Am Chem Soc* **74** 3541 1952, DOI: 10.1021/ja01134a025; Beilstein **2** H 366, **2** II 327, **2** III 924, **2** IV 1133.]

Ethylene dipalmitate [1,2-bis(palmitoyloxy)ethane] [624-03-3] $C_{34}H_{66}O_4$, **M 538.9**, **m 69.1°, 71.2°**. Crystallise the ester from *benzene/MeOH, diethyl ether/MeOH or Me₂CO and dry it in a vacuum desiccator. It forms an inclusion compound with 28.2 mols of urea. [McGreer et al. *J Am Chem Soc* **74** 3541 1952, DOI: 10.1021/ja01134a025; Beilstein **2** H 373, **2** I 166, **2** II 338, **2** III 926, **2** IV 1169.]

Ethylene distearate [1,2-bis(stearoyloxy)ethane] [627-83-8] $C_{38}H_{74}O_4$, **M 595.0**, **m 74.4-75°, 75.3°, 77°**. Crystallise the ester from *benzene/MeOH, diethyl ether/MeOH or Me₂CO and dry it in a vacuum desiccator. It forms an inclusion compound with 31 mols of urea. [McGreer et al. *J Am Chem Soc* **74** 3541 1952, DOI: 10.1021/ja01134a025; Beilstein **2** H 380, **2** II 354, **2** III 1021, **2** IV 1223.]

Ethylene glycol [107-21-1] $C_2H_6O_2$, **M 62.1**, **m -13°**, **b 68°/4mm**, **197.9°/760mm**, **d**₄²⁰ **1.0986**, **n**_D¹⁵ **1.43312**, **n**_D²⁵ **1.43056**, **pK**₁²⁵ **10.6**. It is very *hygroscopic*, and also likely to contain higher diols. Dry it with CaO, CaSO₄, MgSO₄ or NaOH and distil it under vacuum. Dry further by reaction with a small amount of sodium under N₂ (to remove moisture), reflux for several hours and distil. The distillate is then passed through a column of Linde type 4A molecular sieves and finally distil under nitrogen, from more molecular sieves. Then fractionally distil

it. [Beilstein 1 IV 2369.] **Ethylene glycol diacetate** [111-55-7] $C_6H_{10}O_4$, M 146.2, m -41° , b 190.1° , 79-81°/11mm, d_{25}^{25} 1.4188, n_D^{20} 1.4150, should be dried with $CaCl_2$, filtered (excluding moisture), and fractionally distil it under reduced pressure. [Beilstein 2 IV 217 1541.]

Ethylene glycol-bis(2-aminoethylether)-N,N,N,N'-tetraacetic acid (EGTA, Egtazic acid) [67-42-5] $C_{14}H_{24}N_2O_{10}$, M 380.4, m $>245^\circ$ (dec), pK_1^{20} 1.15 (2.40), pK_2^{20} 2.40 (2.50), pK_3^{20} 8.40 (8.67), pK_4^{20} 8.94 (9.22). Dissolve EGTA in aqueous NaOH, precipitate it by adding aqueous HCl, wash it with water and dry at 100° in vacuo. [Beilstein 4 IV 2483.] it is a useful buffer.

Ethylene glycol dibutyl ether [112-48-1] $C_{10}H_{22}O_2$, M 174.3, b $78-80^\circ/16mm$, 200-201°/760mm, d_4^{20} 1.105, n_D^{20} 1.42. Shake the ether with aqueous 5% Na_2CO_3 , dry with $MgSO_4$ and store it with chromatographic alumina to prevent peroxide formation. [Beilstein 1 III 2083, 1 IV 2382.]

Ethylene glycol diethyl ether (1,2-diethoxyethane) [629-14-1] $C_6H_{14}O_2$, M 118.2, m -74° , b $121.5^\circ/atm$, d_4^{20} 0.842, n_D^{20} 1.392. After refluxing for 12 hours, a mixture of the ether (2L), concentrated HCl (27ml) and water (200ml) is added with slow passage of nitrogen. The solution is cooled, and KOH pellets are added slowly and with shaking until no more dissolves. The organic layer is decanted, treated with some KOH pellets and again decanted. It is then refluxed with, and distilled from sodium immediately before use. Alternatively, after removal of peroxides by treatment with activated alumina, the ether is refluxed in the presence of the blue ketyl formed by sodium-potassium alloy with benzophenone, then distilled. [Beilstein 1 H 468, 1 II 519, 1 III 2078, 1 IV 2379.]

Ethyl formate [109-94-4] $C_3H_6O_2$, M 74.1, m -80° , b $54.2^\circ/atm$, d_4^{20} 0.921, d_4^{30} 0.909, n_D^{20} 1.35994, n_D^{25} 1.3565. Free acid or alcohol is removed by standing the ester over anhydrous K_2CO_3 , with occasional shaking, then decanting and distilling from P_2O_5 . Alternatively, the ester can be kept over CaH_2 for several days, then distilled from fresh CaH_2 . It cannot be dried with $CaCl_2$ because it reacts rapidly with the ester to form a crystalline compound. [Beilstein 2 IV 23.]

Ethyl iodide (iodoethane) [75-03-6] C_2H_5I , M 156.0, m -108° , b $72.4^\circ/atm$, d_4^{20} 1.933, n_D^{15} 1.5682, n_D^{25} 1.5104. Drying the iodide with P_2O_5 is unsatisfactory, and with $CaCl_2$ it is incomplete. It is probably best to dry it with sodium wire and distil [Hammond et al. *J Am Chem Soc* 82 704 1960, DOI: 10.1021/ja01488a051]. Exposure of ethyl iodide to light leads to rapid decomposition, with the liberation of iodine. Free iodine can be removed by shaking with several portions of dilute aqueous $Na_2S_2O_3$ (until the colour is discharged), followed by washing with water, drying (with $CaCl_2$, then sodium), and distilling. The distilled ethyl iodide is stored, over some mercury, in a dark bottle away from direct sunlight. Other purification procedures include passage through a 60cm column of silica gel, followed by distillation, and treatment with elemental bromine, extraction of free halogen with $Na_2S_2O_3$ solution, followed by washing with water, drying and distilling. Free iodine and HI have also been removed by direct distillation through a LeBel-Henninger column containing copper turnings. Purification by shaking with alkaline solutions, and storage over silver, are reported to be unsatisfactory. [Beilstein 1 IV 163.]

Ethyl isobutyrate [97-62-1] $C_6H_{12}O_2$, M 116.2, b $110^\circ/atm$, d_4^{20} 0.867, n_D^{20} 1.388. Wash the ester with aqueous 5% Na_2CO_3 , then with saturated aqueous $CaCl_2$. Dry it over $CaSO_4$ and distil. [Beilstein 1 IV 846.]

Ethyl isocyanate [109-90-0] C_3H_5NO , M 71.1, b $559.8^\circ/759mm$, 59-61°/760mm, 60-63°/~760mm, d_4^{20} 0.9031, n_D^{20} 1.3808. Fractionate the isocyanate through an efficient column preferably in an inert atmosphere and store it in aliquots in sealed tubes [Bieber *J Am Chem Soc* 74 4700 1952, DOI: 10.1021/ja01138a501; Slocombe et al. *J Am Chem Soc* 72 1888 1950, DOI: 10.1021/ja01161a009]. [Beilstein 4 IV 402.]

Ethyl isovalerate [108-64-5] $C_7H_{14}O_2$, M 130.2, m -99° , b $134.7^\circ/atm$, d_4^{20} 0.8664, n_D^{20} 1.39621, n_D^{25} 1.3975. Wash the ester with aqueous 5% Na_2CO_3 , then saturated aqueous $CaCl_2$. Dry it over $CaSO_4$ and distil. [Beilstein 2 IV 898.]

Ethyl levulinate (4-oxopentanoic acid ethyl ester) [539-88-8] $C_7H_{12}O_3$, M 144.2, m 37.2° , b $106-108^\circ/2mm$,

138.8°/8mm, 203-205°/atm, d_4^{20} 1.012, n_D^{20} 1.423. Stir the ester with Na_2CO_3 and charcoal, filter and distil. It is freely soluble in H_2O and EtOH [IR, NMR: Sterk *Monatsh Chem* **99** 1770 1968, DOI:10.1007/BF00904307; Thomas & Schuette *J Am Chem Soc* **53** 2324 1931, DOI: 10.1021/ja01357a043; Cox & Dodds *J Am Chem Soc* **55** 3391 1933, DOI: 10.1021/ja01335a059]. [*Beilstein* **3** IV 1562.]

Ethyl malonate monoamide [7597-56-0] $\text{C}_5\text{H}_9\text{NO}_3$, **M 131.1, m 47-50°, 49.5-50°, 50°, b 130-135°/2mm.** The amide crystallises from Et_2O , or by slow evaporation of an aqueous solution as colourless crystals [Snyder & Elston *J Am Chem Soc* **76** 3039 1954, DOI: 10.1021/ja01640a060; McElvain & Schroeder *J Am Chem Soc* **71** 45 1949, DOI: 10.1021/ja01169a014; Rising et al. *J Biol Chem* **89** 1 1930, <http://www.jbc.org/content/89/1/1>]. [*Beilstein* **2** IV 1887.]

Ethyl methacrylate [97-63-2] $\text{C}_6\text{H}_{10}\text{O}_2$, **M 114.2, b 59°/100mm, 118-119°/atm, d_4^{20} 0.915, n_D^{20} 1.515.** Wash the ester successively with 5% aqueous NaNO_2 , 5% NaHSO_3 , 5% NaOH , then water. Dry it over MgSO_4 , add 0.2% (w/w) of phenyl- β -naphthylamine, and distil it through a short Vigreux column [Schultz *J Am Chem Soc* **80** 1854 1958, DOI: 10.1021/ja01541a020]. [*Beilstein* **2** IV 1523.]

Ethyl methyl ether [540-67-0] $\text{C}_3\text{H}_8\text{O}$, **M 60.1, b 7°/760mm, d^0 0.725, n^4 1.3420.** Dry the ether with CaSO_4 , pass it through an alumina column (to remove peroxides), then fractionally distil it while collecting fractions in receivers kept below 0°. It has a high solubility in H_2O . [*Beilstein* **1** H 314, **1** I 158, **1** II 311, **1** III 1288, **1** IV 1314.] **HIGHLY FLAMMABLE.**

3-Ethyl-2-methyl-2-pentene [19780-67-7] C_8H_{16} , **M 112.1, b 109°/757mm, 114.5°/760mm, d_4^{20} 0.72468, n_D^{20} 1.4124.** Purify it by preparative GLC on a column of 20% squalene on Chromosorb P at 70°. Alternatively, fractionate it under an inert atmosphere. It forms an azeotrope with methoxyethanol. [*Beilstein* **1** H 222, **1** III 8471, **1** IV 890.]

Ethyl nitroacetate [626-35-7] $\text{C}_4\text{H}_7\text{NO}_4$, **M 133.1, b 42-43°/0.2mm, 71-72°/3mm, 93-96°/9mm, 194-195°/atm, d_4^{20} 1.1953, n_D^{20} 1.4260, pK^{25} 5.82.** Purify the ester by repeated distillation. IR has ν_{max} at 1748 (CO_2), 1570 and 1337 (NO_2), and 800cm^{-1} [Haszeldine *J Chem Soc* 2525 1953, DOI: 10.1039/JR9530002525]. The **hydrazine salt** crystallises from 95% EtOH or MeOH as yellow crystals **m 104-105°** [Ungnade & Kissinger *J Org Chem* **22** 1662 1957, DOI: 10.1021/jo01363a034; Emmons & Freeman *J Am Chem Soc* **77** 4391 1955, DOI: 10.1021/ja01621a060]. [*Beilstein* **2** IV 537.]

Ethyl propionate [105-37-3] $\text{C}_5\text{H}_{10}\text{O}_2$, **M 102.1, b 99.1°/atm, d_4^{20} 0.891, n^{15} 1.38643, n_D^{20} 1.38394.** Treat the ester with anhydrous CuSO_4 and distil it under nitrogen. [*Beilstein* **2** IV 705.]

Ethyl pyruvate [617-35-6] $\text{C}_5\text{H}_8\text{O}_3$, **M 116.1, m -50°, b 44-45°/10mm, 56°/20mm, 69-71°/42mm, 63°/23mm, 155.5°/760mm, d_4^{20} 1.047, n_D^{20} 1.4052.** Shake the ester with 10ml portions of saturated aqueous CaCl_2 solution (removes ethyl acetate) and the organic layer is removed by centrifugation, decantation and filtration, and is distilled under reduced pressure. Purification of small quantities is carried out *via* the bisulfite adduct: the ester (2.2ml) is shaken with saturated NaHSO_3 (3.6ml), chilled in a freezing mixture when crystals separate rapidly (particularly if seeded). After 5 minutes EtOH (10ml) is added and the crystals are filtered off, washed with EtOH and Et_2O and dried. Yield *ca* 3g of *bisulfite adduct*. Then treat the adduct (16g) with saturated aqueous MgSO_4 (32ml) and 40% formaldehyde (5ml) and shake, whereby the ester separates as an oil which is extracted with Et_2O . The extract is dried (MgSO_4), filtered, evaporated and the residue is distilled (**b 56°/20mm**), and then redistilled (**b 147.5°/750mm**) to give 5.5g of pure ester. [Cornforth *Org Synth Coll Vol* **4** 467 1963, DOI: 10.15227/orgsyn.031.0059; *Beilstein* **3** IV 1513.]

Ethyl stearate [111-61-5] $\text{C}_{20}\text{H}_{40}\text{O}_2$, **M 312.5, m 33°, b 213-215°/15mm.** The solid portion is separated from the partially solid starting material, then recrystallised twice from EtOH , dried by azeotropic distillation with *benzene, and fractionally distilled through a spinning-band column at low pressure [Welsh *Trans Faraday Soc* **55** 52 1959, DOI: 10.1039/TF9595500052]. [*Beilstein* **2** IV 1218.]

Ethyl thiocyanate (ethyl rhodanide) [542-90-5] $\text{C}_3\text{H}_5\text{NS}$, **M 87.1, b 144-145°/atm, d_4^{20} 1.011, n_D^{20} 1.462.**

Fractionally distil the ester at atmospheric pressure. [*Beilstein* 2 IV 1218.] (**CARE LACHRYMATOR.**)

Ethyl thioglycolate (ethyl 2-mercaptoacetate) [623-51-8] $\text{C}_4\text{H}_8\text{O}_2\text{S}$, M 120.2, b 50-51°/10mm, 55°/17mm, 62.5-64°/22mm, 67-68°/24mm, 155-158°/atm, d_4^{20} 1.096, n_D^{20} 1.457. Dissolve the thioglycolate in Et_2O , wash with H_2O , dry it over Na_2SO_4 , filter, evaporate and distil the residue under reduced pressure [Bredereck et al. *Chem Ber* 90 1837 1957, DOI: 10.1002/cber.19570900923]. The **Ni complex** [$\text{Ni}(\text{SCH}_2\text{CO}_2\text{Et})_2$], when recrystallised twice from EtOH, gives crystals which become black when dried in a vacuum over H_2SO_4 , m 104-105° [Draney & Cefola *J Am Chem Soc* 76 1975 1954, DOI: 10.1021/ja01636a081]. [*Beilstein* 3 H 255, 3 IV 616]

N-Ethylthiourea [625-53-6] $\text{C}_3\text{H}_8\text{N}_2\text{S}$, M 104.2, m 110°. Crystallise the thiourea from EtOH, MeOH or ether. [*Beilstein* 4 IV 374.]

Ethyl trichloroacetate [515-84-4] $\text{C}_4\text{H}_5\text{Cl}_3\text{O}_2$, M 191.4, b 100-100.5°/30mm, 168°/atm, d_4^{20} 1.383, n_D^{20} 1.453. Shake the ester with saturated aqueous Na_2CO_3 (three times), aqueous 50% CaCl_2 (three times), saturated aqueous NaCl (twice), then distil it over CaCl_2 , and redistil it under reduced pressure. [*Beilstein* 2 IV 514.]

Ethyl trifluoroacetate [383-63-1] $\text{C}_4\text{H}_5\text{F}_3\text{O}_2$, M 142.1, b 61.3°/750, 60-62°/atm, 62-64°/755mm, d_4^{20} 1.191, n_D^{20} 1.30738. It has been prepared by treating sodium trifluoroacetate with Et_2SO_4 , or the acid with excess of EtOH and a small amount of H_2SO_4 , distilling the azeotropic mixture of ester and EtOH, removing the latter with CaCl_2 , filtering and fractionally distilling [Henne et al *J Am Chem Soc* 69 1819 1947, DOI: 10.1021/ja01199a075]. Fractionate it through a long Vigreux column. IR has ν_{max} at 1800 (CO_2) and 1000 (OCO) cm^{-1} [Fuson et al. *J Chem Phys* 20 1627 1952, DOI: org.virtual.anu.edu.au/10.1063/1.1700229; Bergman *J Org Chem* 23 476 1958, DOI: 10.1021/jo01097a039]. [*Beilstein* 2 II 186, 2 III 427, 2 IV 463.]

Ethyl 4,4,4-trifluoroacetoacetate (ethyl 3-oxo-4,4,4-trifluorobutyrate) [372-31-6] $\text{C}_6\text{H}_7\text{F}_3\text{O}_3$, M 184.1, b 47-49°/25mm, 129-130°/atm, 131.8°/atm, d_4^{25} 1.259, n_D^{20} 1.375. The ester was prepared by the method of Swarts [*Bull Acad Roy Belg* 13 175 1927] in which ethyl trifluoroacetate (1mol) was added slowly to a stirred suspension of NaOEt (1mol) in EtOH (exothermic reaction), followed EtOAc (1mol), and refluxed overnight. An improved purification was by treatment with a concentrated aqueous solution of NaHSO_4 (very small molar excess), followed by addition of a clear aqueous solution of $\text{Cu}(\text{OAc})_2$, and the organic layer was distilled off. The Cu derivative was filtered off and recrystallised from EtOH, washed with Et_2O to give green crystals of the **copper chelate** m 189° [UV has λ_{max} at 220nm (ϵ 11,000) and 270nm (ϵ 21,000), 3.0mg % in EtOH]. The dry Cu chelate was suspended in Et_2O (some dissolved) and H_2S was bubbled through until precipitation of CuS was complete. The CuS was filtered off, and the filtrate was carefully fractionation (to avoid losses) to provide the ester in 54% overall yield. Its UV has λ_{max} at 238nm (ϵ 3000) and 288nm (ϵ 400), in cyclohexane [Breslow et al. *J Am Chem Soc* 68 100, 1946, DOI: 10.1021/ja01205a032; Henne et al. *J Am Chem Soc* 69 1819 1947, DOI: 10.1021/ja01199a075; Haszeldine et al. *J Chem Soc* 609 1951, DOI: 10.1039/JR9510000609; *Beilstein* 3 II 425, 3 III 1206, 3 IV 1548].

Ethyl trifluoromethanesulfonate [425-75-2] $\text{C}_3\text{H}_5\text{F}_3\text{O}_3\text{S}$, M 178.1, b 115°/atm, 118-120°/atm, d_4^{20} 1.378, n_D^{20} 1.336. The ester reacts slowly with H_2O and aqueous alkali. If its IR has no OH bands ($\sim 3000 \text{ cm}^{-1}$) then purify it by redistillation. If OH bands are present, then dilute with dry Et_2O and shake (carefully) with aqueous NaHCO_3 until effervescence ceases, then wash with H_2O and dry (MgSO_4), filter, evaporate and distil the residue under a slight vacuum then at atmospheric pressure in a N_2 atmosphere. **It is a powerful alkylating agent, and the fumes are very toxic — perform all operations in an efficient fume cupboard.** [Gramstad & Haszeldine *J Chem Soc* 173 1956, DOI: 10.1039/JR9560000173; Howells & McCown *Chem Rev* 77 69 1977, DOI: 10.1021/cr60305a005; *Beilstein* 3 IV 34.]

S-Ethyl trifluorothioacetate [383-64-2] $\text{C}_4\text{H}_5\text{F}_3\text{OS}$, M 158.1, b 88-90°/atm, 90.5°/760mm, d_4^{20} 1.255, n_D^{20} 1.372. If IR is free of OH bands then fractionally distil it; otherwise dilute the thio-ester with dry Et_2O , wash with 5% KOH and H_2O , dry over MgSO_4 and fractionate it through an efficient column [Hauptschein et al. *J Am Chem Soc* 74 4005 1952, DOI: 10.1021/ja01136a010]. [*Beilstein* 2 IV 567.] *Powerful obnoxious odour.*

Ethyl vinyl ether (ethoxyethylene) [109-92-2] $\text{C}_4\text{H}_8\text{O}$, **M 72.1**, **m** -116° , **b** $35.5^\circ/\text{atm}$, **d**₄²⁰ **0.755**. It usually contains polymerisation inhibitors (usually amines, e.g. triethanolamine) which can be removed by fractional distillation. Redistil it from sodium. [Beilstein 1 IV 2049.] **LACHRYMATORY**.

Fluoroacetamide [640-19-7] $\text{C}_2\text{H}_2\text{F}_3\text{NO}$, **M 77.1**, **m** 108° . Crystallise fluoroacetamide from chloroform and dry it in a vacuum. **TOXIC** [Beilstein 2 IV 454.]

Formaldehyde [50-00-0] CH_2O , **M 30.0**, **m** -92° , **b** $-79.6^\circ/20\text{mm}$, $-19.5^\circ/760\text{mm}$, **d**₄²⁰ **0.815**, **pK**²⁵ **13.27 (hydrate)**. Technical aqueous formaldehyde (**formalin**) solution commonly contains added MeOH (8-10%) to inhibit oxidation to formic acid. As a rough guide the **d**₄¹⁸ vs [concentration in g of HCHO/100ml (100g) of aqueous solution] is as follows: 1.0054 [2.24 (2.23)], 1.0126 [4.66 (4.60)], 1.0311 [11.08 (10.74)], 1.0410 [14.15 (13.59)], 1.0568 [19.89 (18.82)], 1.0719 [25.44 (23.73)], 1.0853 [30.17 (27.80)], 1.1057 [37.72 (34.11)] and 1.1158 [41.87 (37.53)]; values in curved brackets are from alternative determinations. [Marvel & Porter *Org Synth Coll Vol I* 377 1941, DOI: 10.15227/orgsyn.009.0058]. If pure formaldehyde is required, add KOH solution (1 mole KOH: 100 moles HCHO) to ~37% by weight of aqueous formaldehyde solution (**formalin**), or evaporate formalin to dryness, to give **paraformaldehyde polymer** $\{\text{HO}(\text{CH}_2\text{O})_n\text{H}$, [30525-89-4] **m** **120-170°** depending on *n* as a white solid which, after washing with water, is dried in a vacuum desiccator over P_2O_5 or H_2SO_4 . Formaldehyde is regenerated by heating the paraformaldehyde to 120° under vacuum, or by decomposing it with barium peroxide. The *monomer*, a colourless flammable gas, is passed through a glass-wool filter cooled to -48° in a CaCl_2 /ice mixture to remove particles of polymer, then dried by passage over P_2O_5 and either condensed in a bulb immersed in liquid nitrogen or absorbed in ice-cold conductivity water. The gas or aqueous solutions have *pungent suffocating odours*, are **LACHRYMATORY** and **SUSPECTED CARCINOGEN**; handle carefully. Formalin is a disinfectant and a preservative of dead animal and plant tissues. [Beilstein 1 IV 3017.]

Formaldehyde dimethyl acetal (dimethoxymethane, methylal, formal) [109-87-5] $\text{C}_3\text{H}_8\text{O}_2$, **M 76.1**, **m** -108° , **b** $41-42^\circ/736\text{mm}$, $41-43^\circ/\text{atm}$, $42-46^\circ/\text{atm}$, **d**₄²⁰ **0.8608**, **n**_D²⁰ **1.35335**. It is a volatile flammable liquid which is soluble in three parts of H_2O , and is readily hydrolysed by acids. Purify it by shaking with an equal volume of 20% aqueous NaOH, stand for 20 minutes, dry over fused CaCl_2 , filter and fractionally distil it through an efficient column. Store it over molecular sieves. [Buehler et al. *Org Synth Coll Vol 3* 468 1955, DOI: 10.15227/orgsyn.020.0059; Rambaud & Besserre *Bull Soc Chim Fr* 45 1955, IR: Wilmshurst *Can J Chem* 36 285 1958, DOI: 10.1139/v58-040; Beilstein 1 IV 3026.]

Formaldehyde dimethyl mercaptal (bis-[methylthio]methane) [1618-26-4] $\text{C}_3\text{H}_8\text{S}_2$, **M 108.2**, **b** $44-47^\circ/13\text{mm}$, $45.5^\circ/18\text{mm}$, $148-149^\circ/\sim 760\text{mm}$, **d**₄²⁰ **1.0594**, **n**_D²⁰ **1.5322**. **Work in an efficient fume cupboard as the substance may contain traces (or more) of methylmercaptan which has a very bad odour and is toxic.** Dissolve the mercaptal in Et_2O , shake it with aqueous alkalis then dry it over anhydrous K_2CO_3 , filter and distil it over K_2CO_3 under a stream of N_2 . If the odour is very strong, then allow all gas effluents to bubble through 5% aqueous NaOH solution which is then treated with dilute KMnO_4 in order to oxidise MeSH to odourless products. Its UV has λ_{max} at 238 nm (log ϵ 2.73) [Fehnel & Carmack *J Am Chem Soc* 71 84 1949, DOI: 10.1021/ja01169a025; Fehér & Vogelbruch *Chem Ber* 91 996 1958, DOI: 10.1002/cber.19580910521; Böhme & Marz *Chem Ber* 74 1667 1941, DOI: 10.1002/cber.19410741009]. Oxidation with aqueous KMnO_4 yields **bis-(methylsulfonyl)methane** which has **m** **142-143°** [Fieccchi et al. *Tetrahedron Lett* 1681 1967, DOI: 10.1016/S0040-4039(00)90698-1]. [Beilstein 1 IV 3088.]

Formamide [75-12-7] CH_3NO , **M 45.0**, **f** 2.6° , **b** $103^\circ/9\text{mm}$, $210.5^\circ/760\text{mm}(\text{dec})$, **d**₄²⁰ **1.13**, **n**_D²⁰ **1.44754**, **n**²⁵ **1.44682**. Formamide is easily hydrolysed by acids and bases. It also reacts with peroxides, acid halides, acid anhydrides, esters and (on heating) alcohols, while strong dehydrating agents convert it to a nitrile. It is very *hygroscopic*. Commercial material often contains acids and ammonium formate. Verhoek [*J Am Chem Soc* 58 2577 1956, DOI: 10.1021/ja01303a059] added some bromothymol blue to formamide and then neutralised it with NaOH before heating to $80-90^\circ$ under reduced pressure to distil off ammonia and water. The amide is again neutralised and the process is repeated until the liquid remained neutral on heating. Sodium formate is added, and the formamide is concentrated under reduced pressure at $80-90^\circ$. The distillate is again neutralised and re-

distilled. It is then fractionally crystallised in the absence of CO₂ and water by partial freezing. Formamide (specific conductance $2 \times 10^{-7} \text{ ohm}^{-1} \text{ cm}^{-1}$) of low water content is dried by passage through a column of 3A molecular sieves, then deionized by treatment with a mixed-bed ion-exchange resin loaded with H⁺ and HCONH⁻ ions (using sodium formamide in formamide) [Notley & Spiro *J Chem Soc (B)* 362 1966, DOI: 10.1039/J29660000362]. [*Beilstein* 2 IV 45.]

Formamidine acetate [3473-63-0] CH₄N₂. C₂H₄O₂, M 104.1, m 159-161°(dec), 164°(dec), pK_{Est} ~12. Unlike the hydrochloride below, the acetate salt is not hygroscopic. Prepared from (EtO)₃CH in AcOH by passing NH₃ gas through at 125-130° for 20 to 30 minutes when the acetate starts to crystallise out. Cool, filter the crystals off, wash them with cold EtOH and dry them *in vacuo*. It is recrystallised from a small volume of acetic acid, by addition of EtOH, and the crystals are washed with cold EtOH then Et₂O and dried in a vacuum. It is stable to storage and can be used directly in place of free formamidine. It is less hygroscopic than the hydrochloride below. [Taylor & Rabinovitz *Biochemical Preparations* 5 102 1957, Taylor and Ehrhart *J Am Chem Soc* 82 3138 1960, DOI: 10.1021/ja01497a039; Taylor et al. *Org Synth* 46 39 1966, DOI: 10.15227/orgsyn.046.0039; *Beilstein* 2 IV 82.]

Formamidine hydrochloride [6313-33-3] CH₄N₂. HCl, M 80.1, m 76-78°, 78°, 84-87°, 85-87°(dec), pK_{Est} ~12. The hydrochloride is hygroscopic unlike the acetate salt (see preceding entry). It has been prepared by catalytic reduction of freshly prepared cyanamide in 9% aqueous HCl (to form the hydrochloride) or 12% of aqueous H₂SO₄ (to form the hemisulfate, see below) with 1.4% of Pd/C and hydrogen. The reduction is usually complete in 10 minutes. The hygroscopic *hydrochloride*, obtained in 92% yield, is recrystallised from MeOH and is stored in a vacuum.

Formamidine hemisulfate [77269-36-4; 77269-37-5] CH₄N₂. 0.5 H₂SO₄, M 93.1, has m 156-158° after recrystallisation from MeOH. **Formamidine picrate** CH₄N₂. C₅H₃N₃O₇, has m 246-247° after recrystallisation from hot H₂O, and can be readily converted to the *hydrochloride*, m 78°. [Odo et al. *J Org Chem* 22 1715 1957, DOI: 10.1021/jo01363a620; Shirai et al. *J Org Chem* 23 100 1958, DOI: 10.1021/jo01095a608; Taylor et al. *Org Synth* 46 39 1966, DOI: 10.15227/orgsyn.046.0039; *Beilstein* 3 H 90, 2 I 38, 2 III 125, 2 IV 82.]

Formamidine sulfinic acid (thiourea-S-dioxide) [1758-73-2] CH₄N₂O₂S, M 108.1, m 124-126°(dec). Dissolve it in five parts of aqueous 1:1% NaHSO₃ at 60-63° (charcoal), then allow it to crystallise slowly, with agitation, at 10°. Filter and dry it immediately at 60° [Koniecki & Linch *Anal Chem* 30 1134 1958, DOI: 10.1021/ac60138a041; Havel & Kluttz *Synth Commun* 4 389 1974, DOI: 10.1080/00397917408064100]. [*Beilstein* 3 I 36, 3 IV 145.]

Formic acid [64-18-6] CH₂O₂, M 46.0 (anhydrous), f 8.3°, b 25°/40mm, 100.7°/760mm, d₄²⁰ 1.22, n_D²⁵ 1.37140, n_D²⁵ 1.36938, pK₂₅ 3.74. Anhydrous formic acid can be obtained by direct fractional distillation under reduced pressure, the receiver being cooled in ice-water. The use of P₂O₅ or CaCl₂ as dehydrating agents is unsatisfactory. Reagent grade 88% formic acid can be dried satisfactorily by refluxing with phthalic anhydride for 6 hours and then distilling it. *Alternatively*, if it is left in contact with freshly prepared anhydrous CuSO₄ for several days about one half of the water is removed from 88% formic acid; distillation then removes the remainder. Boric anhydride (prepared by melting boric acid in an oven at a high temperature, cooling in a desiccator, and powdering) is a suitable dehydrating agent for 98% formic acid; after prolonged stirring with the anhydride the formic acid is distilled under vacuum. Formic acid can be further purified by fractional crystallisation using partial freezing. [*Beilstein* 2 IV 3.]

N-Formyl tert-butylamine (N-tert-butylformamide) [2425-74-3] C₅H₁₁NO, M 101.2, m 16°, b 48°/0.2mm, 78-83°/9mm, 135-136°/107mm, 202°/760mm, d₄²⁵ 0.903, n_D²⁰ 1.4330. If the IR indicates some hydrolysis, then dissolve it in Et₂O, wash it with 20% aqueous Na₂CO₃, dry it (MgSO₄), filter and fractionate it. Collect the fraction that solidifies on cooling and recrystallise it from Et₂O at low temperature if necessary. [Emmons *J Am Chem Soc* 79 5739 1957, DOI: 10.1021/ja01578a043; *Beilstein* 4 III 324, 4 IV 661.]

N-Formyl ethylamine (N-ethylformamide) [627-45-2] C₃H₇NO, M 73.1, b 29°/0.5mm, 176-179°/758mm,

d_4^{20} 0.950, n_D^{20} 1.4346. If the IR is good, then distil it and collect the middle fraction and redistil if necessary; otherwise proceed as for the previous amide. [Erickson *J Org Chem* **20** 1569 1955, DOI: 10.1021/jo01128a016; *Beilstein* **4** H 109, **4** I 352, **4** II 601, **4** III 207, **4** IV 346.]

Formyl hydrazine (formic acid hydrazide) [624-84-0] $\text{CH}_4\text{N}_2\text{O}$, **M 60.1, m 54°, 54-57°, $pK_{\text{est}} \sim 2.5$.** Recrystallise it from EtOH and dry it *in vacuo*. Store below 10°; it may disproportionate on storage to 1,2-diformyl hydrazine and hydrazine. It forms a blue $[\text{Cu}(\text{CH}_4\text{N}_2\text{O})]\text{SO}_4$ salt with CuSO_4 . [*Beilstein* **2** H 93, **2** III 127, **2** IV 85.]

Formyloxy acetonitrile (cyanomethyl formate) [150760-95-5] $\text{C}_3\text{H}_3\text{NO}_2$, **M 85.1, b 62-64°/12mm, 172-173°/atm, d_4^{25} 0.903, n_D^{20} 1.4330.** Purify it by fractional distillation and redistilling the middle fraction. It is useful for the formylation of alcohols and amines. The ^{13}C NMR has δ (CDCl_3) at 47.87, 114.47 and 159.46 ppm. [Deutsch & Niclas *Synth Commun* **23** 1561 1993, DOI: 10.1080/00397919308011251; Ducek et al. *Synthesis* **37** 1996, DOI: 10.1055/s-1996-4153.] [*Beilstein* **3** H 243.]

Fumaraldehyde bis(dimethyl acetal) (trans-1,1,4,4-tetramethoxybut-2-ene) [6068-62-8] $\text{C}_8\text{H}_{16}\text{O}_4$, **M 176.2, b 100-103°/15mm, 101-103°/25mm, d_4^{20} 1.011, n_D^{20} 1.425.** Dry it over fused CaCl_2 and distil it *in vacuo*. The maleic (*cis*) isomer has **b 112°/11mm**, and d^{23} 0.932 and n_D^{25} 1.4243. [Zeik & Heusner *Chem Ber* **90** 1869 1957, Clauson-Kaas et al. *Acta Chem Scand* **9** 111 1955, DOI: 10.3891/acta.chem.scand.09-0111; Clauson-Kaas *Acta Chem Scand* **6** 569 1952, DOI: 10.3891/acta.chem.scand.06-0569; *Beilstein* **1** IV 3754.]

Fumaric (trans-but-2-ene-1,4-dioic) acid [110-17-8] $\text{C}_4\text{H}_4\text{O}_4$, **M 116.1, m 289.5-291.5°(sealed tube), pK_1^{25} 3.10, pK_2^{25} 4.60 (4.38).** Crystallise it from hot M HCl or water and dry it at 100°. [*Beilstein* **2** IV 2202.]

Geraniol (trans-3,7-dimethyl-2,6-octadien-1-ol) [106-24-1] $\text{C}_{10}\text{H}_{18}\text{O}$, **M 154.3, b 114-115°/11-12mm, 230°, d_4^{20} 0.879, n_D^{20} 1.4766.** Purify geraniol by ascending chromatography or by thin layer chromatography on plates of kieselguhr G with acetone/water/liquid paraffin (130:70:1) as solvent system. Hexane/ethyl acetate (1:4) is also suitable. Also purify it by GLC on a silicone-treated column of Carbowax 20M (10%) on Chromosorb W (60-80 mesh). [Porter *Pure Appl Chem* **20** 449 1969, DOI: 10.1351/pac196920040449.] Store it in full, tightly sealed containers in the cool and protect from light. It has a pleasant odour of geraniums. [*Beilstein* **1** IV 2277.]

Glutaraldehyde [111-30-8] $\text{C}_5\text{H}_8\text{O}_2$, **M 100.1, b 71°/10mm, as 50% aqueous solution.** Likely impurities are oxidation products-acids, semialdehydes and polymers. It can be purified by repeated washing with activated charcoal (Norit) followed by vacuum filtration, using 15-20g charcoal/100ml of glutaraldehyde solution. Distil it at 60-65°/15mm, discarding the first 5-10%, then dilute with an equal volume of freshly distilled water at 70-75°, using magnetic stirring under nitrogen. The solution is stored at low temperature (3-4°), in a tightly stoppered container, and protected from light. Standardise by titration with hydroxylamine. [Anderson *J Histochem Cytochem* **15** 652 1967, DOI: 10.1177/15.11.652; *Beilstein* **1** IV 3659.] **TOXIC.**

Glutaric acid [110-94-1] $\text{C}_5\text{H}_8\text{O}_4$, **M 132.1, m 97.5-98°, pK_1^{25} 4.35, pK_2^{25} 5.40.** Crystallise the acid from *benzene, CHCl_3 , distilled water or *benzene containing 10% (w/w) of diethyl ether. Dry it under vacuum. [*Beilstein* **2** IV 1934.]

dl-Glyceraldehyde [56-82-6] $\text{C}_3\text{H}_6\text{O}_3$, **M 90.1, m 145°, distils (bath temperature 140-150°) at ~0.8mm.** Crystallise it from EtOH/diethyl ether. Its solubility in H_2O at 18° is 3g/100ml. It is insoluble in hydrocarbon solvents, and forms an **osazone m 138°**. The **D(R+)-enantiomer** [453-17-8] is a syrup (~70% + H_2O) with $[\alpha]_D^{25}$ +14, +8.7 (c 2, H_2O) and the **dimethyl acetal** has **b 124-127°/14mm** and $[\alpha]_D^{15}$ +21 (c 18, H_2O). The **L(S-)-enantiomer** [497-09-6] has $[\alpha]_D^{25}$ -14.3 (c 5, H_2O) and mutarotates from $[\alpha]_D^{20}$ -13.8 to -7.8 in 8 days [Baer & Fischer *J Am Chem Soc* **61** 761 1939, DOI: 10.1021/ja01873a001]; and the **dimethyl acetal** has **b 126-129°/17-20mm** and $[\alpha]_D^{15}$ -20.9 (c 9.2, H_2O). [cf. Witzemann *Org Synth Coll Vol* **2** 305 1943, DOI: 10.15227/orgsyn.011.0050; Perlin *Methods in Carbohydrate Chemistry* **1** 61 1962, *Beilstein* **1** H 845, **1** H 427, **1** II 888, **1** III 3282, 2391, **1** IV 4114.]

dl-Glyceraldehyde Dimer [(±) 3,6-dihydroxy-1,4-dioxane-2,5-dimethanol] [51795-26-7, 26793-98-6] $C_6H_{12}O_6$, M 180.2, m 144-145°. Recrystallise from EtOH (m 120-121°). [Fischer & Baer *Chem Ber* **65** 345 1932, DOI: 10.1002/cber.19320650244; *Beilstein* 1 H 845, 1 IV 4418.]

Glycerol [56-81-5] $C_3H_8O_3$, M 92.1, m 18.2°, b 182°/20mm, 290°/760mm, d_4^{20} 1.261, n_D^{25} 1.47352, pK^{25} 14.4. Glycerol is dissolved in an equal volume of *n*-butanol (or *n*-propanol, amyl alcohol or liquid ammonia) in a water-tight container, cooled and seeded while slowly revolving in an ice-water slurry. The crystals are collected by centrifugation, then washed with cold acetone or isopropyl ether. [Hass & Patterson *Ind Eng Chem (Anal Ed)* **33** 615 1941, DOI: 10.1021/ie50377a015.] Coloured impurities can be removed from substantially dry glycerol by extraction with 2,2,4-trimethylpentane. Alternatively, glycerol can be decolorised and dried by treatment with activated charcoal and alumina, followed by filtering. Glycerol can be distilled at 15mm in a stream of dry nitrogen, and stored in a desiccator over P_2O_5 . Crude glycerol can be purified by digestion with concentrated H_2SO_4 and then saponification with a lime paste, re-acidification with H_2SO_4 , filtration, treatment with an anion exchange resin and fractional distillation in a vacuum. [*Beilstein* 1 IV 2751.]

Glyceryl trinitrate (TNG, trinitroglycerine, nitroglycerin, 1,2,3-trinitroxypropane, 'Blasting Oil', Transiderm-nitro, Nitrolingual, and many other trade names) [55-63-0] $C_3H_5N_3O_9$, M 227.1, m 14°, b 50°/atm (dec, explosive), d^{20} 1.60g/cm³, n_D^{15} 1.474. Sobrero [*Justus Liebigs Ann Chem* **64** 398 1847, DOI: 10.1002/jlac.18480640364; Williamson *Justus Liebigs Ann Chem* **92** 305 1854, DOI: 10.1002/jlac.18540920309] was first to prepare the ester by dissolving syrupy glycerin dropwise into a cold mixture of H_2SO_4 (2 parts) and HNO_3 (1 part) until no more reaction occurred. It was then poured into cold H_2O whereby an oil separated which was washed several times with H_2O by decantation, dissolved in EtOH and the oil separated on adding H_2O and repeating the process. The olive-yellow oil was dissolved in Et₂O (separated from any aqueous layer) and evaporated (**TAKE GREAT CARE HERE AS THIS IS VERY EXPLOSIVE**) to give pure oil which was dried over H_2SO_4 . It detonated on attempted heating. Alternatively, it was prepared by nitrating glycerol with white fuming HNO_3 and keeping the temperature as low as possible. **NO ATTEMPT SHOULD BE MADE TO DISTIL IT** at atmospheric pressure. If it has to be distilled then very small volumes could be transferred from one bulb to another very cold one under vacuum and great precautions should be taken (its vapour pressure at 20°/atm is 0.00026mm).

The tri-nitrate ester is a pale yellow oil which explodes on concussion, is volatile and is sweet smelling. It should not be allowed to come into contact with skin because it is readily absorbed, and on **no account** should it be inhaled as it causes nausea and throbbing headaches (**use a well ventilating fume hood**). Its solubility (%w/w) is 0.13 in H_2O , 0.83 in CS_2 , 5.6 in MeOH and 25.0 in EtOH; and it is very soluble in most organic solvents except for light petroleum and glycerol.

TNG is an ingredient of explosives used by the military and in the construction industry. Medicinally, it is a potent vasodilator which increases blood flow and widely used in patients with heart conditions, e.g. angina and heart failure [Di Carlo 'Nitroglycerin Revisited: Chemistry, Biochemistry, Interactions' *Drug Metabolism Reviews* **4** 1 1975, DOI: 10.3109/03602537508993747. PMID 812687.]

Glycolic (α-hydroxyacetic) acid [79-14-1] $C_2H_4O_3$, M 76.1, m 81°, pK^{25} 3.62. Crystallise it from diethyl ether. [*Beilstein* 3 IV 571.]

Guanidine [113-00-8] CH_5N_3 , M 59.1, m 47.5-48.5°, 48-49°, ~50°, pK^{25} 13.6. Crystallise it from water/EtOH under nitrogen. It is very deliquescent and absorbs CO_2 from the air readily. [Davis *Org Synth Coll Vol* **1** 302 1941, DOI: 10.15227/orgsyn.007.0046; Jones *Trans Faraday Soc* **55** 524 1959, DOI: 10.1039/TF9595500524; *Beilstein* 3 H 82, 3 I 39, 3 II 69, 3 III 154, 3 IV 148.]

Guanidine carbonate [593-85-1] $CH_5N_3 \cdot H_2CO_3$, M 90.1, m 197°, 230°. Crystallise it from MeOH. [*Beilstein* 3 H 86, 3 I 41, 3 II 72, 3 III 161, 3 IV 152.]

Guanidine hydrochloride [50-01-1] $CH_5N_3 \cdot HCl$, M 95.5, m 181-183°. Crystallise the hydrochloride from hot methanol by chilling to about -10°, with vigorous stirring. The fine crystals are filtered through fritted glass, washed with cold (-10°) methanol, dried at 50° under vacuum for 5 hours. (The product is purer than that obtained by crystallisation at room temperature from methanol by adding large amounts of diethyl ether.)

[Kolthoff et al. *J Am Chem Soc* **79** 5102 1957, DOI: 10.1021/ja01576a007; *Beilstein* **3** III 160, **3** IV 150.]

Heptadecanoic acid (margaric) [506-12-7] $C_{17}H_{34}O_2$, **M 270.5**, **m 60-61°, 61.3°, b 227°/100mm**, **pK_{Est} ~4.9**. Crystallise this C-17 (odd number of carbon atoms) acid from MeOH or petroleum ether (colourless plates). Its solubilities at 20° are: H₂O (0.00042), AcOH (1.2), MeOH (2.5), 95% EtOH (4.17), *C₆H₆ (9.3) and Et₂O (very soluble). [Occurrence in butterfat: Hansen et al. *Nature* **179** 98 1957, DOI:10.1038/179098a0; isolation from shark liver oil: Morice & shortland *Biochem J* **61** 453 1955, DOI: 10.1042/bj0610453; from degradation of stearic acid: Levene & West *J Biol Chem* **16** 475 1914; synthesis: Kaufmann & Stamm *Chem Ber* **91** 2121 1958, DOI: 10.1002/cber.19580911018; *Beilstein* **2** IV 1193.] The triglyceride, **glyceryl 1,2,3-triheptadecanoate (triheptadecanoin)** [2438-40-6] $C_{54}H_{104}O_6$, **M 849.4**, has **m 64°, 64-67°, b 786.8°/760mm**, and is crystallised from CHCl₃/petroleum ether or aqueous EtOH. Store under N₂ at -20°. [Gas chromatographic separation of triglycerides: Kail et al. *J Chromatogr Sci* **50** 934 2012, DOI: 10.1093/chromsci/bms093; *Beilstein* **2** III 986.]

1-Heptadecanol [1454-85-9] $C_{17}H_{36}O$, **M 256.5**, **m 54°**. Crystallise it from acetone. [*Beilstein* **1** IV 1884.]

Heptafluoro-2-iodopropane [677-69-0] C_3F_7I , **M 295.9**, **b 39°/735mm, 41°/760mm**, **d₄⁰ 2.1306, n_D²⁰ 1.3281**. Purify it by gas chromatography on a triacetin (glyceryl triacetate) column, followed by bulb-to-bulb distillation at low temperature. Store it over Cu powder to stabilise it. UV has ν_{max} at 271nm (ϵ 240) in petroleum ether (b 60-80°). [Haszeldine *J Chem Soc* 1764, 3761 1953, DOI: 10.1039/JR9530001764; DOI: 10.1039/JR9530003761; *Beilstein* **1** III 255, **1** IV 225.]

n-Heptaldehyde [111-71-7] $C_7H_{14}O$, **M 114.2**, **m -43°, b 40.5°/12mm, 152.8°/760mm**, **d₄²⁰ 0.819, n_D²⁵ 1.4130**. Dry n-heptaldehyde with CaSO₄ or Na₂SO₄ and fractionally distil it under reduced pressure. More extensive purification is by precipitation as the bisulfite compound (formed by adding the aldehyde to saturated aqueous NaHSO₃) which is filtered off and recrystallised from hot H₂O. The crystals, after being filtered and washed well with H₂O, are hydrolysed by adding 700ml of aqueous Na₂CO₃ (12.5% w/w of anhydrous Na₂CO₃) per 100g of aldehyde. The aldehyde is then steam distilled off, separated, dried with CuSO₄ and distilled under reduced pressure in a slow stream of nitrogen. [McNesby & Davis *J Am Chem Soc* **76** 2148 1954, DOI: 10.1021/ja01637a031; *Beilstein* **1** H 695, **1** I 357, **1** II 750, **1** III 2844, **1** IV 3314.]

n-Heptaldoxime [629-31-2] $C_7H_{15}NO$, **M 129.2**, **m 53-55°**. Separate the *cis*(Z) and *trans*(E) oximes by liquid chromatography through a silica gel column and eluting with petroleum ether (b 40-65°)/EtOAc (50:10) at a flow rate of 2-3.4 ml/sec where the *trans*-isomer comes through first and is a liquid with n_D²² 1.38512, followed by the *cis*-isomer which is a solid, and crystallises from 60% aqueous EtOH with **m 55°**. They are identified by TLC on 0.2mm silica gel G by eluting with *C₆H₆/EtOAc (50/10) and visualising with I₂ vapour: the *trans*-isomer has R_F 0.6 and the *cis*-isomer has R_F 0.5 [Pejkovic-Tadic et al. *J Chromatogr* **21** 239 1966, DOI: 10.1016/S0021-9673(01)91297-X; Emmons & Pagano *J Am Chem Soc* **77** 4557 1955, DOI: 10.1021/ja01622a036]. [*Beilstein* **1** H 698, **1** I 358, **1** II 752, **1** III 2850.]

n-Heptane [142-18-5] C_7H_{16} , **M 100.2**, **m -90°, b 98.4°/atm**, **d₄²⁰ 0.684, n_D²⁰ 1.38765, n_D²⁵ 1.38512**. Pass it through a silica gel column which greatly reduces the ultraviolet absorption of n-heptane. (The silica gel is previously heated to 350° before use.) For more extensive purification, heptane is shaken with successive small portions of concentrated H₂SO₄ until the lower (acid) layer remains colourless. The heptane is then washed successively with water, aqueous 10% Na₂CO₃, water (twice), and dried with CaSO₄, MgSO₄ or CaCl₂. It is distilled from sodium. n-Heptane can be distilled azeotropically with methanol, then the methanol is washed out with water and, after drying, the heptane is redistilled. Other purification procedures include passage through activated basic Al₂O₃, drying with CaH₂, storage with sodium, and stirring with 0.5N KMnO₄ in 6N H₂SO₄ for 12 hours after treatment with concentrated H₂SO₄. Carbonyl-containing impurities have been removed by percolation through a column of impregnated Celite made by dissolving 0.5g of 2,4-dinitrophenylhydrazine in 6ml of 85% H₃PO₄ by grinding together, then adding 4ml of distilled water and 10g Celite. [Schwartz & Parks *Anal Chem* **33** 1396 1961, DOI: 10.1021/ac60178a036; *Beilstein* **1** IV 376.]

Hept-1-ene [592-76-7] C_7H_{14} , **M 98.2**, **m** -119°, **b** 93°/771mm, **d**₄²⁰ 0.698, **n**_D²⁰ 1.400. Distil hept-1-ene from sodium, then carefully fractionally distilling it using an 18-in gauze-packed column. It can also be purified by azeotropic distillation with EtOH. It usually contains the 2- and 3-isomers as impurities. These can be removed by gas chromatography using a Carbowax column at 70°. [*Beilstein* 1 IV 857.]

n-Heptyl alcohol (1-heptanol) [111-70-6] $C_7H_{16}O$, **M 116.2**, **m** -36°, **b** 175.6°/atm, **d** 0.825, **n**_D²⁰ 1.425. Shake the alcohol with successive lots of alkaline $KMnO_4$ until the colour persists for 15 minutes, then dry it with K_2CO_3 or CaO, and fractionally distil it. [*Beilstein* 1 IV 1731.]

n-Heptylamine [111-68-2] $C_7H_{17}N$, **M 115.2**, **b** 155°/atm, **d**₄²⁰ 0.775, **n**_D²⁰ 1.434, **pK**²⁵ 10.66. Dry it in over KOH pellets for 24 hours, then decant it and fractionally distil it. Store away from CO_2 . [*Beilstein* 4 IV 734.]

n-Heptyl bromide [629-04-9] $C_7H_{15}Br$, **M 179.1**, **m** -58°, **b** 70.6°/19mm, 180°/760mm, **d**₄²⁰ 1.140, **n**_D²⁰ 1.45. Shake it with conc H_2SO_4 , wash with water, dry it with K_2CO_3 , and fractionally distil. [*Beilstein* 1 IV 391.]

Hexachloro-1,3-butadiene (perchlorobutadiene) [87-68-3] C_4Cl_6 , **M 260.8**, **m** -22° to -19°, **b** 144.1°/100mm, 210-212°/760mm, **d**₄²⁰ 1.683, **n**_D²⁰ 1.5556. Wash the diene with four or five 1/10th volumes of MeOH (or until the yellow colour has been extracted), then stir it for 2 hours with H_2SO_4 , wash it with distilled water until neutral and filter it through a column of P_2O_5 . Distil it under reduced pressure through a packed column. [Rutner & Bauer *J Am Chem Soc* 82 298 1960, DOI: 10.1021/ja01487a012; *Beilstein* 1 IV 998.]

Hexachloroethane [67-72-1] C_2Cl_6 , **M 236.7**, **m** 187°. Steam distil it, then crystallise it from 95% EtOH. Dry it in the dark under vacuum. [*Beilstein* 1 IV 148.]

Hexacosane (C-26) [630-01-3] $C_{26}H_{54}$, **M 366.7**, **m** 56.4°, **b** 169°/0.05mm, 205°/1mm, 262°/15mm. Distil hexacosane under vacuum and recrystallise it from diethyl ether. [*Beilstein* 1 IV 583.]

Hexacosanoic acid (cerotic acid, cerotinic acid) [506-46-7] $C_{26}H_{52}O_2$, **M 396.7**, **m** 86-87°, 88-89°, **pK**_{Est} ~4.9. Crystallise the acid from EtOH, aqueous EtOH and petroleum ether/ Me_2CO . [*Beilstein* 2 IV 1310.]

n-Hexadecane (Cetane) [544-76-3] $C_{16}H_{34}$, **M 226.4**, **m** 18.2°, **b** 105°/0.1mm, 287°/atm, **d**₄²⁰ 0.773, **n**_D²⁰ 1.4345, **n**²⁵ 1.4325. Pass cetane through a column of silica gel and distil it under vacuum in a column packed with Pyrex helices. Store it over silica gel. It also crystallises from acetone, or is fractionally crystallised by partial freezing. [*Beilstein* 1 IV 537.]

1,14-Hexadecanedioic acid (Thaspic acid) [505-54-4] $C_{16}H_{30}O_4$, **M 286.4**, **m** 126°, **pK**_{Est(1)} ~4.5, **pK**_{Est(2)} ~5.5. Recrystallise thaspic acid from EtOH, ethyl acetate or $*C_6H_6$. [*Beilstein* 2 IV 2162.]

Hexadecanoic acid (palmitic acid) [57-10-3] $C_{16}H_{32}O_2$, **M 256.4**, **m** 62-63°, **b** 215°/15mm, **pK**²⁵ 6.46 (50% aqueous EtOH), 5.0 (H_2O). Purify palmitic acid by slow (overnight) recrystallisation from hexane. Some samples are also crystallised from acetone, EtOH or EtOAc. The crystals are kept in air to lose solvent, or are pumped dry of solvent on a vacuum line. [Iwahashi et al. *JCS Faraday Trans 1* 81 973 1985, DOI: 10.1039/F19858100973; **pK**: White *J Am Chem Soc* 72 1859 1950, DOI: 10.1021/ja01160a530; *Beilstein* 2 IV 1157.]

1,5-Hexadiene [592-42-7] C_6H_{10} , **M 82.2**, **m** -141°, **b** 59.6°/atm, **d**₄²⁰ 0.694, **n**_D²⁰ 1.4039. Distil 1,5-hexadiene from $NaBH_4$ and store under Ar or N_2 . [*Beilstein* 1 IV 1013.]

Hexafluoroacetone [684-16-2, 34202-69-2 ($3H_2O$)] C_3F_6O , **M 166.1**, **m** -43-45° (sealed capillary) -129°, **b** -26°, -28°, (trihydrate **m** 18-21°, **b** 55-56°/80mm). Dehydrate hexafluoroacetone by passing the vapours over P_2O_5 . Ethylene is removed by passing the dried vapours through a tube containing Pyrex glass wool moistened with conc H_2SO_4 . Further purification is by low temperature distillation using Warde-Le Roy stills and Dry Ice. Store it in the dark at -78° or at liquid nitrogen temperature. Henne et al. used *Engler* distillation [Henne et al. *J Am Chem Soc* 72 3577 1960, DOI: 10.1021/ja01164a070; Holmes & Kutschke *Trans Faraday Soc* 58 333

1962, DOI: 10.1039/TF9625800333; *Beilstein* 1 IV 3215 3210.]

Hexafluoroacetylacetone (1,1,1,5,5,5-hexafluoro-2,4-pentanedione, hfacac) [1522-22-1] $C_5H_2F_6O_2$, M 208.1, b 63-65°/atm, 68°/736mm, 70-70.2°/760mm, 68-71°/atm, d_4^{20} 1.490, n_D^{20} 1.333. Hfacac has been prepared from a mixture of Na wire (0.34mole) covered with dry Et_2O and CF_3CO_2Et (0.27mole see [383-63-1]) at 0°, to which was added trifluoroacetone (0.27mol, see [421-50-1]) at -78°. The mixture was then set aside overnight at ~25°, and any trifluoroacetone which distilled out was made to condense back into the reaction mixture which was stirred at ~25° until all the Na dissolved [Haszeldine et al. *J Chem Soc* 609 1951, DOI: 10.1039/JR9510000609; compare with Henne et al. *J Am Chem Soc* 69 1819 1947, DOI: 10.1021/ja01199a075]. The reddish-brown mixture was treated with an excess of 2N H_2SO_4 and extracted with ether. The organic layer was dried (Na_2SO_4), and distilled to give hfacac (20g, 37%). By using NaOEt instead of Na metal, Henne and coworkers obtained a 72% yield of hfacac, which gave the *copper chelate* as bright grass green crystals (see below) [Henne et al. *J Am Chem Soc* 69 1819 1947, DOI: 10.1021/ja01199a075]. Hfacac (100g) is purified by shaking twice with 98% H_2SO_4 (300ml, *use protective clothing*) until it is completely dispersed, and set aside overnight. The anhydrous facac is separated and distilled through a glass-helices packed column with fraction b 70.2-70.5°/atm (~20g, analytically pure) being retained for metal complex studies [Buckingham et al. *Aust J Chem* 20 281 1967, DOI:10.1071/CH9670281]. The oil readily forms a solid white stable *covalent dihydrate* [$CF_3-C(OH)_2-CH_2-C(OH)_2-CF_3$] which is caused by the strong electron-withdrawing effect of the fluorine atoms. The dihydrate has no UV absorption spectrum; compare with λ_{max} ($CHCl_3$) 273nm (ϵ 7,800) for the anhydrous diketone. The dihydrate decomposes at ~90°. The hydrate (10g) can also be dehydrated by heating with anhydrous $CaSO_4$ (Drierite, 30g) and distilling; the distillate is treated with more $CaSO_4$ and redistilled (see above). When the distillate is treated with aqueous NaOH and heated, the dihydrate crystallises on cooling. The *Cu (II) complex* is more easily prepared from the anhydrous than from the hydrated diketone which needs to be dehydrated in the reaction medium, and has m 135° (after sublimation or crystallisation from CCl_4). Store hfacac as the dihydrate; but the anhydrous diketone should be kept in an anhydrous atmosphere. [Gilman et al. *J Am Chem Soc* 78 2790 1956, DOI: 10.1021/ja01593a038; Belford et al. *J Inorg Nucl Chem* 2 11 1956, DOI:10.1016/0022-1902(56)80100-0; *Beilstein* 1 III 3123, 1 IV 3681.]

Hexafluoroethane (CFC-116) [76-16-4] C_2F_6 , M 138.0, m -100°, b -79°/atm. Purify it for pyrolysis studies by passing through a copper vessel containing CoF_3 at ca 270°, and held for 3 hours in a bottle with a heated (1300°) platinum wire. It is then fractionally distilled in a vacuum line. [Steunenberg & Cady *J Am Chem Soc* 74 4165 1962, DOI: 10.1021/ja01136a062; *Beilstein* 1 III 132.]

Hexafluoropropene (hexafluoropropylene, perfluoropropene, F-propene) [116-15-4] C_3F_6 , M 150.0, m -153°, -156.2°, b -28°/atm, -29.4°/atm, d_4^{20} 1.583. F-Propene is a gas that is available commercially in brass cylinders and is very **corrosive, toxic**, attacks skin and tissue membranes. It is prepared by treating $CFCl_2CFCICF_3$ with Zn in boiling $EtOH$ (3 days), but is faster under pressure at 100°. The gas is best handled in a vacuum line and should be scrubbed over caustic alkali in a brass tube, as any HF present in it will attack glass. [Henne & Waalkes *J Am Chem Soc* 68 496 1946, DOI: 10.1021/ja01207a041; Henne & Hinkamp *J Am Chem Soc* 67 1194 1945, DOI: 10.1021/ja01223a051; *Beilstein* 1 III 697, 1 IV 735.]

1,1,1,3,3,3-Hexafluoropropan-2-ol (HFIP) [920-66-1] $C_3H_2F_6O$, M 168.1, m -4°, b 57-58°/760mm, d_4^{20} 1.4563, n_D^{22} 1.2750, pK^{25} 9.30. Distil it from 3A molecular sieves, retaining the middle fraction. It has been prepared by reduction of *hexafluoroacetone* in tetrahydrofuran (THF). In this case hexafluoropropanol forms a stable **1:1 THF complex** which distils at 99-100°/760mm (n_D^{25} 1.3283). The complex is decomposed by mixing with 20% oleum and distilling in a vacuum; the distillate is redistilled to give pure hexafluoropropan-2-ol with b 59°/760mm. The 1H NMR shows a doublet at 4.52ppm ($J_{H,H} = 2Hz$). The *benzoyl* derivative, [10315-85-2] M 272.1, has m 53.9° after crystallisation from pentane at -50°, and its IR has ν_{max} at 1760 cm^{-1} . [Middleton & Lindsey *J Am Chem Soc* 86 4948 1964, DOI: 10.1021/ja01076a041; Urry et al. *J Org Chem* 32 347 1967, DOI: 10.1021/jo01288a021.] It has very high peptide solubilising properties, alone or with CH_2Cl_2 [use as a solvent: Narita et al. *Bull Chem Soc Jpn* 61 281 1988, DOI: org/10.1246/bcsj.61.281; Halverson et al. *Biochemistry* 29 2639 1990, DOI: 10.1021/bi00463a003.] It is **CORROSIVE, causes severe eye irritation**.

Hexamethylenediamine (1,6-diaminohexane) [124-09-4] $C_6H_{16}N_2$, M 116.2, m 42°, b 46-47°/1mm, 84.9°/

9mm, 100°/20mm, 204-205°/760mm, pK_1^{25} 10.24, pK_2^{25} 11.02. Crystallise it in a stream of nitrogen. It sublimes in a vacuum. [*Beilstein* 4 IV 1320.]

Hexamethylenediamine dihydrochloride [6055-52-3] $C_6H_{16}N_2 \cdot HCl$, M 189.2, m 248°, 256-257°. Crystallise the salt from water or EtOH. [*Beilstein* 4 IV 1320.]

Hexamethylene glycol (1,6-hexanediol) [629-11-8] $C_6H_{14}O_2$, M 118.2, m 41.6°, 43-45°, b 134°/10mm, 250°, n_D^{20} 1.458. Fractionally crystallise it from its melt or from water. Distil it *in vacuo*. [*Beilstein* 1 IV 2556.]

***n*-Hexane** [110-54-3] C_6H_{14} , M 86.2, m -95°, b 68.7°/atm, d_4^{20} 0.660, n_D^{25} 1.37486, n_D^{25} 1.37226. Purify as for *n*-heptane. Modifications include the use of chlorosulfonic acid or 35% fuming H_2SO_4 instead of concentrated H_2SO_4 in washing the alkane, and final drying and distilling from sodium hydride. Unsaturated impurities can be removed by shaking the hexane with nitrating acid (58% H_2SO_4 , 25% concentrated HNO_3 , 17% water, or 50% HNO_3 , 50% H_2SO_4), then washing the hydrocarbon layer with concentrated H_2SO_4 , followed by H_2O , drying, and distilling over sodium or *n*-butyl lithium. It can also be purified by distillation under nitrogen from sodium benzophenone ketyl solubilised with tetraglyme. Also purify it by passage through a silica gel column followed by distillation [Kajii et al. *J Phys Chem* 91 2791 1987, DOI: 10.1021/j100295a029]. It is a **FLAMMABLE** liquid and a possible nerve toxin. [*Beilstein* 1 IV 338.]

Rapid purification: Distil, discarding the first forerun and store over 4A molecular sieves.

(±)-1,2-Hexanediol [6920-22-5] $C_6H_{14}O_2$, M 118.2, b 96-98°/1mm, 118.4-118.5°/13mm, 214-215°/760mm, d_4^{20} 0.951, n_D^{20} 1.442. Fractionally distil it, preferably in a vacuum. Alternatively, dissolve it in Et_2O , dry with K_2CO_3 then Na_2SO_4 , filter, evaporate and distil it in a vacuum. The ***bis*-4-nitrobenzoyl** derivative has m 101.5-102.5°. [Rudloff *Can J Chem* 36 486 1958, DOI:10.1139/v58-069; *Beilstein* 1 I 251, 1 III 2200, 1 IV 2554.]

1-Hexene [592-41-6] C_6H_{12} , M 84.2, b 63°/atm, d_4^{20} 0.674, n_D^{20} 1.388. Purify it by stirring over Na/K alloy for at least 6 hours, then fractionally distil it from sodium under nitrogen. [*Beilstein* 1 IV 828.]

***cis*-2-Hexene** [7688-21-3] C_6H_{12} , M 84.2, b 68-70°/atm, d_4^{20} 0.699, n_D^{20} 1.399. Purify it as for 1-hexene above. [*Beilstein* 1 IV 833.]

***trans*-2-Hexene** [4050-45-7] M 84.2, m -98.5 to -98°, b 65-67°/atm, n_D^{20} 1.390. Purify it as for 1-hexene above. [*Beilstein* 1 IV 834.]

***cis*-3-Hexene** [7642-09-3] C_6H_{12} , M 84.2, b 66-68°/atm, d_4^{20} 0.681, n_D^{20} 1.395. Purify it as for 1-hexene above. [*Beilstein* 1 IV 830.]

***trans*-3-Hexene** [13269-52-8] C_6H_{12} , M 84.2, b 67-69°/atm, d_4^{20} 0.678, n_D^{20} 1.393. Purify it as for 1-hexene above. [*Beilstein* 1 IV 837.]

***n*-Hexyl alcohol (1-hexanol)** [111-27-3] $C_6H_{14}O$, M 102.2, m -52°, b 157.5°/atm, d_4^{20} 0.818, n_D^{15} 1.4198, n_D^{25} 1.4158. The commercial material usually contains other alcohols which are difficult to remove. A suitable method is to esterify with hydroxybenzoic acid, recrystallise the ester and saponify. [Olivier *Recl Trav Chim Pays-Bas* 55 1027 1936, DOI:10.1002/recl.19360551207.] Drying agents include K_2CO_3 and $CaSO_4$, followed by filtration and distillation. (Some decomposition to the olefin occurs when Al amalgam is used as drying agent at room temperature, even if the amalgam is removed prior to distillation.) If the alcohol is required anhydrous, the redistilled material can be refluxed with the appropriate alkyl phthalate or succinate, as described under *Ethanol*. [*Beilstein* 1 IV 1694.]

***n*-Hexylamine** [111-26-2] $C_6H_{15}N$, M 101.2, m -23°, b 131°/atm, d_4^{20} 0.765, n_D^{20} 1.419, pK^{25} 10.64. Dry with, and fractionally distil the hexylamine from, KOH or CaH_2 . Store away from CO_2 . [*Beilstein* 4 IV 709.]

***n*-Hexyl bromide** [111-25-1] $C_6H_{13}Br$, M 165.1, m -85°, b 87-88°/90mm, 155°/743mm, d_4^{20} 1.176, n_D^{20} 1.448. Shake the bromide with H_2SO_4 , wash with water, dry (K_2CO_3) and fractionally distil. Store in the dark.

[*Beilstein* 1 IV 352.]

***n*-Hexyl methacrylate** [142-09-6] $C_{10}H_{18}O_2$, **M 170.3**, **b 65-66°/4mm**, **88-88.5°/14mm**, **203°/atm**, **d₄²⁰ 0.8849**, **n_D²⁰ 1.4320**. Purify it as for *methyl methacrylate*. [IR Walton & Hughes *J Am Chem Soc* **79** 3985 1957, DOI: 10.1021/ja01572a008; *Beilstein* 2 III 1288, 2 IV 1527.]

Hexyltrimethylammonium bromide [2650-53-5] $C_9H_{22}BrN$, **M 224.3**, **m 186°**. Recrystallise it from acetone. It is extremely *hygroscopic*, store accordingly. [McDowell and Kraus *J Am Chem Soc* **73** 2170 1951, DOI: 10.1021/ja01149a074; *Beilstein* 4 IV 710.]

1-Hexyne [693-02-7] C_6H_{10} , **M 82.2**, **m -132°**, **b 12.5°/75mm**, **71°/760mm**, **d₄²⁰ 0.7156**, **n_D²⁰ 1.3989**. Distil it from NaBH₄ to remove peroxides. Stand over sodium for 24 hours, then fractionally distil it under reduced pressure. Also dry it by repeated vacuum transfer into freshly activated 4A molecular sieves, followed by vacuum transfer onto Na/K alloy, and stirring for 1 hour before fractionally distilling. [*Beilstein* 1 IV 1006.]

2-Hexyne [764-35-2] C_6H_{10} , **M 82.2**, **b 83.8°/760mm**, **d₄²⁰ 0.73146**, **n_D²⁰ 1.41382**. Purify as for 1-hexyne above. [*Beilstein* 1 IV 1009.]

3-Hexyne [928-49-4] C_6H_{10} , **M 82.2**, **b 81°/760mm**, **d₄²⁰ 0.7231**, **n_D²⁰ 1.4115**. Purify as for 1-hexyne above. [*Beilstein* 1 IV 1009.]

(±)-5-Hexyn-3-ol (4-hydroxy-1-hexyne) [19780-84-8] $C_6H_{10}O$, **M 98.1**, **b 58-59°/25mm**, **73-76°/60mm**, **d₄²⁰ 0.8918**, **n_D²⁰ 1.4437**. Purify the hexynol by fractionation in a vacuum. The *carbamoyl* derivative (prepared by reaction with COCl₂/toluene followed by NH₃) is crystallised by dissolving in the minimum volume of toluene and adding excess of petroleum ether (b 40-60°), and has **m 70-71°**. [Läuger et al. *Helv Chim Acta* **42** 2379 1959, DOI: 10.1002/hlca.19590420710; *Beilstein* 1 IV 2235.]

Hydrazine *N,N'*-dicarboxylic acid diamide [110-21-4] $C_2H_6NO_2$, **M 116.1**, **m 245-246°(dec)**, **248°**. Crystallise the diamide from water, wash the crystals with EtOH then Et₂O and dry in vacuum over P₂O₅. It does **not** decompose on drying at 110°/48 hours. Its solubility in H₂O is 1% at 0°. [Audrieth & Mohr *Inorg Synth* **4** 26 1953, *Beilstein* 3 I 116, 3 I 56, 3 II 95, 3 III 229.]

3-Hydroxy-2-butanone (acetoin) [513-86-0] $C_4H_8O_2$, **M 88.1**, **m 15°**, **b 144-145°**, [**m 100-105° dimer**]. Wash acetoin with EtOH until colourless, then with diethyl ether or acetone to remove biacetyl. Dry it in air by suction and dry further in a vacuum desiccator. [*Beilstein* 1 IV 3991.]

(±)-α-Hydroxy-γ-butyrolactone [19444-84-9, *S*(-)- 733-52-4, 52079-23-9; *R*(±) 56881-90-4] $C_4H_6O_3$, **M 102.1**, **b 84°/0.2mm**, **133°/10mm**, **d₄²⁰ 1.310**, **n_D²⁰ 1.4656**. It has been purified by repeated fractionation and forms a colourless liquid. It has to be distilled at high vacuum; otherwise it will dehydrate. The *acetoxo* derivative has **b 94°/0.2mm**. The *S*-enantiomer [52079-23-9] has **d₄²⁰ 1.309**, **n_D²⁰ 1.467**, **[α]_D²⁵ -82** (c 2, MeOH), **[α]_D²⁵ -68** (c 1.15, CHCl₃). [NMR: Daremon & Rambaud *Bull Soc Chim Fr* 294 1971, Schmitz et al. *Chem Ber* **108** 1010 1975, DOI: 10.1002/cber.19751080405; *Beilstein* 18/I V 3.]

12-Hydroxydodecanoic acid (12-hydroxylauric acid) [505-95-3] $C_{12}H_{24}O_3$, **M 216.3**, **m 86-88°**, **pK_{Est} ~4.8**. Recrystallise the acid from toluene. [Sadownik et al. *J Am Chem Soc* **108** 7789 1986, DOI: 10.1021/ja00284a050]. [*Beilstein* 3 III 658.]

***N*-[2-Hydroxyethyl]ethylenediamine [2-(2-aminoethylamino)ethanol]** [111-41-1] $C_4H_{12}N_2O$, **M 104.1**, **b 91.2°/5mm**, **238-240°/752mm**, **d₄²⁰ 1.030**, **n_D²⁰ 1.485**, **pK₁²⁰ 3.75**, **pK₂²⁰ 9.15**. Distil the amine twice through a Vigreux column. Redistil it from solid NaOH, then from CaH₂. *Alternatively*, it can be converted to the dihydrochloride and recrystallised from water. It is then dried, mixed with excess of solid NaOH and the free base is distilled from the mixture. It is finally redistilled from CaH₂. [Drinkard et al. *J Am Chem Soc* **82** 2992 1960, DOI: 10.1021/ja01497a004; *Beilstein* 4 IV 1558.]

***N*-[2-Hydroxyethyl]ethylenediaminetriacetic acid (HEDTA)** [150-39-0] $C_{10}H_{18}N_2O_7$, **M 278.3, m 212-214°(dec), pK_1^{20} 2.51, pK_2^{20} 5.31, pK_3^{20} 9.86.** Crystallise HEDTA from warm H_2O , after filtering, by addition of 95% EtOH and allowing to cool. The crystals, collected on a sintered-glass funnel, are washed three times with cold absolute EtOH, then again crystallised from H_2O . After leaching with cold H_2O , the crystals are dried at 100° under vacuum. [Spedding et al. *J Am Chem Soc* **78** 34 1956, DOI: 10.1021/ja01582a007; *Beilstein* **4** IV 2449.] It is useful buffer.

***N*-Hydroxyethyliminodiacetic acid (HIMDA, ethanol diglycine)** [93-62-9] $C_6H_{11}NO_5$, **M 177.2, m 181°(dec), pK_1^{25} 2.16, pK_2^{25} 8.72, pK_3^{25} 13.7 (OH).** Crystallise HIMDA from water. [*Beilstein* **4** IV 2432.]

2-Hydroxyethylimino-tris(hydroxymethyl)methane (MONO-TRIS) [7343-51-3] $C_6H_{15}NO_4$, **M 165.2, m 91°, pK_{Est} ~9.8.** Crystallise it twice from EtOH. Dry it under vacuum at 25°. The pure **hydrochloride** was isolated as a syrup [Pierce & Wotiz *J Am Chem Soc* **66** 879 1944, DOI: 10.1021/ja01234a010.] [*Beilstein* **4** III 858.] It is useful buffer.

2-Hydroxyethyl methacrylate [868-77-9] $C_6H_{10}O_3$, **M 130.1, b 67°/3.5mm, d_4^{20} 1.071, n_D^{20} 1.452.** Dissolve the ester in water and extract with *n*-heptane to remove ethylene glycol dimethacrylate (checked by gas-liquid chromatography and by NMR) and distil it twice under reduced pressure [Strop et al. *J Phys Chem* **80** 694 1976, DOI: 10.1021/j100548a008]. [*Beilstein* **2** IV 1530.]

***dl*-2-Hydroxy-2-methylbutyric acid** [3739-30-8] $C_5H_{10}O_3$, **M 118.1, m 72-73°, pK^{25} 3.73.** Crystallise the acid from *benzene, and sublime it at 90°. [*Beilstein* **3** H 324.] **IRRITANT.**

***dl*-2-Hydroxy-3-methylbutyric (α -hydroxyisovaleric) acid** [4026-18-0] $C_5H_{10}O_3$, **M 118.1, m 86°, pK_{Est} ~3.9.** Crystallise the acid from ether/pentane. [*Beilstein* **3** IV 618.] **IRRITANT.**

***R*(-)- γ -Hydroxymethyl- γ -butyrolactone** [52813-63-5] $C_5H_8O_3$, **M 116.1, b 101-102°/0.048mm, d_4^{20} 1.2238, n_D^{20} 1.471, $[\alpha]_{546}^{20}$ -38, $[\alpha]_D^{20}$ -33 (c 3, EtOH), $[\alpha]_D^{30}$ -53.5° (c 3, EtOH).** Purify it by column chromatography on Silica gel 60 (Merck 70-230 mesh) and elute with 7% EtOH/73% $CHCl_3$. Its IR (film) has ν_{max} at 3400 (OH), 1765 (C=O) and 1180 (COC) cm^{-1} . [Eguchi & Kakuta *Bull Chem Soc Jpn* **47** 1704 1974, DOI: org/10.1246/bcsj.47.1704; IR and NMR: Ravid et al. *Tetrahedron* **34** 1449 1978, DOI: 10.1016/0040-4020(78)80164-1; *Beilstein* **3** III 620, **18** V/1 13]

3-Hydroxy-3-methylglutaric acid (Meglutol) [503-49-1] $C_6H_{10}O_5$, **M 162.1, m 99-102°, 108-109°, 100°, $pK_{Est(1)}$ ~4.0, $pK_{Est(2)}$ ~5.0.** Recrystallise the acid from diethyl ether/hexane and dry it under a vacuum at 60° for 1 hour. [*Beilstein* **3** IV 1166.]

4-Hydroxy-4-methyl-2-pentanone (diacetone alcohol) [123-42-2] $C_6H_{12}O_2$, **M 116.2, b 166°, d_4^{20} 0.932, n_D^{20} 1.4235, n_D^{25} 1.4213.** The pentanone loses water when heated. It can be dried with $CaSO_4$, then fractionally distilled under reduced pressure. [*Beilstein* **1** IV 4023.]

2-Hydroxy-2-methylpropionic acid (α -hydroxyisobutyric acid, 2-methylactic acid) [594-61-6] $C_4H_8O_3$, **M 104.1, m 79°, b 114°/12mm, 212°/760mm, pK^{25} 3.78.** Distil the acid in steam, crystallise it from Et_2O or *benzene, dry it under vacuum, or sublime it at 50°. [*Beilstein* **3** IV 782.]

(\pm)-2-Hydroxyoctanoic acid (2-hydroxycaprylic acid) [617-73-2, 6482-96-2 *non-specific*] $C_8H_{16}O_3$, **M 160.2, m 69.5°, b 160-165°/10mm, pK_{Est} ~3.7.** Crystallise the (\pm)-acid from EtOH/petroleum ether or ether/ligroin. The (\pm)-racemic and *laevo* 2-hydroxy acids were prepared by a general procedure of anodic coupling in a cooled electrolytic cell with Platinum-foil electrodes in methanolic solution using a current of 1 amp and at 50 Volts, and run until the solution became neutral (~ 10–15 hrs). Thus a mixture of *ethyl O-acetylmalic acid mono- α -ester* (racemic or chiral, only the isomer with the free un-esterified γ -carboxy group reacts) and *n-hexanoic acid* decarboxylate and couple to give the respective (\pm)-**2-hydroxyoctanoic acid** or, when the mono ester derived from *S*(-)-*malic acid* is used, the chiral ***S*(+)-2-hydroxyoctanoic acid** [70267-27-5] which, after distillation followed by recrystallisation from isopropyl ether (fine needles) has **m 70.2-70.6°, $[\alpha]_D^{19}$**

+6.2 (*l* 0.5, *c* 9.2, CHCl₃), [α]_D¹⁹ +1.5 (*l* 0.5, *c* 6.8, EtOH), [α]_D¹⁹ +3.0 (*l* 0.5, *c* 4.27, H₂O containing 10–15% of MeOH). *S*(-)-2-hydroxyoctanoic acid sodium salt, obtained by titration with one mol of aqueous 0.185 N NaOH, has [α]_D¹⁹ -18 (*l* 0.5, *c* 5.05, 50% aqueous EtOH), and [α]_D¹⁹ -10.3 (*l* 0.5, *c* 10.1, H₂O). *Methyl S*(+)-2-hydroxyoctanoate [85549-54-8] C₉H₁₈O₃, M 174.2, has *b* 118°/22mm, *d*₄¹⁶ 0.9686, *n*_D¹⁸ 1.4342, [α]_D¹⁶ +2.80 (neat), [α]_D¹⁶ +11 (*l* 0.5, *c* 10.03, CHCl₃). [Horn & Pretorius *J Chem Soc* 1460 1954, DOI: 10.1039/JR9540001460.] *R*(-)-2-hydroxyoctanoic acid has [30117-44-3]. [*Beilstein* 3 IV 873.]

N-Hydroxysuccinimide [6066-82-6] C₄H₅NO₃, M 115.1, *m* 96-98°, *pK*_a²⁵ 6.0. Recrystallise from EtOH/ethyl acetate [Manesis & Goodman *J Org Chem* 52 5331 1987, DOI: 10.1021/jo00233a006]. [*Beilstein* 21/9 V 498.]

(±)-2-Hydroxy-*n*-tetradecanoic acid (α-hydroxymyristic acid) [2507-55-3] C₁₄H₂₈O₃, M 244.4, *m* 81-82°, *pK*_{Est} ~3.7. Crystallise the acid from chloroform or twice from MeOH (*m* 85.8-86.6°) [Horn & Pretorius *J Chem Soc* 1460 1954, DOI: 10.1039/JR9540001460; Chibnall et al. *Biochem J* 30 100 1936, DOI: 10.1042/bj0300100; *Beilstein* 3 H 361, 3 I 130, 3 II 246, 3 III 660, 3 IV 921].

R(-)-2-Hydroxy-*n*-tetradecanoic acid [26632-17-7] C₁₄H₂₈O₃, M 244.4, *m* 88-2-88.5°, [α]_D²⁰ -3.1 (CHCl₃). Crystallise the acid from CHCl₃, or first from Me₂CO, then hexane at -30°. Obtained by liquid-liquid extraction (counter-current distribution of liquids) of the *methyl ester* isolated from wool wax, hydrolysed and recrystallised. *Methyl R*(-)-2-hydroxy-*n*-tetradecanoate [149948-89-0] C₁₅H₃₀O₃, M 258.4, was purified by distillation, *b* 133.5-136.5°/1mm, followed by recrystallisation from Et₂O then Me₂CO at low temperature to provide a white powder with *m* 34.8-35.0°, [α]_D²⁰ -3.6 (*c* 9, CHCl₃). [Horn & Pretorius *J Chem Soc* 1460 1954, DOI: 10.1039/JR9540001460; Horn et al. *J Chem Soc* 177 1954, DOI: 10.1039/JR9540000177; *Beilstein* 3 III 660.] The *S*(+)-*enantiomer* [67253-09-2] was isolated from *Salmonella* polysaccharide. *Methyl S*(+)-2-hydroxy-*n*-tetradecanoate has [149948-88-9], and the *racemic methyl ester* has [130823-84-6].

Iminodiacetic acid [142-73-4] C₄H₇NO₄, M 133.1, *m* 225°(dec), *pK*₁²⁵ 2.50, *pK*₂²⁵ 9.40. Recrystallise iminodiacetic acid from water and dry it in a vacuum over P₂O₅. [*Beilstein* 4 IV 2428.]

Iodoacetamide [144-48-9] C₂H₄INO₂, M 185.0, *m* 93-94°, 92-95°. It has been prepared from chloroacetamide and NaI in Me₂CO [Hellstrom *Z physic Chem [A]* 157 246 1931]. Crystallise it from water CHCl₃, or CCl₄. It is used for tagging proteins. [Gurd *Methods Enzymol* 25 424 1972, DOI: 10.1016/S0076-6879(72)25038-8; *Beilstein* 2 H 223, 2 III 487, 2 IV 537.]

Iodoacetic acid [64-69-7] C₂H₃IO₂, M 186.0, *m* 78°, *pK*_a²⁵ 3.19. Crystallise it from petroleum ether (*b* 60-80°) or CHCl₃/CCl₄. Soluble in H₂O. [*Beilstein* 2 IV 534.]

2-Iodobutane (sec-butyl iodide) [513-48-4] C₄H₉I, M 184.0, *m* -104°, *b* 120°/atm, *d*₄²⁰ 1.50, *n*_D²⁵ 1.4973. Purify the iodide by shaking with concentrated H₂SO₄, then washing it with water, aqueous Na₂SO₃ and again with water. Dry (MgSO₄) and distil. Alternatively, pass it through a column of activated alumina before distillation, or treat with bromine, followed by extraction of the free halogen with aqueous Na₂S₂O₃, thoroughly washing with H₂O, drying and distilling. Store over Ag powder and distilled before use. [*Beilstein* 1 IV 272.]

Iodoform [75-47-8] CHI₃, M 393.7, *m* 119°, *d*₄²⁵ 4g/ml. Crystallise it from MeOH, EtOH or EtOH/EtOAc. It is steam volatile. It is soluble (in g/100ml) in cold EtOH (1.7), boiling EtOH (6.3), cold CHCl₃ (10.0), Et₂O (13.3), glycerol (1.2), CS₂ (33.3), olive oil (2.9), freely soluble in *C₆H₆ and Me₂CO slightly soluble in petroleum ether and very slightly soluble in H₂O. It is a disinfectant. [*Beilstein* 1 IV 97.]

***N*-Iodosuccinimide** [516-12-1] C₄H₄INO₂, M 225.0, *m* 200-201°. Crystallise it from dioxane/CCl₄. It is soluble in Me₂CO, MeOH, and dioxane but almost insoluble in CCl₄ or Et₂O and is decomposed by H₂O. It is decomposed by light. It is an iodinating agent and iodates arenes in triflic acid. [Olah et al *J Org Chem* 58 3194 1993, DOI: 10.1021/jo00063a052; *Beilstein* 21/9 V 544.]

Isoamyl acetate (1-butyl-3-methyl acetate, isopentyl acetate) [123-92-2] C₇H₁₄O₂, M 130.2, *m* -78°, *b*

142°/atm, d_4^{20} 0.871, n_D^{20} 1.40535. Dry the acetate with finely divided K_2CO_3 and fractionally distil it. [Beilstein 2 IV 157.]

Isoamyl alcohol (3-methyl-1-butanol, 1-butyl-3-methyl alcohol) [123-51-3] $C_5H_{12}O$, M 88.2, m -117°, b 128°/750mm, 132°/760mm, d^{15} 0.8129, n^{15} 1.4085, n_D^{20} 1.4075. Dry the alcohol by heating with CaO and fractionally distilling, then heating with BaO and redistilling. *Alternatively*, boil it with concentrated KOH solution, wash it with dilute H_3PO_4 , and dry it with K_2CO_3 , then anhydrous $CuSO_4$, before fractionally distilling it. If very dry alcohol is required, the distillate is refluxed with the appropriate alkyl phthalate or succinate as described for *ethanol*. It is separated from 2-methyl-1-butanol by fractional distillation, fractional crystallisation and preparative gas chromatography. [Beilstein 1 IV 1677.]

Isoamyl bromide (1-butyl-3-methyl bromide) [107-82-4] $C_5H_{11}Br$, M 151.1, f -112°, b 119.2°/ 737mm, d_4^{20} 1.208, n 1.444. Shake the bromide with concentrated H_2SO_4 , wash with water, dry with K_2CO_3 and fractionally distil it. [Beilstein 1 IV 328.]

Isoamyl chloride (1-butyl-3-methyl chloride) [107-84-6] $C_5H_{11}Cl$, M 106.6, b 99°/734mm, d_4^{20} 0.8704, n_D^{20} 1.4084. Shake the chloride vigorously with 95% H_2SO_4 until the acid layer no longer becomes coloured during 12 hours, then wash it with water, saturated aqueous Na_2CO_3 , and more water. Dry it with $MgSO_4$, filter and fractionally distil it. *Alternatively*, a stream of oxygen containing 5% of ozone is passed through the chloride for a time, three times longer than is necessary to cause the first coloration of starch iodide paper by the exit gas. Subsequent washing of the liquid with aqueous $NaHCO_3$ hydrolyses the ozonides and removes organic acids. After drying and filtering, the isoamyl chloride is distilled. [Chien & Willard *J Am Chem Soc* 75 6160 1953, DOI: 10.1021/ja01120a016; Beilstein 1 IV 287.]

Isoamyl ether [diisopentyl ether, di-(1-butyl-3-methyl) ether] [544-01-4] $C_{10}H_{22}O$, M 158.3, b 173.4°/atm, d_4^{20} 0.778, n_D^{20} 1.40850. This is a mixture of 2- and 3-methylbutyl ether. It is purified by refluxing with sodium for 5 hours, then it is distilled under reduced pressure, to remove alcohols. Isoamyl ether can also be dried with $CaCl_2$ and fractionally distilled from P_2O_5 . [Beilstein 1 IV 1682.]

Isobutane (2-methylpropane) [75-28-5] C_4H_{10} , M 58.1, m -160°, b -10.2°/atm, d_4^{20} 0.557. Olefins and moisture can be removed by passage at 65° through a bed of silica-alumina catalyst which has previously been evacuated at about 400°. *Alternatively*, water and CO_2 can be removed by passage through P_2O_5 , then asbestos impregnated with NaOH. Treatment with anhydrous $AlBr_3$ at 0° then removes traces of olefins. Inert gases can be separated by freezing the isobutane at -195° and evacuating out the system. [Beilstein 1 IV 282.]

Isobutene (2-methylpropene, isobutylene) [115-11-7] C_4H_8 , M 56.1, b -6.6°/760mm. Dry isobutene by passage through anhydrous $CaSO_4$ at 0°. Purify it further by freeze-pump-thaw cycles and trap-to-trap distillation. [Beilstein 1 IV 796.]

Isobutyl bromide (1-bromo-2-methylpropane) [78-77-3] C_4H_9Br , M 137.0, b 91.2°/atm, d_4^{20} 1.260, n_D^{20} 1.437. Partially hydrolyse it to remove any tertiary alkyl halide, fractionally distil it, then wash it with concentrated H_2SO_4 , water and aqueous K_2CO_3 , then redistil it from dry K_2CO_3 . [Dunbar & Hammett *J Am Chem Soc* 72 109 1950, DOI: 10.1021/ja01157a030; Beilstein 1 IV 294.]

Isobutyl chloride (1-chloro-2-methylpropane) [513-36-0] C_4H_9Cl , M 92.6, m -131°, 68.8°/760mm, d_4^{20} 0.877, n_D^{20} 1.398. Use the same methods as described under *isoamyl chloride* above. [Beilstein 1 IV 287.]

Isobutyl chloroformate [543-27-1] $C_5H_9ClO_2$, M 136.6, b 123-127°/atm, 128.8°/atm, d_4^{20} 1.053, n_D^{20} 1.4070. It can be dried over $CaCl_2$ and fractionated at atmospheric pressure while keeping moisture out. Its purity can be checked by conversion to the *phenyl urethane* derivative with $PhNCO$ [Saunders et al. *J Am Chem Soc* 73 3796 1951, DOI: 10.1021/ja01152a069.] Its IR (film) has ν_{max} at 1780cm^{-1} . [Thompson & Jameson *Spectrochim Acta* 13 236 1959, Röse *Justus Liebigs Ann Chem* 205 227 1880, DOI: 10.1002/jlac.18802050208]. [Beilstein 3 IV 26.]

Isobutyl formate [542-55-2] $C_5H_{10}O_2$, M 102.1, b 98.4°/atm, d_4^{20} 0.885, n_D^{20} 1.38546. Wash the formate with saturated aqueous $NaHCO_3$, in the presence of saturated aqueous $NaCl$ solution until no further reaction occurs, then with saturated aqueous $NaCl$, dry ($MgSO_4$) and fractionally distil it. [Beilstein 2 H 21, 2 I 18, 2 II 30, 2 III 41, 2 IV 29.]

Isobutyl iodide (1-iodo-2-methylpropane) [513-38-2] C_4H_9I M 184.0, b 83°/250mm, 120°/760mm, d_4^{20} 1.60, n_D^{20} 1.495. Shake the iodide with concentrated H_2SO_4 , and wash it with water, aqueous Na_2SO_3 , and water, dry with $MgSO_4$ and distil it. Alternatively, pass it through a column of activated alumina before distillation. Store it under nitrogen with some mercury in a brown bottle or in the dark. [Beilstein 1 IV 299.]

Isobutyl vinyl ether [109-53-5] $C_6H_{12}O$, M 100.2, m -112°, b 108-110°/atm, d_4^{20} 0.768, n_D^{20} 1.398. Wash the ether three times with equal volumes of aqueous 1% $NaOH$, dry it with CaH_2 , reflux it with sodium for several hours, then fractionally distil it from sodium. [Beilstein 1 IV 2054.]

Isobutyraldehyde (2-methylpropionaldehyde) [78-84-2] C_4H_8O , M 72.1, m -65°, b 62.0°/atm, d_4^{20} 0.789, n_D^{20} 1.377. Dry isobutyraldehyde with $CaSO_4$ and use it immediately after distillation under nitrogen because of the great difficulty in preventing oxidation. It can be purified through its acid bisulfite derivative. [Beilstein 1 IV 3262.]

Isobutyramide (2-methylpropionamide) [563-83-7] C_4H_9NO , M 87.1, m 128-129°, b 217-221°/760mm. Crystallise the amide from acetone, $*C_6H_6$, $CHCl_3$, $EtOAc$ or water, then dry it under vacuum over P_2O_5 or 99% at H_2SO_4 , or at 70°/3 hours in a desiccator. Sublime it under vacuum. [Kent & McAlvain *Org Synth Coll Vol* 3 490 1955, DOI: 10.15227/orgsyn.025.0058; Beilstein 2 H 293, 2 I 129, 2 II 262, 2 III 654, 2 IV 852.]

Isobutyric acid (2-methylpropionic acid) [79-31-2] $C_4H_8O_2$, M 88.1, b 78°/34mm, 154-154.5°/760mm, d_4^{20} 0.949, n_D^{20} 1.393, pK^{25} 4.60. Distil the acid from $KMnO_4$, then redistil it from P_2O_5 . [Beilstein 2 H 288, 2 I 126, 2 II 257, 2 III 637, 2 IV 843.]

Isobutyronitrile (2-methylpropionitrile, isopropyl cyanide) [78-82-0] C_4H_7N , M 69.1, m -72°, b 103.6°/760mm, d_4^{25} 0.7650, n_D^{20} 1.378. Shake the nitrile with concentrated HCl (to remove isonitriles), then with water and aqueous $NaHCO_3$. After a preliminary drying with silica gel or Linde type 4A molecular sieves, it is shaken or stirred with CaH_2 until hydrogen evolution ceases, then decanted and distilled from P_2O_5 (not more than 5g/L, to minimise gel formation) or Drierite (b 101-103°/760mm). Finally it is refluxed with, and slowly distilled from CaH_2 (5g/L), taking precautions to exclude moisture. [Beilstein 2 H 294, 2 I 129, 2 II 263, 2 III 655, 2 IV 853.]

Isonitrosoacetone (anti-pyruvic aldehyde-1-oxime) [31915-82-9] $C_3H_5NO_2$, M 87.1, m 65-67°, 69°, pK^{25} 8.3. Crystallise isonitrosoacetone from $*C_6H_6$, ether/petroleum ether or CCl_4 . It sublimes at 90-100° (water bath temperature)/0.05mm [Kahovec & Kohlrausch *Chem Ber* 75 1541 1942, DOI: 10.1002/cber.19420751219]. It forms an iron and a Cu^{2+} salt (m 170° dec). [Beilstein 1 H 763, 1 I 396, 1 II 822, 1 III 3092, 1 IV 3632.]

(±)-Isononane (3,3,4-trimethylhexane) [34464-40-9] C_9H_{20} , M 128.3, b 142°/760mm, 142°/760mm, d_4^{20} 0.7454, n_D^{20} 1.4178. Isononane is passed through columns of activated silica gel and basic alumina (activity 1) and distilled under high vacuum from Na/K alloy. [Beilstein 1 III 517, 1 IV 462.]

Isopentyl formate [110-45-2] $C_6H_{12}O_2$, M 116.2, b 27°/10mm, 123-123.6°/760mm, 123-124°/atm, d_4^{20} 0.8713, n_D^{20} 1.391. The colourless liquid ester is soluble in 300 volumes of H_2O and is soluble in common organic solvents. It is purified by repeated distillation using an efficient column at atmospheric pressure. [Beilstein 2 H 22, 2 I 18, 2 II 31, 2 III 43, 2 IV 30.]

Isoprene (2-methyl-1,3-butadiene) [78-79-5] C_5H_8 , M 68.1, b 34.5-35°/762mm, d_4^{20} 0.681, n_D^{25} 1.4225. Reflux it with sodium then distil it from sodium or $NaBH_4$ under nitrogen, and pass it through a column containing KOH , $CaSO_4$ and silica gel. *tert*-Butylcatechol (0.02% w/w) is added, and the isoprene is stored in this way until redistilled before use. The inhibitor (*tert*-butylcatechol) in isoprene can be removed by several

washings with dilute NaOH and water. The isoprene is then dried over CaH_2 , distilled under nitrogen at atmospheric pressure, and the fraction distilling at 32° is collected. Store it under nitrogen at -15° . [*Beilstein* 1 H 252, 1 IV 1001.]

Isopropenyl acetate (1-methylvinyl acetate, 2-acetoxyprene) [108-22-5] $\text{C}_5\text{H}_8\text{O}_2$, M 100.1, b $96.7^\circ/749\text{mm}$, $97.2\text{--}97.4^\circ/757\text{mm}$, d_4^{25} 0.909, n_D^{20} 1.4010. Prepared by reacting ketene with a mixture of Me_2CO and polyphosphoric acid [Wacher US Patent 2 867 653 1957], fractionally distilling through a 75cm packed column, and collecting the fraction with b $96.7^\circ/749\text{mm}$. It forms an azeotrope with H_2O . Its equivalent weight is determined by hydrolysis in 90% EtOH containing twice the required amount of NaOH and titrating the excess of alkali; which should give an equivalent weight of ~ 100.1 . Its heat of hydrolysis has been determined [Sunner *Acta Chem Scand* 11 1757 1957, DOI: 10.3891/acta.chem.scand.11-1757]. It is a good source the $=\text{CMe}_2$ group, as in the preparation of Meldrum's acid (see [2033-24-1]), because it is the activated enol ester of acetone. Its FT-IR (neat) has ν_{max} at 1757.3, 1673.0, 1431.4, 1372.5, 1253.1, 1198.2, 1027.9, 900.0 and 871.4 cm^{-1} ; the ^1H NMR (60MHz, CDCl_3 , TMS) has δ at 1.92 (s, 3H, ester Me), 2.13 (s, 3H, isoPr-Me) and 7.00 (d, $J = \sim 6\text{Hz}$, 2H, $=\text{CH}_2$); and the ^{13}C NMR (15MHz, CDCl_3 , TMS) has δ at 19.54, 21.05, 102.05, 152.94 and 169.04. [*Beilstein* 2 III 278, 2 IV 179.]

Isopropanol (propan-2-ol) [67-63-0] $\text{C}_3\text{H}_8\text{O}$, M 60.1, m -89.5° , b $82.5^\circ/\text{atm}$, d_4^{20} 0.783, $n_D^{25.8}$ 1.3739, pK^{25} 17.1. Isopropyl alcohol is prepared commercially by dissolution of propene in H_2SO_4 , followed by hydrolysis of the sulfate ester. Major impurities are water, lower alcohols and oxidation products such as aldehydes and ketones. Purification of isopropanol follows substantially the same procedure as for *n*-propyl alcohol. Isopropanol forms a constant-boiling mixture, b 80.3° , with water. Most of the water can be removed from this 91% isopropanol by refluxing with CaO (200g/L) for several hours, then distilling. The distillate can be dried further with CaH_2 , magnesium ribbon, BaO, CaSO_4 , calcium, anhydrous CuSO_4 or Linde type 5A molecular sieves. Distillation from sulfanilic acid removes ammonia and other basic impurities. Peroxides [indicated by liberation of iodine from weakly acid (HCl) solutions of 2% KI] can be removed by refluxing with solid stannous chloride or with NaBH_4 then the alcohol is fractionally distilled. To obtain isopropanol containing only 0.002M of water, sodium (8g/L) is dissolved in material dried by distillation from CaSO_4 . Isopropyl benzoate (35ml) is then added and, after refluxing for 3 hours, the alcohol is distilled through a 50-cm Vigreux column [Hine & Tanabe *J Am Chem Soc* 80 3002 1958, DOI: 10.1021/ja01545a026]. Other purification steps for isopropanol include refluxing with solid aluminium isopropoxide, refluxing with NaBH_4 for 24 hours, and removing acetone by treatment with, and distillation from 2,4-dinitrophenylhydrazine. Peroxides re-form in isopropanol if it is kept for several days in contact with air. [*Beilstein* 1 IV 1461.]

Isopropyl acetate [108-21-4] $\text{C}_5\text{H}_{10}\text{O}_2$, M 102.1, m -73° , b $88.4^\circ/\text{atm}$, d_4^{20} 0.873, n_D^{20} 1.3773. Wash the acetate with 50% aqueous K_2CO_3 (to remove acid), then with saturated aqueous CaCl_2 (to remove any alcohol). Dry it with CaCl_2 and fractionally distil it. [*Beilstein* 2 IV 141.]

Isopropyl acrylamide [2210-25-5] $\text{C}_6\text{H}_{11}\text{NO}$, M 113.2, m $60\text{--}63^\circ$, b $89\text{--}92^\circ/2\text{mm}$, $110\text{--}115^\circ/15\text{mm}$. Fractionate the amide under reduced pressure, and recrystallise the solid distillate from hexane (m 59°), $^*\text{C}_6\text{H}_6$ (m 62°) or $^*\text{C}_6\text{H}_6/\text{hexane}$ (m $62\text{--}63^\circ$). Store it with 0.05% of 4-*tert*-butylcatechol. It is used for making water soluble swellable hydrogels. [*Beilstein* 4 IV 517.]

Isopropyl bromide (2-bromopropane) [75-26-3] $\text{C}_3\text{H}_7\text{Br}$, M 123.0, m -89° , b $0^\circ/69.2\text{mm}$, $59.4^\circ/760\text{mm}$, d_4^{20} 1.31, n_D^{15} 1.42847, n_D^{20} 1.4251. Wash the bromide with 95% H_2SO_4 (concentrated acid partially oxidised it) until a fresh portion of acid did not become coloured after several hours, then with water, aqueous NaHSO_3 , aqueous 10% Na_2CO_3 and again with water. (The H_2SO_4 can be replaced by concentrated HCl.) Prior to this treatment, isopropyl bromide has been purified by bubbling a stream of oxygen containing 5% ozone through it for 1 hour, followed by shaking with 3% hydrogen peroxide solution, neutralising with aqueous Na_2CO_3 , washing with distilled water and drying. Alternatively, it has been treated with elemental bromine and stored for 4 weeks, then extracted with aqueous NaHSO_3 and dried with MgSO_4 . After the acid treatment, isopropyl bromide can be dried with Na_2SO_4 , MgSO_4 or CaH_2 , and fractionally distilled. [*Beilstein* 1 IV 208.]

Isopropyl chloride (2-chloropropane) [75-29-6] $\text{C}_3\text{H}_7\text{Cl}$, M 78.5, m -118° , b $34.8^\circ/\text{atm}$, d_4^{20} 0.864, n_D^{20}

1.3779, n²⁵ 1.3754. Purify the chloride with 95% H₂SO₄ as described for *isopropyl bromide*, then dry with MgSO₄, P₂O₅ or CaH₂, and fractionally distil it from Na₂CO₃ or CaH₂. *Alternatively*, a stream of oxygen containing *ca* 5% ozone is passed through the chloride for about three times as long as is necessary to obtain the first coloration of starch iodide paper by the exit gas, and the liquid is then washed with NaHCO₃ solution to hydrolyse ozonides and remove organic acids before drying and distilling. [*Beilstein* 1 IV 191.]

Isopropyl ether (diisopropyl ether) [108-20-3] C₆H₁₄O, M 102.2, m -85°, b 68.3°/atm, d₄²⁰ 0.719, n_D²⁰ 1.3688, n²⁵ 1.36618. Common impurities are water and peroxides [detected by the liberation of iodine from weakly acid (HCl) solutions of 2% KI]. Peroxides can be removed by shaking with aqueous Na₂SO₃ or with acidified ferrous sulfate (0.6g FeSO₄ and 6ml concentrated H₂SO₄ in 110ml of water, using 5-10g of solution per L of ether), or aqueous NaBH₄ solution. The ether is then washed with water, dried with CaCl₂ and distilled. *Alternatively*, refluxing with LiAlH₄ or CaH₂, or drying with CaSO₄, then passage through an activated alumina column, can be used to remove water and peroxides. Other dehydrating agents used with isopropyl ether include P₂O₅, sodium amalgam and sodium wire. (The ether is often stored in brown bottles, or in the dark, with sodium wire.) Bonner and Goishi (*J Am Chem Soc* 83 85 1961, DOI: 10.1021/ja01462a018) treated isopropyl ether with dilute sodium dichromate/sulfuric acid solution, followed by repeated shaking with a 1:1 mixture of 6M NaOH and saturated KMnO₄. The ether is washed several times with water, dilute aqueous HCl, and water, with a final washing with, and storage over, ferrous ammonium sulfate acidified with H₂SO₄. Blaustein and Gryder (*J Am Chem Soc* 79 540 1957, DOI: 10.1021/ja01560a012), after washing with alkaline KMnO₄, then water, treated the ether with ceric nitrate in nitric acid, and again washed it with water. Hydroquinone is added before drying with CaCl₂ and MgSO₄, and refluxing with sodium amalgam (108g Hg/100g Na) for 2 hours under nitrogen. The distillate (nitrogen atmosphere) is made 2 x 10⁻⁵M in hydroquinone to inhibit formation of peroxides (which is negligible if the ether is stored in the dark). Catechol (pyrocatechol) and resorcinol are alternative inhibitors. [*Beilstein* 1 IV 1471.]

Isopropyl iodide (2-iodopropane) [75-30-9] C₃H₇I, M 170.0, m -90°, b 88.9°/atm, d₄²⁰ 1.70, n_D²⁰ 1.4987. Treat the iodide with bromine, followed by extraction of free halogen with aqueous Na₂S₂O₃ or NaHSO₃, washing with water, drying (MgSO₄ or CaCl₂) and distilling. (The treatment with bromine is optional.) Other purification methods include passage through activated alumina, or shaking with copper powder or mercury to remove iodine, drying with P₂O₅ and distilling. Washing with concentrated H₂SO₄ or concentrated HCl (to remove any alcohol), water, aqueous Na₂SO₃, water and aqueous Na₂CO₃ has also been used. Treatment with silica gel causes some liberation of iodine. Distillations should be carried out at slightly reduced pressure. Purified isopropyl iodide is stored in the dark in the presence of a little mercury. [*Beilstein* 1 IV 223.]

Isopropyl methyl ether [598-53-8] C₄H₁₀O, M 74.1, b 32.5°/777mm, d₁₅¹⁵ 0.724, n_D²⁰ 1.3576. Purify the ether by drying with CaSO₄, passing through a column of alumina (to remove peroxides) and fractional distillation. [*Beilstein* 1 H 362, 1 II 381, 1 III 1458, 1 IV 1471.]

Isovaleric acid (3-methylbutyric acid) [502-74-2] C₅H₁₀O₂, M 102.1, m -29°, b 176.5°/762mm, d₄²⁰ 0.927, n_D¹⁵ 1.4064, n_D²⁰ 1.40331, pK²⁵ 4.77. Dry the acid (Na₂SO₄), then fractionally distil. [*Beilstein* 2 IV 895.]

Itaconic acid (2-propen-1,2-dicarboxylic acid) [97-65-4] C₅H₆O₄, M 130.1, m 165-166°, pK₁²⁵ 3.63, pK₂²⁵ 5.00. Crystallise itaconic acid from EtOH, EtOH/water or EtOH/*benzene. [*Beilstein* 2 IV 2228.]

Itaconic anhydride (2-propen-1,2-dicarboxylic anhydride) [2170-03-8] C₅H₄O₃, M 112.1, m 66-68°, 67-68°, 68°, b 114-115°/12mm, 139-140°/30mm. Crystallise the anhydride from CHCl₃/petroleum ether. It can be distilled under reduced pressure. Distillation at atmospheric pressure, or prolonged distillation causes rearrangement to citraconic anhydride (2-methylmaleic anhydride). If the material (as seen in the IR spectrum) contains much free acid, then heat with acetyl chloride or SOCl₂, evaporate and distil at as high a vacuum as possible. The crude anhydride deposits crystals of itaconic acid on standing probably due to hydrolysis by H₂O — store it in sealed ampoules under dry N₂. [Shriner et al. *Org Synth Coll Vol* 2 368 1943, DOI: 10.15227/orgsyn.011.0070; IR: Nagai *Bull Chem Soc Jpn* 37 369 1964, DOI:org/10.1246/bcsj.37.369; Kelly & Segura *J Am Chem Soc* 56 2497 1934, DOI: 10.1021/ja01326a503; *Beilstein* 17/11 V 66.]

Kerosene [8008-20-6] (mixture of hydrocarbons) **b** ~175-225°/atm, ~190-250°/atm, **d**₄²⁰ 0.75-0.82, **n**_D²⁰ 1.443. Stir it with concentrated H₂SO₄ until a fresh portion of acid remains colourless, then wash with water, dry with solid KOH and distil it in a Claisen flask. For more complete drying, the kerosene can be refluxed with Na, and distilled from Na.

Ketene [463-51-4] C₂H₂O, **M** 42.0, **m** - 678°, -150°, **b** -56°/atm, -41°/atm, **d**₄²⁰ 1.093, **n**_D²⁰ 1.441. Ketene is prepared by pyrolysis of acetic anhydride. Purify it by passing through a trap at -75° and collecting in a liquid-nitrogen-cooled trap. Ethylene is removed by evacuating it in an isopentane-liquid-nitrogen slush pack at -160°. Store it at room temperature in a suitable container in the dark or better at -80°, but do not store it under pressure as it may **EXPLODE**. It is a strong **IRRITANT** when inhaled and is as **POISONOUS** as phosgene. See diketene in 'Heterocyclic Compounds', in this Chapter. [Hurd *Org Synth Coll Vol I* 330 1941, DOI: 10.15227/orgsyn.004.0039; Andreades & Carlson *Org Synth Coll Vol 5* 679 1973, DOI: 10.15227/orgsyn.045.0050.]

Ketene diethyl acetal (1,1-diethoxyethene) [2678-54-8] C₆H₁₂O₂, **M** 116.2, **b** 85-87°/200mm, 124-126°/atm, **d**₄²⁵ 0.87760, **n**_D²⁵ 1.4101. It can be prepared by the removal of HI from diethyl iodoacetal with *tert*-BuOK in ~52% yield. The ketene diethyl acetal is distilled off using a Widmer column preferably under vacuum. It is then fractionated again under a vacuum. Distilling is possible at atmospheric pressure but the clear water-like liquid distillate slowly becomes turbid on standing. When distilled under a vacuum the distillate takes a much longer time to become turbid. It should be stored in a dry atmosphere. It reacts rapidly with H₂O or EtOH to liberate heat and form EtOAc or ethyl orthoacetate. It polymerises on heating. [Beyerstedt & McElvain *J Am Chem Soc* **58** 529 1936, DOI: 10.1021/ja01294a039; also cf. McElvain & McShane Jr. *J Am Chem Soc* **74** 2662 1952, DOI: 10.1021/ja01130a057.] [*Beilstein* **1** III 2948, **1** IV 3420.]

Ketene dimethyl acetal (1,1-dimethoxyethane) [922-69-0] C₄H₈O₂, **M** 88.1, **b** 89°/atm, **d**₄²⁵ 0.9294, **n**_D²⁵ 1.4050, 1.3962. It can be prepared from anhydrous pinacol and sodium which was made to react with dimethyl chloroacetal and the dimethyl ketene acetal formed is distilled off. It is then refractionated to constant refractive index to give pure acetal with **b** 89-89.5°/740mm. [McElvain & McShane Jr. *J Am Chem Soc* **74** 2662 1952, DOI: 10.1021/ja01130a057; *Beilstein* **1** III 2947, **1** IV 3420.]

L(+)-Lactic acid (S(+)-2-hydroxypropionic acid) [79-33-4] C₃H₆O₂, **M** 90.1, **m** 52.8°, **b** 105°/0.1mm, **[α]_D²⁰** +3.82 (H₂O), -13.5 (c 2.5, 1.5M NaOH), **pK**³¹ 3.83. Purify lactic acid by fractional distillation at 0.1mm pressure, followed by fractional crystallisation from diethyl ether/isopropyl ether (1:1, dried with sodium). [Borsook et al. *J Biol Chem* **102** 449 1933, <http://www.jbc.org/content/102/2/449.citation>.] The solvent mixture, *benzene/diethyl ether (1:1) containing 5% petroleum ether (b 60-80°) has also been used. Most salts of (+)-lactic acid are *levorotatory* accounting for the negative rotation of alkaline solutions. It tends to form varying amounts of intermolecular esterification. [Brin *Biochemical Preparations* **3** 61 1953, *Beilstein* **3** IV 633.]

Lanthanide shift reagents A variety of these reagents are available commercially, and they are generally quite stable and should not deteriorate on long storage in a dry state and in the absence of light. [See J.R. Campbell *Aldrichimica Acta* **4** 55 1971, G.R. Sullivan in *Top Stereochem* (Eliel & Allinger Eds) J Wiley & Sons Vol 10 287 1978, T.C. Morrill Ed. *Lanthanide Shift Reagents* Deerfield Beach Florida 1986, ISBN 0895731193.]

Lauraldehyde (1-dodecanal, dodecyl aldehyde) [112-54-9] C₁₂H₂₄O, **M** 184.3, **b** 99.5-100°/3.5mm, 185°/100mm, **d**₄²⁰ 0.831, **n**_D²⁰ 1.435, **n**_D^{24.7} 1.4328. Convert lauraldehyde to the bisulfite addition compound by shaking with saturated aqueous NaHSO₃ for 1 hour. The precipitate is filtered off, washed with ice cold water, EtOH and ether, then decomposed with aqueous Na₂CO₃. The aldehyde is extracted into diethyl ether which, after drying and evaporating, gives an oil which is fractionally distilled under vacuum. [*Beilstein* **1** IV 3380.]

Lauric acid (1-dodecanoic acid) [143-07-7] C₁₂H₂₄O₂, **M** 200.3, **m** 44.1°, **b** 141-142°/0.6-0.7mm, 225°/100mm, **d**₄²⁵ 0.883, **pK**²⁰ 5.3. Distil the acid in a vacuum. Also crystallise it from absolute EtOH, or from acetone at -25°. Alternatively, purify it via its methyl ester (**b** 140.0°/15mm), as described for capric acid (see

[334-48-5]). It has also been purified by zone melting. [cf. *Beilstein* 1 III 2913, 3 IV 1082]

Lauroyl peroxide (di-[dodecanoyl] peroxide) [105-74-8] $C_{24}H_{46}O_4$, M 398.6, m 53-54°. Crystallise it from *n*-hexane or *benzene and store it below 0°. Potentially **EXPLOSIVE**. [*Beilstein* 2 IV 1102.]

Z-Maleamic acid (cis-maleic acid monoamide) [557-24-4] $C_4H_5NO_3$, M 115.1, m 172°, 172-173°(dec), 178-180°, $pK_{Est} \sim 2.65$. Crystallise it from EtOH. [*Beilstein* 2 H 752, 2 II 646, 2 III 1927, 2 IV 1927.] **IRRITANT**.

Maleic acid (toxic acid, cis-butenedioic acid) [110-16-7] $C_4H_4O_4$, M 116.1, m 143.5°, $pK_1^{25} 1.91$, $pK_2^{25} 6.33$. Crystallise the acid from acetone/petroleum ether (b 60-80°) or hot water. Dry it at 100°. [*Beilstein* 2 H 748, 2 I 303, 2 II 641, 2 III 1911, 2 IV 2199.]

Maleic anhydride see furan-2,5-dione; **Maleic hydrazide** see pyridazine-3,6-diol, and **Maleimide** See pyrrol-2,5-dione all in 'Heterocyclic Compounds', in this Chapter.

Maleuric acid (Z-N-carbamoylmaleamic acid) [105-61-3] $C_5H_6N_2O_4$, M 158.1, m 158.5°(dec), 167-168°(dec), $pK_{Est} \sim 2$. Crystallise the acid from hot water. Dry it at $\sim 100^\circ$ over H_2SO_4 . It is prepared from maleic anhydride and urea, and is not soluble in most solvents. In H_2O it has λ_{max} at 235nm with ϵ of 8720. The *methyl ester*, m 113-114°, is soluble in hot H_2O , MeOH, EtOH, Me_2CO and dioxane. The *butyl ester*, m 96-98°, is soluble in EtOH, Me_2CO , $CHCl_3$ and $*C_6H_6$. [Batt et al. *J Am Chem Soc* 76 3663 1954, DOI: 10.1021/ja01643a018.] *Beilstein* 3 H 68, 3 III 314, 3 IV 124.]

dl-Malic acid (2-hydroxysuccinic acid) [617-48-1 and 6915-15-7] $C_4H_6O_5$, M 134.1, m 128-129°, 131-132°, $pK_1^{25} 3.46$, $pK_2^{25} 5.10$. Crystallise the acid from acetone, then from acetone/ CCl_4 , or from ethyl acetate by adding petroleum ether (b 60-70°). Dry it at 35° under 1mm pressure to avoid formation of the anhydride. It is soluble (in g/100ml at 20°) in MeOH (82.7), H_2O (55.8), EtOH (45.3), dioxane (22.7), Me_2CO (17.8), Et_2O (0.8) and $*C_6H_6$ (~ 0) [*Beilstein* 3 IV 1124.]

L-Malic acid [S(-)-2-hydroxysuccinic acid] [97-67-6] $C_4H_6O_5$, M 134.1, m 104.5-106°, $[\alpha]_D^{20} -2.3$ (c 8.5, H_2O), $[\alpha]_D^{20} -30$ (c 5.5, pyridine), $pK_1^{25} 3.46$, $pK_2^{25} 5.10$. Crystallise *S*-malic acid (charcoal) from ethyl acetate/petroleum ether (b 55-56°), keeping the temperature below 65°. Or dissolve it by refluxing in fifteen parts of anhydrous diethyl ether, decant, concentrate to one-third volume and crystallise it at 0°, repeatedly to constant melting point. The **D-enantiomer (R(+)-2-hydroxysuccinic acid)** [636-61-3] has identical properties except that the optical rotations are positive. [*Beilstein* 3 IV 1123.]

Malonamide (malondiamide) [108-13-4] $C_3H_6N_2O_2$, M 102.1, m 170°, 172-175°. Crystallise the amide from water. [*Beilstein* 2 IV 1887.]

Malonic acid [141-82-2] $C_3H_4O_4$, M 104.1, m 136°(dec). $pK_1^{25} 2.58$, $pK_2^{25} 5.69$. Crystallise malonic acid from *benzene/diethyl ether (1:1) containing 5% of petroleum ether (b 60-80°), wash with diethyl ether, then recrystallise it from H_2O or acetone. Dry it under vacuum over concentrated H_2SO_4 . It sublimes in high vacuum. It is soluble (in g/100ml at room temperature) in H_2O (15.4), MeOH (10), EtOH (5), *iso*-Pr (3.3), pyridine (1.4) and Et_2O (0.8). [*Beilstein* 2 IV 1874.]

Malononitrile (dicyanomethane) [109-77-3] $C_3H_2N_2$, M 66.1, m 32-34°, b 109°/20mm, 113-118°/25mm, 220°/760mm. Crystallise the nitrile from water, EtOH, *benzene or chloroform. Distil it in a vacuum from, and store over, P_2O_5 . [Bernasconi et al. *J Am Chem Soc* 107 3612 1985, DOI: 10.1021/ja00298a035; Grootenhuis *J Am Chem Soc* 109 8044 1987, DOI: 10.1021/ja00260a016; *Beilstein* 2 IV 1892.]

Meproamate [2,2-di(carbamoyloxymethyl)pentane] [57-53-4] $C_9H_{18}N_2O_4$, M 218.3, m 104-106°. Crystallise it from hot water [solubility is 0.34% (w/w) at 20°, and 0.79% (w/w) at 37°], aqueous EtOH (m 104-105.5°) or xylene (m 104.1-105.3°). It is soluble in most organic solvents. Its aqueous solutions are neutral,

and it is stable in dilute acid and alkaline solutions. It can be an addictive depressant drug. [Ludwig & Piech *J Am Chem Soc* **73** 5779 1951, DOI: 10.1021/ja01156a086; *Beilstein* **3** IV 73.]

2-Mercaptoethanol [60-24-2] C_2H_6OS , **M 78.1**, **b** 44°/4mm, 53.5°/10mm, 58°/12mm, 68°/20mm, 78.5°/40mm, 96-97° (92°)/100mm, 157°/748mm, d_4^{20} 1.114, n_D^{20} 1.500, pK^{25} 9.72 (9.43). Purify it by distilling in a vacuum. Distilling at atmospheric pressure causes some oxidation and should be done in an inert atmosphere. [Woodward *J Chem Soc* 1892 1948, DOI: 10.1039/JR9480001892.] It has a foul odour, is irritating to the eyes, nose and skin — should be handled in an efficient **fume cupboard**. It is miscible with H_2O , EtOH, Et_2O and $*C_6H_6$ and the UV has λ_{max} at 235nm. The **2,4-dinitrophenyl thioether** has **m 101-102°** (from EtOH or aqueous MeOH) [Grogan et al. *J Org Chem* **20** 50 1955, DOI: 10.1021/jo01119a009]. [*Beilstein* **1** IV 2428.]

Mesaconic acid (methyfummaric acid) [498-24-8] $C_5H_6O_4$, **M 130.1**, **m** 204-205°, **b** 204-205°/atm (dec), pK_1^{25} 3.02, pK_2^{25} 4.82. Crystallise it from H_2O , EtOH (needles) or EtOAc (platelets) and sublime it *in vacuo*. Its solubility in H_2O , is 2.7% w/w at 18° and 118% w/w at the boiling point; in 90% EtOH its solubility is 31% w/w at 17° and 96% w/w at the boiling point; it is soluble in Et_2O but sparingly in $CHCl_3$, CS_2 and light petroleum. It has been prepared by reaction of citraconic anhydride with nitric acid. The **dimethyl ester** $C_7H_{10}O_4$ has **b** 204-206°/atm, d_4^{20} 1.1201, n_D^{20} 1.4557. The **diethyl ester** $C_9H_{14}O_4$ has **b** 229°/atm, d_4^{20} 1.0467, n_D^{20} 1.4494. [Shriner et al. *Org Synth* **11** 74 1931, DOI: 10.15227/orgsyn.011.0074; Katakis et al. *JCS Dalton Trans* 1491 1986, DOI: 10.1039/DT9860001491]. [*Beilstein* **2** H 763, **2** I 308, **2** II 650, **2** III 1934, **2** IV 2231.]

Mesityl oxide (4-methyl-3-penten-2-one) [141-79-7] $C_6H_{10}O$, **M 98.2**, **m** -53°, **b** 57°/55mm, 128-129°/745mm, 112°/760mm, n_D^{24} 1.4412, **d** 0.854, pK^{20} -5.36 (H_0 scale, aqueous H_2SO_4). Purify it by distillation, preferably in a vacuum or *via* the **semicarbazone** (**m** 165°) which is decomposed to pure ketone. The **2,4-dinitrophenylhydrazone** (**m** 205-206°) crystallises from EtOH. [Johnson *J Am Chem Soc* **73** 5888 1951, DOI: 10.1021/ja01156a530; Johnson *J Am Chem Soc* **75** 2720 1953, DOI: 10.1021/ja01107a051; Erskine & Waight *J Chem Soc* 3425 1960, DOI: 10.1039/JR9600003425; *Beilstein* **1** H 736, **1** I 382, **1** II 793, **1** III 2995, **1** IV 3471.]

α -Methacraldehyde (methacrolein) [78-85-3] C_4H_6O , **M 68.1**, **m** -81°, **b** 68.4°/atm, d_4^{20} 0.849, n_D^{20} 1.416. Fractionally distil it under nitrogen through a short Vigreux column. Store it in sealed ampoules. (Slight polymerisation may occur.) [*Beilstein* **1** IV 3455.]

Methacrylamide [79-39-0] C_3H_7NO , **M 85.1**, **m** 111-112°. Crystallise the amide from $*benzene$ or ethyl acetate and dry it under vacuum at room temperature. [*Beilstein* **2** IV 1538.]

Methacrylic acid (2-methylpropenoic acid) [79-41-4] $C_4H_6O_2$, **M 86.1**, **m** 12-16°, **b** 72°/14mm, 160°/760mm, 163°/atm d_4^{20} 1.015, n_D^{20} 1.431, pK^{25} 4.65. Aqueous methacrylic acid (90%) is saturated with NaCl (to remove the bulk of the water), then the organic phase is dried with $CaCl_2$ and distilled under vacuum. Polymerisation inhibitors should be added to the corrosive distillate and include 0.25% *p*-methoxyphenol, 0.1% hydroquinone, or 0.05% *N,N'*-diphenyl-*p*-phenylenediamine. [*Beilstein* **2** IV 1518.]

Methacrylic anhydride [760-93-0] $C_8H_{10}O_3$, **M 154.2**, **b** 65°/2mm, d_4^{20} 1.040, n_D^{20} 1.454. Distil the anhydride at 2mm pressure, immediately before use, in the presence of hydroquinone. [*Beilstein* **2** IV 1537.]

Methacrylonitrile [126-98-7] C_4H_5N , **M 67.1**, **m** -35.8°, **b** 90.3°/atm, d_4^{20} 0.800, n_D^{20} 1.4007, n_D^{30} 1.3954. Wash it with saturated aqueous $NaHSO_3$ (to remove inhibitors such as *p*-*tert*-butylcatechol), 1% NaOH in saturated NaCl and then with saturated NaCl. Dry it with $CaCl_2$ and fractionally distil it under nitrogen to separate it from impurities such as methacrolein and acetone. [*Beilstein* **2** IV 1537.]

Methacryloyl chloride [920-46-7] C_4H_5ClO , **M 104.5**, **m** -60°, **b** 95-96°/760mm, 98.4°/772mm, d_4^{25} 1.076, n_D^{20} 1.4432. Purify the ester by fractional distillation. If it contains the acid (OH bands in the IR) then add redistilled $SOCl_2$ (with cooling) and cuprous chloride (ca to 2%), reflux the mixture gently for 1 hour and fractionate it through a 1 metre column packed with glass helices. Redistillation then provides the acid chloride

in high purity as a colourless liquid. It is necessary to keep the apparatus moisture free (use CaCl_2 tubes). Stabilise it with 0.05% of 2,6-di-*tert*-butyl-4-methylphenol. [Lal & Green *J Org Chem* **20** 1030 1955, DOI: 10.1021/jo01365a013; *Beilstein* **2** IV 1537.]

Methane [74-82-8] CH_4 , **M 16.0, m -184° , b $-164^\circ/760\text{mm}$, $-130^\circ/6.7\text{atmospheres}$, d_4^{20} 0.554** (cf. d_4^{20} 1.00 for air). Dry methane by passing over CaCl_2 and P_2O_5 , then through a Dry-ice trap and fractionally distil it from a liquid-nitrogen trap. Oxygen can be removed by prior passage in a stream of hydrogen over reduced copper oxide at 500° , and higher hydrocarbons can be removed by chlorinating about 10% of the sample: the hydrocarbons, chlorides and HCl are readily separated from the methane by condensing the sample in the liquid nitrogen trap and fractionally distilling it. Methane has also been washed with concentrated H_2SO_4 , then solid NaOH and then 30% NaOH solution. It is dried with CaCl_2 , then P_2O_5 , and condensed in a trap at liquid air temperature, then transferred to another trap cooled in liquid nitrogen. CO_2 , O_2 , N_2 and higher hydrocarbons can be removed from methane by adsorption on charcoal. [Eiseman & Potter *J Res Nat Bur Stand* **58** 213 1957, DOI: org/10.6028/jres.058.028; *Beilstein* **1** IV 3.] **HIGHLY FLAMMABLE.**

Methanesulfonic acid [75-75-2] $\text{CH}_3\text{O}_3\text{S}$, **M 96.1, m 20° , b $134.5\text{--}135^\circ/3\text{mm}$, d_4^{20} 1.483, n_D^{20} 1.432, pK^{25} -1.86 (-1.2).** Dry the acid, either by azeotropic removal of water with *benzene or toluene, or by stirring 20g of P_2O_5 with 500ml of the acid at 100° for 0.5 hours. Then distil it under vacuum and fractionally crystallise it by partial freezing. Sulfuric acid, if present, can be removed by prior addition of $\text{Ba}(\text{OH})_2$ to a dilute solution, filtering off the BaSO_4 and concentrating under reduced pressure; and is sufficiently pure for most applications. [*Beilstein* **4** IV 10.]

Methanesulfonyl chloride [124-63-0] $\text{CH}_3\text{ClO}_2\text{S}$, **M 114.5. b $55^\circ/11\text{mm}$, d_4^{20} 1.474, n_D^{20} 1.452.** Distil the sulfonyl chloride from P_2O_5 under vacuum. It is a strong **IRRITANT**. [*Beilstein* **4** IV 27.]

Methanol (methyl alcohol) [67-56-1] CH_4O , **M 32.0, m -98° , b $64.5^\circ/\text{atm}$, $64.7^\circ/\text{atm}$, d^{15} 0.79609, d^{25} 1.32663, n^{15} 1.33057, n^{25} 1.32663, pK^{25} 15.5.** Almost all methanol is now obtained synthetically. Likely impurities are water, acetone, formaldehyde, ethanol, methyl formate and traces of dimethyl ether, methylal, methyl acetate, acetaldehyde, carbon dioxide and ammonia. Most of the water (down to about 0.01%) can be removed by fractional distillation. Drying with CaO is unnecessary and wasteful. Anhydrous methanol can be obtained from 'absolute' material by passage through Linde type 4A molecular sieves, or by drying with CaH_2 , CaSO_4 , or with just a little more sodium than required to react with the water present, in all cases the methanol is then distilled. Two treatments with sodium reduces the water content to about $5 \times 10^{-5}\%$. [Friedman et al. *J Am Chem Soc* **83** 4050 1961, DOI: 10.1021/ja01480a023.] Lund and Bjerrum [*Chem Ber* **64** 210 1931, DOI: 10.1002/cber.19310640204] warmed clean dry magnesium turnings (5g) and iodine (0.5g) with 50-75ml of 'absolute' methanol in a flask until the iodine disappeared and all the magnesium was converted to the methoxide. Up to 1L of methanol was added and, after refluxing for 2-3 hours, it was distilled off, excluding moisture from the system. Redistillation from tribromobenzoic acid removes basic impurities and traces of magnesium oxides, and leaves conductivity-quality material. The method of Hartley and Raikes [*J Chem Soc* **127** 524 1925, DOI: 10.1039/CT9252700524] gives a slightly better product. This consists of an initial fractional distillation, followed by distillation from aluminium methoxide, and then ammonia and other volatile impurities are removed by refluxing for 6 hours with freshly dehydrated CuSO_4 (2g/L) while dry air is passed through: the methanol is finally distilled. (The aluminium methoxide is prepared by warming with aluminium amalgam (3g/L) until all the aluminium has reacted. The amalgam is obtained by warming pieces of sheet aluminium with a solution of HgCl_2 in dry methanol.) This treatment also removes aldehydes.

If acetone is present in the methanol, it is usually removed prior to drying. Bates, Mullaly and Hartley [*J Chem Soc* 401 1923, DOI: 10.1039/CT9232300401] dissolved 25g of iodine in 1L of methanol and then poured the solution, with constant stirring, into 500ml of M NaOH. Addition of 150ml of water precipitated iodoform. The solution was allowed to stand overnight, filtered, then boiled under reflux until the odour of iodoform disappeared, and fractionally distilled. (This treatment also removes formaldehyde.) Morton and Mark [*Ind Eng Chem (Anal Edn)* **6** 151 1934] refluxed methanol (1L) with furfural (50ml) and 10% NaOH solution (120ml) for 6-12 hours, the refluxing resin carries down with it the acetone and other carbonyl-containing impurities. The alcohol was then fractionally distilled. Evers and Knox [*J Am Chem Soc* **73** 1739 1951, DOI: 10.1021/ja01148a090], after refluxing 4.5L of methanol for 24 hours with 50g of magnesium, distilled off 4L of

it, which they then refluxed with AgNO_3 for 24 hours in the absence of moisture or CO_2 . The methanol was again distilled, shaken for 24 hours with activated alumina before being filtered through a glass sinter and distilled under nitrogen in an all-glass still. Material suitable for conductivity work was obtained.

Variations of the above methods have also been used. For example, a sodium hydroxide solution containing iodine has been added to methanol and, after standing for 1 day, the solution has been poured slowly into about a quarter of its volume of 10% AgNO_3 , shaken for several hours, then distilled. Sulfanilic acid has been used instead of tribromobenzoic acid in Lund and Bjerrum's method. A solution of 15g of magnesium in 500ml of methanol has been heated under reflux, under nitrogen, with hydroquinone (30g), before degassing and distilling the methanol, which was subsequently stored with magnesium (2g) and hydroquinone (4g per 100ml). Refluxing for about 12 hours removes the bulk of the formaldehyde from methanol: further purification has been obtained by subsequent distillation, refluxing for 12 hours with dinitrophenylhydrazine (5g) and H_2SO_4 (2g/L), and again fractionally distilling. [Beilstein 1 IV 1227.]

Rapid purification: Methanol purification is the same as for ethanol. Another simple purification procedure consists of adding 2g of NaBH_4 to 1.5L methanol, gently bubbling argon through it and refluxing for a day at 30° , then adding 2g of freshly cut sodium (washed with methanol) and refluxing for 1 day before distilling. The middle fraction is taken. [Jou & Freeman *J Phys Chem* 81 909 1977, DOI: 10.1021/j100524a021.] **VERY TOXIC SHOULD NOT BE SWALLOWED.**

Methoxyacetic acid [625-45-6] $\text{C}_3\text{H}_6\text{O}_3$, M 90.1, m $7-9^\circ$, b $97^\circ/13-14\text{mm}$, 202-204°/atm, d_4^{20} 1.175, n_D^{20} 1.417, pK^{25} 3.57. Fractionally crystallise the acid by repeated partial freezing, then fractionally distil it under vacuum through a vacuum-jacketed Vigreux column ~20cm long. [Beilstein 3 IV 574.]

Methoxyamine hydrochloride (O-methylhydroxylamine HCl) [593-56-6] $\text{CH}_5\text{NO} \cdot \text{HCl}$, M 83.5, m $151-152^\circ$, pK^{25} 4.60. Crystallise the hydrochloride from absolute EtOH or EtOH by addition of diethyl ether. [Kovach et al. *J Am Chem Soc* 107 7360 1985, DOI: 10.1021/ja00311a024; Beilstein 1 IV 1252.]

2-Methoxyethanol (methylcellosolve) [109-86-4] $\text{C}_3\text{H}_8\text{O}_2$, M 76.1, m -85° , b $124.4^\circ/\text{atm}$, d_4^{20} 0.964, n_D^{20} 1.4017, pK^{25} 14.8. Peroxides can be removed by refluxing with stannous chloride or by filtration under slight pressure through a column of activated alumina. 2-Methoxyethanol can be dried with K_2CO_3 , CaSO_4 , MgSO_4 or silica gel, then distilled from sodium. Aliphatic ketones (and water) can be removed by making the solvent 0.1% in 2,4-dinitrophenylhydrazine and allowing to stand overnight with silica gel before fractionally distilling. [Beilstein 1 IV 2375.]

2-Methoxyethoxymethylchloride (MEMCl) [3970-21-6] $\text{C}_4\text{H}_9\text{ClO}_2$, M 124.6, b $50-52^\circ/13\text{mm}$, 140-145°(dec)/atm, d_4^{20} 1.092, n_D^{20} 1.427. Possible impurities are methoxyethanol (b $124^\circ/\text{atm}$), HCHO and HCl which can be removed below the boiling point of MEMCl. Purify MEMCl by fractional distillation in a vacuum. If too impure, prepare it from methoxyethanol (152g) and *s*-trioxane (66g) by bubbling a stream of dry HCl (with stirring) until a clear mixture is obtained. Dilute with pentane (900ml), dry (3 hours over 100g MgSO_4 , at 5°), evaporate and the residue is distilled in a vacuum. It is **MOISTURE SENSITIVE** and **TOXIC**. The **MEM.NEt₃·Cl** salt, prepared by reaction with 1.3 equivalents of Et_3N (16 hours/ 25°) and dried in a vacuum, has m $58-61^\circ$, and is moisture sensitive. [Corey et al. *Tetrahedron Lett* 809 1976, DOI: 10.1021/ie50367a005; Yoshimatsu et al. *J Org Chem* 59 1011 1994, DOI: 10.1021/jo00084a017; Greene & Wuts *Protective Groups in Organic Synthesis* 3rd edn, J Wiley & Sons NY 1991.] **CARCINOGEN.**

2-Methoxyethylamine [109-85-3] $\text{C}_3\text{H}_9\text{NO}$, M 75.1, b $94^\circ/\text{atm}$, d_4^{20} 0.874, n_D^{20} 1.407, pK^{25} 9.40. An aqueous 70% solution of the amine is dehydrated by azeotropic distillation with *benzene or methylene chloride and the amine is distilled twice from zinc dust. Store it in a tight container as it absorbs CO_2 from the atmosphere. [Beilstein 4 IV 1411.]

N-Methylacetamide [79-16-3] $\text{C}_3\text{H}_7\text{NO}$, M 73.1, m 30° , b $70-71^\circ/2.5-3\text{mm}$, pK_1^{25} -3.70, pK_2^{25} -0.42. Fractionally distil it under vacuum, then fractionally crystallise it twice from its melt. Likely impurities include acetic acid, methyl amine and H_2O . For a detailed purification procedure, see Knecht and Kolthoff, *Inorg Chem* 1 195 1962, DOI: 10.1021/ic50002a002. Although N-methylacetamide is commercially available it is often extensively contaminated with acetic acid, methylamine, water and an unidentified impurity. The recommended

procedure is to synthesise it in the laboratory by direct reaction. The gaseous amine is passed into hot glacial acetic acid, to give a partially aqueous solution of methylammonium acetate which is heated to *ca* 130° to expel water. Chemical methods of purification such as extraction by petroleum ether, treatment with H₂SO₄, K₂CO₃ or CaO can be used but are more laborious.

Tests for purity include the Karl Fischer titration for water; this can be applied directly. Acetic acid and methylamine can be detected polarographically.

In addition to the above, purification of *N*-methylacetamide can be achieved by fractional freezing, including zone melting, repeated many times, or by vacuum distillation under reduced pressures. For details of zone melting techniques, see Knecht in *Recommended Methods for Purification of Solvents and Tests for Impurities*, Coetzee Ed. Pergamon Press 1982. [Beilstein 4 IV 176.]

Methyl acetate [79-20-9] C₃H₆O₂, M 74.1, m -98°, b 56.7-57.2°/atm, d₄²⁰ 0.934, n_D²⁰ 1.36193, n_D²⁵ 1.3538, pK²⁰ -7.28 (H₀ scale, aqueous H₂SO₄). Methanol in methyl acetate can be detected by measuring its solubility in water. At 20°, the solubility of methyl acetate in water is *ca* 35g per 100ml, but 1% MeOH confers complete miscibility. Methanol can be removed by conversion to methyl acetate, by refluxing for 6 hours with acetic anhydride (85ml/L), followed by fractional distillation. Acidic impurities can be removed by shaking with anhydrous K₂CO₃ and distilling. An alternative treatment is with acetyl chloride, followed by washing with concentrated NaCl and drying with CaO or MgSO₄. (Solid CaCl₂ cannot be used because it forms a crystalline addition compound.) Distillation from copper stearate destroys peroxides. Free alcohol or acid can be eliminated from methyl acetate by shaking with strong aqueous Na₂CO₃ or K₂CO₃ (three times), then with aqueous 50% CaCl₂ (three times), saturated aqueous NaCl (twice), drying with K₂CO₃ and distilling it from P₂O₅. [Beilstein 2 IV 122.]

Methyl acetimidate hydrochloride [14777-27-6] C₃H₇NO. HCl, M 109.6, m 93-95°, 105°(dec), pK_{Est} ~ 5.5. Crystallise the imidate from methanol by adding dry ether to a ratio of 1:1 and cooling at 0°. Filter off the crystals in a cold room, wash them with methanol/ether (1:2), then dry in a vacuum. [Hunter & Ludwig *J Am Chem Soc* 84 3491 1962, DOI: 10.1021/ja00877a016.] The *free base* has b 90-91°/765mm, d₄²⁰ 0.867, n_D²⁰ 1.403. [Hunter & Ludwig *Methods Enzymol* 25 585 1972, DOI: 10.1016/S0076-6879(72)25058-3; Beilstein 2 IV 181.]

Methyl acrylate [96-33-3] C₄H₆O₂, M 86.1, m -75°, b 80°/atm, d₄²⁰ 0.9535, n_D²⁰ 1.4040. Wash the ester repeatedly with aqueous NaOH until free from inhibitors (such as hydroquinone), then wash it with distilled water, dry (CaCl₂) and fractionally distil it under reduced pressure in an all-glass apparatus. Seal it under nitrogen and store it at 0° in the dark. [Bamford & Han *JCS Faraday Trans 1* 78 855 1982, DOI: 10.1039/F19827800855; Beilstein 2 IV 1457.]

Methylamine (gas) [74-89-5] CH₅N, M 31.1, m -93°, b -7.55°/719mm, pK²⁵ 10.62. Dry the amine with sodium or BaO. It is commercially available in metal cylinders. It is a strong organic base. [Beilstein 4 IV 118.]

Methylamine hydrochloride [593-51-1] CH₅N. HCl, M 67.5, m 231.8-233.4°, b 225-230°/15mm, pK²⁵ 10.62. Crystallise the salt from *n*-butanol, absolute EtOH or MeOH/CHCl₃. Wash it with CHCl₃ to remove traces of dimethylamine hydrochloride. Dry it under vacuum first with H₂SO₄ then P₂O₅. It is deliquescent; store it in a desiccator over P₂O₅. [Beilstein 4 IV 122.]

Methyl bromide [74-83-9] CH₃Br, M 94.9, m -94°, b 3.6°/atm. Purify it by bubbling through conc H₂SO₄, followed by passage through a tube containing glass beads coated with P₂O₅. Also purify it by distillation from AlBr₃ at -80°, by passage through a tower of KOH pellets and by partial condensation. [Beilstein 1 IV 68.]

2-Methylbutane (isopentane) [78-78-4] C₅H₁₂, M 72.2, b 27.9°/atm, d₄²⁰ 0.621, n_D²⁰ 1.35373, n_D²⁵ 1.35088. Stir isopentane for several hours in the cold with concentrated H₂SO₄ (to remove olefinic impurities), then wash it with H₂O, aqueous Na₂CO₃ and H₂O again. Dry it with MgSO₄ and fractionally distil it using a Todd column packed with glass helices. Material transparent down to 180nm is obtained by distilling from sodium wire, and passing through a column of silica gel which had previously been dried in place at 350° for 12 hours before use. [Potts *J Chem Phys* 20 809 1952, DOI: 10.1063/1.1700572; Beilstein 1 IV 320.]

2-Methyl-1-butanol [137-32-6, *RS*(±) 34713-94-5, *S*(-) 1565-80-6] $C_5H_{12}O$, *M* 88.2, *b* 130°(*RS*), *m* -70°, 128.6°(*S*), $[\alpha]_D^{25}$ -5.8 (neat), d_4^{20} 0.809, n_D^{52} 1.4082. Reflux the butanol with CaO, distil, reflux with magnesium and again fractionally distil it. A small sample of highly purified material is obtained by fractional crystallisation after conversion into a suitable ester such as the trinitrophthalate or the 3-nitrophthalate. The latter is converted to the cinchonine salt in acetone and recrystallised from $CHCl_3$ by adding pentane. The salt is saponified, extracted with ether, and fractionally distilled. [Terry et al. *J Chem Eng Data* **5** 403 1960, DOI: 10.1021/je60008a002; *Beilstein* **1** IV 1666.]

2-Methyl-2-butanol (tert-pentyl alcohol) [75-85-4] $C_5H_{12}O$, *M* 88.2, *m* -12°, *b* 50°/60mm, 66°/165mm, 102°/atm, d_4^{20} 0.805, n_D^{20} 1.405. Reflux it with magnesium, then fractionally distil it with high reflux ratio. It forms azeotropic mixtures with many organic liquids. [Cottle & Power *J Am Chem Soc* **58** 2267, DOI: 10.1021/ja01302a052; Urry et al. *J Am Chem Soc* **76** 450 1954, DOI: 10.1021/ja01631a037.] [*Beilstein* **1** H 388, **1** I 195, **1** II 422, **1** III 1622, **1** IV 1668.]

3-Methyl-1-butanol (iso-pentyl alcohol) [123-51-3] $C_5H_{12}O$, *M* 88.2, *m* -117°, *b* 130°/atm, 132°/760mm, d_4^{25} 0.812, n_D^{25} 1.406. It is a major component of fusel oil. Reflux it with magnesium, then fractionally distil it with high reflux ratio. It was synthesised from the Grignard reagent, 2-propane magnesium bromide, and ethylene oxide in 85% yield, and the α -naphthylurethane had *m* 67° [Huston & Agett *J Org Chem* **06** 123 1941, DOI: 10.1021/jo01201a012]. It is an **IRRITANT**. The *acetate ester* [123-92-2] $C_7H_{14}O_2$, *M* 130.1, on the other hand is a liquid with a fruity odour and has *b* 142°/atm, d_4^{15} 0.876, n_D^{21} 1.3997. [*Beilstein* **1** H 392, **1** I 196, **1** II 426, **1** III 1633, **1** IV 1677.]

3-Methyl-2-butanol [598-75-4] $C_5H_{12}O$, *M* 88.2, *b* 111.5°/atm, d_4^{20} 0.807, n_D^{20} 1.4095, n_D^{25} 1.4076. Reflux it with magnesium, then fractionally distil it. [*Beilstein* **1** IV 1675.]

3-Methyl-2-butanone (methyl isopropyl ketone) [563-80-4] $C_5H_{10}O$, *M* 86.1, *m* -92°, *b* 93-94°/752mm, d_4^{20} 0.818, *n* 1.410, pK^{25} -7.1 (aqueous H_2SO_4). Reflux the ketone with a little $KMnO_4$. Fractionate it through a spinning-band column, dry with $CaSO_4$ and distil it. [*Beilstein* **1** IV 3287.]

2-Methyl-2-butene see **amylene** above.

2-Methyl-3-butyne-2-amine (1,1-dimethylpropargylamine, 3-amino-3-methyl-1-butyne) [2978-58-7] C_5H_9N , *M* 83.1, *b* 79-80°/760mm, d_4^{25} 0.790, n_D^{25} 1.4183, pK_{Est} ~8.0. Dissolve the amine in Et_2O , dry over anhydrous K_2CO_3 , filter, evaporate and distil (preferably under N_2). Store it away from CO_2 . The *hydrochloride* [2978-59-8] has *m* 234° (from $EtOH/Et_2O$). The *benzoyl* derivative has *m* 152-153° (from $EtOH$). [Hennion & Teach *J Am Chem Soc* **75** 1653 1953, DOI: 10.1021/ja01103a040; Hennion & DiGiovanna *J Org Chem* **30** 2645 1965, DOI: 10.1021/jo01019a033.]

Methyl *n*-butyrate [623-42-7] $C_5H_{10}O_2$, *M* 102.1, *b* 102.3°/760mm, d_4^{20} 0.898, n_D^{20} 1.389. Treat the ester with anhydrous $CuSO_4$, then distil it under dry nitrogen. [*Beilstein* **2** IV 786.]

***S*-(+)-2-Methylbutyric acid** [1730-91-2] $C_5H_{10}O_2$, *M* 102.1, *b* 64°/2mm, 78°/15mm, 90-94°/23mm, 174-175°/atm, d_4^{20} 0.938, n_D^{20} 1.406, $[\alpha]_{546}^{20}$ +23, $[\alpha]_D^{20}$ +19.8 (neat), $[\alpha]_D^{13}$ +18.3 (c 6, $EtOH$), pK^{25} 4.76 (for *RS*). Purify the acid by distilling it *in vacuo* [Sax & Bergmann *J Am Chem Soc* **77** 1910 1955, DOI: 10.1021/ja01612a065; Doering & Aschner *J Am Chem Soc* **75** 393 1953, DOI: 10.1021/ja01098a040]. The *methyl ester* is formed by addition of diazomethane and has *b* 112-115°/760mm, $[\alpha]_D^{27}$ +21.1 (c 1.7, $MeOH$). [*Beilstein* **2** IV 888.] The racemate, ***RS*-2-methylbutyric acid** [116-53-0] has *b* 102-103°/atm, d_4^{25} 0.986, n_D^{20} 1.405. [*Beilstein* **2** 305.]

Methyl carbamate (urethylane) [598-55-0] $C_2H_5NO_2$, *M* 75.1, *m* 54.4-54.8°, 56-58°, *b* 176-177°/~760mm. Crystallise the carbamate from *benzene or distil it. [*Beilstein* **3** H 21, **3** IV 37.]

Methyl chloride [74-87-3] CH_3Cl , *M* 50.5, *m* -97°, *b* -24.1°/atm, d_4^{25} 0.915g/ml. Bubble methyl chloride through a sintered-glass disc dipped into conc H_2SO_4 , then wash it with water, condense it at low temperature

and fractionally distil it. It has been distilled from AlCl_3 at -80° . *Alternatively*, pass it through towers containing AlCl_3 , soda-lime and P_2O_5 , then condense and fractionally distil it. Use low temperature techniques. Store it as a gas. [Beilstein 1 IV 28.]

Methyl chloroacetate [96-34-4] $\text{C}_3\text{H}_5\text{ClO}_2$, M 108.5, m -33° , b $129\text{--}130^\circ/\text{atm}$, d_4^{20} 1.230, n_D^{20} 1.423. Shake the ester with saturated aqueous Na_2CO_3 (three times), aqueous 50% CaCl_2 (three times), saturated aqueous NaCl (twice), dry (Na_2SO_4) and fractionally distil it. **Very toxic**. [Beilstein 2 IV 480.]

R-(+) Methyl 2-chloropropionate [77287-29-7] $\text{C}_4\text{H}_7\text{ClO}_2$, M 122.6, b $49\text{--}50^\circ/35\text{mm}$, $78\text{--}80^\circ/120\text{mm}$, $132\text{--}134^\circ/760\text{mm}$, d_4^{20} 1.152, n_D^{20} 1.417, $[\alpha]_D^{20}$ +26 (19.0) (neat). Purify the ester by repeated distillation [Walker *J Chem Soc* 67 914 1895, DOI: 10.1039/CT8956700914; Walden *Chem Ber* 28 1293 1985, see also Gless *Synth Commun* 16 633 1986, DOI: 10.1080/00397918608057732]. [Beilstein 2 H 248.] The racemate **RS-methyl 2-chloropropionate** [17639-93-9] has b $132\text{--}134^\circ/760\text{mm}$, d_4^{25} 1.075, n_D^{20} 1.417.

Methyl cyanoacetate [105-34-0] $\text{C}_4\text{H}_5\text{NO}_2$, M 99.1, f -13° , b $115^\circ/36\text{mm}$, $200.4\text{--}200.9^\circ/761\text{mm}$, d_4^{20} 1.128, n_D^{20} 1.420. Purify the ester by shaking with 10% Na_2CO_3 solution, wash well with water, dry with anhydrous Na_2SO_4 , and distil it. [Beilstein 2 H 584, 2 I 253, 2 II 530, 2 III 1628, 2 IV 1889.]

Methyl cyanoformate [17640-15-2] $\text{C}_3\text{H}_3\text{NO}_2$, M 85.1, b $81^\circ/47\text{mm}$, $97^\circ/751\text{mm}$, $100\text{--}101^\circ/760\text{mm}$, $104\text{--}107^\circ/\text{atm}$, d_4^{20} 1.072, n_D^{20} 1.37378. Purify the ester by fractionation through a 45cm glass helices packed column or a 30cm spinning band column. [Sheppard *J Org Chem* 27 3756 1962, DOI: 10.1021/jo01058a003.] It has been distilled through a short Vigreux column, and further purified by recrystallisation from Et_2O at -40° as white crystals which melt at room temperature. NMR: δ 4.0 (CH₃), and the IR has ν_{max} at 2250 (CN) and 1750 (CO) cm^{-1} . [Childs & Weber *J Org Chem* 41 3486 1976, DOI: 10.1021/jo00883a041; Beilstein 2 III 1587, 2 IV 1889.]

Methyl decanoate (methyl caprate) [110-42-9] $\text{C}_{11}\text{H}_{22}\text{O}_2$, M 186.3, m -14° to -11° , b $114^\circ/15\text{mm}$, $224^\circ/760\text{mm}$, d_4^{20} 0.874, n_D^{20} 1.426. Pass the ester through alumina before use and distil in a vacuum. [Beilstein 2 IV 1044.]

N-Methyldiethanolamine [MDEA, *N,N*-bis(hydroxyethyl)methylamine, 2,2'-methyliminodiethanol] [105-59-9] $\text{C}_5\text{H}_{13}\text{NO}_2$, M 119.2, b $75\text{--}77^\circ/0.5\text{mm}$, $115^\circ/5\text{mm}$, $131^\circ/10\text{mm}$, $141\text{--}142^\circ/18\text{mm}$, $246\text{--}248^\circ/\text{atm}$, d_4^{25} 1.038, n_D^{20} 1.469, pK^{25} 8.57, pK^{35} 8.31, pK^{45} 8.13, pK^{60} 7.87. Purify MDEA by fractional distillation preferably under vacuum in a stream of N_2 , and store it under N_2 . The colourless distillate darkens in air and absorbs CO_2 under pressure [Goodridge *Trans Faraday Soc* 51 1703 1955, DOI: 10.1039/TF9555101703]. It is a tertiary base, hence it does not form an *N*-carbamate salt like primary and secondary amines, but **does** form a *carbonate salt* in the presence of carbonic acid (i.e. $\text{CO}_2 + \text{H}_2\text{O} = \text{H}_2\text{CO}_3$). It is soluble in H_2O and EtOH , but slightly soluble in Et_2O . The *hydrochloride* is very hygroscopic; the *O,O'*-diacetyl ester has b $110^\circ/4\text{mm}$, $133^\circ/15\text{mm}$, b $251^\circ/\text{atm}$, the *O,O'*-bis(4-nitrobenzoyl) ester has m $112\text{--}113^\circ$, the *Reineckate salt* has m 168° , the *picrate* crystallises from EtOAc with m $95\text{--}96^\circ$, and with a strong aqueous solution of AuCl_3/HCl the *chloroaurate salt*, m $101\text{--}102^\circ$, is formed which crystallises from H_2O . With PtCl_4/HCl , the *chloroplatinate salt* is obtained, forming orange-yellow rhombic crystals upon recrystallisation from a concentrated aqueous solution which sinter at 145° and decompose at $148\text{--}150^\circ$. The IR spectrum in CCl_4 has C-H str at ν_{max} (ϵ mole $^{-1}$ L.cm $^{-1}$) 2779 (93), 2844 (81), 2877 (99) and 2948 (107) cm^{-1} . [Hanby & Rydon *J Chem Soc* 513 1947, DOI: 10.1039/JR9470000513; Hill & Meakins *J Chem Soc* 760 1958, DOI: 10.1039/JR9580000760; Knorr & Matthes *Chem Ber* 31 1069 1898, DOI: 10.1002/cber.189803101193; Beilstein 4 H 284, 4 II 729, 4 III 692, 4 IV 1517.]

Methyl dodecanoate (methyl laurate) [111-82-0] $\text{C}_{13}\text{H}_{26}\text{O}_2$, M 214.4, m 5° , b $141^\circ/15\text{mm}$, d_4^{20} 0.870, n_D^{50} 1.4199. Pass the ester through alumina before use, and distil it in a vacuum. [Beilstein 2 IV 1090.]

N-Methyleneaminoacetonitrile (MAAN) [109-82-0] $(\text{C}_3\text{H}_4\text{N}_2)_3 = \text{C}_9\text{H}_{12}\text{N}_6$, M 68.1, m 129° . The reaction involving ammonia, formaldehyde and cyanide provided a crystalline substance originally thought to be $\text{CH}_2=\text{NCH}_2\text{CN}$. However, it turned out to be a trimeric form (see further). After washing the crude product thoroughly with cold H_2O , drying at $50\text{--}60^\circ$ and recrystallising from 95% EtOH a 52% yield of colourless cryst-

als **m 129°** (*orthorhombic*) were obtained which turned out to be a *trimeric form*. From the ethanolic mother liquors a *second trimeric form* with **m 86°** (monoclinic crystals) was isolated in a relative proportion of ~8%. These could be recrystallised from Me₂CO. Their reactivities, however are different. It crystallises nicely from H₂O but reacts with it with considerable loss of material. It is sparingly soluble in *C₆H₆. It is an inhibitor of bone growth. [Johnson & Rinehart *J Am Chem Soc* **46** 768, 1653 1924, DOI: 10.1021/ja01668a031; Adams & Langley *Org Synth Coll Vol I* 355 1941, DOI: 10.15227/orgsyn.004.0047.]

Methyl ether (dimethyl ether) [115-10-6] C₂H₆O, **M 46.1**, **m -141°**, **b -63.5°/96.5mm**, **-24°/~760mm**, **d₄²⁵ 1.918/L** (at 1 atmosphere relative to air as 1). Dry methyl ether by passing over alumina and then BaO, or over CaH₂, followed by fractional distillation at low temperatures. Its solubility is 37ml per ml of H₂O at 18°, and it is very soluble in EtOH and Et₂O. [Beilstein **1** IV 1245.]

N-Methyl ethylamine hydrochloride [624-60-2] C₃H₉N. HCl, **M 95.6**, **m 126-130°**, **pK²⁵ 10.9** (free base). Crystallise the hydrochloride from absolute EtOH or diethyl ether. Dry it *in vacuo*. [Beilstein **4** H 94.]

N-Methyl formamide [123-39-7] C₂H₅NO, **M 59.1**, **m -3.5°**, **b 100.5°/25mm**, **d₄²⁰ 1.005.**, **n 1.4306** Dry it over molecular sieves for 2 days, then distil it under reduced pressure through a column packed with glass helices. Fractionally crystallise it by partial freezing and the solid portion is distilled in a vacuum. [Beilstein **4** IV 170.]

Methyl formate [107-31-3] C₂H₄O₂, **M 60.1**, **m -100°**, **b 31.5°/atm, 34°/atm**, **d₄²⁰ 0.971**, **n¹⁵ 1.34648**, **n_D²⁰ 1.34332**. Wash the formate with strong aqueous Na₂CO₃, dry it with solid Na₂CO₃ and distil it from P₂O₅. (Procedure removes free alcohol or acid.) [Beilstein **2** IV 20.]

2-Methylglutaric acid [18069-17-5] C₆H₁₀O₄, **M 146.1**, **m 79°**, **80-82°**, **b 214-215°/22mm**, **pK₁²⁵ 4.36**, **pK₂²⁵ 5.37**. Crystallise the acid from distilled water, then dry it under vacuum over concentrated H₂SO₄. [Beilstein **2** IV 1989.]

3-Methylglutaric acid [626-51-7] C₆H₁₀O₄, **M 146.1**, **m 87°**, **pK₁²⁵ 4.35**, **pK₂²⁵ 5.44**. Crystallise the acid from distilled water, then dry it under vacuum over concentrated H₂SO₄. [Beilstein **2** IV 1992.]

Methylglyoxal [78-98-8] C₃H₄O₂, **M 72.1**, **b ca 72°/760mm**. Commercial 30% (w/v) aqueous solution is diluted to about 10% and distilled twice, taking the fraction boiling below **50°/20mm**. (This treatment does not remove lactic acid). It is *hygroscopic* and polymerises readily. The aqueous solution is more stable probably because it forms a hydrate, i.e. the 1,1-diol. [Beilstein **1** IV 3631.]

Methylglyoxal 1,1-dimethyl acetal (1,1-dimethoxyacetone, pyruvaldehyde-1-dimethyl acetal) [6342-56-9] C₅H₁₀O₃, **M 118.1**, **b 40°/10mm, 63-66°/35mm, 141°/760mm, 144-147°/atm**, **d₄²⁵ 0.976**, **n_D²⁴ 1.4480**, **n_D²⁵ 1.4102**. Purify by careful fractionation, preferably under reduced pressure. If however it is too impure (cf. IR and NMR) then add at least an equal volume of absolute MeOH, and saturate the solution with dry HCl. Any precipitated salt is filtered off, the filtrate is allowed to stand at room temperature in a stoppered vessel. The mixture is neutralised (to litmus) with methanolic NaOMe, the precipitated salts are filtered off and the filtrate is evaporated under diminished pressure. The residue is mixed with an equal volume of *C₆H₆, the *C₆H₆—immiscible layer is removed and the benzene solution is distilled. [Royals & Robinson III *J Am Chem Soc* **78** 4161 1956, DOI: 10.1021/ja01597a083; Braude & Evans *J Chem Soc* 3324 1955, DOI: 10.1039/JR9550003324.] [Beilstein **1** I 395, **1** IV 3631.]

2-Methylhexane (isoheptane) [591-76-4] C₇H₁₆, **M 100.2**, **m -118°**, **b 90.1°/atm**, **d₄²⁰ 0.678**, **n_D²⁰ 1.38485**, **n_D²⁰ 1.38227**. Purify it by azeotropic distillation with MeOH, then wash it with water (to remove the MeOH), dry it over type 4A molecular sieves and distil it. [Beilstein **1** IV 397.]

3-Methylhexane [589-34-4] C₇H₁₆, **M 100.2**, **m -119°**, **b 91.9°/atm**, **d₄²⁰ 0.687**, **n_D²⁰ 1.38864**, **n_D²⁰ 1.38609**. Purify it as for 2-methylhexene. [Beilstein **1** IV 399.]

Methyl hexanoate (methyl caproate) [106-70-7] $\text{C}_7\text{H}_{14}\text{O}_2$, M 130.2, m -71° , b $52^\circ/15\text{mm}$, $150^\circ/760\text{mm}$, d_4^{20} 0.885, n_D^{20} 1.410. Pass it through alumina and distil it before use. [Beilstein 2 IV 921.]

Methylhydrazine [60-34-4] CH_6N_2 , M 46.1, b $87^\circ/745\text{mm}$, d_4^{20} 0.876, n_D^{20} 1.436, pK^{30} 7.87. Dry with BaO, then distil it in a vacuum. Store it under nitrogen away from CO_2 . [Beilstein 4 IV 3322.]

Methyl hydrazinocarboxylate [6294-89-9] $\text{C}_2\text{H}_6\text{N}_2\text{O}_2$, M 90.1, m $70-73^\circ$, b $108^\circ/12\text{mm}$. To remove impurities, the material is melted and pumped under vacuum until the vapours are spectroscopically pure [Caminati et al. *J Am Chem Soc* 108 4364 1986, DOI: 10.1021/ja00275a023]. Distil it in a vacuum. [Beilstein 3 I 46.]

Methyl iodide [74-88-4] CH_3I , M 141.9, m -64° , b $42.8^\circ/\text{atm}$, d_4^{20} 2.281, n_D^{20} 1.5315. Methyl iodide deteriorates rapidly with liberation of iodine if exposed to light. It is usually purified by shaking it with dilute aqueous $\text{Na}_2\text{S}_2\text{O}_3$ or NaHSO_3 until colourless, then washing with water, dilute aqueous Na_2CO_3 , and more water, drying with CaCl_2 and distilling. It is stored in a brown bottle away from sunlight in contact with a small amount of mercury, powdered silver or copper. (Prolonged exposure of mercury to methyl iodide forms methylmercuric iodide.) Methyl iodide can be dried further using CaSO_4 or P_2O_5 . An alternative purification is by percolation through a column of silica gel or activated alumina, then distillation. The solution can be degassed by using repeated freeze-pump-thaw cycles. Cu metal can be added as stabiliser. [Beilstein 1 IV 87.]

O-Methylisourea hydrogen sulfate (2-methylpseudourea sulfate, O-methylisourea bisulfate) [29427-58-5] $\text{C}_2\text{H}_6\text{N}_2\text{O} \cdot 0.5 \text{H}_2\text{SO}_4$, M 172.2, m $114-118^\circ$, 119° . Recrystallise the salt from MeOH/Et₂O (327g of salt dissolved in 1L of MeOH and 2.5L of Et₂O is added) [Fearing & Fox *J Am Chem Soc* 76 4382 1954, DOI: 10.1021/ja01646a033]. The *picrate* has m 192° [Odo et al. *J Org Chem* 23 1319 1958, DOI: 10.1021/jo01103a020]. [Beilstein 3 IV 143.]

N-Methyl maleimide [930-88-1] $\text{C}_5\text{H}_5\text{NO}_2$, M 111.1, m $94-96^\circ$. Crystallise the imide three times from diethyl ether. Dry it *in vacuo*. [Beilstein 21/10 V 5.]

Methylmalonic acid (2-methylpropanedioic acid) [516-05-2] $\text{C}_4\text{H}_6\text{O}_4$, M 118.1, m $135^\circ(\text{dec})$, pK_1^{25} 3.05, pK_2^{25} 5.76. The acid crystallises as the *hydrate* from water. [Beilstein 2 IV 1932.]

Methyl methacrylate [80-62-6] $\text{C}_5\text{H}_8\text{O}_2$, M 100.1, f -50° , b $46^\circ/100\text{mm}$, $100^\circ/\sim 760\text{mm}$, d_4^{20} 0.937, n_D^{20} 1.4144. Wash the ester twice with aqueous 5% NaOH (to remove inhibitors such as hydroquinone) and twice with water. Dry it with CaCl_2 , Na_2CO_3 , Na_2SO_4 or MgSO_4 , then with CaH_2 under nitrogen under reduced pressure. The distillate is stored at low temperatures and redistilled before use. Prior to distilling, inhibitors such as hydroquinone (0.004%), β -naphthylamine (0.2%) or di- β -naphthol are sometimes added. Also purify it by boiling with aqueous H_3PO_4 solution and finally with saturated NaCl solution. It is dried for 24 hours over anhydrous CaSO_4 , distilled at 0.1mm Hg at room temperature, and stored at -30° [Albeck et al. *JCS Faraday Trans 1* 1 1488 1978, DOI: 10.1039/F19787401488]. [Beilstein 2 II 398, 2 III 1279, 2 IV 1519.]

Methyl methanesulfonate [66-27-3] $\text{C}_2\text{H}_6\text{O}_3\text{S}$, M 110.3, b $59^\circ/0.6\text{mm}$, $96-98^\circ/19\text{mm}$, $202-203^\circ/\text{atm}$, d_4^{20} 1.300, n_D^{20} 1.4140. Purify the ester by careful fractionation and collecting the middle fraction. Suspected **CARCINOGEN**. (Note that MeSO_3H has b $167-167.5^\circ/10\text{mm}$ and methanesulfonic anhydride has b $138^\circ/10\text{mm}$)—both are possible impurities. [Beilstein 4 IV 11.]

S-Methyl methanethiolsulfonate [2949-92-0] $\text{C}_2\text{H}_6\text{O}_2\text{S}_2$, M 126.2, b $69-71^\circ/0.4\text{mm}$, $96-97^\circ/4.5\text{mm}$, $104-105^\circ/10\text{mm}$, $119^\circ/16\text{mm}$, d_4^{20} 1.226, n_D^{20} 1.515. Purify it by fractional distillation under reduced pressure, the IR has ν_{max} at $1350, 750 \text{ cm}^{-1}$. [Slusarchyk et al. *J Org Chem* 38 943 1973, DOI: 10.1021/jo00945a023; Beilstein 4 IV 31.]

Methyl nitrate [598-58-3] CH_3NO_3 , M 77.0, b $5^\circ/50\text{mm}$, explodes at $65^\circ/760\text{mm}$, d_5^{15} 1.2322, d_5^{15} 1.2167, d_5^{25} 1.2032. Wash MeONO_2 once with H_2O then again with H_2O containing a few drops of concentrated NaOH to keep it slightly alkaline (litmus). Dry the ester over anhydrous CaCl_2 , decant it and use it directly. It is poss-

ible to distil it under a vacuum with slow and gentle heating, as a sudden rise in temperature can cause decomposition with copious release of nitrous fumes (use extreme precautions and protection). The middle fraction can then be subjected to several freeze-pump-thaw cycles. **TAKE ALL NECESSARY SAFETY PRECAUTIONS as the VAPOUR CAN EXPLODE ON HEATING EVEN IN THE ABSENCE OF OXYGEN.** [Black & Babers *Org Synth Col Vol* **2** 412 1943, DOI: 10.15227/orgsyn.019.0064.] [Beilstein **1** H 284, **1** I 141, **1** II 273, **1** III 1201, **1** IV 1254.]

Methyl nitrite [624-91-9] CH_3NO_2 , **M 61.0, b -18°/atm, -17°/atm, d_{15}^{15} (liquid) 0.991.** Condense MeONO in a liquid nitrogen trap. Distil the greenish liquid under vacuum (preferably in a vacuum line), into the first trap containing dry Na_2CO_3 to free it from acid impurities then into further Na_2CO_3 and fused CaCl_2 traps before collection at -78°. It has been distilled through columns that are surrounded by Et_2O /Dri-ice cooled to -30°. [Leermakers & Ramsperger *J Am Chem Soc* **54** 1838 1932, DOI: 10.1021/ja01344a015; Thompson & Purkis *Trans Faraday Soc* **32** 674 1936, DOI: 10.1039/TF9363200674; Beilstein **1** H 284, **1** I 141, **1** II 273, **1** III 1201, **1** IV 1253.] **CARCINOGEN.**

2-Methyl-2-nitro-1,3-propanediol [77-49-6] $\text{C}_4\text{H}_9\text{NO}_4$, **M 135.1, m 145°, 147-148°, 149-150°, 150.6°.** Crystallise it from *n*-butanol or Me_2CO (**m 150.6°**). It decomposes on attempted distillation at 10mm. Its solubility in H_2O is 80g/100ml at 20°. It can be distilled at 10mm pressure (take precautions). [Gabriel *Ind Eng Chem* **32** 887 1940, DOI: 10.1021/ie50367a005.] [Beilstein **1** H 489, **1** II 547, **1** III 2190, **1** IV 2537.]

2-Methyl-2-nitro-1-propanol [76-39-1] $\text{C}_4\text{H}_9\text{NO}_3$, **M 119.1, m 87-88°, 93-96°, b 60-61°/1mm, 75°/2mm, 94-95°/10mm, d_{25}^{25} 1.303, n_D^{25} 1.4425.** Distil it under vacuum and/or crystallise it from petroleum ether or MeOH . [Astle & Abbott *J Org Chem* **21** 1228 1956, DOI: 10.1021/jo01117a007; Kambe & Yasuda *Bull Soc Chem Jpn* **41** 1444 1968, DOI:org/10.1246/bcsj.41.1444; Beilstein **1** H 378, **1** III 1546, **1** IV 1604.]

2-Methyloctane [3221-61-2] C_9H_{20} , **M 128.3, f.p -80.1°, b 141.6°/727.5mm, 142.8°/760mm, d_4^{20} 0.7107, n_D^{20} 1.40285.** It is present in petroleum and is a major constituent in *Hypericum perforatum*. Take it through a silica gel column and distil it. If too crude then wash sequentially with 70% H_2SO_4 , 85% H_2SO_4 , then concentrated H_2SO_4 (handle with necessary protection from the strong acid) until the hydrocarbon liquid is almost free from the yellow colour. Cool the organic layer then wash with ice cold H_2O , twice with 25% NaOH solution, dry over Na , then carefully fractionate from Na-K alloy using high reflux ratios (30—60:1). [Whitmore & Southgate *J Am Chem Soc* **60** 2571 1938, DOI: 10.1021/ja01278a003; Whitmore & Orem *J Am Chem Soc* **60** 2573 1938, DOI: 10.1021/ja01278a004, and for fractionating columns see Whitmore & Lux *J Am Chem Soc* **54** 3448 1932, DOI: 10.1021/ja01347a071]. [Beilstein **1** III 507, **1** IV 454]

3-Methyloctane [2216-33-3] C_9H_{20} , **M 128.3, m b -108°/atm, 142-144°/760mm, d_4^{20} 0.719, n_D^{20} 1.407.** It was isolated from a petroleum fraction [Jamison, Lesslie and Turner *J Inst Petr* **75** 595 1949]. Take it through a silica gel column and distil it. Alternatively, purify it like the preceding 2-methyloctane. [Klassen & Ross *J Phys Chem* **91** 3668 1987, DOI: 10.1021/j100297a041; Beilstein **1** III 507, **1** IV 455]

S(+)-3-Methyloctane C_9H_{20} , **M 128.3, b 32-42°/10mm, n_D^{20} 1.4080, 144°/atm, n_D^{20} 1.4081, $[\alpha]_{5461}^{19}$ +8.39, $[\alpha]_{5896}^{19}$ +7.54 (c 9.8, CHCl_3).** Prepared by catalytic reduction of optically active 6-methyloct-3-ene-1-ol using $\text{Pd/BaSO}_4/\text{H}_2$ in Et_2O overnight (complete H_2 uptake; i.e. apparently the OH is replaced by H before reduction of the double bond — allylic OH hydrogenolysis) whose absolute configuration at the chiral centre had been correlated with that of *S*(-)-glyceraldehyde. Removal of catalyst, evaporation, followed by two distillations gave pure chiral hydrocarbon. [Crombie & Harper *J Chem Soc* 2685 1950, DOI: 10.1039/JR9500002685; Beilstein **1** IV 455.] **R(-)-3-Methyloctane** C_9H_{20} , **M 128.3, has b 143°/atm, n_D^{25} 1.4052, $[\alpha]_D^{27}$ -8.5 (CHCl_3),** Levene & Marker *J Biol Chem* **91** 77 (102) 1931, <http://www.jbc.org/content/91/1/77.citation>].

Methyl octanoate (methyl caprylate) [111-11-5] $\text{C}_9\text{H}_{18}\text{O}_2$, **M 158.2, b 83°/15mm, 193-194°/760mm, d^{25} 0.877g/ml, n_D^{20} 1.419.** Pass the ester through alumina and distil it before use. [Beilstein **2** IV 986.]

Methyl oleate (methyl *cis*-9-octadecenoate) [112-62-9] $\text{C}_{19}\text{H}_{36}\text{O}_2$, **M 296.5, f -19.9°, b 217°/16mm, d_4^{20}**

0.874, n_D^{20} 1.4522. Purify the oleate by fractional distillation under reduced pressure, and by low temperature crystallisation from acetone. Store it in the dark under N_2 . [Beilstein 2 IV 1649.]

Methylpentane (mixture of isomers) C_6H_{14} . Pass the mixture through a long column of activated silica gel (or alumina) and collect material that is transparent down to 200nm in the UV.

2-Methylpentane (isohexane) [107-83-5] C_6H_{14} , M 86.2, m -154° , b 60.3° , d_4^{20} 0.655, n_D^{20} 1.37145, n_D^{25} 1.36873. Purify it by azeotropic distillation with MeOH, followed by washing out the MeOH with water, drying ($CaCl_2$, then sodium), and distilling it. [Forziati et al. *J Res Nat Bur Stand* 36 129 1946, DOI: org/10.6028/jres.036.005; Beilstein 1 IV 358.]

3-Methylpentane [96-14-0] C_6H_{14} , M 86.2, b $63.3^\circ/atm$, d_4^{20} 0.664, n_D^{20} 1.37652, n_D^{25} 1.37384. Purify it by azeotropic distillation with MeOH, as for 2-methylpentane. Purify it for ultraviolet spectroscopy by passing it through columns of silica gel or alumina activated by heating for 8 hours at 210° under a stream of nitrogen. Alternatively, treat it with conc (or fuming) H_2SO_4 , then wash it with water, aqueous 5% NaOH, water again, then dry ($CaCl_2$, then sodium), and distil it through a long, glass helices-packed, column. [Beilstein 1 IV 363.]

RS-2-Methyl-2,4-pentanediol (MPD, pinecon, hexylene glycol) [107-41-5] $C_6H_{14}O_2$, M 118.2, m -40° , b $97^\circ/10mm$, $102^\circ/12mm$, $107.5-108.5^\circ/25mm$, $135-136^\circ/40mm$, $197^\circ/760mm$, d^{25} 0.922g/ml, n_D^{20} 1.4265. It is present in *Santolina rosmarinifolia*. Dry the diol with Na_2SO_4 , then CaH_2 , and fractionally distil it under reduced pressure through a helices packed column, taking precautions to avoid absorption of water. It is soluble in H_2O , EtOH, Et_2O , pentane and hexane. It is a reagent for characterising aldehydes by forming cyclic acetates. A 20% solution can be used for the extraction of Boron. The *RS-diacetate* [99113-75-4] $C_{10}H_{18}O_4$, M 202.2 is a liquid with m -40° , b $95^\circ/15mm$, $92^\circ/18mm$. It has very harmful vapours; use efficient fume cupboard as it irritates eyes, skin throat and lungs, causing headaches and nausea and is a CNS depressant. The *bis-4-nitrobenzoyl ester* forms pale yellow flakes from aqueous EtOH, m $155-157^\circ$. The *R(-) enantiomer* [99210-90-9] has b $108-110^\circ/6mm$, $[\alpha]_D^{25}$ -14.7 (neat 87% ee). The *S(+)-enantiomer* [99210-91-0] has b $87-90^\circ/6mm$, $[\alpha]_D^{25}$ $+15$ (c 1, EtOH, 92% ee). [For optical resolutions using yeast or porcine pancreatic lipases in nearly anhydrous organic solvents see Kirchner et al. *J Am Chem Soc* 107 7072 1985, DOI: 10.1021/ja00310a052.] [Beilstein 1 H 486, 1 I 252, 1 II 552, 1 III 2206, 1 IV 2565.]

2-Methyl-1-pentanol [105-30-6] $C_6H_{14}O$, M 102.2, b $65-66^\circ/60mm$, $146-147^\circ/760mm$, d_4^{20} 0.827, n_D^{20} 1.420. Dry the 1-pentanol with Na_2SO_4 and distil it. [Beilstein 1 IV 1713.]

4-Methyl-2-pentanol (isobutyl methyl carbinol) [108-11-2] $C_6H_{14}O$, M 102.2, m -90° , b $47^\circ/10mm$ $131-132^\circ/atm$ d_4^{20} 0.810, n_D^{20} 1.413. Wash the 2-pentanol with aqueous $NaHCO_3$, dry and distil it. Purify it further by converting it to the phthalate ester by adding 120ml of dry pyridine and 67g of phthalic anhydride per mole of alcohol, purifying the ester and steam distilling it in the presence of NaOH. The distillate is extracted with ether, and the extract is dried and fractionally distilled. The *phenylurethane* has m 143° , and the *2,4-dinitrobenzoate* has m $60.5-61.5^\circ$. *R(-)-4-Methyl-2-pentanol* [16404-54-9] has b $132^\circ/atm$, d_4^{25} 0.802, n_D^{20} 1.4630, $[\alpha]_D^{22}$ -18.5 (c 1, $CHCl_3$), -20.8 (neat) and the *1-naphthylurethane* has m $86-87^\circ$, $[\alpha]_D^{22}$ -3.7 (EtOH). *S(+)-4-methyl-2-pentanol* [14898-80-7] has b $63-64^\circ/45mm$, $138^\circ/atm$, d_4^{25} 0.802, n_D^{20} 1.4790, $[\alpha]_D^{22}$ $+18.0$ (c 1, $CHCl_3$), $+21.4$ (neat). [Levine & Walti *J Biol Chem* 94 367 1931, <http://www.jbc.org/content/94/2/373.citation>; for optical resolution via the brucine salts of the *o*-carboxybenzoate ester followed by acidification and hydrolysis see Kenyon & Strauss *J Chem Soc* 2153 1949, DOI: 10.1039/JR9490002153; for chiral synthesis of the alcohols via reduction of the respective ketone using $\{LiAlH(OR')\}$ (*N*-methylephedrine) see Vigneron & Jacquet *Tetrahedron* 32 939 1976, DOI: 10.1016/0040-4020(76)85053-3; and for absolute configuration see Bell *Aust J Chem* 32 2625 1979, DOI: 10.1071/CH9792625; Beilstein 1 H 410, 1 I 203, 1 II 440, 1 III 1669, 1 IV 1717.]

3-Methyl-3-pentanol carbamate (Emylcamate) [78-28-4] $C_7H_{15}NO_2$, M 145.2, m $56-58.5^\circ$, b $24^\circ/0.7mm$, $35^\circ/10mm$. It crystallises in needles with the odour of camphor from 30% aqueous EtOH, and is dried over a dessicant. It distils at a low temperature in a vacuum. Its solubility in H_2O at room temperature is 0.4g/100mL. This drug reduces anxiety. [Beilstein 1 IV 1773.]

4-Methyl-2-pentanone (methyl isobutyl ketone) [108-10-1] $C_6H_{12}O$, M 100.2, m -80° , b $115.7^\circ/760\text{mm}$, d_4^{20} 0.801, n_D^{20} 1.3958, n_D^{25} 1.3938. Reflux the ketone with a little $KMnO_4$, wash it with aqueous $NaHCO_3$, dry with $CaSO_4$ and distil it. Acidic impurities are removed by passage through a small column of activated alumina. [Beilstein 1 IV 3305.]

2-Methyl-1-pentene [763-29-1] C_6H_{12} , M 84.2, m -126° , b $61.5-62^\circ/\text{atm}$, d_4^{20} 0.680, n_D^{20} 1.395. Water is removed, and formation of peroxides is prevented by several vacuum distillations of 2-methyl-1-pentene from sodium. It is stored with sodium-potassium alloy. [Beilstein 1 IV 841.]

4-Methyl-1-pentene [691-37-2] C_6H_{12} , M 84.2, b $61.5-62^\circ/\text{atm}$, d^{25} 0.680g/ml, n_D^{20} 1.395. Water is removed, and formation of peroxides is prevented by several vacuum distillations of 2-methyl-1-pentene from sodium. It is stored with sodium-potassium alloy. [Beilstein 1 IV 846.]

cis-4-Methyl-2-pentene [691-38-3] C_6H_{12} , M 84.2, m -134.4° , b $57.7-58.5^\circ/\text{atm}$, d_4^{20} 0.672, n_D^{20} 1.388. Dry the *cis*-pentene with CaH_2 , and distil it. [Beilstein 1 IV 841.]

trans-4-Methyl-2-pentene [674-76-0] C_6H_{12} , M 84.2, m -140.8° , b $58.5^\circ/\text{atm}$, d_4^{20} 0.669, n_D^{20} 1.389. Dry the *trans*-isomer with CaH_2 , and distil it. [Beilstein 1 IV 844.]

2-Methylpropane-1,2-diamine (1,2-diamino-2-methylpropane) [811-93-8] $C_4H_{12}NO_2$, M 88.2, b $47-48^\circ/17\text{mm}$, $50^\circ/20\text{mm}$, $59^\circ/20\text{mm}$ (mono hydrate?), pK_I^{25} 6.25 (6.18), pK_2^{25} 9.82 (9.42). Dry the diamine with sodium for 2 days, then distil it from sodium under reduced pressure. The *dihydrochloride* [15444-85-6] m 306° (dec) forms plates from aqueous EtOH. The anhydrous *picrate* has m 241° . It complexes with Cu^{2+} , Cr^{3+} , Co^{2+} and Ni^{2+} . [Bosnich et al. *Aust J Chem* 19 2045 1966, DOI:10.1071/CH9662045; Basolo et al. *J Am Chem Soc* 75 1478 1953, DOI: 10.1021/ja01102a507.] [Beilstein 4 H 264, 4 II 707, 4 III 581, 4 IV 1306.]

2-Methylpropane-1-thiol (isobutylmercaptan) [513-44-0] $C_4H_{10}S$, M 90.2, b $41.2^\circ/142\text{mm}$, $88.5^\circ/760\text{mm}$, n_D^{25} 1.43582, $pK_{\text{Est}} \sim 10.8$. Dissolve the thiol in EtOH, and add to 0.25M $Pb(OAc)_2$ in 50% aqueous EtOH. The precipitated lead mercaptide is filtered off, washed with a little EtOH, and impurities are removed from the molten salt by steam distillation. After cooling, dilute HCl is added dropwise to the residue, and the mercaptan is distilled directly from the flask. Water is separated from the distillate, and the mercaptan is dried (Na_2CO_3) and distilled under nitrogen. [Mathias *J Am Chem Soc* 72 1897 1950, DOI: 10.1021/ja01161a012; Beilstein 1 H 378, 1 I 191, 1 II 412, 1 III 1565, 1 IV 1605.]

2-Methylpropane-2-thiol (tert-butylmercaptan) [75-66-1] $C_4H_{10}S$, M 90.2, m -0.5° , b $61.6^\circ/701\text{mm}$, $66^\circ/760\text{mm}$, d_4^{25} 0.79426, n_D^{25} 1.41984, pK^{25} 11.22. Dry the thiol for several days over CaO , then distil it from CaO . Purify it as for 2-methylpropane-1-thiol above. [Beilstein 1 H 383, 1 II 416, 1 III 1589, 1 IV 1634.]

2-Methyl-1-propanol (isobutanol) [78-83-1] $C_4H_{10}O$, M 74.1, m -108° , b $107.9^\circ/760\text{mm}$, d_4^{20} 0.804, n_D^{15} 1.39768, n_D^{25} 1.3939. Isobutanol is dried by refluxing with CaO and BaO for several hours, followed by treatment with calcium or aluminium amalgam, then fractional distilling it from sulfanilic or tartaric acids. More exhaustive purifications involve formation of phthalate or borate esters. Heating it with phthalic anhydride gives the *acid phthalate* which, after crystallisation to constant melting point (m 65°) from petroleum ether, is hydrolysed with aqueous 15% KOH. The alcohol is distilled off as the water azeotrope and dried (K_2CO_3 , then anhydrous $CuSO_4$), and finally magnesium turnings, followed by fractional distillation. [Hückel & Ackermann *J Prakt Chem* 136 15 1933.] The borate ester is formed by heating the dried alcohol for 6 hours in an autoclave at $160-175^\circ$ with a quarter of its weight of boric acid. After fractional distillation under vacuum, the ester is hydrolysed by heating for a short time with aqueous alkali and the alcohol is dried with CaO and distilled. [Michael et al. *J Am Chem Soc* 38 653 1916, DOI: 10.1021/ja02260a018.] Alternatively, dry the alcohol with K_2CO_3 , $CaSO_4$ or $CaCl_2$, filter and fractionally distil it. For further drying, the redistilled alcohol can be refluxed with the appropriate alkyl phthalate or succinate as described under *ethanol*. [Beilstein 1 IV 1588.]

Methyl propiolate [922-67-8] $C_4H_4O_2$, M 84.1, b $100^\circ/\text{atm}$, $102^\circ/\text{atm}$, $103-105^\circ/\text{atm}$, d_4^{20} 0.945, n_D^{20} 1.4080. Purify the propiolate by fractional distillation and collecting the middle fraction, note that propiolic acid

has a higher boiling point [144°(dec)/760mm]. [*Beilstein* 2 IV 1688.] **LACHRYMATORY**.

N-Methylpropionamide [1187-58-2] C_4H_9NO , **M 87.1**, **f -30.9°**, **m -43°**, **b 103°/12-13mm**, **d₄²⁰ 0.934**, **n_D²⁵ 1.4356**. The amide is a colourless, odourless, neutral liquid at room temperature with a high dielectric constant. The amount of water present can be determined directly by Karl Fischer titration, GLC and NMR have been used to detect unreacted propionic acid. Commercial material of high quality is available, probably from the condensation of anhydrous methylamine with 50% excess of propionic acid. Rapid heating to 120-140° with stirring favours the reaction by removing water either directly or as the ternary xylene azeotrope. The quality of the distillate improves during the distillation.

N-Methylpropionamide can be dried over CaO. Water and unreacted propionic acid are removed as their xylene azeotropes. It is then distilled in a vacuum. Material used as an electrolyte solvent (specific conductance less than $10^{-6} \text{ ohm}^{-1} \text{ cm}^{-1}$) is obtained by fractional distillation under reduced pressure, and storage over BaO or molecular sieves because it readily absorbs moisture from the atmosphere on prolonged storage. [*Hoover Pure Appl Chem* 37 579 1974, DOI: org.virtual.anu.edu.au/10.1351/pac197437040579; *Recommended Methods for Purification of Solvents and Tests for Impurities*, Coetzee Ed., Pergamon Press, 1982, *Beilstein* 4 IV 183.]

Methyl propionate [554-12-1] $C_4H_8O_2$, **M 88.1**, **m -88°**, **b 79.7°/atm**, **d²⁵ 0.915 g/ml**, **n_D²⁰ 1.376**. Wash the ester with saturated aqueous NaCl, then dry it with Na_2CO_3 and distil it from P_2O_5 . (This removes any free acid and alcohol.) It has also been dried with anhydrous $CuSO_4$. [*Beilstein* 2 IV 704.]

Methyl n-propyl ether [557-17-5] $C_4H_{10}O$, **M 74.1**, **b 39°/743mm**, **38.8°/atm**, **d₄²⁰ 0.736**, **n_D¹⁴ 1.3602**, **pK²⁵ -3.79 (aqueous H_2SO_4)**. Dry it with $CaSO_4$, then pass the ether through a column of alumina (to remove peroxides) and fractionally distil it. Its solubility in H_2O at 25° is 5% v/v. It is a **FLAMMABLE** anaesthetic. [Bennett & Phillip *J Chem Soc* 1930 1928, DOI: 10.1039/JR9280001930; *Beilstein* 1 H 354, 1 I 178, 1 II 367, 1 III 1413, 1 IV 1421.]

Methyl n-propyl ketone (pentan-2-one) [107-87-9] $C_5H_{10}O$, **M 86.1**, **m -78°**, **b 102.4°/atm**, **d₄²⁰ 0.807**, **n_D²⁰ 1.3903**. Purify the ketone by refluxing it with a little $KMnO_4$, dry it with $CaSO_4$ and distil it. It can be converted to its bisulfite addition compound by shaking with excess saturated aqueous $NaHSO_3$ at room temperature, cooling to 0°, filtering, washing with diethyl ether and drying. Steam distillation of the adduct gives a distillate from which the ketone is recovered, washed with aqueous $NaHCO_3$ and distilled water, dried (K_2CO_3) and fractionally distilled. **IRRITANT**. [Waring & Garik *J Am Chem Soc* 78 5198 1956, DOI: 10.1021/ja01601a019; *Beilstein* 1 IV 3271.] 5198 1956,

RS-3-Methyl-1-pentyn-3-ol (Meparfynol) [77-75-8] $C_6H_{10}O$, **M 98.1**, **m -30.6°**, **b 20°/6.5mm**, **50°/37mm**, **121-122°/atm**, **d₄²⁰ 0.8688**, **n_D²⁰ 1.4318**. Purify by removing H_2O with Drierite and distillation in a vacuum or at atmospheric pressure. The acrid smelling liquid has a burning taste. Its solubility in H_2O is 12.8% w/v and it is very soluble in most organic solvents. [Hurd & McPhee *J Am Chem Soc* 69 239 1947, DOI: 10.1021/ja01194a018; *Beilstein* 1 IV 2242.]

RS-3-Methyl-1-pentyn-3-ol carbamate (Meparfynol carbamate,) [302-66-9] $C_7H_{11}NO_2$, **M 141.2**, **m 55.8-57°**, **56-58°**, **120-121°/16mm**. Crystallise it from $*C_6H_6$, hexane, ether/petroleum ether or cyclohexane. Its solubility in H_2O is 1.6% w/v. [McLamore, P'an & Bawley *J Org Chem* 20 1379 1955, DOI: 10.1021/jo01127a014; *Beilstein* 1 IV 65.] It is a sedative.

Methyl stearate (methyl octadecanoate) [122-61-8] $C_{19}H_{38}O_2$, **M 298.5**, **m 41-43°**, **b 181-182°/4mm**. Crystallise the ester from petroleum ether or distil it in a vacuum. [*Beilstein* 2 IV 1216.]

RS-2-Methylsuccinic acid (2-methylbutane-1,4-dioic acid) [498-21-5] $C_5H_8O_4$, **M 132.1**, **m 112-112.5**, **111-113°**, **115.0°**, **pK₁²⁵ 3.88**, **pK₂²⁵ 5.35**. Crystallise the acid from water (prisms), Et_2O or $CHCl_3$ /petroleum ether (b 60-80°), and dry it *in vacuo* at 20° over P_2O_5 . The acid can be sublimed at 70-80°/5 x 10^{-4} mm or 150°/5 x 10^{-4} mm unchanged. The **dimethyl ester** [21307-96-0] $C_7H_{14}O_4$, **M 160.1**, is an oil with **f.p -80°**, **b 197°/atm**. [Conn et al. *J Am Chem Soc* 64 1747 1942, DOI: 10.1021/ja01260a001; Duncanson *J Chem Soc* 1753 1952, DOI: 10.1039/JR9520001753; Brown *Org Synth Coll Vol* 3 615 1955, DOI: 10.15227/orgsyn.026.0054.]

[*Beilstein* 2 H 637, 2 I 274, 2 II 568, 2 III 1695, 2 IV 1948.]

***R*(+)-2-Methylsuccinic acid** [3641-51-8] $C_5H_8O_4$, M 132.1, m 111-112°, 114.5°, 115°, $[\alpha]_D^{20} +9.2$ (c 5.1, H_2O), $[\alpha]_D^{20} +11.3$ (c 3.1, H_2O), $[\alpha]_D^{20} +16.5$ (c 4.4, EtOH), +16.88 (c 2.16, EtOH). It was prepared from the (±)-acid by optical resolution of the strychnine salt. It has also been isolated from degradation of terpenes. Recrystallise the acid from water, $*C_6H_6$ or $*C_6H_6/Et_2O$. The **dimethyl ester** [22644-27-5] $C_7H_{14}O_4$, M 160.1, is an oil with b 93-94°/0.03mm, $[\alpha]_D^{17.9} +9.98$ (MeOH). [Duncanson *J Chem Soc* 1753 1952, DOI: 10.1039/JR9520001753; Naps & Johns *J Am Chem Soc* 62 2450 1940, DOI: 10.1021/ja01866a053; McMillan *J Chem Soc* 1823 1959, DOI: 10.1039/JR9590001823.] [*Beilstein* 2 III 1694, 2 IV 1948.]

***S*(-)-2-Methylsuccinic acid** [2174-58-5] $C_5H_8O_4$, M 132.1, m 110-115°, 114.5°, $[\alpha]_D^{20} -9.2$ (c 5.1, H_2O), $[\alpha]_D^{20} -9.9$ (c 13, H_2O), $[\alpha]_D^{20} -15.9$ (EtOH). Crystallise the acid from water, $*C_6H_6$, or $*C_6H_6$ /petroleum ether (b 60-80°). The **dimethyl ester** [63163-08-8] $C_7H_{14}O_4$, M 160.1, is an oil with b 140°/12mm, $[\alpha]_D^{19.4} -9.94$ (MeOH). [*Beilstein* 2 III 1694, 2 IV 1948.]

***RS*-2-Methylsuccinic anhydride (pyrotartaric anhydride)** [4100-80-5] $C_5H_6O_3$, M 114.0, m 33-35°, b 105°/2mm, 118-120°/7mm, 135.78°/24mm, 150°/13mm, 238-240°/atm, 244-248°/atm, $d_4^{25} 1.22$. Reflux the anhydride with acetic anhydride, or $AcCl$ + a little $SOCl_2$, or by boiling with $POCl_3$ (1-3 hrs), evaporate off the solvent in a vacuum, redistil the residue and collect the solid distillate. When recrystallised from $CHCl_3$ it has m 33°. The (±)-anhydride can be sublimed. [*Beilstein* 17 H 414.]

***R*(+)-2-Methylsuccinic anhydride** [83247-83-0] $C_5H_6O_3$, M 114.0, m 64-65°, 67-69°, $[\alpha]_D^{29} +32.1$ (c 2.717, EtOH), +31.3 (c 1.737, $CHCl_3$), $[\alpha]_D^{28} +31.3$ (c 1.280, Me_2CO), $[\alpha]_D^{20} +32.6$ (c 15, dioxane). It is prepared by dehydration of the chiral acid or impure anhydride with Ac_2O , or $AcCl$ + a little $SOCl_2$, or by boiling with $POCl_3$ (1-3 hrs), and evaporating to dryness. The residual solid anhydride is dried *in vacuo* over P_2O_5 or KOH , and recrystallised from $CHCl_3$ /petroleum ether or $*C_6H_6$ /petroleum ether (b 60-80°). [Naps & Johns *J Am Chem Soc* 62 2450 1940, DOI: 10.1021/ja01866a053; McMillan *J Chem Soc* 1823 1959, DOI: 10.1039/JR9590001823.]

***S*(-)-2-Methylsuccinic anhydride** [6973-20-2] $C_5H_6O_3$, M 114.0, m 69-70°, $[\alpha]_D^{20} -26.4$ (undiluted extrapolated), $[\alpha]_D^{30} -32.6$ (c 2, $CHCl_3$), $[\alpha]_D^{28} -31.3$ (c 1.280, Me_2CO), $[\alpha]_D^{31} -32.6$ (c 2, EtOH). Purified as for the preceding *R*-enantiomer and recrystallised from $*C_6H_6$. [Naps & Johns *J Am Chem Soc* 62 2450 1940, DOI: 10.1021/ja01866a053.]

***R*(+)-2-Methylsuccinic acid 4-methyl ester** [81025-83-4] $C_6H_{10}O_4$, M 146.1, b 241-242°/760mm, $d_4^{25} 1.150$, $n_D^{20} 1.4320$, $[\alpha]_D^{20} +11$ (c 1, $CHCl_3$). Purify by distillation [*Beilstein* 2 III 1694, 2 IV 1948.]

***R*(+)-2-Methylsuccinimide** [117307-07-0] $C_5H_7NO_2$, M 113.1, m 78°, $[\alpha]_D^{20} +29.4$ (c 2, $CHCl_3$). The imide is purified by recrystallising from toluene.

***N*-Methylsuccinimide** [1121-07-9] $C_5H_7NO_2$, M 113.1, m 61-70°, 66°, 66-67°, 68-70°, b 234-235°/atm, 225-227°/atm. It crystallises from Et_2O /petroleum ether in needles, and also from hot EtOH or Me_2CO , but distils at 234° at atmospheric pressure without decomposition. It is slightly soluble in H_2O . [*Beilstein* 21 H 373, 21 II 303, 21 III/IV 4544.]

***N*-Methylthioacetamide** [5310-10-1] C_2H_5NS , M 89.1, m 59°. Recrystallise the amide from $*benzene$ or EtOH. [Todd et al. *Chem Ber* 69 217 1936, DOI: 10.1002/cber.19360690139; *Beilstein* 4 I 329, 4 III 124.]

Methyl trifluoromethanesulfonate (methyl triflate) [333-27-7] $C_2H_3F_3O_3S$, M 164.1, b 97-97.5°/736mm, 99°/~760mm, 100-102°/~760mm, $d_4^{20} 1.496$, $n_D^{25} 1.3238$. It is a strong methylating agent but is corrosive and **POISONOUS**. Fractionate it carefully and collecting the middle fraction (use an efficient fume cupboard) and keep away from moisture. It is a **POWERFUL ALKYLATING AGENT** and a strong **IRRITANT**. [IR: Gramstad & Haszeldine *J Chem Soc* 173 1956, DOI: 10.1039/JR9560000173; *J Chem Soc* 4069 1957 DOI: 10.1039/JR9570004069.] **Trifluoromethanesulfonic acid (triflic acid)** [1493-13-6] M 151.1, boils higher (b

162°/atm), has a **pK_a** of 3.10, and is **TOXIC** and hygroscopic. [Hansen *J Org Chem* **30** 4322 1965, DOI: 10.1021/jo01023a511; Kurz & El-Nasr *J Am Chem Soc* **104** 5823 1982, DOI: 10.1021/ja00385a062; *Beilstein* **3** IV 34.]

Methyl vinyl ketone (3-buten-2-one) [78-94-4] **C₄H₆O**, **M 70.1**, **b 34°/120mm**, **62-68°/400mm**, **79-80°/760mm**, **d₄²⁰ 0.845**, **n_D²⁰ 1.413**. It forms an 85% azeotrope with water. After drying with K₂CO₃ and CaCl₂ (with cooling), the ketone is distilled at low pressures. [*Beilstein* **1** IV 3444.]

Methyl vinyl sulfone [3680-02-2] **C₃H₆O₂S**, **M 106.1**, **b 116-118°/20mm**, **d₄²⁰ 1.215**, **n_D²⁰ 1.461**. Pass the sulfone through a column of alumina, then de-gas, distil it in a vacuum line and store it at -190° until required. [*Beilstein* **1** III 1866.]

N-Monobutyl urea [592-31-4] **C₅H₁₂N₂O**, **M 116.2**, **m 96-98°**, **pK_{Est} ~0.2**. Crystallise the urea from EtOH/water, then dry it under vacuum at room temperature. [*Beilstein* **4** I 371, **4** IV 578.]

N-Monoethyl urea [625-52-5] **C₃H₈N₂O**, **M 88.1**, **m 92-95°**, **pK_{Est} ~0.2**. Crystallise the urea from EtOH/water, then dry it under vacuum at room temperature. [*Beilstein* **4** IV 369.]

N-Monomethyl urea [598-50-5] **C₂H₆N₂O**, **M 74.1**, **m 93-95°**, **pK_{Est} ~0.2**. Crystallise the urea from EtOH/water, then dry it under vacuum at room temperature. [*Beilstein* **4** IV 205.]

N-Monopropyl urea [627-06-5] **C₄H₁₀N₂O**, **M 102.1**, **m 107°, 110°**, **pK_{Est} ~0.2**. Crystallise the urea from EtOH or EtOH/Et₂O. [Biovin & Biovin *Can J Chem* **29** 479 1951, DOI:10.1139/v51-056; IR: Biovin & Biovin *Can J Chem* **32** 563 1954, DOI: 10.1139/v54-073; *Beilstein* **4** H 142, **4** III 261, **4** IV 482.]

Mucochloric acid (2,3-dichloro-4-oxo-2-butenic acid) [87-56-9] **C₄H₂Cl₂O₃**, **M 169.0**, **m 124-126°**, **pK²⁵ 4.20**. Crystallise the acid twice from water (charcoal). [*Beilstein* **3** IV 1720.]

trans,trans-Muconic acid (hexa-2,4-dienedioic acid) [3588-17-8] **C₆H₆O₄**, **M 142.1**, **m 300°**, **pK²⁵ 4.51**, for **cis,cis-acid** [1119-72-8] **m 194-195°**, **pK²⁵ 4.49**. Crystallise the diacid from H₂O. [*Beilstein* **2** IV 2298.]

Myristic acid (tetradecanoic acid) [544-63-8] **C₁₄H₂₈O₂**, **M 228.4**, **m 58°**, **b 250°/100mm** **pK²⁰ 6.3 (50% aqueous EtOH)**, **pK_{Est} ~4.9 (H₂O)**. Purify the acid *via* the *methyl ester* (**b 153-154°/10mm**, **n_D²⁵ 1.4350**), as for capric acid. [Trachtman & Miller *J Am Chem Soc* **84** 4828 1962, DOI: 10.1021/ja00883a039.] Also purify it by zone melting. It crystallises from petroleum ether, and is dried in a vacuum desiccator containing shredded wax. [*Beilstein* **2** IV 1126.]

Neopentane (2,2-dimethylpropane, tetramethylmethane) [463-82-1] **C₅H₁₂**, **M 72.2**, **flash point 79.3°**, **m -19.8°**, **b 9.5°/760mm**, **d₄²⁰ 0.6737**, **n_D²⁰ 1.38273**. It was prepared by the reaction of *tert*-butyl chloride and methyl magnesium chloride in toluene at 45-60° in 42-50% yield. The crude hydrocarbon was freed from isobutene by passage over concentrated H₂SO₄ or P₂O₅, and through silica gel. It was then fractionated through a packed column (42 x 1.2 cm) provided with a total condenser and a variable take-off. The jacket of the column was kept at 9-10° and the reflux condenser at 0-3° by circulating cooled water through them. Fractions were collected and their freezing points measured. Those with freezing points close to -19° were pooled, the vapour was scrubbed through 85% H₂SO₄ then 25% KOH, dried with P₂O₅, condensed, and re-fractionated as before. The fraction with **b 9.4°/760mm**, and **m -19.8°** was collected. [Whitmore & Flemming *J Am Chem Soc* **55** 3803 1933, DOI: 10.1021/ja01336a058; Whitmore et al. *J Am Chem Soc* **56** 749 1934, DOI: 10.1021/ja01318a508; *Beilstein* **1** H 141, **1** I 50, **1** II 104, **1** 369, **1** IV 333.]

Nerolidol (3,7,11-trimethyl-1,6,10-dodecatrien-3-ol) **C₁₅H₂₆O₂**, **M 222.4** [*cis/trans* 7212-44-4] **b 122°/3mm**, **d₄²⁰ 0.73**, **n_D²⁰ 1.477**, [*cis* 3790-78-1] **b 70°/0.1mm**, [*trans* 40716-66-3] **b 78°/0.2mm**, **145-146°/2mm**. Purify it by TLC on plates of Kieselguhr G [McSweeney *J Chromatogr* **17** 183 1965, DOI: 10.1016/S0021-9673(00)99850-9] or silica gel impregnated with AgNO₃, using 1,2-CH₂Cl₂/CHCl₃/EtOAc/PrOH (10:10:1:1) as solvent system. Also by GLC on butanediol succinate (20%) on Chromosorb W. Store it under N₂ at ~5° in the

dark. [Beilstein 1 IV 2336.]

Nitrilotriacetic acid [tris(carboxymethyl)amine, NTA, Complexone 1] [139-13-9] $\text{C}_6\text{H}_9\text{NO}_6$, M 191.1, m 247°(dec), pK_1 0.8, pK_2 1.71, pK_3 2.47, pK_4 9.71. Crystallise it from water and dry it at 110° in a vacuum. [Beilstein 4 IV 2441.]

Nitroethane [79-24-3] $\text{C}_2\text{H}_5\text{NO}_2$, M 75.1, m -90°, b 115°/atm, d_4^{20} 1.049, n_D^{20} 1.3920, n_D^{25} 1.39015, pK^{25} 8.60 (8.46, pH equilibrium requires *ca* 5 minutes). Purify it as described for *nitromethane* below. A spectroscopic impurity can be removed by shaking it with activated alumina, decanting and distilling it rapidly. [Beilstein 1 IV 170.]

Nitroguanidine [556-88-7] $\text{CH}_4\text{N}_4\text{O}_2$, M 104.1, m 239°(dec), 246-246.5°(dec), 257°, pK_1^{25} -0.55, pK_2^{25} 12.20. Crystallise it from water (20ml/g). The *nitrate* has m 147°(dec) (prisms, H_2O). [Beilstein 3 H 126, 3 III 236.]

Nitromethane [75-52-5] CH_3NO_2 , M 61.0, f -28.5°, b 101.3°/atm, d_4^{20} 1.13749, d^{30} 1.12398, n_D^{20} 1.3819, n^{30} 1.37730, pK^{25} 10.21. Nitromethane is generally manufactured by gas-phase nitration of methane. The usual impurities include aldehydes, nitroethane, water and small amounts of alcohols. Most of these can be removed by drying with CaCl_2 or by distillation to remove the water/nitromethane azeotrope, followed by drying with CaSO_4 . Phosphorus pentoxide is **not** suitable as a drying agent as it forms a sol in MeNO_2 , and a white solid is deposited in the condenser when it is distilled [Wright et al. *J Chem Soc* 199 1931, DOI: 10.1039/JR9310000199]. In an early preparation, Whitmore and Whitmore [*Org Synth* 3 83 1923 DOI: 10.1002/0471264180.os003.24] synthesised nitromethane from sodium nitrite and sodium chloroacetate. The purified material should be stored in dark bottles, away from strong light, in a cool place. Purifications using extraction are commonly used. For example, Van Looy and Hammett [*J Am Chem Soc* 81 3872 1959, DOI: 10.1021/ja01524a019] mixed about 150ml of concentrated H_2SO_4 with 1L of nitromethane and allowed it to stand for 1 or 2 days. The solvent was washed with water, aqueous Na_2CO_3 , and again with water, then dried for several days with MgSO_4 , filtered again with CaSO_4 . It was fractionally distilled before use. Smith, Fainberg and Winstein [*J Am Chem Soc* 83 618 1961, DOI: 10.1021/ja01464a029] washed it successively with aqueous NaHCO_3 , aqueous NaHSO_3 , water, 5% H_2SO_4 , water and dilute NaHCO_3 . The solvent was dried with CaSO_4 , then percolated through a column of Linde type 4A molecular sieves, followed by distillation from some of this material (in powdered form). Buffagni and Dunn [*J Chem Soc* 5105 1961, DOI: 10.1039/JR9610005105] refluxed it for 24 hours with activated charcoal while bubbling a stream of nitrogen through the liquid. The suspension was filtered, dried (Na_2SO_4) and distilled, then passed through an alumina column and redistilled. It has also been refluxed over CaH_2 , distilled and kept under argon over 4A molecular sieves.

It has been purified by zone melting at low temperature, or by distillation under a vacuum at 0°, subjecting the middle fraction to several freeze-pump-thaw cycles. An impure sample containing higher nitroalkanes and traces of cyanoalkanes was purified (on the basis of its NMR spectrum) by crystallisation from Et_2O at -60° (cooling in Dry-Ice) [Parrett & Sun *J Chem Educ* 54 448 1977, DOI: 10.1021/ed054p448].

Fractional crystallisation is more effective than fractional distillation from Drierite in purifying nitromethane for conductivity measurements. [Coetzee & Cunningham *J Am Chem Soc* 87 2529 1965, DOI: 10.1021/ja01090a001.] Specific conductivities around $5 \times 10^{-9} \text{ ohm}^{-1}\text{cm}^{-1}$ were obtained. [Beilstein 1 IV 100.]

1-Nitropropane [108-03-2] $\text{C}_3\text{H}_7\text{NO}_2$, M 89.1, b 131.4°/atm, d_4^{20} 1.004, n_D^{20} 1.40161, n_D^{25} 1.39936, pK^{25} 8.98. Purify it as for *nitromethane*. [Beilstein 1 IV 229.]

2-Nitropropane [79-46-9] $\text{C}_3\text{H}_7\text{NO}_2$, M 89.1, b 120.3°/atm, d_4^{20} 0.989, n_D^{20} 1.3949, n_D^{25} 1.39206, pK^{25} 7.68. Purify it as for *nitromethane*. [Beilstein 1 IV 230.]

N-Nitrosodiethanolamine (NDELA) [1116-54-7] $\text{C}_4\text{H}_{10}\text{N}_2\text{O}_3$, M 134.4, b 100°/2.6x10⁻⁵mm, 125°/0.01mm, n_D^{20} 1.4849. Purify NDELA by dissolving the amine (0.5g) in 1-propanol (10ml) and 5g of anhydrous Na_2SO_4 added with stirring. After standing for 1-2 hours, it is filtered and passed through a chromatographic column packed with 10ml of AG 50W x 8 (H^+ form 50-100mesh, a strongly acidic cation exchanger). The eluent and washings (50 ml EtOH) are combined and evaporated to dryness at 35°. It has also been extracted with EtOH

from the nitrosation mixture of ethanolamine, filtered and distilled under high vacuum. [Fukuda et al. *Anal Chem* **53** 2000 1981, DOI: 10.1021/ac00236a013; Jones & Wilson *J Chem Soc* 547 1949, DOI: 10.1039/JR9490000547; *Beilstein* **1** III 721, see Spiegelhalder et al. *N-Nitroso Compounds: Occurrence Biological Effects and Relevance in Human Cancer* (eds. O'Neill et al. IARC Scientific Publications No 57; IARC Lyon p943 1984.) *Possible CARCINOGEN*.

Nitrourea [556-89-8] $\text{CH}_3\text{N}_3\text{O}_3$, **M 105.1**, **m 158.4-158.8°(dec)**, **pK²⁰ 2.15**. Crystallise it from EtOH/petroleum ether. Dry it *in vacuo* ~50°. [Ingersoll & Armendt *Org Synth Coll Vol I* 417 1941, DOI: 10.15227/orgsyn.005.0085.]

n-Nonane [111-84-2] C_9H_{20} , **M 128.3**, **m -53°**, **b 150.8°/atm**, **d₄²⁰ 0.719**, **n_D²⁰ 1.40542**, **n_D²⁵ 1.40311**. Fractionally distil n-nonane, then stir it with successive volumes of concentrated H_2SO_4 for 12 hours each until no further coloration is observed in the acid layer. Then wash it with water, dry with MgSO_4 and fractionally distil it. *Alternatively*, it is purified by azeotropic distillation with 2-ethoxyethanol, followed by washing out the alcohol with water, drying and distilling it. [Forziati et al. *J Res Nat Bur Stand* **36** 129 1946, DOI: org/10.6028/jres.036.005; *Beilstein* **1** IV 447.]

Nylon powder. Pellets are purified by dissolving them in ethylene glycol under reflux, then precipitating nylon as a white powder by adding EtOH at 25°. This is washed with EtOH and dried at 100° under vacuum.

n-Octacosane [630-02-4] $\text{C}_{28}\text{H}_{58}$, **M 394.8**, **m 62.5°**, **b 278°/15mm**. Purify it by forming its adduct with urea, washing it and crystallising it from acetone/water. [McCubbin *Trans Faraday Soc* **58** 2307 1962, DOI: 10.1039/TF9625802307.] Crystallise it then from hot filtered isopropyl ether solution (10ml/g). [*Beilstein* **1** IV 588.]

n-Octacosanol (octacosyl alcohol) [557-61-9] $\text{C}_{28}\text{H}_{58}\text{O}$, **M 410.8**, **m 83.4°**, **84°**. It has been prepared from behenic acid (see docosanoic acid above). Recrystallise it from large volumes of Me_2CO . Sublime it at 200-250°/1mm instead of distilling it. It is insoluble in H_2O but soluble in organic solvents, fats, oils and CS_2 . [Bleyberg & Ulrich *Ber* **64** 2504 1931, DOI: 10.1002/cber.19310640929; *Beilstein* **2** IV 1318.]

n-Octadecane [593-45-3] $\text{C}_{18}\text{H}_{38}$, **M 254.5**, **m 28.1°**, **b 173.5°/10mm**, **316.1°/760mm**, **d₄²⁰ 0.7768**, **n_D²⁰ 1.4390**. Crystallise it from acetone and distil it from sodium in a vacuum. [*Beilstein* **1** IV 553.]

1-Octadecene (ODE) [112-88-9] $\text{C}_{18}\text{H}_{37}$, **M 252.5**, **m 95-98°**, **14-16°**, **17-18°**, **b 179°/15mm**, **b 315°/atm**, **d²⁵ 0.789g/ml**, **n_D²⁰ 1.444**. May contain n-octadecene, 2-octyl-1-decene, 2-butyl-1-tetradecene and/or 2-hexyl-1-dodecene. Best purified by fractional distillation through a 50cm column with 2cm diameter packed with 3/32-inch single turn glass helices [Swern et al. *J Am Chem Soc* **68** 1504 1946, DOI: 10.1021/ja01212a034]. [For the use of ODE in 'monolayers on Si(111) hydrogen-terminated surfaces: Effect of substrate doping' see Miramond & Vuillaume *J Appl Phys* **96** 1529 2004, DOI: 10.1063/1.1767984; ODE is used in the synthesis of colloidal quantum dots of high quality CdSe quantum dots, core/shell quantum dots and quantum rods see Asokan et al. *Nanotechnology* **16** 2000 2005, DOI: 10.1088/0957-4484/16/10/004. PMID 20817962; for general synthetic procedure see Lundeen & Hoozer *J Am Chem Soc* **85** 2180 1963, DOI: 10.1021/ja00897a041; *Beilstein* **1** IV 930.]

Octadecyl acetate [822-23-1] $\text{C}_{20}\text{H}_{40}\text{O}_2$, **M 312.5**, **m 32.6°**, **b 166-168°/1mm**, **172-174°/1.5mm**. Distil the ester under high vacuum, then crystallise it from $\text{Et}_2\text{O}/\text{MeOH}$, EtOH (**m 32.8°**) or Me_2CO (**m 32.4°**). Also recorded are **m 30.2°**; and **35.3°** for an α form as well as **38-39°** for a β form. [Phillips & Mumford *J Chem Soc* 1657 1934, DOI: 10.1039/JR9340001657; *Beilstein* **2** H 136, **2** II 147, **2** III 266, **2** IV 171.]

n-Octadecyl alcohol (stearyl alcohol, octadecanol) [112-92-5] $\text{C}_{18}\text{H}_{38}\text{O}$, **M 270.5**, **m 61°**, **b 153-154°/0.3mm**. Crystallise octadecanol from MeOH, or dry Et_2O and C_6H_6 , then fractionally distil it *in vacuo*. Also purify it by column chromatography. Free it from cetyl alcohol by zone refining. [*Beilstein* **1** IV 1888.]

***n*-Octadecylamine (1-aminooctadecane, stearylamine, ODA) [124-30-1] C₁₈H₃₉N, M 269.5, m 49-52°, 50-52°, 52-54°, b 183.0-183.1°/5mm, 232°/32mm, pK²⁵ 10.60.** *n*-Octadecylamine can be prepared from nonadecanoic acid (*n*-nonadecylic acid, [646-30-0]) by Schmidt's hydrazoic acid method whereby the carboxylic acid (15.8g, 53mmol) in *C₆H₆ (500ml) is treated carefully with concentrated H₂SO₄ (30ml), stirred vigorously at 40° and hydrazoic acid (52ml of a 5.3% solution in *C₆H₆, 1.2 equivalents, i.e. 63.3mmol; see [7782-79-8], **POISONOUS** use efficient fumehood) is added slowly. After evolution of N₂ and CO₂ have ceased (~2 hours) the acid layer is poured into cold H₂O whereby octadecylamine sulfate precipitates (~96% yield decomposing at ~200°). [see Briggs et al *J Chem Soc* 61 1942, DOI: 10.1039/JR9420000061.] The free base is obtained by shaking with 10% KOH solution, extracting into Et₂O, drying (K₂CO₃), evaporating and distilling the residue in a vacuum. Its FT-IR (film) has ν_{\max} at 2923.9, 1466.3, 792.4, 720.9 cm⁻¹; its ¹H NMR (300MHz, CDCl₃, TMS) has δ at 0.89 (t, *J* = 7 Hz, 3H, CH₃), 1.01 (s), 1.25 (s, 10H), 1.43 (m, 2H), 2.67 (t, *J* = ~7 Hz, 2H, CH₂-N); and the ¹³C NMR (75MHz, CDCl₃, CHCl₃ standard) has δ at 42.35, 33.99, 31.95, 29.72, 29.56, 29.39, 26.94, 22.71, 14.11. It has also been prepared by the Gabriel Synthesis whereby octadecanol is converted to octadecyl iodide, m 29-32° (by reaction with P₂O₅/KI), then with potassium phthalimide in DMF, to the *N*-*n*-octadecylphthalimide, m 80-81°, followed by hydrazinolysis with 85% NH₂NH₂·H₂O, and isolation as the **hydrochloride** [1838-08-0] **m 162-163°** after crystallisation from EtOH/Et₂O [Wood *J Chem Soc* 3327 1953, DOI: 10.1039/JR9530003313]. Another method of preparation include reduction of stearic amide with LAH in ~60% yield using a Soxhlet extractor with the amide in the thimble and LAH/Et₂O in the boiling flask [Murr & Lester *J Am Chem Soc* 77 1684 1955, DOI: 10.1021/ja01611a089]. The ***N*-acetyl derivative** has **m 84-85°** (EtOH/Et₂O), and the ***N*-methyl derivative** [2439-55-6] has **m 42-46°, b 155°/0.5mm**. [Beilstein 4 II 661, 4 III 431, 4 IV 825.] It is a strong base and should be stored in a dry CO₂-free atmosphere. The base and its derivatives are **skin and eye irritants**, and release histamine.

Octadecyl ether (dioctadecyl ether) [6297-03-6] C₃₆H₇₄O, M 523.0, m 59.4°, n_D⁶⁰ 1.440. Distil the ether in a vacuum, then crystallise it from MeOH/*C₆H₆, MeOH (m 58.5-59.5°) or Me₂CO (m 59.5°). It has an α form **m 57.8°** and a β form **m 40°**. [Beilstein 1 IV 1891.]

Octafluoropropane (profluorane) [76-19-7] C₃F₈, M 188.0, m -183°, b -36°/atm -38°/atm Purify it for pyrolysis studies by passage through a copper vessel containing CoF₃ at about 270°, then fractionally distil it. [Steunenbergh & Cady *J Am Chem Soc* 74 4165 1952, DOI: 10.1021/ja01136a062.] Also purify it by several trap-to-trap distillations at low temperatures. From the reaction of carbon and fluorine several fluorocarbons were obtained and separated by using a low temperature column packed with glass helices at 200mm, and the temperature remained constant within $\pm 1^\circ$. Each fraction was redistilled and their molecular weights were determined from the density balance. Thus obtained were C₃F₈ (188), C₄F₁₀ (234, 238), C₅F₁₂ (277, 278) and C₆F₁₄ (328, 228). The redistilled fractions had relatively sharp melting points. The **Critical Temperature of C₃F₈ is 343.7K**. [Simons & Block *J Am Chem Soc* 59 1407 1937, DOI: 10.1021/ja01286a512]. [Beilstein 1 III 219, 1 IV 189.]

***n*-Octane [111-65-9] C₈H₁₈, M 114.2, m -56.8°, b 19.2°/10mm, 125.6°/760mm, d₄²⁰ 0.704, n_D²⁰ 1.39743, n_D²⁵ 1.39505.** Extract the octane repeatedly with concentrated H₂SO₄ or chlorosulfonic acid, then wash it with water, dry and distil it. Alternatively, purify it by azeotropic distillation with EtOH, followed by washing with water to remove the EtOH, drying and distilling it. For further details, see *n*-heptane. It is also purified by zone melting at low temperatures. [Beilstein 1 H 159, 1 I 60, 1 II 122, 1 III 457, 1 IV 412.]

***RS*-Octane-1,2-diol [1117-86-8] C₈H₁₈O₂, M 146.2, m 30-30.5°, 36°, b 103-105°/0.5mm, 131-132°/10mm.** Distil the diol *in vacuo* and/or recrystallise it from petroleum ether. The **α -naphthylurethane** has **m 112-114°**. [Beilstein 1 III 2217, 1 IV 2590.] ***S*-(-)-Octane-1,2-diol** [87720-91-0] also crystallises from petroleum ether with **m 35-37°** and $[\alpha]_D^{17}$ -4.7 (c 35, EtOH) [Späth et al. *Chem Ber* 66 591 1933, DOI: 10.1002/cber.19330660430]; ***R*-(+)-octane-1,2-diol** [87720-90-9] has similar properties but with a positive optical rotation.

Octane-4,5-diol (dipropyl glycol) [110-98-5] C₈H₁₈O₂, M 134.2, b 109-110°/8mm (mostly *d,l*), d₄²⁰ 1.022, n_D²⁰ 1.441. Fractionally distil the diol below 15mm pressure, using a packed column and taking precautions to avoid absorption of water. [Beilstein 1 H 490, 1 II 556, 1 III 2219, 1 IV 2493.] Note that kilograms of a 1:1 mixture of *d,l*- and *meso*- dipropenylglycol (**b 113.4-114.4°/9mm**) can be obtained in *ca* 67% yield from *trans*-

crotonaldehyde by reaction with Zn—Cu couple in dilute acetic acid. When this glycol is reduced in 95% EtOH with a Pt catalyst (Adam's) it gives a 1:1 mixture of *d,l*- and *meso*- dipropyl glycol in >90% yield. After removal of Pt and evaporation of the solvent through a Hempel column, a pasty solid, m 20-90°, is obtained. Recrystallisation of the pasty solid from petroleum ether gives **meso-dipropyl glycol m 123.5-124.5°**, whose **bis-3,5-dinitrobenzoate** (prepared in dry pyridine, precipitation in the usual way, dissolving the crude ester in warm nitrobenzene and adding EtOH) has **m 200.3-200.4°**. The **bis-phenylurethane** has **m 191.2°**. Evaporation of the petroleum ether filtrate and repeated fractional distillation (using a 90 cm Claisen-Head column packed with glass rings and fitted with a Hopkins condenser) gives pure ***d,l*-dipropyl glycol m 28°, b 109.8-110°/9mm, n_D²⁵ 1.4419**, whose **bis-3,5-dinitrobenzoate** has **m 125-125.3°**. The **bis-phenylurethane** has **m 133.3°**. Studies of the melting points *versus* molar composition of mixtures of *meso* and *d,l* forms displayed a *eutectic* mixture of 5% *meso*- and 95% of *d,l*- dipropyl glycol at 20.0-20.5 °. Also the melting points of the *bis*-3,4-dinitrobenzoates of mixtures of these are indicative of their ratios [Young et al. *J Am Chem Soc* **58** 2274 1936, DOI: 10.1021/ja01302a054]. Separation of *meso* and *d,l* forms can be achieved by chromatography in petroleum ether on Al₂O₃. [Beilstein **1 I** 490, **1 II** 556, **1 III** 2219, **1 IV** 2493.]

Octane-1,8-diol (octamethylene glycol) [629-41-4] C₈H₁₈O₂, M 146.2, m 59-61°, b 172°/20mm. Recrystallise the diol from EtOH and distil it in a vacuum. [Beilstein **1 IV** 2592.]

1-Octanethiol [111-88-6] C₈H₁₈S, M 146.3, b 86°/15mm, 197-200°/760mm, d₄²⁰ 0.8433, n_D²⁰ 1.4540, pK²⁵ 10.72(dilute *t*-BuOH). Pass the thiol through a column of alumina and work under N₂, or Argon. Distil it under N₂ and a vacuum. Store it under N₂, or Argon in the dark. [Fletcher *J Am Chem Soc* **68** 2726 1946, DOI: 10.1021/ja01216a505]. [Beilstein **1 III** 1710, **1 IV** 1767.]

1-Octene [111-66-0] C₈H₁₆, M 112.2, m -101°, b 121°/742mm, d₄²⁰ 0.716, n_D²⁰ 1.4087. Distil 1-octene under nitrogen from sodium which removes water and peroxides. Peroxides can also be removed by percolation through dried, acid washed, alumina. Store it under N₂, or Argon in the dark. [Strukul & Michelin *J Am Chem Soc* **107** 7563 1985, DOI: 10.1021/ja00311a056; Beilstein **1 H** 221, **1 II** 199, **1 IV** 874.]

***trans*-2-Octene [13389-42-9] C₈H₁₆, M 112.2, b 124-124.5°/760mm, d₄²⁰ 0.722, n_D²⁰ 1.4132.** Purify it as for 1-octene above. [Beilstein **1 IV** 879.]

***n*-Octyl alcohol [111-87-5] C₈H₁₈O, M 130.2, m -15°, b 98°/19mm, 195.3°/760mm, d₄²⁰ 0.828, n_D²⁰ 1.43018.** Fractionally distil it under reduced pressure. Dry it with sodium and again fractionally distil or reflux with boric anhydride and re-distilled (b 195-205°/5mm), the distillate being neutralised with NaOH and again fractionally distilled. Also purify it by distillation from Raney nickel and by preparative GLC. [Beilstein **1 IV** 1756.]

***n*-Octylamine (1-aminooctane, caprylamine) [111-86-4] C₈H₁₉N, M 129.2, m -5° to -1°, b 62-64°/11mm 175-177°/745mm, 185-187°/atm, d₄²⁰ 0.7819, d₄²⁵ 0.728, n_D²⁰ 1.4292, pK²⁵ 10.57.** *n*-Octylamine has been prepared from nonanoic acid (pelargonic acid, see [112-05-0]) by Schmidt's hydrazoic acid method as described for octadecylamine above. [Adamson & Kenner *J Chem Soc* 838 1934, DOI: 10.1039/JR9340000838.] The syntheses described below for octadecyl amine can be used for preparing octylamine. A more recent preparation involves formation of tri-*n*-octylborane from 1-octene and BH₃.THF *in situ*, then reaction with NaN₃/HCl to give a 79% yield of octylamine [Kabalka et al. *Organometallics* **6** 1369 1987, DOI: 10.1021/om00149a046]. The free base can be isolated from its salts or the picrate by treatment with strong aqueous alkaline solution, extraction into Et₂O or CH₂Cl₂, drying (K₂CO₃), evaporating and distilling the residue preferably in a vacuum. Solubility in H₂O is ~2%. It is a strong base which readily absorbs CO₂ in moist air and should be stored under N₂. Its FT-IR (film) has ν_{max} at 3371.9, 2924.6, 1617.3, 1467.0, 1378.3, 1072.7, 822.1, 722.9 cm⁻¹; its ¹H NMR (300MHz, CDCl₃, TMS) has δ at 0.89 (t, *J* = ~7 Hz, 3H, CH₃), 1.15 (s), 1.29 (s, 6CH₂), 1.42 (t, *J* = ~7 Hz, NH₂), 2.69 (t, *J* = ~7 Hz, 2H, CH₂-N) ppm; and the ¹³C NMR (300MHz, CDCl₃, CHCl₃ standard) has δ at 42.32, 33.96, 31.88, 29.51, 29.34, 26.94, 22.69, 14.10 ppm. The **picrate** [78498-55-2] crystallises as plates with **m 111.5-112.5°**, the ***N*-acetyl derivative** [7462-62-6] has **b 148-149°/3mm**, and the ***N*-methyl derivative** [2439-54-5] has **b 60-65°/3mm, 75-78°/13mm**. [Beilstein **4 H** 196, **4 I** 386, **4 II** 655, **4 III** 379, **4 IV** 751.] *n*-Octylamine is quite a strong primary base capable of forming a carbamate salt of the type RNHCO₂⁻ RNH₃⁺ in the presence of CO₂ and H₂O (see storage above). The free base and its derivatives are **skin and eye irritants**.

***n*-Octylammonium hexadecanoate** [88020-97-7] $C_{24}H_{51}NO_2$, M 385.7, m 52-53°. Purify it by several recrystallisations from *n*-hexane or ethyl acetate. The solid is then washed with cold anhydrous diethyl ether, and dried *in vacuo* over P_2O_5 . [Beilstein 4 IV 751 for octylamine.]

***n*-Octylammonium octadecanoate** [32580-92-0] $C_{26}H_{55}NO_2$, M 413.7, m 56-57°. Purify it as for the hexadecanoate above.

***n*-Octylammonium tetradecanoate** [17463-35-3] $C_{22}H_{47}NO_2$, M 357.6, m 46-48°. Purify it as the hexadecanoate above.

***n*-Octyl bromide** [111-83-1] $C_8H_{17}Br$, M 193.1, m -55°, b 201.5°/atm, d_4^{20} 1.118, n_D^{25} 1.4503. Shake the bromide with H_2SO_4 , wash it with water, dry with K_2CO_3 and fractionally distil it. [Beilstein 1 IV 422.]

1-Octyne [629-05-0] C_8H_{14} , M 110.2, m -80°, b 76-77°/150mm, 126.2°/760mm, d_4^{20} 0.717, n_D^{25} 1.4159. Distil 1-octyne from $NaBH_4$ to remove peroxides. Fractionate it through a 10inch Widmer column at 125-126°/759mm. [Sletzing & Dawson *J Org Chem* 14 849 1949, DOI: 10.1021/jo01157a018.] [Beilstein 1 III 1005, 1 IV 1034.]

Oleic acid (OLA, *cis*-9-octadecenoic acid, olainic acid) [112-80-1] $C_{18}H_{34}O_2$, M 282.5, m 13-14°, 16°, b 145°/0.1mm, 194-195°/1.2mm, 228-229°/15mm, 360°(dec), d_4^{20} 0.891, n_D^{30} 1.4571, pK^{25} 6.42 (50% aqueous EtOH), pK_{Est} ~4.8 (H_2O). Purify the acid by fractional crystallisation from its melt, followed by molecular distillation at 10^{-3} mm, or by conversion to its methyl ester, the free acid can be crystallised from acetone at -40° to -45° (12ml/g). For purification by the use of lead and lithium salts see Keffler and McLean [*J Soc Chem Ind* 54 176T 1935, DOI: 10.1002/jctb.5000542514]. Purification based on direct crystallisation from acetone is described by Brown and Shinowara [*J Am Chem Soc* 59 6 1937, DOI: 10.1021/ja01280a003; pK White *J Am Chem Soc* 72 1859 1950, DOI: 10.1021/ja01160a530]. [Purification *via* urea adduct see Rubin & Paisley: *Biochem Preparations* 9 113 1962, Synthesis: Robinson & Robinson *J Chem Soc* 127 175 1925, DOI: 10.1039/CT9252700175; Beilstein 2 H 463, 2 I 198, 2 II 429, 2 III 1387, 2 IV 1641.]

Oleyl alcohol [143-28-2] $C_{18}H_{36}O$, M 268.5, m 1-5°, b 182-184°/1.5mm, 207°/13mm, d_4^{20} 0.847, $n_D^{27.5}$ 1.4582. Purify it by fractional crystallisation at -40° from acetone, then distil it under vacuum. [Beilstein 2 IV 2204.]

Oxalic acid ($2H_2O$) [6153-56-6] $C_2H_2O_4$, M 90.0, m 101.5°, [anhydrous 144-62-7] m 189.5°, pK_1^{25} 1.08 (1.37), pK_2^{25} 3.55 (3.80). Crystallise oxalic acid from distilled water. Dry it in a vacuum over H_2SO_4 . The anhydrous acid can be obtained by drying at 100° overnight. [Beilstein 2 IV 1819.]

Oxaloacetic acid [328-42-7] $C_4H_4O_5$, M 132.1, m 160°(decarboxylates), pK_1^{25} 2.22, pK_2^{25} 3.89, pK_3^{25} 13.0. Crystallise it from boiling EtOAc, or from hot Me_2CO /hot $*C_6H_6$. [Beilstein 3 IV 1808.]

2-Oxoglutaric acid (2-oxopentane-1,5-dioic, α -ketoglutaric acid) [328-50-7] $C_5H_6O_5$, M 146.1, m 114°, 115-117°, (pK_{Est} see oxaloacetic acid above). Crystallise the keto-acid repeatedly from Me_2CO /*benzene, EtOAc or ethyl propionate. Dry it *in vacuo*. [Beilstein 3 IV 1813.]

Oxamide [471-46-5] $C_2H_4N_2O_2$, M 88.1, m >320°(dec). Crystallise oxamide from water, grind it and dry it in an oven at 150°. [Beilstein 2 IV 1860.]

Palmitic acid anhydride (hexadecanoic anhydride) [623-65-4] $C_{32}H_{62}O_3$, M 494.9, m 63-64°, 64°, d_4^{82} 0.838, n_D^{68} 1.436. It is moisture sensitive and hydrolyses in water. Purify it by refluxing with acetic anhydride for 1 hour, evaporating and freeing the residue of acetic acid and anhydride by drying the residue at high vacuum and recrystallising from petroleum ether at low temperature. [Beilstein 2 IV 1181.]

Paraffin (oil) [8012-95-1] d_4^{20} 0.880, n_D^{20} 1.482. Treat the oil with fuming H_2SO_4 (care), then wash it with water and dilute aqueous NaOH, then percolate it through activated silica gel.

Paraffin Wax [8012-74-2] C_nH_{2n+2} , d_4^{20} 0.82, n_D^{20} 1.45 (extra light). A colourless (white), odourless wax from petroleum which is commercially available in fractions which solidify at various temperatures, e.g. with **solidification points at 44-46°** (in large white chunks), **50-52°** (white pellets), **54-56°** (white pellets), **58-60°** (white pellets), and **68-70°** (white pellets). Also available are waxes with **melting point ranges of 53-57°, 58-62°, ≥ 65°, and 70-80°**. Melt the wax in the presence of NaOH, wash it with water until all of the base had been removed. The paraffin is allowed to solidify after each wash. Finally, 5g of paraffin is melted by heating it on a water-bath, then shaken for 20-30 minutes with 100ml of boiling water and dry the melt under vacuum.

Pelargonic acid (nonanoic acid) [112-05-0] $C_9H_{18}O_2$, M 158.2, m 15°, b 98.9°/1mm, 225°/760mm, 268-269°/760mm, pK^{25} 4.96. Esterify the acid with ethylene glycol and distil the ester. (This removes dibasic acids as undistillable residues.) The acid is regenerated by hydrolysing the ester in the usual way and is distilled *in vacuo*. [Beilstein 2 IV 1018.]

Pelargononitrile (octyl cyanide) [2243-27-8] $C_9H_{17}N$, M 139.2, m -34°, b 92°/10mm, 224°, d_4^{20} 0.818, n_D^{20} 1.4255. Stir the nitrile with P_2O_5 (~5%), distil it from P_2O_5 and redistil it under a vacuum. IR should have CN but no OH bands. [Beilstein 2 IV 1204.]

Pelargonyl chloride (nonanoyl chloride) [764-85-2] $C_9H_{17}ClO$, M 176.7, b 88°/12mm, 108-110°/22mm, d_4^{20} 0.941, n_D^{20} 1.436. Reflux it with acetyl chloride (~3 volumes) for 1 hour, then distil off AcCl followed by the nonanoyl chloride at ~12mm. It is moisture sensitive and should be stored in sealed ampoules. [Beilstein 2 IV 1023.]

Pentabromoacetone [79-49-2] C_3HBr_5O , M 452.6, m 76°, pK^{25} 8.0 (MeOH), pK_{Est} ~4.6 (H_2O). Crystallise it from Et_2O , EtOH or aqueous EtOH (m 73.2°). Its solubility in H_2O is 0.01mg/100ml. It sublimes at its melting point. [Beilstein 1 H 659, 1 I 345, 1 III 2753, 1 IV 3226.]

Pentachloroethane (pentalin) [76-01-7] C_2HCl_5 , M 202.3, b 69°/37mm, 152.2°/64mm, 162.0°/~760mm, d_4^{20} 1.678, n^{15} 1.50542. Usual impurities include trichloroethylene. It partially decomposes if it is distilled at atmospheric pressure. Drying it with CaO, KOH or sodium is unsatisfactory because of the elimination of the elements of HCl. It can be purified by steam distillation, or by washing with concentrated H_2SO_4 , water, and then aqueous K_2CO_3 , drying with solid K_2CO_3 or $CaSO_4$, and fractionally distilling under reduced pressure. [Beilstein 2 IV 147.]

Pentadecafluoro octanoic acid (perfluorocaprylic acid) [335-67-1] $C_8HF_{14}O_2$, M 414.1, m 54.9-55.6°, b 189°/736mm, pK_{Est} <0. Crystallise the acid from CCl_4 and toluene, and distil it. It forms micelles in H_2O and the solubility is 1% in H_2O . The *acid chloride* has b 129-130°/744mm. The *amide* has m 138°. [Bernett & Zisman *J Phys Chem* 63 1911 1959, DOI: 10.1021/j150581a028; Bro & Sperati *J Polym Sci* 38 289 1959, DOI: 10.1002/pol.1959.1203813401; Beilstein 2 IV 994.]

Pentadecanoic acid [1002-84-2] $C_{15}H_{30}O_2$, M 242.4, m 51-53°, 80°, b 158°/1mm, 257°/100mm, d_4^{80} 0.8424, pK_{Est} ~5.0. Crystallise the acid from Et_2O and distil it *in vacuo*. It is very *hygroscopic*. See the purification of palmitic acid. [Beilstein 2 IV 1147.]

Pentadecanolide (1-oxacyclohexadecan-2-one, pentadecanoic- ω -lactone, 15-hydroxypentadecanoic lactone, exaltolide, Tibetolide) [106-02-5] $C_{15}H_{28}O_2$, M 240.4, m 34-36°, 37-37.5°, 37-38°, b 102-103°/0.03mm, 112-114°/0.2mm, 137°/2mm, 169°/10-11mm, d_4^{40} 0.9401. It has been recrystallised from MeOH (4 parts) at -15°, and distilled under high vacuum. [Hunsdiecker & Erlbach *Chem Ber* 80 129 1947, DOI: 10.1002/cber.19470800207; Galli & Mandolini *Org Synth* 58 98 1978, DOI: 10.15227/orgsyn.058.0098; Demole & Enggist *Helv Chim Acta* 11 2318 1978, DOI: 10.1002/hlca.19780610705; Beilstein 17/9 V 106.]

Penta-1,3-diene [*cis*: 1574-41-0, *trans*: 2004-70-8] C_8H_{14} , M 98.1, m -87°, b 42°/atm, d_4^{20} 0.680, n_D^{20} 1.4316. Distil the diene from $NaBH_4$. Purify it also by preparative gas chromatography. [Reimann et al. *J Am Chem Soc* 108 5527 1986, DOI: 10.1021/ja00278a026; Beilstein 1 IV 994.]

Penta-1,4-diene [591-93-5] C_5H_8 , M 68.1, m -1.5° , b 25.8-26.2°/756mm, d_4^{20} 0.645, n_D^{20} 1.3890. Distil it from $NaBH_4$. Purify it by preparative gas chromatography or distillation and stabilise it with 0.1% of 2,6-di-*tert*-butyl-*p*-cresol. [Reimann et al. *J Am Chem Soc* **108** 5527 1986, DOI: 10.1021/ja00278a026; *Beilstein* **1** IV 998.]

Pentaethylenhexamine (3,6,9,12-tetraazatetradecane-1,4-diamine) [4067-16-7] $C_{10}H_{28}N_6$, M 232.4, d_4^{20} 0.950, n_D^{20} 1.510, pK_1 1.2, pK_2 2.7, pK_3 4.3, pK_4 7.8, pK_5 9.1, pK_6 9.9 (all estimated). Fractionally distil it twice at 10-20mm, the fraction boiling at 220-250° being collected. It can be further purified *via* the *hydrochloride*. Its solution in MeOH (40ml of base in 250ml) is cooled in an ice-bath and concentrated HCl (~50ml) is added dropwise with stirring. The precipitated *hydrochloride* is filtered off, washed with Me_2CO , and Et_2O , then dried in a vacuum desiccator. The free base is then obtained by basification, extraction into Et_2O , drying (NaOH), filtering, evaporating and distilling the residue. It forms a Cu complex $[Cu(C_{10}H_{28}N_6)]^{2+}$. [Jonassen et al. *J Am Chem Soc* **79** 4279 1957, DOI: 10.1021/ja01573a011; *Beilstein* **4** IV 1245.]

2,2,3,3,3-Pentafluoropropan-1-ol (PFPOH) [422-05-9] $C_3H_3F_5O$, M 150.1, b 80°/~760mm, d_4^{20} 1.507, n_D^{20} 1.288, pK^{25} 12.74. Shake the alcohol with alumina for 24 hours, dry with anhydrous K_2CO_3 , and distil it, collect the middle fraction (b 80-81°) and redistil it. [*Beilstein* **1** IV 1438.]

***n*-Pentane** [109-66-0] C_5H_{12} , M 72.2, m -130° , b 36.1°/atm, d_4^{20} 0.626, n_D^{25} 1.35472. Stir the pentane with successive portions of concentrated H_2SO_4 until there is no further coloration during 12 hours, then with 0.5N $KMnO_4$ in 3M H_2SO_4 for 12 hours, wash with water and aqueous $NaHCO_3$. Dry it with $MgSO_4$ or Na_2SO_4 , then P_2O_5 and fractionally distil it through a column packed with glass helices. It is also purified by passage through a column of silica gel, followed by distillation and storage with sodium hydride. An alternative purification is by azeotropic distillation with MeOH, which is subsequently washed out from the distillate (using water), followed by drying and re-distilling. For removal of carbonyl-containing impurities, see *n-heptane*. Also purify it by fractional freezing (*ca* 40%) on a copper coil through which cold air is passed, then wash with concentrated H_2SO_4 and fractionally distil it. [*Beilstein* **1** IV 303.]

Pentane-1-thiol [110-66-7] $C_5H_{12}S$, M 104.2, m -76° , b 122.9°/697.5mm, d_4^{25} 0.8375, pK_{Est} ~10.1. Dissolve the thiol in aqueous 20% NaOH, then extract with a small amount of diethyl ether. The aqueous solution is acidified slightly with 15% H_2SO_4 , and the thiol is distilled out, dried with $CaSO_4$ or $CaCl_2$, and fractionally distilled under nitrogen. [Ellis & Reid *J Am Chem Soc* **54** 1674 1932, DOI: 10.1021/ja01343a067; *Beilstein* **1** IV 1453.]

***RS*-Pentan-2-ol (*sec*-amyl alcohol)** [13403-73-1, 6032-29-7 *non-specific*] $C_5H_{12}O$, M 88.2, b 73-73.5°/735mm, 118-118.5°/749mm, 119.9°/~760mm, d_4^{20} 0.810, n_D^{20} 1.41787, n_D^{25} 1.4052. Reflux it with CaO, distil it, then reflux it with magnesium and again fractionally distil it. Its solubility in H_2O is 16.6% w/v at 20°. The *acetate*, [626-38-0] $C_7H_{14}O_2$, M 130.1, has b 133.5°/atm, and the *1-naphthylurethane* [111994-17-1] crystallises from petroleum ether with m 76° . [Brown & Nakagawa *J Am Chem Soc* **77** 3614 1955, DOI: 10.1021/ja01618a056; Brown & Wheeler *J Am Chem Soc* **78** 2199 1956, DOI: 10.1021/ja01591a049; *Beilstein* **1** IV 1655.]

***R*(-)-Pentan-2-ol** [31087-44-2] $C_5H_{12}O$, M 88.2, b 119-120°/~760mm, d_4^{20} 0.809, n_D^{20} 1.409, $[\alpha]_D^{20}$ -16.1 (neat), -14.6 (c 1.2, EtOH), -14.3 (c 4.8, pentane). Reflux it with CaO, distil it, then reflux it with magnesium and again fractionally distil it. The *R-acetate*, [54638-10-7] $C_7H_{14}O_2$, M 130.1, has b 133°/atm, $[\alpha]_D^{20}$ -17.6 (neat). The *R-1-naphthylurethane* [120055-25-6] crystallises from aqueous EtOH with m 71-73° and has $[\alpha]_D^{20}$ +12.8 (EtOH). [For optical resolution see Jephcote et al. *JCS Perkin Trans I* 1529 1989, DOI: 10.1039/P19890001529; and for the absolute configuration see Lemieux et al. *Can J Chem* **29** 678 1951, DOI: 10.1139/v51-076.] [*Beilstein* **1** II 420.]

***S*(+)-Pentan-2-ol** [26184-62-3] $C_5H_{12}O$, M 88.2, b 118-119°/~760mm, d_4^{20} 0.81, n_D^{20} 1.406, $[\alpha]_D^{25}$ +13 (neat), $[\alpha]_D^{20}$ +13.5 (c 1.3, EtOH). Reflux it with CaO, distil it, then reflux it with magnesium and again fractionally distil it. The *S-acetate*, [5562-90-4] $C_7H_{14}O_2$, M 130.1, has b 130-131°/atm, $[\alpha]_D^{20}$ +17.6 (neat or petroleum ether). The *S-1-naphthylurethane* [120055-35-8] crystallises from aqueous EtOH with m 88-91° and has $[\alpha]_D^{20}$ +13.3 (EtOH). [*Beilstein* **1** IV 1656.]

Pentan-3-ol (diethyl carbinol) [584-02-1] $C_5H_{12}O$, M 88.2, b 113.5-113.7°/738mm, 114-115°/749mm, 116.2°/atm, d_4^{20} 0.819, n_D^{25} 1.4097. Reflux the alcohol with CaO, distil, then reflux it with magnesium and again fractionally distil it. [Brown & Nakagawa *J Am Chem Soc* **77** 3614 1955, DOI: 10.1021/ja01618a056; Brown & Wheeler *J Am Chem Soc* **78** 2199 1956, DOI: 10.1021/ja01591a049; *Beilstein* **1** IV 1662.]

Pentan-3-one see **diethyl ketone** above.

1-Pentene [109-67-1] C_5H_{10} , M 70.1, b 29.9-30.1°/atm, d_4^{25} 0.641, n_D^{20} 1.371. Reflux the mixture with sodium wire, then fractionally distil it twice through a Fenske (glass helices packing) column. [*Beilstein* **1** IV 808.]

Pent-2-ene (mixed isomers) [109-68-2] C_5H_{10} , M 70.1, b 36.4°/atm, d_4^{20} 0.650, n_D^{20} 1.38003, n_D^{25} 1.3839. Reflux the mixture with sodium wire, then fractionally distil it twice through a Fenske (glass helices packing) column. [Norris & Joubert *J Am Chem Soc* **49** 873 1927, DOI: 10.1021/ja01402a032; *Beilstein* **1** IV 815.]

cis-Pent-2-ene [627-20-3] C_5H_{10} , M 70.1, m -180°, b 37.1°/atm, d_4^{20} 0.657, n_D^{25} 1.3798. Prepared by dehydration of 2-pentanol. [Norris *Org Synth Coll Vol I* 430 1941, DOI: 10.1522/orgsyn.007.0076.] Dry it with sodium wire and fractionally distil it, or purify it by azeotropic distillation with MeOH, followed by washing out the MeOH with water, drying and distilling. Also purify it by chromatography through silica gel and alumina [Klassen & Ross *J Phys Chem* **91** 3668 1987, DOI: 10.1021/j100297a041]. [Sherrill & Matlak *J Am Chem Soc* **59** 2134 1937, DOI: 10.1021/ja01290a014; Carr & Stücklen *J Am Chem Soc* **59** 2138 1937, DOI: 10.1021/ja01290a015; Lucas et al. *J Am Chem Soc* **63** 22 1941, DOI: 10.1021/ja01846a005; *Beilstein* **1** IV 814.]

trans-Pent-2-ene [646-04-8] C_5H_{10} , M 70.1, m -140°, b 36.5°/atm, d_4^{20} 0.6482, n_D^{20} 1.3793. It is treated as above and washed with water, dried over anhydrous Na_2CO_3 , and fractionally distilled. The middle cut is purified by two passes of fractional melting. [Sherrill & Matlak *J Am Chem Soc* **59** 2134 1937, DOI: 10.1021/ja01290a014; Carr & Stücklen *J Am Chem Soc* **59** 2138 1937, DOI: 10.1021/ja01290a015; Lucas et al. *J Am Chem Soc* **63** 22 1941, DOI: 10.1021/ja01846a005; *Beilstein* **1** IV 814.]

Pent-1-yne [627-19-0] C_5H_8 , M 68.1, m -106° to -105°, b 40°/760mm, d_4^{25} 0.691, n_D^{20} 1.385. It is stood with, then distilled at low pressure from sodium or $NaBH_4$. [*Beilstein* **1** III 958, **1** IV 990.]

Pent-2-yne [627-21-4] C_5H_8 , M 68.1, m -109°, b 26°/2.4mm, 56.1°/760mm, d_4^{20} 0.710, n_D^{25} 1.4005. It is stood with, then distilled at low pressure from sodium or $NaBH_4$. [*Beilstein* **1** III 958, **1** IV 992.]

Perfluorobutyric acid (heptafluorobutyric acid, HFBA) [375-22-4] $C_4HF_7O_2$, M 214.0, m -17.5°, b 120°/755mm, d_4^{20} 1.651, n_D^{16} 1.295, pK^{25} -0.17. Fractionally distil the acid twice in an Oldershaw column with an automatic vapour-dividing head, the first distillation being in the presence of concentrated H_2SO_4 as a drying agent. (Take care with the hot acid.) [*Beilstein* **2** IV 810.]

Perfluoroheptane (hexadecafluoroheptane) [335-57-9] C_7F_{16} , M 388.1, b 82-84°/atm, 99-101°/atm, d_4^{25} 1.745, n_D^{20} 1.3. Purify it as for *perfluorodimethylhexane*. Other procedures include shaking with H_2SO_4 , washing with water, and drying with P_2O_5 for 48 hours then fractionally distilling. Alternatively, it has been refluxed for 24 hours with saturated acid $KMnO_4$ (to oxidise and remove hydrocarbons), then neutralised, steam distilled, dried with P_2O_5 , and passed slowly through a column of dry silica gel. It has been purified by fractional crystallisation using partial freezing. [*Beilstein* **1** IV 388.]

Perfluoro-n-hexane (tetradecafluorohexane) [355-42-0] C_6F_{14} , M 338.1, m -4°, b 58-60°/atm, d_4^{20} 1.684. Purify the fluorohexane by fractional freezing. The methods described for *perfluoroheptane* should be applicable here. [*Beilstein* **1** IV 348.]

Perfluorononane (eicosafluorononane) [375-96-2] C_9F_{20} , M 488.1, b 126-127°/atm, d_4^{20} 1.80, n_D^{20} 1.275. Purify as for *perfluorodimethylcyclohexane*. [*Beilstein* **1** III 505.]

Perfluoropropyl iodide (heptafluoro-1-iodopropane) [754-34-7] C_3F_7I , M 295.9, b 41°/atm, d_4^{20} 2.13, n_D^{20}

1.339. Purify the fluoro-iodide by fractional distillation. Store it over Cu as stabiliser. [*Beilstein* 1 IV 225.]

Perfluorotributylamine (heptacosafuorotributylamine, PFTBA) [311-89-7] $C_{12}F_{27}N$, M 671.1, b 177.6°/760mm, 178°/760mm, d_4^{20} 1.881, n_D^{20} 1.291, pK_{Est} ~5.0. Purify it as for perfluorodimethylcyclohexane (see [335-27-3]); see also perfluorotripropylamine below [*Haszeldine J Chem Soc* 102 1951, DOI: 10.1039/JR9510000102]. [*Beilstein* 2 IV 819.]

Perfluorotripropylamine (heneicosafuorotripropylamine FTPA) [338-83-0] $C_9F_{21}N$, M 521.1, b 129.5-130.5°/atm, 130°/atm, 129.5-130.5°/atm, d_4^{20} 1.822, n_D^{20} 1.279, pK_{Est} ~5.6. Purify it as for perfluorodimethylcyclohexane (see [335-27-3]) [*Haszeldine J Chem Soc* 102 1951, DOI: 10.1039/JR9510000102; for azeotropes see Simons & Linevsky *J Am Chem Soc* 74 4750 1972, DOI: 10.1021/ja01139a007.] **IRRITANT.**

Petroleum ether (ligroin) [8032-32-4] b 35-60°, d_4^{20} 0.640, n_D^{20} 1.363. Shake it several times with concentrated H_2SO_4 , then 10% H_2SO_4 and concentrated $KMnO_4$ (to remove unsaturated, including aromatic, hydrocarbons) until the permanganate colour persists. Wash it with water, aqueous Na_2CO_3 and again with water. Dry it with $CaCl_2$ or Na_2SO_4 , and distil it. It can be dried further using CaH_2 or sodium wire. Passage through a column of activated alumina, or treatment with CaH_2 or sodium, removes peroxides. For the elimination of carbonyl-containing impurities without using permanganate, see *n-heptane*. These procedures could be used for all fractions of petroleum ethers. See skellysolve below.

Rapid purification: Pass it through an alumina column and fractionally distilling, collecting the desired boiling fraction.

Phorone (2,6-dimethylhepta-2,5-dien-4-one) [504-20-1] $C_9H_{14}O$, M 138.2, m 28°, b 88°/17mm, 197°/743mm. Purify by fractional distillation or distillation and/or repeatedly recrystallise from EtOH. It can be converted to its **2,4-dinitrophenylhydrazone** by adding it to a moderate excess of 0.4% of 2,4-dinitrophenyl hydrazine in 2N-aqueous HCl which precipitates within 24 hours. The derivative can be washed, dried and recrystallised from EtOH, or chromatographed on alumina, eluting with benzene and recrystallising from MeOH to provide red crystals m 118-118.5°. [Braude & Timmons *J Chem Soc* 2000 1950, DOI: 10.1039/JR9500002000; Connolly *J Chem Soc* 338 1944, DOI: 10.1039/JR9440000338; Joseph & Blumenthal *J Org Chem* 24 1371 1959, DOI: 10.1021/jo01091a626; *Beilstein* 1 IV 3564.]

Pimelic acid (heptane-1,7-dioic acid) [111-16-0] $C_7H_{12}O_4$, M 160.2, m 105-106°, m 212°/10mm, pK_1^{25} 4.46, pK_2^{25} 5.58. Crystallise the acid from water or from *benzene containing 5% diethyl ether. [*Beilstein* 1 IV 2003.]

Pinacol (hexahydrate) [6091-58-3 ($6H_2O$), 76-09-5 (anhydrous)] $C_6H_{14}O_2 \cdot 6H_2O$, M 194.3, m 46.5°, b 59°/4mm. Distil pinacol, then crystallise it repeatedly from water. (See also below.) [*Beilstein* 1 IV 2575.]

Pinacol (anhydrous) (2,3-dimethyl-2,3-butanediol) [76-09-5] $C_6H_{14}O_2$, M 118.1, m 41.1°, 171-172°/739mm, b 172°/atm. The hydrate is rendered anhydrous by azeotropic distillation of water with *benzene. Recrystallise it from *benzene or toluene/petroleum ether, absolute EtOH or dry diethyl ether. It recrystallises from water to give the *hexahydrate*. [*Beilstein* 1 IV 2575.]

Pinacolone see *tert-butyl methyl ketone*.

Pinacolone oxime [2475-93-6] $C_6H_{13}NO$, M 115.2, m 75.5-76°, 78°, 78.5-79.5°, 171.6°/748mm. Crystallise the oxime from aqueous EtOH, EtOH (small needles) or petroleum ether (plates). [Markownikoff *Chem Ber* 32 1445 1899, DOI: 10.1002/cber.18990320221; Smith & Adkins *J Am Chem Soc* 60 657 1938, DOI: 10.1021/ja01270a048; Whitmore et al. *J Am Chem Soc* 61 683 1939, DOI: 10.1021/ja01872a043; *Beilstein* 1 H 694, 1 II 750, 1 III 2842, 1 IV 3310.]

Pivalic acid (trimethylacetic acid) [75-98-9] $C_5H_{10}O_2$, M 102.1, m 35.4°, b 71-73°/0.1mm, 163-164°/atm, pK^{25} 5.03. Fractionally distil the acid under reduced pressure, then fractionally crystallise it from its melt. Re-

crystallise it from *benzene. [*Beilstein* 2 IV 908.]

Pivaloyl chloride (trimethylacetyl chloride) [3282-30-2] $\text{C}_5\text{H}_9\text{ClO}$, M 120.6, b 57.6°/150mm, 70.5-71/250mm, 104°/754mm, 104-105°/atm, 105-108°/atm, d_4^{20} 1.003, n_D^{20} 1.4142. First check the IR to see if OH bands are present. If absent, or present in small amounts, then redistil it under a moderate vacuum. If present in large amounts then treat it with oxalyl chloride or thionyl chloride and reflux for 2-3 hours, evaporate and distil the residue. **Strongly LACHRYMATORY - work in a fume cupboard.** Store it in sealed ampoules under N_2 . [Traynham & Battiste *J Org Chem* 22 1551 1957, DOI: 10.1021/jo01363a004; for Grignard reactions: Whitmore et al. *J Am Chem Soc* 63 643 1941, DOI: 10.1021/ja01848a004; *Beilstein* 2 IV 912.]

Polyacrylonitrile [25014-41-9] $(\text{C}_3\text{H}_3\text{N})_n$, M 150,000 (average). Precipitate it from dimethylformamide by addition of MeOH.

Poly(diallyldimethylammonium) chloride [26062-79-3]. Precipitate it from water with acetone, and dry the salt in a vacuum for 24 hours. [Hardy & Shriver *J Am Chem Soc* 107 3823 1985, DOI: 10.1021/ja00299a012.]

Polyethylene [9002-88-4] $\text{H}(\text{CH}_2\text{-CH}_2)_n\text{H}$. It is available in several molecular weight averages (from M_w ~4000 to 6000.000) as well as in low, medium and high density packages. Crystallise it from thiophen-free *benzene and dry it over P_2O_5 under vacuum.

Polymethyl acrylate [9003-21-8]. Precipitate it from a 2% solution in acetone by addition of water.

Polyvinyl acetate [9003-20-7]. Precipitate it from acetone by addition of *n*-hexane.

Polyvinyl chloride [9002-81-2]. Precipitate it from cyclohexanone by addition of MeOH.

Propane [74-98-6] C_3H_8 , M 44.1, m -189.7°, b -42.1°/760mm, d_4^{20} 0.5005, n_D^{20} 1.2898. Purify propane by bromination of the olefinic contaminants. Propane is treated with bromine for 30 minutes at 0°. Unreacted bromine is quenched, and the propane is distilled through two -78° traps and collected at -196° [Skell et al. *J Am Chem Soc* 108 6300 1986, DOI: 10.1021/ja00280a030]. It autoignites at 450° and the flash point is -104°. It is highly **FLAMMABLE** and is available in metal cylinders. [*Beilstein* 1 H 103, 1 I 33, 1 II 71, 1 III 204, 1 IV 175.]

Propane-1,2-diamine (propylenediamine) [78-90-0] $\text{C}_3\text{H}_{10}\text{N}_2$, M 74.1, b 120.5°/~760mm, d_4^{20} 0.868, n_D^{20} 1.446, pK_1^{25} 6.61, pK_2^{25} 9.82. Purify this hygroscopic diamine by azeotropic distillation with toluene. Then redistil it. Store it in a CO_2 free atmosphere. **SKIN IRRITANT**. [Horton et al. *Anal Chem* 27 269 1955, DOI: 10.1021/ac60098a025; for optical resolutions *via* the *d*-tartrate salts or the Co(II) complexes see Dwyer et al. *J Am Chem Soc* 81 290 1959, DOI: 10.1021/ja01511a009, see also Smith et al. *J Am Chem Soc* 96 2908 1974, DOI: 10.1021/ja00816a041; *Beilstein* 4 IV 1255.]

***R*(-)-propane-1,2-diamine** [6852-78-4] $\text{C}_3\text{H}_{10}\text{N}_2$, has b 120.5°/~760mm, $[\alpha]_D^{25}$ -35 (c 0.97, * C_6H_6) and is purified by distillation. The optical rotation is inverted in the ***R*(+)-propane-1,2-diamine dihydrochloride** [19777-67-4] $\text{C}_3\text{H}_{10}\text{N}_2 \cdot 2\text{HCl}$, M 147.1, which has m 241-244°, $[\alpha]_D^{25}$ +4.0 (c 20, H_2O) [*Beilstein* 4 IV 1255].

***S*(+)-propane-1,2-diamine** [15967-73-3] $\text{C}_3\text{H}_{10}\text{N}_2$, has b 120.5°/~760mm, $[\alpha]_D^{25}$ +34.8 (c 0.97, * C_6H_6) and is purified by distillation, and the ***S*(-)-propane-1,2-diamine dihydrochloride** [19777-66-3] $\text{C}_3\text{H}_{10}\text{N}_2 \cdot 2\text{HCl}$, M 147.1, has m 227-229, $[\alpha]_D^{25}$ -4.0 (c 20, H_2O) [*Beilstein* 4 258, 4 IV 1255].

Propane-1,3-diamine (trimethylenediamine) [109-76-2] $\text{C}_3\text{H}_{10}\text{N}_2$, M 74.1, m -12°, b 140°/~760mm, d_4^{25} 0.888, n_D^{20} 1.458, pK_1^{25} 8.29, pK_2^{25} 10.30. Purify like the preceding diamine. [*Beilstein* 4 IV 1255.] The **dihydrochloride** [10517-44-9] $\text{C}_3\text{H}_{10}\text{N}_2 \cdot 2\text{HCl}$, M 147.1, has m 243°, 246-250°. [*Beilstein* 4 H 261.]

(±)-Propane-1,2-diol (propyleneglycol) [57-55-6] $\text{C}_3\text{H}_8\text{O}_2$, M 76.1, m -60°, b 45.5°/1.0mm, 70.8°/5.0mm, 83.2°/10mm, 96.4°/20mm, 104°/32mm, 111.2°/40mm, 119.9°/60mm, 132.0°/100mm, 149.7°/200mm, 168.1°/

400mm, 188.2°/atm, d_4^{20} 1.040, n_D^{20} 1.433. Dry the diol over Na_2SO_4 , decant and distil it under reduced pressure. It is soluble in H_2O , Me_2CO , CHCl_3 , many oils, is used as an antifreeze in dairies and breweries; and is a substitute for glycerol or ethylene glycol. [Beilstein 1 IV 2468.]

***R*-(*-*)- and *S*-(*+*)- Propane-1,2-diol (*R*- and *S*- propylene glycol) [*R*(*-*) 4254-14-2; *S*(*+*) 4254-15-3] $\text{C}_3\text{H}_8\text{O}_2$, M 76.1, b 78.5°/10mm, 94-96°/14mm, 186-188°/765mm, d_4^{20} 1.036, n_D^{20} 1.432, $[\alpha]_D^{25}$ (-) or (+) 17.5 (neat).** The laevo *R*-enantiomer can be obtained from reduction of hydroxyacetone by yeast [Levene & Walti *Org Synth Coll Vol* 2 545 1943, DOI: 10.15227/orgsyn.010.0084], and the dextro *S*-enantiomer is obtained by reduction of *S*-(*+*)-lactic acid. Thus (+)-lactic acid (33.4g, 370mmol) in dry THF (200ml) is added dropwise to a suspension of LAH (32.1g, 850mmol) in dry THF (500ml) at 0° during 2 hours, allowed to warm to 25°, refluxed for 2 hours, cooled, then quenched by careful addition of ice-cold H_2O (32ml), followed by 4N NaOH (32ml) and again H_2O (96ml). The white precipitate is coagulated by refluxing the mixture for 0.5 hours, filtered, washed with THF, the white cake is slurred with hot THF (4 x ~100ml), the combined THF layers are evaporated *in vacuo* (~40°), and the residual oil is distilled through a vacuum jacketed Vigreux column under vacuum to give the *S*-diol as a colourless liquid (23.9g, 85%). [Fryzuk & Bosnich *J Am Chem Soc* 100 5491 1978, DOI: 10.1021/ja00485a037; Beilstein 1 IV 2468.]

Propane-1,3-diol [504-63-2] $\text{C}_3\text{H}_8\text{O}_2$, M 76.1, m -27°, b 110-122°/12mm, d_4^{20} 1.053, $n_D^{18.5}$ 1.4398. Dry this diol with K_2CO_3 and distil it under reduced pressure. More extensive purification involves conversion with benzaldehyde to 2-phenyl-1,3-dioxane (m 47-48°) which is subsequently decomposed by shaking with 0.5M HCl (3ml/g) for 15 minutes and standing overnight at room temperature. After neutralisation with K_2CO_3 , the benzaldehyde is removed by steam distillation and the diol is recovered from the remaining aqueous solution by continuous extraction with CHCl_3 for 1 day. The extract is dried with K_2CO_3 , the CHCl_3 is evaporated and the diol is distilled. [Foster et al. *Tetrahedron* 16 177 1961, DOI: 10.1016/0040-4020(61)80068-9; Beilstein 1 IV 2493.] The **bis-*p*-tosylate [5469-66-9] $\text{C}_{12}\text{H}_{20}\text{O}_6\text{S}_2$, M 384.5**, prepared in the usual way (see below), has m 90-92° [Beilstein 11 II 49].

***S*-(*-*)-Propane-1,2-diol bis(*p*-toluenesulfonate) [60434-71-1] $\text{C}_{12}\text{H}_{20}\text{O}_6\text{S}_2$, M 384.5, m 68-70° (62°), $[\alpha]_D^{25}$ -20° (c 1, CHCl_3),** is prepared in much the same way as for *S,S*-butane-2,3-diol bis-tosylate (see above) from *S*-(*+*)-propane-1,2-diol (310mmol) in dry pyridine (30ml) and toluene-*p*-sulfonyl chloride (700mmol, recrystallised from hexane) in dry pyridine (135ml) at 0° during 0.5 hours, then at 25° for 17 hours. The crude ester is purified by dissolving in CH_2Cl_2 and cyclohexane is added to cloud point at 40°, then allowed to crystallise at 25° when more cyclohexane is added and set aside (5°/12 hours) to give the *bis*-tosylate as white feathery crystals which are dried over CaCl_2 *in vacuo* (114g, 95%). [Fryzuk & Bosnich *J Am Chem Soc* 100 5491 1978, DOI: 10.1021/ja00485a037.] This *S*-(*-*)-2,3-ditosylate provides the chiral ligand *R*-(*+*)- [cf. 15629-92-2], which involves inversion of configuration at the chiral centre.

Propane-1-thiol [107-03-9] $\text{C}_3\text{H}_8\text{S}$, M 76.1, m -113°, b 65.3°/702mm, d_4^{25} 0.83598, n_D^{25} 1.43511, pK^{20} 10.82. Purify the thiol by dissolving it in aqueous 20% NaOH, extracting with a small amount of *benzene and steam distilling until clear. After cooling, the solution is acidified slightly with 15% H_2SO_4 , and the thiol is distilled out, dried with anhydrous CaSO_4 or CaCl_2 , and fractionally distilled under nitrogen. [Mathias & Filho *J Phys Chem* 62 1427 1958, DOI: 10.1021/j150569a018.] Also purify it by liberating the mercaptan by adding dilute HCl to the residue remaining after steam distilling. After direct distillation from the flask, and separation of the water, the mercaptan is dried (Na_2SO_4) and distilled under nitrogen. [Beilstein 1 IV 1449.]

Propane-2-thiol (Isopropyl mercaptan) [75-33-2] $\text{C}_3\text{H}_8\text{S}$, M 76.1, m -131°, b 49.8°/696mm, d_4^{25} 0.80895, n_D^{25} 1.42154, pK^{25} 10.86. Purify it as for propane-1-thiol above. [Beilstein 1 IV 1498.]

Propargyl alcohol (2-propyn-1-ol) [107-19-7] $\text{C}_3\text{H}_4\text{O}$, M 56.1, m -53°, b 54°/57mm, 113.6°/760mm, d_4^{20} 0.947, n_D^{20} 1.432. The commercial material contains a stabiliser. An aqueous solution of propargyl alcohol can be concentrated by azeotropic distillation with butanol or butyl acetate. Dry it with K_2CO_3 and distil it under reduced pressure, in the presence of about 1% succinic acid, through a glass helices-packed column. [Beilstein 1 IV 2214.]

Propargyl chloride (3-chloropropyne) [624-65-7] $\text{C}_3\text{H}_5\text{Cl}$, **M 74.5**, **b 58°/760mm**, **65°/760mm**, d_4^{20} **1.03**, n_D^{20} **1.435**. Purify the chloride by fractional distillation at atmospheric pressure. Note that a possible impurity is **propargyl alcohol** which has **b 114-115°/~760mm** (see above). [Henry *Chem Ber* **8** 398 1875, DOI: 10.1002/cber.187500801129.] **HIGHLY TOXIC** and **FLAMMABLE**. [Beilstein **1** IV 963.]

Propene (propylene) [115-07-1] C_3H_6 , **M 42.1**, **m -185.2°**, **b -47.8°/750mm**, d_4^{20} **0.519**, n^{71} **1.357**. Purify it by freeze-pump-thaw cycles and trap-to-trap distillation. [Beilstein **1** IV 725.]

β -Propiolactone see oxetan-2-one in ‘Heterocyclic Compounds’, in this Chapter.

Propionaldehyde (propanal) [123-38-6] $\text{C}_3\text{H}_6\text{O}$, **M 58.1**, **m -81°**, **b 48.5-48.7°/atm**, d_4^{20} **0.804**, n_D^{20} **1.3733**, n_D^{25} **1.37115**. Dry the aldehyde with CaSO_4 or CaCl_2 , and fractionally distil it under nitrogen or in the presence of a trace of hydroquinone (to retard oxidation). Blacet and Pitts [*J Am Chem Soc* **74** 3382 1952, DOI: 10.1021/ja01133a049] repeatedly distilled the middle fraction in a vacuum until it no longer gave a solid polymer when cooled to -80°. It is stored with CaSO_4 . **IRRITANT**. The *oxime* [627-39-4] $\text{C}_3\text{H}_7\text{NO}$, **M 73.0**, distils at **77°/100mm**, **130-131°/atm**, the distillate solidifies and the solid crystallises from EtOH with **m 40°**. It is a colorimetric reagent for Pd(II). [Beilstein **1** H 629, **1** I 333, **1** II 687, **1** III 1682, **1** IV 3165.] The **2,4-dinitrophenylhydrazone** [725-00-8] has **m 142-148°**. [Hurd & Meinert *Org Synth Coll Vol* **2** 541 1943, DOI: 10.1021/orgsyn.012.0064.] The **cyanohydrin (3-hydroxybutyronitrile)** [4476-02-2] $\text{C}_4\text{H}_7\text{NO}$, **M 85.1** has **b 62-64°/2mm**, **96-98°/15mm**, **105°/20mm**, d_4^{20} **0.962**, n_D^{20} **1.414**, and is purified by fractional distillation, taking precautions as some HCN (**POISONOUS**) may be liberated (work in an efficient fume cupboard). The *cyanohydrin* forms a **2,4-dinitrobenzoate** with **m 83-84°**. [Stevens et al. *J Am Chem Soc* **76** 2695 1954, DOI: 10.1021/ja01639a029; Beilstein **3** H 305, **3** II 217, **3** III 564, **3** IV 757.]

Propionamide [79-05-0] $\text{C}_3\text{H}_7\text{NO}$, **M 73.1**, **m 79.8-80.8°**, pK^{24} **-0.9** (H_0 scale, aqueous H_2SO_4). Crystallise it from acetone, *benzene, CHCl_3 , water or acetone/water, then dry it in a vacuum desiccator over P_2O_5 or concentrated H_2SO_4 . [Beilstein **2** H 243, **2** I 108, **2** II 223, **2** III 542, **2** IV 725.]

Propionic acid [79-09-4] $\text{C}_3\text{H}_6\text{O}_2$, **M 74.1**, **m -24° to -23°**, **b 141°/atm**, d_4^{20} **0.992**, n_D^{20} **1.3865**, n_D^{25} **1.3843**, pK_1^{25} **-6.8** (H_0 scale, aqueous H_2SO_4), pK_2^{25} **4.88**. Dry the acid with Na_2SO_4 or by fractional distillation, then redistil after refluxing with a few crystals of KMnO_4 . An alternative purification uses conversion to the ethyl ester, fractional distillation and hydrolysis. [Bradbury *J Am Chem Soc* **74** 2709 1952, DOI: 10.1021/ja01131a005]. Propionic acid can also be heated for 0.5 hour with an amount of benzoic anhydride equivalent to the amount of water present (in the presence of CrO_3 as catalyst), followed by fractional distillation. [Chan & Israel *J Chem Soc* 196 1960, DOI: 10.1039/JR9600000196; Beilstein **2** IV 695.]

Propionic anhydride [123-62-6] $\text{C}_6\text{H}_{10}\text{O}_3$, **M 130.2**, **m -43°**, **b 67°/18mm**, **168°/780mm**, d_4^{20} **1.407**, n_D^{20} **1.012**. Shake the anhydride with P_2O_5 for several minutes, then distil. [Beilstein **2** IV 722.]

Propionitrile (ethyl cyanide) [107-12-0] $\text{C}_3\text{H}_5\text{N}$, **M 55.1**, **m -93°**, **b 97.2°/atm**, d_4^{20} **1.407**, n^{15} **1.36812**, n^{30} **1.36132**. Shake the nitrile with dilute HCl (20%), or with concentrated HCl until the odour of isonitrile has gone, then wash with water, and aqueous K_2CO_3 . After a preliminary drying with silica gel or Linde type 4A molecular sieves, it is stirred with CaH_2 until hydrogen evolution ceases, then decant and distil from P_2O_5 (not more than 5g/L, to minimise gel formation). Finally, it is refluxed with, and slowly distilled from CaH_2 (5g/L), taking precautions to exclude moisture. [Beilstein **2** IV 728.]

***n*-Propyl acetate** [109-60-4] $\text{C}_5\text{H}_{10}\text{O}_2$, **M 102.1**, **m -95°**, **b 101.5°/atm**, d_4^{20} **0.887**, n_D^{20} **1.38442**, pK^{25} **-7.18** (H_0 scale, aqueous H_2SO_4). Wash the ester with saturated aqueous NaHCO_3 until neutral, then with saturated aqueous NaCl . Dry it with MgSO_4 and fractionally distil it. [Beilstein **2** IV 138.]

***n*-Propyl alcohol (1-propanol)** [71-23-8] $\text{C}_3\text{H}_8\text{O}$, **M 60.1**, **m -127°**, **b 97.2°/atm**, d_4^{25} **0.79995**, n_D^{20} **1.385**, pK^{25} **16.1**. The main impurities in *n*-propyl alcohol are usually water and 2-propen-1-ol, reflecting the commercial production by hydration of propene. Water can be removed by azeotropic distillation either directly (azeotrope contains 28% water) or by using a ternary system, e.g. by also adding *benzene. Alternatively, for

removal of gross amounts of water, reflux over CaO for several hours is desirable, followed by distillation and a further drying. To obtain more nearly anhydrous alcohol, suitable drying agents are firstly NaOH, CaSO₄ or K₂CO₃, then CaH₂, aluminium amalgam, magnesium activated with iodine, or a small amount of sodium. *Alternatively*, the alcohol can be refluxed with *n*-propylsuccinate or phthalate in a method similar to the one described under EtOH. Allyl alcohol is removed by adding bromine (15ml/L), and then fractionally distilling from a small amount of K₂CO₃. Propionaldehyde, also formed in the bromination, is removed as the 2,4-dinitrophenylhydrazone. *n*-Propyl alcohol can be dried down to 20 ppm of water by passage through a column of pre-dried molecular sieves (type 3 or 4A, heated for 3 hours at 300°) in a current of nitrogen. Distillation from sulfanilic or tartaric acids removes impurities.

Albrecht [*J Am Chem Soc* **82** 3813 1960, DOI: 10.1021/ja01500a004] obtained spectroscopically pure material by heating with charcoal to 50-60°, filtering and adding 2,4-dinitrophenylhydrazine and a few drops of concentrated H₂SO₄. After standing for several hours, the mixture is cooled to 0°, filtered and distilled in a vacuum. Gold and Satchell [*J Chem Soc* 1938 1963, DOI: 10.1039/JR9630001938] heated *n*-propyl alcohol with 3-nitrophthalic anhydride at 76-110° for 15 hours, then recrystallised the resulting ester from H₂O, *benzene/petroleum ether (b 100-120°)(3:1), and *benzene. The ester was hydrolysed under reflux with aqueous 7.5M NaOH for 45 minutes under nitrogen, followed by distillation (also under nitrogen). The fraction with **b** 87-92° is dried with K₂CO₃ and stirred under reduced pressure in the dark over 2,4-dinitrophenylhydrazine, then freshly distilled. Also purify it by adding 2g NaBH₄ to 1.5L of alcohol, gently flushing with argon and refluxing for 1 day at 50°. Then 2g of freshly cut sodium (washed with propanol) is added and refluxed for one day, and finally distilled, taking the middle fraction [Jou & Freeman *J Phys Chem* **81** 909 1977, DOI: 10.1021/j100524a021]. [*Beilstein* **1** IV 1413.]

***n*-Propylamine** [107-10-8] C₃H₉N, **M 59.1**, **m -83°**, **b 48.5°/atm**, **d₄²⁰ 0.716**, **n_D²⁰ 1.38815**, **pK²⁵ 10.69**. Distil the amine from zinc dust, under reduced pressure, in an atmosphere of nitrogen. [*Beilstein* **4** IV 464.]

***n*-Propyl bromide** [106-94-5] C₃H₇Br, **M 123.0**, **m -110°**, **b 71.0°/atm**, **d₄²⁰ 1.354**, **n_D¹⁵ 1.43695**, **n_D²⁵ 1.43123**. Likely contaminants include *n*-propyl alcohol and isopropyl bromide. The simplest purification procedure uses drying with MgSO₄ or CaCl₂ (with or without a preliminary wash of the bromide with aqueous NaHCO₃, then water), followed by fractional distillation away from bright light. Chien and Willard [*J Am Chem Soc* **79** 4872 1957, DOI: 10.1021/ja01575a013] bubbled a stream of oxygen containing 5% ozone through *n*-propyl bromide for 1 hour, then stirred it with 3% hydrogen peroxide solution, neutralised it with aqueous Na₂CO₃, washed it with distilled water and dried it. This was followed by vigorous stirring with 95% H₂SO₄ until fresh acid did not discolour within 12 hours. The propyl bromide was separated, neutralised, washed, dried with MgSO₄ and fractionally distilled. The centre cut was stored in the dark. Instead of ozone, Schuler and McCauley [*J Am Chem Soc* **79** 821 1957, DOI: 10.1021/ja01561a010] added bromine and stored it for 4 weeks, the bromine then being extracted with aqueous NaHSO₃ before the sulfuric acid treatment was applied and finally distilled. Further purification is by preparative gas chromatography on a column packed with 30% SE-30 (General Electric ethylsilicone rubber) on 42/60 Chromosorb P at 150° and 40psi, using helium as carrier gas. [Chu *J Chem Phys* **41** 226 1964, DOI: 10.1063/1.1725626; *Beilstein* **1** IV 205.]

***n*-Propyl chloride** [540-54-5] C₃H₇Cl, **M 78.5**, **m -123°**, **b 46.6°/atm**, **d₄²⁰ 0.890**, **n_D²⁰ 1.3880**. Dry the chloride with MgSO₄ and fractionally distil. It can be more extensively purified using extraction with H₂SO₄ as for *n*-propyl bromide. *Alternatively*, Chien and Willard [*J Am Chem Soc* **75** 6160 1953, DOI: 10.1021/ja01120a016] passed a stream of oxygen containing about 5% ozone through the *n*-propyl chloride for three times as long as was needed to cause the first coloration of starch iodide paper by the exit gas. After washing with aqueous NaHCO₃ to hydrolyse ozonides and remove organic acids, the chloride was dried with MgSO₄ and fractionally distilled. [*Beilstein* **1** IV 189.]

Propylene glycol diacetate (1,2-diacetoxypropane, propylene diacetate) [623-84-7] C₇H₁₂O₄, **M 160.2**, **b 84-85°/12mm**, **191°/760mm**, **d₄²⁵ 1.050**, **n_D²⁰ 1.414**. Wash the ester with aqueous NaHCO₃ in the presence of solid NaCl, dry it with MgSO₄ and fractionally distil it. [*Beilstein* **2** H 142, **2** II 156, **2** III 312, **2** IV 220.]

Propylene glycol 1-methylether (1-methoxy-2-propanol) [RS(±) 107-98-2, R(-) 4984-22-9, S(+) 26550-55-0] C₄H₁₀O₂, **M 90.1**, **b 118.5°/740mm**, **119-120°/atm for RS**, **131-132°/atm for R or S**, **d₄²⁵ 0.922**, **n_D²⁰ 1.403**.

Wash the ethers with aqueous NaHCO_3 in the presence of solid NaCl , dry them with MgSO_4 and fractionally distil them. The *RS*-acetate [108-65-6] **M 132.2** has **b 145-146°/atm**, and the *3,5*-dinitrobenzoate has **m 95°** (from ligroin). The *RS*-alcohol was converted into the *o*-carboxyphenylbenzoate (oil) and the optical resolution of its brucine salt in Me_2CO was achieved. The free acid phthalates were released (as oils), then hydrolysed with aqueous NaOH (2 equivalents) and the alcohols distilled off as azeotropes. Pure alcohols were obtained by saturating the azeotropes with K_2CO_3 , drying the alcohol layer from CaO and distilling.

The *R*(-) and *S*(+) enantiomers have **b 119-120°/atm**, $[\alpha]_{\text{D}}^{20}$ - and + **3.3 or 3.8 (neat)**, **20.5 (c 10, H_2O)**, **20.1 (c 10, CHCl_3)**, **21.1 (c 10, $^*\text{C}_6\text{H}_6$)**, and **4.1 (c 10, EtOH)**. [Inoue et al. *Macromol Khem* **90** 131 1966, DOI: 10.1002/macp.1966.020900112; Butcher & Westheimer *J Am Chem Soc* **77** 2420 1955, DOI: 10.1021/ja01614a018.] [*Beilstein* **1** II 536, **1** III 2146, **1** IV 2471.]

***n*-Propyl ether (dipropyl ether)** [111-43-3] **$\text{C}_6\text{H}_{14}\text{O}$, M 102.2**, **m -123°**, **b 90.1°/atm**, **d_4^{20} 0.740**, **n_{D}^{15} 1.38296**, **n_{D}^{20} 1.3803**, **pK^{25} -4.40 (aqueous H_2SO_4)**. Purify the ether by drying with CaSO_4 , by passage through an alumina column (to remove peroxides), and by fractional distillation. [*Beilstein* **1** III 2146, **1** IV 1422.]

Propyl formate [110-74-7] **M 88.1**, **m -93°**, **b 81.3°/atm**, **d_4^{20} 0.9058**, **n_{D}^{20} 1.3779**. Distil the formate, then wash it with saturated aqueous NaCl , and with saturated aqueous NaHCO_3 in the presence of solid NaCl , dry it with MgSO_4 and fractionally distil it. [*Beilstein* **2** IV 26.]

***n*-Propyl iodide (1-iodopropane)** [107-08-4] **$\text{C}_3\text{H}_7\text{I}$, M 170.0**, **m -101°**, **b 38°/80mm**, **102.5°/atm**, **d_4^{20} 1.745**, **n_{D}^{20} 1.5041**. It should be distilled first under reduced pressure to avoid decomposition. Dry the iodide with MgSO_4 or silica gel and fractionally distil it. Store it under nitrogen with mercury in a brown bottle. Prior to distillation, free iodine can be removed by shaking with copper powder or by washing with aqueous $\text{Na}_2\text{S}_2\text{O}_3$ and drying. Alternatively, the *n*-propyl iodide can be treated with bromine, then washed with aqueous $\text{Na}_2\text{S}_2\text{O}_3$, dried and distilled. See also *n*-butyl iodide. [*Beilstein* **1** IV 222.]

2-Propylpentanoic acid (Valproic acid, VPA, di-*n*-propylacetic acid) [99-66-1] **$\text{C}_8\text{H}_{16}\text{O}_2$, M 144.2**, **b 120-121°/14mm**, **219.5°/760mm**, **220°/atm**, **d_4^{20} 0.91**, **n_{D}^{20} 1.425**, **pK^{25} 4.61**. Purify VPA in the same way as for propionic acid using KMnO_4 (see [79-09-4]). It distils as a colourless oil with a characteristic smell and is soluble in most organic solvents. Its solubility in H_2O is 1.3mg/ml at room temperature. Its FT-IR (film) has ν_{max} at 3572.9, 2969.9, 2885.0, 1771.8, 1465.6, 1388.2, 1102.6, 738.5, 586.5 cm^{-1} ; its ^1H NMR (300MHz, CDCl_3 , TMS) has δ at 0.92 (t, $J = 7\text{Hz}$, 2 CH_3), 1.40 (m, 6H), 1.60 (m, 2H), 2.29 (m, 1H), 12.2 (brs, OH); and the ^{13}C NMR (75MHz, CDCl_3 , CHCl_3 standard) has δ at 183.41, 45.20, 34.37, 20.59, 13.99.

The *ethyl ester* [17022-31-0] **M 172.2** has **b 183°/760mm**. It forms a **1:1 Na salt (sodium valproate, Depakin among other trade names)** [1069-66-5] **M 166.2** as a white, crystalline, odourless, hygroscopic solid with the following solubilities in g/ml: 2.5 in H_2O , 0.66 in EtOH, 0.2 in MeOH, and is insoluble in organic solvents. It also forms a **2:1 Na salt [sodium hydrogen bis(2-propylpentanoate), valproate semisodium]** [76584-70-8] **M 310.4**, a **Magnesium salt (Depamag)**, and the pro-drug amide, **Valpromide (Depramide)** [2430-27-5] **M 143.2**, **m 123°**, **122-124°**, **125-126°**, obtained as needles by heating the acid with NH_3 at 180°, is a bitter, crystalline white powder [*Beilstein* **2** IV 695]. These are all commercially available drugs for the treatment of epilepsy, bipolar disorders and migraine. They are **toxic** at high concentrations. [Wiemann et al. *Bull Soc Chim Fr* 189 1958, Nau et al. *Pharmacol Toxicol* **69** 310 1991, DOI: 10.1111/j.1600-0773.1991.tb01303.x; Löscher **58** 31 1999, DOI: 10.1016/S0301-0082(98)00075-6; *Beilstein* **2** H 350.]

***n*-Propyl propionate** [106-36-5] **$\text{C}_6\text{H}_{12}\text{O}_2$, M 116.2**, **m -76°**, **b 122°/atm**, **d_4^{20} 0.881**, **n 1.393**. Treat the ester with anhydrous CuSO_4 , then distil it under nitrogen. [*Beilstein* **2** IV 707.]

Propyne [74-99-7] **C_3H_4 , M 40.1**, **m -101.5°**, **b -23.2°/760mm**, **d^{50} 0.7062**, **n^{40} 1.3863**. Purify it by preparative gas chromatography. [*Beilstein* **1** 246.]

Pyruvic acid (2-oxopropionic acid) [127-17-3] **$\text{C}_3\text{H}_4\text{O}_3$, M 88.1**, **m 13°**, **b 21°/0.1mm**, **57.8°/10mm**, **85°/40mm**, **165°(dec)/atm**, **pK^{25} 2.39 (2.60)**. Distil it twice under high vacuum, then fractionally crystallise it by partial freezing. It polymerises with decomposition on standing unless pure and in a sealed container. [Howard & Frazer *Org Synth Coll Vol* **1** 475 1941, DOI: 10.15227/orgsyn.004.0063; *Beilstein* **3** IV 1505.]

Ricinoleic acid (*R* 12-hydroxy-*cis*-9-octadecenoic acid, 12-hydroxyoleic acid) [141-22-0] $C_{18}H_{34}O_3$, M 298.5, m 7-8° (α -form), 5.0° (γ -form), n_D^{20} 1.4717, $pK_{Est} \sim 4.5$. Purify it as the *methyl acetylricinoleate* [Rider *J Am Chem Soc* **53** 4130 1931, DOI: 10.1021/ja01362a031], fractionally distilling it at 180-185°/0.3mm, then 87g of this ester is hydrolysed by refluxing with KOH (56g), water (25ml), and MeOH (250ml) for 10 minutes. The free acid is separated on acidification and extraction into Et₂O, dried (Na₂SO₄), filtered, evaporated and the residue is crystallised from acetone at -50°, and distilled in small batches, b 180°/0.005mm. The *R*-enantiomer has m 5.5°, b 245°/10mm and $[\alpha]_D^{26} +7.2$ (c 5, Me₂CO). [Bailey et al. *J Chem Soc* 3027 1957, DOI: 10.1039/JR9570003027; *Beilstein* **3** IV 1026, 1207.]

Sebacic acid (1,10-decanedioic acid) [111-20-6] $C_{10}H_{18}O_4$, M 202.3, m 134.5°, b 232°/10mm, 234°/15mm, 273°/50mm, 294.5°/100mm, pK_1^{25} 4.58, pK_2^{25} 5.54. Purify sebacic acid *via* the disodium salt which, after crystallisation from boiling water (charcoal), is then converted to the free acid. The free acid is crystallised repeatedly from hot distilled water or from Me₂CO/petroleum ether (in plates), and is dried under vacuum. It sublimes slowly at its melting point. It is slightly soluble in hydrocarbon and chlorinated solvents; its solubility in Et₂O is ~0.1% (20°), and in H₂O in %w/v it is 0.004% (0°), 0.10% (20°), 0.42% (65°) and 2.0% (100°). When castor oil is heated at 40° with 40% aqueous NaOH or KOH for 1hr, cooled, the upper layer is separated, mixed with solid NaOH or KOH and heated in a Cu or Fe basin until the odour of octanol vanishes (3hrs), poured into cold H₂O, acidified with HCl and collected to give sebacic acid (20-40% yield) [Dominques et al. *J Chem Ed* **29** 446 1952, DOI: 10.1021/ed029p446]. [*Beilstein* **2** IV 2078.]

Sebacic acid monomethyl ester [818-88-2] $C_{11}H_{20}O_4$, M 216.1, m 40-41°, 42-43°, b 148°/0.025mm, 157°/1mm, 169-171°/4mm, 208°/20mm. Obtained by partial hydrolysis of dimethyl sebacate (17.22 moles, m 26.5-27.5°) with KOH (13.1 moles) in MeOH and isolation of the mono-ester. The crude mono-ester was fractionated in several batches at very low pressure through a 120 cm perforated-disk column (5 theoretical plates) specially designed to permit a minimum pressure gradient (i.e. minimise the unavoidable formation of the diester which is the most volatile component — keep reflux ratio low). Recrystallise the ester from Me₂CO/petroleum ether or petroleum ether at low temperature and distil it in a vacuum. The viscous liquid ester distillate solidifies. **9-Methoxycarbonylnonanoyl chloride** obtained by usual treatment with SOCl₂ distils as a water white oil at 143°/3.5mm (96% yield) which solidified on cooling, m 16.5-17°. When dissolved in petroleum ether (b 35-60°) and saturated with dry NH₃, the *ester-amide*, m 74.5-76° (also 77.4°) is obtained in 83% yield after recrystallisation from MeOH-Me₂O (4:1). The amide provided **9-methoxycarbonylnonanonitrile** (95% yield) after further treatment with SOCl₂ (45°, 6 hrs) as a water-white oil which distils at 131-132°/1.6 mm (also b 178°/14mm) and solidifies readily, m 3-4°. [Soffer et al. *J Am Chem Soc* **69** 1684 1947, DOI: 10.1021/ja01199a035.] [*Beilstein* **2** IV 608.]

Sebaco-1,10-diamidine dihydrochloride (decane-1,10-diamidine dihydrochloride) [5578-81-4 *free base*] $C_{10}H_{22}N_4 \cdot 2HCl$, M 271.2, m 175-175.5°. The solubility of the salt in H₂O is ~60%. Recrystallise it from a small volume of H₂O, or dissolve in EtOH and add Et₂O. It has antimicrobial activity [Fuller *Biochem J* **36** 548 1942, DOI: 10.1042/bj0360548] and is a rabbit liver amine oxidase inhibitor [Blaschko & Duthie *Biochem J* **39** 347 1945, DOI: 10.1042/bj0390347]. The *picrate* crystallises from Et₂O/pyridine with m 175-175.5°. [Broom *J Pharmacol Exp Therap* **57** 81 1936, Lee et al. *J Med Chem* **38** 3053 1995, DOI: 10.1021/jm00016a008; *Beilstein* **2** III 1817.]

Sebaconitrile (decanedinitrile) [1871-96-1] $C_{10}H_{16}N_2$, M 164.3, m 8°, b 127°/0.2mm, n_D^{20} 1.4479. Mix the nitrile with P₂O₅ (10% by wt), distil the dinitrile from it, then redistil it. [Kaufman & Whittaker *J Chem Phys* **24** 1104 1956, DOI: 10.1063/1.1742690; *Beilstein* **2** H 720, **2** II 610, **2** III 1817, **2** IV 2089.]

Sebacoyl chloride (sebacyl chloride, 1,10-decanedioyl dichloride) [111-19-3] $C_{10}H_{16}Cl_2O_2$, M239.1, -5 to -3°, b 168°/12mm, d_4^{25} 1.121, n_D^{20} 1.468. Check for OH bands in the IR spectrum. If absent the distil carefully under a vacuum; but if present then some hydrolysis must have occurred. In this case add excess of SOCl₂ and boil gently until liberation of HCl and SO₂ stops. Evaporate and distil. Note that it is an acid chloride and is reactive towards H₂O and protic solvents. Store in tight containers, or better in sealed tubes. **Eye and skin irritant**. [*Beilstein* **2** IV 2088.] It readily forms the *diamide* [1740-54-1] $C_{10}H_{20}N_2O_2$, M 200.2, with ammonia

and has **m 210°**.

Semicarbazide hydrochloride (hydrazine carboxamide hydrochloride) [563-41-7] $C_{10}H_{16}N_2O \cdot HCl$, **M 111.5**, **m 173°(dec)**, **175°(dec)**, **pK²⁴ 3.66**. Crystallise the salt from aqueous 75% EtOH and dry it under vacuum over $CaSO_4$. Alternatively, crystallise it from a mixture of 3.6 mole % MeOH and 6.4 mole % of water. [Kovach et al. *J Am Chem Soc* **107** 7360 1985, DOI: 10.1021/ja00311a024.] It IR has ν_{max} at 700, 3500 cm^{-1} [Ingersoll et al. *Org Synth Coll Vol I* 485 1941, DOI: 10.15227/orgsyn.005.0093; Davison & Christie *J Chem Soc* 3389 1955, DOI: 10.1039/JR9550003389; Thiele & Stange *Chem Ber* **27** 31 1894, DOI: 10.1002/cber.18940270109; pK: Bartlett *J Am Chem Soc* **54** 2853 1923, DOI: 10.1021/ja01346a027]. The **free base** crystallises as prisms from absolute EtOH, **m 96°**. [Curtius & Heidenreich *Chem Ber* **27** 55 1894, DOI: 10.1002/cber.18940270114; *Beilstein* **3** IV 177.] **TOXIC ORALLY**, possible **CARCINOGEN** and **TERATOGEN**.

Senecialdehyde (3,3-dimethylacraldehyde, 3-methyl-2-butenal) [107-86-8] C_5H_8O , **M 84.1**, **b 78-80°/70mm**, **133-135°/atm**, **d₄²⁰ 0.911**, **n_D²⁰ 1.428**. This flammable oil oxidises readily and should be fractionated under N_2 . It should be stored under N_2 and/or vacuum. The UV has λ_{max} at 235.5nm. The **semicarbazone** has **m 221-222°** (from MeOH) and the **2,4-dinitrophenylhydrazone** has **m 184-185°** (from MeOH). [Forbes & Shilton *J Org Chem* **24** 436 1959, DOI: 10.1021/jo01085a617; *Beilstein* **1** III 2990, **1** IV 3464.]

Skellysolve A is essentially *n*-pentane, **b 28-30°/atm**,
Skellysolve B is essentially *n*-hexane, **b 60-68°/atm**,
Skellysolve C is essentially *n*-heptane, **b 90-100°/atm**,
Skellysolve D is mixed heptanes, **b 75-115°/atm**,
Skellysolve E is mixed octanes, **b 100-140°/atm**,
Skellysolve F is petroleum ether, **b 30-60°/atm**,
Skellysolve G is petroleum ether, **b 40-75°/atm**,
Skellysolve H is hexanes and heptanes, **b 69-96°/atm**,
Skellysolve L is essentially octanes, **b 95-127°/atm**. For methods of purification, see **petroleum ether**.

Solanone [S(+)-trans-2-methyl-5-isopropyl-1,3-nonan-8-one] [1937-54-8] $C_{13}H_{22}O$, **M 194.3**, **b 60°/1mm**, **n_D²⁰ 1.4755**, **[α]_D²⁰ +14° (neat)**. Purify solanone by high vacuum distillation and store it in sealed ampules [Kohda & Sato *JCS Chem Commun* 951 1981, DOI: 10.1039/C39810000951]. It has UV (hexane) at λ_{max} 230nm (ϵ 11,800). The **semicarbazone** crystallises from aqueous EtOH or toluene with **m 160.5-161.5°**. [Johnson et al. *J Org Chem* **30** 2918 1965, DOI: 10.1021/jo01020a009.]

Sorbic acid (2,4-hexadienoic acid) [110-44-1] $C_6H_8O_2$, **M 112.1**, **m 134°**, **pK²⁵ 4.76**. Crystallise the acid from water. Dry it in air or in a desiccator over P_2O_5 . [*Beilstein* **2** IV 1701.]

Stearic acid (octadecanoic acid) [57-11-4] $C_{18}H_{36}O_2$, **M 284.5**, **m 71.4°, 72°**, **b 144-145°/27mm**, **383°/760mm**, **d₄²⁰ 0.911**, **n_D²⁰ 1.428**, **pK²⁵ 4.78**. Crystallise stearic acid from acetone, acetonitrile, EtOH (5 times), aqueous MeOH, ethyl methyl ketone or petroleum ether (b 60-90°), or by fractional precipitation by dissolving in hot 95% EtOH and pouring into distilled water, with stirring. The precipitate, after washing with distilled water, is dried under vacuum over P_2O_5 . It has also been purified by zone melting and partial freezing. [Tamai et al. *J Phys Chem* **91** 841 1987, DOI: 10.1021/j100288a017; *Beilstein* **2** IV 1206.]

Suberic acid (hexane-1,6-dicarboxylic acid) [505-48-6] $C_8H_{14}O_4$, **M 174.2**, **m 141-142°**, **b 230°/15mm**, **pK₁²⁵ 4.12**, **pK₂²⁵ 5.40**. Crystallise it from acetone. It sublimes at 300° without decomposition. [*Beilstein* **2** IV 2028.]

Succinamic acid (succinic acid amide) [638-32-4] $C_6H_7NO_3$, **M 117.1**, **m 155°, 156-157°**, **pK²⁵ 4.54**. Crystallise the amide from Me_2CO or H_2O and dry it in a vacuum. It is not very soluble in MeOH. It is converted into succinimide above 200°. [*Beilstein* **2** H 614.]

Succinamide [110-14-5] $\text{C}_4\text{H}_8\text{N}_2\text{O}_2$, M 116.1, m 262-265°(dec). Crystallise it from water. [Beilstein 2 IV 1922.]

Succinic acid [110-15-6] $\text{C}_4\text{H}_6\text{O}_4$, M 118.1, m 185-185.5°, pK_1^{25} 4.21, pK_2^{25} 5.72. Wash it with diethyl ether. Crystallise it from acetone, distilled water, or *tert*-butanol. Dry it under vacuum over P_2O_5 or concentrated H_2SO_4 . Also purify it by conversion to the disodium salt which, after crystallisation from boiling water (charcoal), is treated with mineral acid to regenerate the succinic acid. The acid is then recrystallised and dried in a vacuum. [Beilstein 2 H 606, 2 IV 1908.]

Succinic anhydride (dihydro-2,5-furandione) [108-30-5] $\text{C}_4\text{H}_4\text{O}_3$, M 100.1, m 119-120°, b 261°/atm. Crystallise the anhydride from redistilled acetic anhydride or CHCl_3 , then filter, wash with diethyl ether and dry it in a vacuum. [Beilstein 17 H 606, 17 V 6.]

Succinimide (2,5-pyrrolidindione) [123-56-8] M 99.1, m 124-125°, b 285-290°/atm, pK^{25} 9.62. Crystallise the imide from EtOH (1ml/g) or water. [Beilstein 21 H 369, 21/9 V 438.]

Succinonitrile (1,2-dicyanoethane) [110-61-2] $\text{C}_4\text{H}_4\text{N}_2$, M 80.1, m 57.9°, b 108°/1mm, 267°/760mm. Purify the nitrile by vacuum sublimation, and/or crystallisation from acetone. [Beilstein 2 H 615, 2 IV 1923.]

D(-)-Tartaric acid (2S,3S-threarc acid [147-71-7] $\text{C}_4\text{H}_6\text{O}_6$, M 150.1, m 169.5-170°, 170-171° (2S,3S-form, natural) $[\alpha]_{546}^{20}$ -15 (c 10, H_2O), $[\alpha]_{546}^{20}$ -12 (c 20, H_2O); m 208°, 210-212°, (2RS,3RS-form, DL [133-37-9]), pK_1^{25} 3.03, pK_2^{25} 4.46, pK_3^{25} 14.4. Crystallise the acid from distilled H_2O or *benzene/diethyl ether containing 5% of petroleum ether (b 60-80°) (1:1). Soxhlet extraction with diethyl ether has been used to remove an impurity absorbing at 265nm. It has also been crystallised from absolute EtOH/hexane and dried in a vacuum for 18 hours [Kornblum & Wade *J Org Chem* 52 5301 1987, DOI: 10.1021/jo00233a001]. The *non-natural* 2R,3S(+) tartaric acid [87-69-4] has m 171-172°, $[\alpha]_{546}^{20}$ +12 (c 20, H_2O). [Beilstein 3 IV 1229.]

meso-Tartaric acid [2RS,3SR(±)-tartaric acid] [147-73-9] $\text{C}_4\text{H}_6\text{O}_6$, M 150.1, m 165-166°; monohydrate [5990-63-6] $\text{C}_4\text{H}_6\text{O}_6 \cdot \text{H}_2\text{O}$, M 168.1, m 139-141°; pK_1^{25} 3.17, pK_2^{25} 4.91. Crystallise it from water (solubility in H_2O at 20° is 125%w/v), wash it with cold MeOH and dry it at 60° under vacuum. This form is optically inactive and was prepared by racemising (isomerising) L-tartaric acid in boiling alkali [Holleman *Org Synth Coll Vol I* 497 1941, DOI: 10.15227/orgsyn.006.0082]. [Beilstein 3 IV 1218.]

Tetra-*n*-amylammonium bromide (tetra-*n*-pentylammonium bromide) [866-97-7] $\text{C}_{20}\text{H}_{44}\text{N} \cdot \text{Br}$, M 378.5, m 100-101°. Crystallise it from petroleum ether, *benzene or acetone/ether mixtures and dry in vacuum at 40-50° for 2 days. It is used in ion-paired chromatography (Sagara et al. *J Chromatogr* 328 289 1985, DOI:10.1016/S0021-9673(01)87399-4). [Beilstein 4 IV 677.]

Tetra-*n*-amylammonium iodide [2498-20-6] $\text{C}_{20}\text{H}_{44}\text{N} \cdot \text{I}$, M 425.5, m 135-137°. Crystallise the iodide from EtOH and dry it at 35° under a vacuum. It has also been purified by dissolving in acetone and precipitating by adding diethyl ether, and drying at 50° for 2 days. [Beilstein 4 IV 677.]

1,1,2,2-Tetrabromoethane [79-27-6] $\text{C}_2\text{H}_2\text{Br}_4$, M 345.7, f 0.0°, b 119°/15mm, 243.5°/atm, d_4^{20} 2.965, n_D^{20} 1.63533. Wash it successively with conc H_2SO_4 (three times) and H_2O (three times), dry it with K_2CO_3 and CaSO_4 and distil it in a vacuum or at ~760mm. [Beilstein 1 IV 162.]

Tetra-*n*-butylammonium bromide [1643-19-2] $\text{C}_{16}\text{H}_{36}\text{N} \cdot \text{Br}$, M 322.4, m 102-106°, 119.6°. Recrystallise the salt from *benzene (5ml/g) at 80° by adding hot *n*-hexane (three volumes) and allowing to cool. Dry it over P_2O_5 or $\text{Mg}(\text{ClO}_4)_2$, under vacuum. The salt is *very hygroscopic*. It can also be crystallised from ethyl acetate or dry acetone by adding diethyl ether and dried *in vacuo* at 60° for 2 days. It has been crystallised from acetone by addition of diethyl ether. It is so *hygroscopic* that all manipulations should be carried out in a dry-box. It has been purified by precipitation from a saturated solution in dry CCl_4 on addition of cyclohexane or by recrystallisation from ethyl acetate, then heating in vacuum to 75° in the presence of P_2O_5 . [Symons et al. *JCS*

Faraday Trans 1 **76** 2251 1980, DOI: 10.1039/F19807602251.] It also recrystallises from CH_2Cl_2 /diethyl ether and is dried in a vacuum desiccator over P_2O_5 . [Blau & Espenson *J Am Chem Soc* **108** 1962 1986, DOI: 10.1021/ja00268a039; *Beilstein* **4** IV 657.]

Tetra-*n*-butylammonium chloride [1112-67-0] $\text{C}_{16}\text{H}_{36}\text{N} \cdot \text{Cl}$, **M 277.9, m 15.7°**. Crystallise the chloride from acetone by addition of diethyl ether. It is very **hygroscopic** and forms crystals with $34\text{H}_2\text{O}$ [37451-68-6] and **m 41-44°**. It is used in ion-paired chromatography (Sagara et al. *J Chromatogr* **328** 289 1985, DOI:10.1016/S0021-9673(01)87399-4). [*Beilstein* **4** IV 557.]

Tetra-*n*-butylammonium hexafluorophosphate [3109-63-5] $\text{C}_{16}\text{H}_{36}\text{N} \cdot \text{PF}_6$, **M 387.5, m 239-241°, 244-246°**. Recrystallise it from saturated EtOH/water and dry it for 10 hours in a vacuum at 70°. Also recrystallise it three times from absolute EtOH and dry it for 2 days in a drying pistol under a vacuum at boiling toluene temperature [Bedard & Dahl *J Am Chem Soc* **108** 5933 1986, DOI: 10.1021/ja00279a044]. It is a stable supporting electrolyte in organic solvents [Baiser in *Organic Electrochemistry* M. Dekker NY p228 1973.]

Tetra-*n*-butylammonium hydrogen sulfate [32503-27-8] $\text{C}_{16}\text{H}_{36}\text{N} \cdot \text{HSO}_4$, **M 339.5, m 171-172°**. The sulfate crystallises from Me_2CO . It has been used as a phase transfer catalyst (see ‘Catalysts, Part 1’ in Chapter 5).

Tetra-*n*-butylammonium iodide [311-28-4] $\text{C}_{16}\text{H}_{36}\text{N} \cdot \text{I}$, **M 369.4, m 141-143°, 146°**. Crystallise the iodide from toluene/petroleum ether (see entry for the corresponding bromide), acetone, ethyl acetate, EtOH/diethyl ether, nitromethane, aqueous EtOH or water. Dry it at room temperature under a vacuum. It has also been dissolved in MeOH/acetone (1:3, 10ml/g), filtered and allowed to stand at room temperature to evaporate to *ca* half its original volume. Distilled water (1ml/g) is then added, and the precipitate is filtered off and dried. It can also be dissolved in acetone, precipitated by adding ether and dried in a vacuum at 90° for 2 days. It has also been recrystallised from CH_2Cl_2 /petroleum ether or hexane, or anhydrous methanol and stored in a vacuum desiccator over H_2SO_4 . [Blau & Espenson *J Am Chem Soc* **108** 1962 1986, DOI: 10.1021/ja00268a039; *Beilstein* **4** IV 558.]

Tetra-*n*-butylammonium nitrate [1941-27-1] $\text{C}_{16}\text{H}_{36}\text{N} \cdot \text{NO}_3$, **M 304.5, m 116-118°, 119°**. Crystallise it from *benzene (7ml/g), EtOH or EtOAc (**m 121-122°**); dry it in a vacuum over P_2O_5 at 60° for 2 days. [*Beilstein* **4** IV 558.]

Tetra-*n*-butylammonium perchlorate [1923-70-2] $\text{C}_{16}\text{H}_{36}\text{N} \cdot \text{NClO}_4$, **M 341.9°, m 210°(dec)**. Crystallise the perchlorate from EtOH, ethyl acetate, *n*-hexane or diethyl ether/acetone mixture, or hot CH_2Cl_2 . Dry it in a vacuum at room temperature over P_2O_5 for 24 hours. [Anson et al. *J Am Chem Soc* **106** 4460 1984, DOI: 10.1021/ja00328a028; Ohst & Kochi *J Am Chem Soc* **108** 2897 1986, DOI: 10.1021/ja00271a019; Collman et al. *J Am Chem Soc* **108** 2916 1986, DOI: 10.1021/ja00271a021; Blau & Espenson *J Am Chem Soc* **108** 1962 1986, DOI: 10.1021/ja00268a039; Gustowski et al. *J Am Chem Soc* **108** 7553 1986, DOI: 10.1021/ja00284a019; Ikezawa & Kutal *J Org Chem* **52** 3299 1987, DOI: 10.1021/jo00391a022; *Beilstein* **4** IV 557.]

Tetra-*n*-butylammonium picrate [914-45-4] $\text{C}_{16}\text{H}_{36}\text{N} \cdot \text{C}_6\text{H}_2\text{N}_3\text{O}_7$, **M 490.6, m 106.5-107°**. Crystallise the picrate from EtOH. Dry it in a vacuum desiccator over P_2O_5 . [*Beilstein* **6** II 271, **4** III 292.]

Tetra-*n*-butylammonium tetrabutylborate ($\text{Bu}_4\text{N}^+ \text{Bu}_4\text{B}^-$) [23231-91-6] $\text{C}_{32}\text{H}_{72}\text{BN}$, **M 481.7, m 106-109°, 155°, 161.8°**. Dissolve it in MeOH or acetone, and crystallise by adding distilled water. It also crystallises from EtOH or EtOAc. Dry it in a vacuum at 70°. It has also been successively recrystallised from isopropyl ether, isopropyl ether/acetone (50:1) and isopropyl ether/EtOH (50:1) for 10 hours, then isopropyl ether/acetone for 1 hour, and dried at 65° under reduced pressure for 1 week. [Kondo et al. *JCS Faraday Trans 1* **76** 812 1980, DOI: 10.1039/F19807600812; *Beilstein* **4** III 293, **4** III 558.]

Tetra-*n*-butylammonium tetrafluoroborate [429-42-5] $\text{C}_{16}\text{H}_{36}\text{N} \cdot \text{BF}_4$, **M 329.3, m 155-161°, 161.8°, 161-163°, $\text{pK}^{25} -4.9$ (for HBF_4)**. Recrystallise it from H_2O , aqueous EtOH or from EtOAc by cooling in Dry-ice. Also recrystallise it from ethyl acetate/pentane or dry acetonitrile. Dry it at 80° under vacuum. [Detty & Jones *J Am Chem Soc* **109** 5666 1987, DOI: 10.1021/ja00253a017; Hartley & Faulkner *J Am Chem Soc* **107** 3436

1985, DOI: 10.1021/ja00298a007.] The **acetate** has **m 118±2°** (from BuCl), the **bromide** has **m 118°** (from EtOAc), and the **nitrate** has **m 120°** (from *C₆H₆). [Witschonke & Kraus *J Am Chem Soc* **69** 2472 1947, DOI: 10.1021/ja01202a067; Wheeler & Sandstedt *J Am Chem Soc* **77** 2024 1955, DOI: 10.1021/ja01612a102; *Beilstein* **4** IV 558.]

1,1,2,2-Tetrachloro-1,2-difluoroethane [72-12-0] C₂Cl₂F₄, **M 203.8**, **f 26.0°**, **b 92.8°/760mm**, **d₄²⁵ 1.6252**, **n_D²⁵ 1.4130**. Purify it as for trichlorotrifluoroethane. [*Beilstein* **1** III 165, **1** III 146.]

sym-Tetrachloroethane (acetylene tetrachloride) [79-34-5] C₂H₂Cl₄, **M 167.9**, **m -43°**, **b 62°/100mm**, **146.2°/atm**, **d₄²⁰ 1.588**, **n_D¹⁵ 1.49678**. Stir the ethane, on a steam-bath, with concentrated H₂SO₄ until a fresh portion of acid remains colourless. The organic phase is then separated, distilled in steam, dried (CaCl₂ or K₂CO₃), and fractionally distilled in a vacuum. [*Beilstein* **1** IV 144.]

Tetrachloroethylene (PCE, pechloroethylene) [127-18-4] C₂Cl₄, **M 165.8**, **b 62°/80mm**, **121.2°**, **d₄¹⁵ 1.63109**, **d₄²⁰ 1.623**, **n_D¹⁵ 1.50759**, **n_D²⁰ 1.50566**. It decomposes under similar conditions to CHCl₃, to give phosgene and trichloroacetic acid. Inhibitors of this reaction include EtOH, diethyl ether and thymol (effective at 2-5 ppm). Tetrachloroethylene should be distilled under a vacuum (to avoid formation of phosgene) and stored in the dark out of contact with air. It can be purified by washing with 2M HCl until the aqueous phase no longer becomes coloured, then with water, drying with Na₂CO₃, Na₂SO₄, CaCl₂ or P₂O₅, and fractionally distilling just before use. 1,1,2-Trichloroethane and 1,1,1,2-tetrachloroethane can be removed by counter-current extraction with EtOH/water. [*Beilstein* **1** IV 715.]

Tetracosane (C₂₄) [646-31-1] C₂₄H₅₀, **M 338.7**, **m 49-52°**, **54°**, **b 243-244°/15mm**, **391°/atom**. Crystallise it from diethyl ether and/or distil it under high vacuum. [*Beilstein* **1** IV 578.]

Tetracosanoic (lignoceric) acid [557-59-5] C₂₄H₄₈O₂, **M 368.7**, **m 84°**, **87.5-88°**, **pK_{Est} ~5.0**. Crystallise the acid from acetic acid, Me₂CO, toluene, petroleum ether/Me₂CO or *C₆H₆/Me₂CO. [*Beilstein* **2** IV 1301.]

Tetracyanoethylene (TCNE, ethylenetetracarbonitrile) [670-54-2] C₆N₄, **M 128.1**, **m 197-199°**, **199-200°(sealed tube)**. Recrystallise it from chlorobenzene, dichloroethane, or dichloromethane [Hall et al. *J Org Chem* **52** 5528 1987, DOI: 10.1021/jo00234a006]. Note that its solubility in dry chlorobenzene increases rapidly near the boiling point of the solvent. Therefore crystals should be extracted with boiling chlorobenzene which should be dry. Store it at 0° in a desiccator over NaOH pellets. (It slowly evolves HCN on exposure to moist air **CARE**.) Also purify it by repeated sublimation at 120-130°/0.5mm. Pure product with barely a trace of yellow colour is obtained by placing TCNE (35g) in a glass thimble and covering it with activated wood charcoal chips (4-8 mesh). The mouth of the thimble is covered with coarse grade filter paper which is held in place by wiring. The thimble is then placed in a sublimator, and sublimation is carried out at 1-2 mm with bath temperature at 175-200°. TCNE is recovered in 80-90% yield as a colourless, hard crystalline mass with **m 201-202°(sealed tube)**. [Frey et al. *J Am Chem Soc* **107** 748 1985, DOI: 10.1021/ja00290a002; Traylor & Miksztal *J Am Chem Soc* **109** 2778 1987, DOI: 10.1021/ja00243a033; Fatiadi *Synthesis* 249 1986, DOI: 10.1055/s-1986-31584; Fatiadi *Synthesis* 749 1987, DOI: 10.1055/s-1987-28074; Fatiadi *Synthesis* 959 1987, DOI: 10.1055/s-1987-28140; Carboni *Org Synth Coll Vol* **4** 877 1963, DOI: 10.15227/orgsyn.039.0064; *Beilstein* **2** IV 2450.] It is a π-acid which forms π-complexes with polycyclic hydrocarbons, e.g. with pyrene where the π-complex has an equilibrium constant of 29.5 [Melby et al. *J Am Chem Soc* **84** 3374 1962, DOI: 10.1021/ja00876a029; Acker & Hertler *J Am Chem Soc* **84** 3370 1962, DOI: 10.1021/ja00876a028].

Tetradecane (C₁₄) [629-59-4] C₂H₃₀, **M 198.4**, **m 6°**, **b 122°/10mm**, **252-254°/atm**, **d₄²⁰ 0.763**, **n_D²⁰ 1.429**. Wash it successively with 4M H₂SO₄ and water. Dry it over MgSO₄ and distil it several times under reduced pressure [Poë et al. *J Am Chem Soc* **108** 5459 1986, DOI: 10.1021/ja00278a015]. It is used as a standard in gas chromatography. [*Beilstein* **1** H 171, **1** IV 520.]

1-Tetradecanol (myristyl alcohol) [112-72-1] C₁₄H₃₀O, **M 214.4**, **m 35-39°**, **39-39.5°**, **b 160°/10mm**, **170-173°/20mm**, **289°/atm**. Crystallise the alcohol from aqueous EtOH or distil preferably in a vacuum. It has also been purified by zone melting. [*Beilstein* **1** IV 1864.]

Tetradecyl ether (di-tetradecyl ether) [5412-98-6] $C_{28}H_{58}O$, M 410.7, m 43.5°, 45°, b 457°/760mm, d₄²⁰ 0.8117. Distil the ether under a vacuum and then crystallise it repeatedly from MeOH/*benzene. It also crystallises from MeOH alone (m 31.2°, 33°, and 44.4° also reported), or Me₂CO (m 43.5°). [Di Giacomo & Smyth *J Am Chem Soc* **78** 2027 1956, DOI: 10.1021/ja01591a001; *Beilstein* **1** IV 1865.]

Tetradecyltrimethylammonium bromide (myristyl trimethylammonium bromide) [1119-97-7] $C_{17}H_{38}N$. Br, M 336.4, m 244-245°, 244-249°, 245-250°. Crystallise the bromide from Me₂CO or a mixture of Me₂CO and >5% MeOH or Me₂CO and EtOH. Wash it with diethyl ether and dry it in a vacuum oven at 60°. It is a cationic detergent with micellar average weight 27,000, aggregation number 80 and CMC 4-5mM at 20-25°. Its solubility is 1g/5g H₂O. [Dearden & Woolley *J Phys Chem* **91** 2404 1987, DOI: 10.1021/j100293a040; Shelton et al. *J Am Chem Soc* **68** 753 1946, DOI: 10.1021/ja01209a012; *Beilstein* **4** III 419, **4** IV 813.]

Tetraethoxymethane See tetraethyl orthocarbonate below.

Tetraethylammonium bromide (TEA bromide) [71-91-0,] $C_8H_{20}N$. Br, M 210.2, m 269°(dec), 284°(dec), 285°(dec). Recrystallise the bromide from EtOH, CHCl₃ or diethyl ether, or, recrystallise it from acetonitrile and dry it over P₂O₅ under reduced pressure for several days. It also recrystallises from EtOH/diethyl ether (1:2), EtOAc, water or boiling MeOH/acetone (1:3) or by adding an equal volume of acetone and allowing to cool. Dry it at 100° *in vacuo* for 12 days, and store over P₂O₅. [*Beilstein* **4** IV 332.]

Tetraethylammonium chloride hydrate (TEA chloride) [68696-18-4 (H₂O), 56-34-8 (anhydrous)] $C_8H_{20}N$. Cl, M 165.7 (anhydrous), m dec >200°. Crystallise the chloride from EtOH by adding diethyl ether, from warm water by adding EtOH and diethyl ether, from dimethylacetamide or from CH₂Cl₂ by addition of diethyl ether. Dry it over P₂O₅ in vacuum for several days. It also crystallises from acetone/CH₂Cl₂/hexane (2:2:1) [Blau & Espenson *J Am Chem Soc* **108** 1962 1986, DOI: 10.1021/ja00268a039; White & Murray *J Am Chem Soc* **109** 2576 1987, DOI: 10.1021/ja00243a005]. [*Beilstein* **4** IV 332.]

Tetraethylammonium hexafluorophosphate [429-07-2] $C_8H_{20}N$. PF₆, M 275.2, m >300°, 331°(dec), pK₁²⁵ ~ 0.5, pK₂²⁵ 5.12 (for fluorophosphoric acid H₂PO₃F). Dissolve the salt (0.8g) in hot H₂O (3.3ml) and cool to crystallise. Yield of prisms is 0.5g. Its solubility in H₂O is 8.1g/L at 19° [Lange & Müller *Chem Ber* **63** 1058 1930, DOI: 10.1002/cber.19300630510]. [*Beilstein* **4** III 199.]

Tetraethylammonium iodide [68-05-3] $C_8H_{20}N$. I, M 257.2, m 302°, >300°(dec). Crystallise the iodide from acetone/MeOH, EtOH/water, dimethylacetamide or ethyl acetate/EtOH (19:1). Dry it under a vacuum at 50° and store it over P₂O₅. [*Beilstein* **4** IV 332.]

Tetraethylammonium perchlorate [2567-83-1] $C_8H_{20}N$. ClO₄, M 229.7, m 345°(dec). Crystallise the perchlorate repeatedly from water, aqueous MeOH, acetonitrile or acetone, and dry it at 70° under a vacuum for 24 hours. [Cox et al. *J Am Chem Soc* **106** 5965 1984, DOI: 10.1021/ja00332a035; Liu et al. *J Am Chem Soc* **108** 1749 1986, DOI: 10.1021/ja00268a006; White & Murray *J Am Chem Soc* **109** 2576 1987, DOI: 10.1021/ja00243a005.] It has also been crystallised twice from ethyl acetate/95% EtOH (2:1) [Lexa et al. *J Am Chem Soc* **109** 6464 1987, DOI: 10.1021/ja00255a036]. [*Beilstein* **4** IV 332.]

Tetraethylammonium picrate [741-03-7] $C_8H_{20}N$. C₆H₂N₃O₇, M 342.1, m >300°(dec). Purify it by successive crystallisations from water or 95% EtOH followed by drying in vacuum at 70°. [*Beilstein* **4** IV 332.]

Tetraethylammonium tetrafluoroborate [429-06-1] $C_8H_{20}N$. BF₄, M 217.1, m 235°, 356-367°, 275-277°, 289-291°, pK₂₅ -4.9 (for HBF₄). Dissolve the salt in hot MeOH, filter and add Et₂O. It is soluble in ethylene chloride [Thompson & Kraus *J Am Chem Soc* **69** 1016 1947, DOI: 10.1021/ja01197a012; Wheeler & Sandstedt *J Am Chem Soc* **77** 2025 1955, DOI: 10.1021/ja01612a103]. It has also been recrystallised three times from a mixture of ethyl acetate/hexane (5:1) or MeOH/petroleum ether, then stored at 95° for 48 hours under vacuum [Hartley & Faulkner *J Am Chem Soc* **107** 3436 1985, DOI: 10.1021/ja00298a007; Huang et al. *Anal Chem* **58** 2889 1986, DOI: 10.1021/ac00126a070]. It is used as a supporting electrolyte. [*Beilstein* **4** IV 333.]

Tetraethylammonium tetraphenylborate [12099-10-4] $C_8H_{20}N.C_{24}H_{20}B$, M 449.4. Recrystallise the borate from aqueous acetone. Dry it in a vacuum oven at 60° for several days. *Similarly for the propyl and butyl homologues.* [Beilstein 4 IV 333.]

Tetraethylene glycol dimethyl ether (tetraglyme) [143-24-8] $C_{10}H_{22}O_5$, M 222.3, m -30°, b 105°/1mm, 275-276°/atm, d_4^{20} 1.010, n_D^{20} 1.435. Stand the ether over CaH_2 , $LiAlH_4$ or sodium, and distil it when required. [Beilstein 1 IV 2404.]

Tetraethylenepentamine (TEPA, tetrene) [112-57-2] $C_8H_{23}O_5$, M 189.3, m -40°, b 169-171°/0.05mm, 340°/atm, d_4^{20} 0.999, n_D^{20} 1.506, pK_1^{25} 2.98, pK_2^{25} 4.72, pK_3^{25} 8.08, pK_4^{25} 9.10, pK_5^{25} 9.68. Distil the amine under vacuum. Also purify *via* its *penta hydrochloride*, *nitrate* or *sulfate*. Jonassen, Frey and Schaafsma [*J Phys Chem* 61 504 1957, DOI: 10.1021/j150550a033] cooled a solution of 150g of the base in 300ml of 95% EtOH, and added dropwise 180ml of concentrated HCl, keeping the temperature below 20°. The white precipitate was filtered off, crystallised three times from EtOH/water, then washed with diethyl ether and dried by suction. Reilley & Holloway [*J Am Chem Soc* 80 2917 1958, DOI: 10.1021/ja01545a001], starting with a similar solution cooled to 0°, added slowly (keeping the temperature below 10°) a solution of 4.5g-moles of HNO_3 in 600ml of aqueous 50% EtOH (also cooled to 0°). The precipitate was filtered by suction, recrystallised five times from aqueous 5% HNO_3 , then washed with acetone and absolute EtOH and dried at 50°. [For purification *via* the sulfate see Reilley and Vavoulis (*Anal Chem* 31 243 1959, DOI: 10.1021/ac60146a026), and for an additional purification step using the Schiff base with benzaldehyde see Jonassen et al. *J Am Chem Soc* 79 4279 1957, DOI: 10.1021/ja01573a011]. [Beilstein 4 IV 1244.] The **TEPA pentahydrochloride** [4961-41-5] $C_8H_{23}O_5 \cdot 5HCl$, M 371.6 has m 280°(dec).

Tetraethyl 1,1,2,2-ethanetetracarboxylate [632-56-4] $C_{14}H_{22}O_8$, M 318.3, m 73-74°, 73-75°. Recrystallise the ester twice from EtOH by cooling to 0°. [Mochizuki et al. *Bull Chem Soc Jpn* 64 1750 1991, DOI: org/10.1246/bcsj.64.1750; Weinges et al. *Angew Chem Int Ed* 20 960 1981, DOI: 10.1002/anie.198109601; Beilstein 2 IV 2415.]

Tetraethyl orthocarbonate (ethyl orthocarbonate, tetraethoxy ethane) [78-09-1] $C_9H_{20}O_4$, M 192.3, b 59.6-60°/14mm, 158°/atm, 159°/atm, 160-161°/atm, d_4^{20} 0.9186, n_D^{20} 1.3932. Likely impurities are hydrolysis products. Shake the orthocarbonate with brine (saturated NaCl, dilute with a little Et_2O if amount of material is small) and dry ($MgSO_4$). The organic layer is filtered, evaporated, and the residue is distilled through a helices packed fractionating column with a total reflux partial take-off head. All distillations can be done at atmospheric pressure in an inert atmosphere (e.g. N_2). [Roberts & McMahon *Org Synth Coll Vol* 4 457 1963, DOI: 10.15227/orgsyn.032.0068; Connolly & Dyson *J Chem Soc* 828 1937, DOI: 10.1039/JR9370000827; Tieckelmann & Post *J Org Chem* 13 265 1948, DOI: 10.1021/jo01160a014; for review see Kantlehner et al. *Justus Liebig's Ann Chem* 507 1982, DOI: 10.1002/jlac.198219820313; Beilstein 3 IV 6.]

2,2,3,3-Tetrafluoropropan-1-ol [76-37-9] $C_3H_4F_4O$, M 132.1, b 106-106.5°/~760mm, d_4^{25} 1.48 g/ml, n_D^{20} 1.321, pK^{25} 12.74. Tetrafluoro-1-propanol (450ml) is added to a solution of $NaHSO_3$ (2.25g) in water (90ml), shaken vigorously and set aside for 24 hours. The fraction distilling at or above 99° is refluxed for 4 hours with 5-6g of KOH and rapidly distilled, followed by a final fractional distillation. [Kosower & Wu *J Am Chem Soc* 83 3142 1961, DOI: 10.1021/ja01475a034.] Alternatively, shake the alcohol with alumina for 24 hours, dry it overnight with anhydrous K_2CO_3 and distil it, taking the middle fraction (b 107-108°). [Beilstein 1 IV 1438.]

Tetra-*n*-heptylammonium bromide [4368-51-8] $C_{28}H_{60}N.Br$, M 490.7, m 88.9-89.1°. Crystallise the bromide from *n*-hexane, then dry it in a vacuum oven at 70°. [Goodrich et al. *J Am Chem Soc* 72 4411 1950, DOI: 10.1021/ja01166a020; Beilstein 4 IV 736.]

Tetra-*n*-heptylammonium iodide [3535-83-9] $C_{28}H_{60}N.Cl$, M 537.7, m 102-103°. Recrystallise the iodide from EtOH or aqueous EtOH. [Erikson et al. *J Org Chem* 25 849 1960, DOI: 10.1021/jo01075a616; Beilstein 4 IV 736 for triheptylamine.]

Tetra-*n*-hexylammonium bromide [4328-13-6] $C_{24}H_{52}N.Br$, M 434.6, m 99-100°. Wash the bromide with

ether, and dry it in a vacuum at room temperature for 3 days.

Tetra-*n*-hexylammonium chloride [5922-92-9] $C_{24}H_{52}N.Cl$, M 390.1, m 111-113°. Crystallise the chloride from EtOH.

Tetra-*n*-hexylammonium iodide [2138-24-1] $C_{24}H_{52}N.I$, M 481.6, m 99-101°, 102-103°. Wash the iodide with diethyl ether and dry it at room temperature *in vacuo* for 3 days. It is soluble in CH_2Cl_2 . [Eriksen et al. *J Org Chem* **25** 849 1960, DOI: 10.1021/jo01075a616; *Beilstein* **4** IV 711 for trihexylamine.]

Tetrahexylammonium perchlorate [4656-81-9] $C_{24}H_{52}N.ClO_4$, M 454.1, m 104-106°. Crystallise the salt from acetone and dry it *in vacuo* at 80° for 24 hours.

Tetrakis(dimethylamino)ethylene (TDAE) [996-70-3] $C_{10}H_{24}N_4$, M 300.2, b 60°/1mm, d_4^{20} 0.861, n_D^{20} 1.4817, $pK_{Est(1)} < 0$, $pK_{Est(2)} < 0$, $pK_{Est(3)} \sim 1.5$, $pK_{Est(4)} \sim 5.1$. Impurities include tetramethylurea, dimethylamine, tetramethylethane-diamine and tetramethyloxamide. It is washed with water while being flushed with nitrogen to remove dimethylamine, dried over molecular sieves, then passed through a silica gel column (previously activated at 400°) under nitrogen. De-gas it in a vacuum line by distillation from a trap at 50° to one at -70°. Finally, it is stirred over sodium-potassium alloy for several days. [Holroyd et al. *J Phys Chem* **89** 4244 1985, DOI: 10.1021/j100266a019; Wiberg *Angew Chem Int Ed* **7** 766 1968, DOI: 10.1002/anie.196807661; *Beilstein* **4** IV 167.]

Tetramethylammonium bromide [64-20-0] $C_4H_{12}N.Br$, M 154.1, sublimes with dec >230°. Crystallise the bromide from EtOH, EtOH/diethyl ether, MeOH/acetone, water or from acetone/MeOH (4:1) by adding an equal volume of acetone. It is dried at 110° under reduced pressure or at 140° for 24 hours. [*Beilstein* **4** IV 145.]

Tetramethylammonium chloride (TMA) [75-57-0] $C_4H_{12}N.Cl$, M 109.6, m >230°(dec). Crystallise the chloride from EtOH, EtOH/ $CHCl_3$, EtOH/diethyl ether, acetone/EtOH (1:1), isopropanol or water. Traces of the free amine can be removed by washing with $CHCl_3$. [*Beilstein* **4** IV 145.]

Tetramethylammonium hexafluorophosphate [558-32-7] $C_4H_{12}N.PF_6$, M 219.1, m >300°, d_4^{25} 1.617, $pK_1^{25} \sim 0.5$, pK_2^{25} 5.12 (for fluorophosphoric acid H_2PO_3F). The salt (0.63g) is recrystallised from boiling H_2O (76ml), yielding pure (0.45) $Me_4N.PF_6$ after drying at 100°. It is a good supporting electrolyte. [Lange & Müller *Chem Ber* **63** 1058 1930, DOI: 10.1002/cber.19300630510; *Beilstein* **4** III 110.]

Tetramethylammonium hydroxide ($5H_2O$) [10424-65-4 ($5H_2O$), 75-59-2 (aqueous solution)] $C_4H_{12}N.OH$, M 181.2, m 63°, 65-68°, 67-70°. It is freed from chloride ions by passage through an ion-exchange column (e.g. Amberlite IRA-400, prepared in its OH^- form by passing 2M NaOH until the effluent is free from chloride ions, then washed with distilled H_2O until neutral). A modification, to obtain carbonate-free hydroxide, uses the method of Davies and Nancollas [*Nature* **165** 237 1950, DOI:10.1038/165237b0]. An aqueous solution (1.0 ± 0.02 M in H_2O) and a methanolic solution (25% w/w) are also commercially available. [*Beilstein* **4** H 50, **4** IV 145.]

Tetramethylammonium iodide [75-58-1] $C_4H_{12}N.I$, M 201.1, m >230°(dec), >300°(dec). Crystallise the iodide from water or 50% EtOH, EtOH/diethyl ether, ethyl acetate, or from acetone/MeOH (4:1) by adding an equal volume of acetone. Dry it in a vacuum desiccator. [*Beilstein* **4** IV 146.]

Tetramethylammonium nitrate [1941-24-8] $C_4H_{12}N.NO_3$, M 136.2, m >300°, 410°. Recrystallise the nitrate from EtOH and dry at 110° in an air oven. [Coats & Taylor *J Chem Soc* 1495 1936, DOI: 10.1039/JR9360001495; *Beilstein* **4** III 113, **4** IV 147.]

Tetramethylammonium perchlorate [2537-36-2] $C_4H_{12}N.ClO_4$, M 123.6, m >300°, pK^{25} -2.4 to -3.1 (for $HClO_4$). Crystallise it twice from H_2O and dry it at 110° in an air oven. It is insoluble in most organic solvents. [Mead et al. *J Chem Soc* 1207 1933, DOI: 10.1039/JR9330001207; Coats & Taylor *J Chem Soc* 1495 1936, DOI: 10.1039/JR9360001495.]

Tetramethylammonium tetrphenylborate [15525-13-0] $C_4H_{12}N$, $C_{24}H_{20}B$, **M 393.3**. Recrystallise it from acetone, acetone/ CCl_4 and from acetone/1,2-dichloroethane. Dry it over P_2O_5 in a vacuum, or in a vacuum oven at 60° for several days. [Beilstein 4 IV 145.]

***N,N,N',N'*-Tetramethylethylenediamine (TMEDA, TMED)** [110-18-9] $C_6H_{16}N_2$, **M 116.2, m -55°, b 122°/atm, d_4^{20} 1.175, n_D^{25} 1.4153, pK_1^{25} 5.90, pK_2^{25} 9.14**. Dry TMEDA partially with molecular sieves (Linde type 4A), then distil it in a slight vacuum from butyl lithium. This treatment removes all traces of primary and secondary amines and water. [Hay et al. *JCS Faraday Trans 1* **68** 1 1972, DOI: 10.1039/F19726800001.] Or dry it with KOH pellets, reflux for 2 hours with one-sixth its weight of *n*-butyric anhydride (to remove primary and secondary amines) and fractionally distil it. Reflux it with fresh KOH, and distil it under nitrogen. [Cram & Wilson *J Am Chem Soc* **85** 1245 1963, DOI: 10.1021/ja00892a008.] It was also distilled from Na. Store it sealed under N_2 . The **dipicrate** has **m 263°(dec)**. [Beilstein 4 H 250, 4 I 415, 4 II 690, 4 III 512, 4 IV 1172.]

Tetramethylethylenediamine dihydrochloride [7677-21-8] $C_6H_{16}N_2 \cdot 2HCl$, **M 198.2, m ~300°**. Crystallise the salt from 98% EtOH/concentrated HCl. It is *hygroscopic*. [Knorr *Chem Ber* **37** 3507 1904, DOI: 10.1002/cber.190403703177; Beilstein 4 IV 1172.]

1,1,3,3-Tetramethylguanidine [80-70-6] $C_5H_{13}N_3$, **M 115.2, b 52-54°/11mm, 159-160°/atm, 162-163°/atm, d_4^{20} 0.917 n_D^{20} 1.470, pK^{25} 13.6**. Reflux it over granulated BaO, then fractionally distil it. Protect it from CO_2 and H_2O as it is a strong base. [Beilstein 4 IV 227.]

Tetramethyl orthocarbonate (methyl orthocarbonate, tetramethoxy methane) [1850-14-2] $C_5H_{12}O_4$, **M 136.2, m -5.6°, -5°, -2°, b 113.5°/760mm, 113.5-114°/755mm, 112-114°/atm, d_4^{20} 1.0202, n_D^{20} 1.3860**. Purify it in the same way as for tetraethyl orthocarbonate. [Smith *Acta Chem Scand* **10** 1006 1956, DOI: 10.3891/acta.chem.scand.10-1006; Tieckelmann & Post *J Org Chem* **13** 265 1948, DOI: 10.1021/jo01160a014; Kantlehner et al. *Synthesis* **73** 1977, DOI: 10.1055/s-1977-24283; Fatiadi *Synthesis* 749 1987, DOI: 10.1055/s-1987-28074; Fatiadi *Synthesis* 959 1987, DOI: 10.1055/s-1987-28140; Beilstein 3 IV 4.]

2,6,10,14-Tetramethylpentadecane (pristane, norphytane) [1921-70-6] $C_{19}H_{40}$, **M 268.5, b 68° (bath temp)/0.004mm, 158°/10mm, 296°/atm, d_4^{20} 0.7827, n_D^{20} 1.4385**. Purify pristane by shaking it with concentrated H_2SO_4 (**care**, if amount of pristane is too small then it should be diluted with petroleum ether *not* Et_2O which is quite soluble in H_2SO_4), then H_2O (**care**, as it may heat up in contact with concentrated H_2SO_4), dry ($MgSO_4$), evaporate and distil it over Na. It is a lubricant, which was isolated from shark liver oil, petroleum crude oils and wool wax; and was synthesised from phytol [7541-49-3] [Sørensen & Sørensen *Acta Chem Scand* **3** 939 1949, DOI: 10.3891/acta.chem.scand.03-0939]. [Beilstein 1 III 570.]

Tetramethylthiuram disulfide [bis-(dimethylthiocarbamyl)disulfide, Thiram] [137-26-8] $C_6H_{12}N_2S_4$, **M 240.4, m 146-148°, 155-156°**. Crystallise thiram (three times) from boiling $CHCl_3$, then recrystallise it from boiling $CHCl_3$ by adding EtOH dropwise to initiate crystallisation, and allow it to cool. Finally, it is precipitated from cold $CHCl_3$ by adding EtOH (which retains the monosulfide in solution). [Ferington & Tobolsky *J Am Chem Soc* **77** 4510 1955, DOI: 10.1021/ja01622a020; Beilstein 4 IV 242.]

1,1,3,3-Tetramethyl urea [632-22-4] $C_5H_{12}N_2O$, **M 116.2, m -1.2°, b 62-63°/12mm, 175.2°/760mm, d_4^{20} 0.969, n_D^{20} 1.453**. Dry it over BaO and distil it under nitrogen. It denatures proteins in H_2O . [Elbaum & Herskovits *Biochemistry* **13** 1268 1974, DOI: 10.1021/bi00703a033; Kane *Anal Biochem* **53** 350 1973, DOI: 10.1016/0003-2697(73)90081-X; Beilstein 4 IV 225.]

Tetranitromethane (TNM) [509-14-8] CN_4O_8 , **M 196.0, m 14.2°, b 46°/36mm, 21-23°/23mm, 126°/760mm, d_4^{20} 1.640, n_D^{20} 1.438**. Shake tetranitromethane with dilute NaOH, wash (H_2O), steam distil, dry with Na_2SO_4 and fractionally crystallise it by partial freezing. The melted crystals are dried with $MgSO_4$ and fractionally distilled under reduced pressure. *Alternatively*, shake it with a large volume of dilute NaOH until no absorption attributable to the *aci*-nitro anion (from mono- di- and tri- nitromethanes) is observable in the water. Then wash it with distilled water, and distil it at room temperature by passing a stream of air or nitrogen through the liquid and condensing it in a trap at -80°. It can be dried with $MgSO_4$ or Na_2SO_4 , fractionally recrystallised from the

melt, and fractionally distilled under reduced pressure. [Liang *Org Synth Coll Vol* **3** 803 1955, DOI: 10.15227/orgsyn.021.0105; *Beilstein* **4** H 80, **4** I 21, **4** II 45, **4** III 116, **4** IV 107.] **Potentially explosive (when impure e.g. with toluene), toxic, carcinogenic.**

Tetra-*n*-propylammonium bromide [1941-30-6] $\text{C}_{12}\text{H}_{28}\text{N} \cdot \text{Br}$, **M 266.3, m >280°(dec).** Crystallise it from ethyl acetate/EtOH (9:1), acetone or MeOH. Dry it at 110° under reduced pressure. [*Beilstein* **4** IV 471.]

Tetra-*n*-propylammonium iodide [631-40-3] $\text{C}_{12}\text{H}_{28}\text{N} \cdot \text{I}$, **M 313.3, m >280°(dec).** Purify the iodide by crystallising it from EtOH, EtOH/diethyl ether (1:1), EtOH/water or aqueous acetone. Dry it at 50° under a vacuum and store it over P_2O_5 in a vacuum desiccator. Keep it away from light. [*Beilstein* **4** IV 472.]

Tetra-*n*-propylammonium perchlorate [15780-02-6] $\text{C}_{12}\text{H}_{28}\text{N} \cdot \text{ClO}_4$, **M 285.8, m 238-240°, 239-241°, $\text{pK}^{25}_{\text{a}}$ -2.4 to -3.1 (for HClO_4).** Purify it by recrystallisation from H_2O or MeCN/ H_2O (1:4.v/v), and dry it in an oven at 60° for several days, or in a vacuum over P_2O_5 at 100°. [Walden & Hilgert *Z Phys Chem* **165** 245 1933, Walden & Birr *Z Phys Chem* **144** 281 1929, Walden & Busch *Z Phys Chem* **140** 97 1929, *Beilstein* **4** II 628.]

Thioacetamide [62-55-5] $\text{C}_2\text{H}_5\text{NS}$, **M 75.1, m 112-113°, $\text{pK}^{25}_{\text{a}}$ 13.4.** Crystallise the amide from absolute diethyl ether or *benzene. Dry it at 70° in a vacuum and store it over P_2O_5 at 0° under nitrogen. (*It develops an obnoxious odour on storage*, and the absorption at 269nm decreases, hence it should be freshly recrystallised before use). [*Beilstein* **2** IV 565.]

Thiodiglycollic acid (2,2'-dithioacetic acid) [123-93-3] $\text{C}_4\text{H}_6\text{O}_4\text{S}$, **M 150.2, m 129°, 128-131°, pK^{25}_1 3.15 (3.24), pK^{25}_2 4.13 (4.56).** Crystallise the acid from water. It forms Cu, Pb, Hg and Ag salts used for their detection. [*Beilstein* **3** IV 612.]

3,3'-Thiodipropionic acid (bis[2-carboxyethyl]sulfide) [111-17-1] $\text{C}_6\text{H}_{10}\text{O}_4\text{S}$, **M 178.2, m 134°, pK^{25}_1 3.84, pK^{25}_2 4.66.** Crystallise the sulfide from water (very soluble at 100°, but 3.7% at 26°). It is soluble in EtOH and Me_2CO . It is an antioxidant. [*Beilstein* **3** IV 735.]

Thioformamide [115-08-2] CH_3NS , **M 61.0, m 29°, 32.0-33.8°, $\text{pK}_{\text{Est}} \sim 12.4$.** Crystallise thioformamide from EtOAc, Et_2O or ether/petroleum ether. The *monohydrate* is a yellow oil soluble in many organic solvents. Its UV has λ_{max} at 263nm (ϵ 2500) in MeOH. [Erlenmeyer & Menzi *Helv Chim Acta* **31** 2065 1948, DOI: 10.1002/hlca.19480310722.] *Alternatively*, dissolve it in Et_2O to separate it from any formamide and/or polymers, filter, evaporate and recrystallise the residue from EtOAc at Dry-ice temperature [Londergan et al. *J Am Chem Soc* **75** 4456 1953, DOI: 10.1021/ja01114a018]. Store it in Et_2O solution over P_2O_5 . [Cousineau & Secristill *J Org Chem* **44** 4351 1979, DOI: 10.1021/jo01338a023; *Beilstein* **2** H 95, **2** I 39, **2** III 128, **2** IV 92.]

Thioglycollic acid (mercaptoacetic acid) [68-11-1] $\text{C}_2\text{H}_4\text{O}_2\text{S}$, **M 92.1, m -16°, b 95-96°/8mm, d^{20}_4 1.326, n^{20}_D 1.505, pK^{25}_1 3.42, pK^{25}_2 10.20.** Mix the acid with an equal volume of *benzene; the *benzene is then distilled off to dehydrate the acid. After heating to 100° to remove most of the *benzene, the residue is distilled under vacuum and stored in sealed ampoules at 3°. [Eshelman et al. *Anal Chem* **32** 844 1960, DOI: 10.1021/ac60163a036; *Beilstein* **3** IV 1130.]

RS-Thiomalic (mercaptosuccinic) acid [70-49-5] $\text{C}_4\text{H}_6\text{O}_4\text{S}$, **M 150.2, m 153-154°, 155-157°, pK^{25}_1 3.64 (3.17), pK^{25}_2 4.64 (4.67), pK^{25}_3 10.37 (10.52).** Dissolve the acid in water and extract it several times with diethyl ether to remove impurities. The aqueous solution gave the acid on freeze-drying. [*Beilstein* **3** IV 472.]

Thiosemicarbazide [79-19-6] $\text{CH}_5\text{N}_3\text{S}$, **M 91.1, m 181-183°, pK^{25}_1 1.88, pK^{25}_2 12.81.** Crystallise thiosemicarbazide from H_2O (solubility is 20.3% w/w at 80°). The *hydrochloride* has **m 190-191°(dec, 184° also reported)**. It forms salts with heavy metals. [*Beilstein* **3** H 195, **3** I 79, **3** II 134, **3** III 315, **3** IV 374.]

Thiourea [62-56-6] $\text{CH}_4\text{N}_2\text{O}$, **M 76.1, m 179°, $\text{pK}^{20}_{\text{a}}$ -1.19 (aqueous H_2SO_4).** Crystallise thiourea from absolute EtOH, MeOH, acetonitrile or water. Dry it under vacuum over H_2SO_4 at room temperature. [*Beilstein* **3** IV 342.]

Tiglic acid (*trans*-2,3-dimethylacrylic acid) [80-59-1] $C_5H_8O_2$, M 100.1, m 63.5-64°, b 198.5°/atm, 95°/11mm, pK¹⁸ 4.96. Crystallise it from water. It is steam volatile and is soluble in organic solvents. [Beilstein 2 IV 1552.]

***trans*-Traumatic acid** (2-dodecene-1,12-dioic acid) [6402-36-4] $C_{12}H_{20}O_4$, M 228.3, m 165-166°, 150-160°/0.001mm, pK_{Est(1)}²⁰ ~4.2, pK_{Est(2)}²⁰ ~4.6. Crystallise the acid from EtOH, acetone or glyme. The *bis*-4-phenylphenacyl ester has m 144-145° (from EtOH). [Beilstein 2 III 1978, 2 IV 2279.]

1,2,3-Triaminopropane trihydrochloride [free base 21291-99-6] $C_8H_{11}N_3$, M 198.7, m 250° (sintering at 100°), pK₁²⁰ 3.72, pK₂²⁰ 7.95, pK₃²⁰ 9.59. Crystallise the trihydrochloride from EtOH or H₂O. The *free base* decomposes at 190°/760mm, but has b 92-93°/9mm without decomposition. [Beilstein 4 H 274, 4 III 630.]

Tribromochloromethane [594-15-0] $CBBr_3Cl$, M 287.2, m 55°, b 158-159.5°/~760mm, 160°/~760mm. Melt it, wash it with aqueous Na₂S₂O₃, dry it with BaO and fractionally crystallise from its melt. It also crystallises from EtOH and distils at atmospheric pressure. [Miller & Smyth *J Chem Phys* 24 816 1956, DOI: 10.1063/1.17426; Beilstein 1 H 68, 1 II 35, 1 III 91, 1 IV 85.]

Tribromofluoromethane [353-54-8] $CBBr_3F$, M 270.7, m -73.6°, b 106-107°/~760mm, d₄²⁵ 2.765, n_D²⁰ 1.525. It has been prepared from dry CBr₄ (1mole), SbF₃ (0.5 mole), and Br₂ (0.05 mole) by heating at 120-130° during 1 hr whereby CBr₃F distils out as it is formed. The reaction is completed by further heating at 150-160° for 0.5 hr. The crude product is washed with 5% aqueous tartaric acid, 10% aqueous sodium sulfite, 2% aqueous NaOH, then H₂O, dried (P₂O₅) and distilled to give CBr₃F (65-75%), b 106-107°/atm. Some CBr₂F₂ (~5-10%) with b 23.5°/atm is also formed and distils first. [Birchall & Haszeldine *J Chem Soc* 13 1959, DOI: 10.1039/JR9590000013.] Cu is added as a stabiliser [Hellmann et al. *J Am Chem Soc* 79 5654 1957, DOI: 10.1021/ja01578a018; Beilstein 1 IV 87.]

Tri-*n*-butylamine (TBA) [102-82-9] $C_{12}H_{27}N$, M 185.4, m -70°, b 68°/3mm, 120°/44mm, 216°/atm, d₄²⁰ 0.7788, n_D²⁰ 1.4294, pK²⁵ 9.93. Purify the amine by fractional distillation from sodium under reduced pressure. Pegolotti and Young [*J Am Chem Soc* 83 3251 1961, DOI: 10.1021/ja01476a018] heated the amine overnight with an equal volume of acetic anhydride, in a steam bath. The amine layer was separated and heated with water for 2 hours on the steam bath (to hydrolyse any remaining acetic anhydride). The solution was cooled, solid K₂CO₃ was added to neutralise any acetic acid that had been formed, and the amine was separated, dried (K₂CO₃) and distilled at 44mm pressure. Davis and Nakshbendi [*J Am Chem Soc* 84 2085 1926, DOI: 10.1021/ja00870a017] treated the amine with one-eighth of its weight of benzenesulfonyl chloride in aqueous 15% NaOH at 0-5°. The mixture was shaken intermittently and allowed to warm to room temperature. After a day, the amine layer was washed with aqueous NaOH, then water and dried with KOH. (This treatment removes primary and secondary amines.) It was further dried with CaH₂ and distilled under vacuum. [Beilstein 4 IV 554.]

Tri-*n*-butylammonium hydrobromide [37026-85-0] $C_{12}H_{27}N \cdot HBr$, M 308.3, m 75.2-75.9°. Crystallise the hydrobromide from ethyl acetate. [Beilstein 4 H 157, 4 III 292, 4 IV 555.]

Tri-*n*-butylammonium nitrate [33850-87-2] $C_{12}H_{27}N \cdot HNO_3$, M 304.5. Crystallise the nitrate from mixtures of *n*-hexane and acetone (95:5). Dry it over P₂O₅ in a vacuum. [Beilstein 4 H 157, 4 I 371, 4 II 633, 4 III 291, 4 IV 554.]

Tri-*n*-butylammonium perchlorate [14999-66-7] $C_{12}H_{27}N \cdot HClO_4$, M 285.5. Recrystallise the perchlorate from *n*-hexane. (Potentially explosive.) [Beilstein 4 IV 554.]

Tricarballic acid (propane-1,2,3-tricarboxylic acid) [99-14-9] $C_6H_8O_6$, M 176.1, m 156-161°, 166°, pK₁²⁵ 3.47, pK₂²⁵ 4.54, pK₃²⁵ 5.89. Crystallise the acid from diethyl ether. [Beilstein 2 IV 2366.]

Trichloroacetamide [594-65-0] $C_2H_2Cl_3NO$, M 162.4, m 139-141°, b 238-240°/atm. Its solution in xylene is dried with P₂O₅, then fractionally distilled. [Beilstein 2 IV 520.]

Trichloroacetic acid (TCA) [76-03-9] $\text{C}_2\text{HCl}_3\text{O}_2$, M 163.4, m $59.4\text{--}59.8^\circ$, b $196^\circ/\text{atm}$, $\text{pK}^{25}_{\text{a}}$ 0.51. Purify this strong acid (care) by fractional crystallisation from its melt, then crystallise it repeatedly from dry *benzene and store it over concentrated H_2SO_4 in a vacuum desiccator. It can also be crystallised from CHCl_3 or cyclohexane, and dried over P_2O_5 or $\text{Mg}(\text{ClO}_4)_2$ in a vacuum desiccator. Trichloroacetic acid can be fractionally distilled under reduced pressure from MgSO_4 . Layne, Jaffé and Zimmer [*J Am Chem Soc* **85** 435 1963, DOI: 10.1021/ja00887a014] dried trichloroacetic acid in *benzene by distilling off the *benzene-water azeotrope, then crystallised the acid from the remaining *benzene solution. Manipulations should be carried out under N_2 . [Toxic vapours, use a well ventilated fume cupboard.] [Beilstein **2** IV 508.]

Trichloroacetonitrile [545-06-2] $\text{C}_2\text{Cl}_3\text{N}$, M 144.4, m -42° , -44° , b $84.6^\circ/741\text{mm}$, $85.8\text{--}86^\circ/764\text{mm}$, $85.7^\circ/760\text{mm}$, d_4^{20} 1.441, n_D^{25} 1.4409. It is prepared by mixing trichloroacetamide (150g, obtained from ethyl trichloroacetate and ammonia) with an equal weight of P_2O_5 and heating at $\sim 200^\circ$ in an oil bath, and a further amount of P_2O_5 is added before distilling the nitrile off. The distillate is then carefully fractionated, and the fraction b $85.8\text{--}86^\circ/764\text{mm}$ is collected (yield 70-75%), and should contain $<0.02\%$ of amide: cf. IR (film) has a nitrile band at 2250 cm^{-1} , and is almost free from bands at ~ 1735 (CO) and >2502 (NH) cm^{-1} . The liquid shows no signs of change after several months, even with traces of moisture, provided that it is kept in away from light. [Davies & Jenkin *J Chem Soc* 2374 1954, DOI: 10.1039/JR9540002374, and references therein, Carpenter *J Org Chem* **27** 2085 1962, DOI: 10.1021/jo01053a043; Beilstein **2** H 212, **2** I 95, **2** II 201, **2** III 477, **2** IV 524]. **It is lachrymatory, a skin and eye irritant, and has been used as a fumigant.**

It is a useful reagent for selectively esterifying phosphoric acid, e.g. with benzyl alcohol in the presence of Et_2N it gives monobenzylphosphate with H_2PO_4 [Cramer & Baldauf *Chem Ber* **92** 370 1959, DOI: 10.1002/cber.19590920219], and converts a monophosphoric ester into a symmetrical pyrophosphate in the presence of pyridine [Cramer et al. *Justus Liebigs Ann Chem* **654** 180 1962, DOI: 10.1002/jlac.19626540118]. It is readily converted into its trichloroacetimidate esters by reaction with allylic alcohols in CH_2Cl_2 , in the presence of DBU at $\sim 0^\circ$ to ambient temperatures [Anderson & Overman *J Am Chem Soc* **125** 12412 2003, DOI: 10.1021/ja037086r; Kirsch et al. *Org Lett* **9** 911 2007, DOI: 10.1021/ol070110b]; similarly prepared trichloroacetimidates of allylic alcohols were shown to undergo ether-directed Pd(II)-catalysed aza-Claisen rearrangements [Jamieson & Sutherland *Tetrahedron* **63** 2132 2007 DOI:10.1016/j.tet.2006.12.067], and bis-trichloroacetimidates from 2-aminopropane-1,3-diols yielded dihydro-oxazines through an acid catalysed cyclisation [Rondot et al. *Org Lett* **9** 247 2007, DOI: 10.1021/ol0627202].

1,1,1-Trichloroethane [71-55-6] $\text{C}_2\text{H}_3\text{Cl}_3$, M 133.4, f -32.7° , b $74.0^\circ/\text{atm}$, d_4^{20} 1.337, n_D^{20} 1.4385. Wash it successively with concentrated HCl (or concentrated H_2SO_4), aqueous 10% K_2CO_3 (Na_2CO_3), aqueous 10% NaCl, dry it with CaCl_2 or Na_2SO_4 , and fractionally distil it. It can contain up to 3% dioxane as preservative. This is removed by washing successively with 10% aqueous HCl, 10% aqueous NaHCO_3 and 10% aqueous NaCl, and distilling over CaCl_2 before use. **TOXIC VAPOURS.** [Beilstein **1** IV 138.]

1,1,2-Trichloroethane [79-00-5] $\text{C}_2\text{H}_3\text{Cl}_3$, M 133.4, f -36.3° , b $113.6^\circ/\text{atm}$, d_4^{20} 1.435, n_D^{20} 1.472. Purify the chloroethane as for 1,1,1-trichloroethane above. **TOXIC VAPOURS.** [Beilstein **1** IV 139.]

Trichloroethylene (TCE) [79-01-6] C_2HCl_3 , M 131.4, m -84.8° , b $-12.4^\circ/10\text{mm}$, $20^\circ/60\text{mm}$, $48^\circ/200\text{mm}$, $67^\circ/400\text{mm}$, $86.9^\circ/760\text{mm}$, $87.2^\circ/\text{atm}$, d_4^{20} 1.463, n_D^{21} 1.4767. Trichloroethylene undergoes decomposition in a similar way as CHCl_3 , giving HCl, CO, COCl_2 and organic products. It reacts with KOH, NaOH and 90% H_2SO_4 , and forms azeotropes with water, MeOH, EtOH, and acetic acid. It is purified by washing successively with 2M HCl, water and 2M K_2CO_3 , then dried with K_2CO_3 and CaCl_2 , then fractionally distilled before use. It has also been steam distilled from 10% $\text{Ca}(\text{OH})_2$ slurry, most of the water being removed from the distillate by cooling to -30° to -50° and filtering off the ice through chamomile skin: the trichloroethylene is then fractionally distilled at 250mm pressure and collected in a blackened container. **ANAESTHETIC, TOXIC VAPOURS.** [Carlisle & Levine *Ind Eng Chem (Anal Ed)* **24** 1164 1932, DOI: 10.1021/ie50274a019; Beilstein **1** IV 712.]

1,1,2-Trichloro-1,2,2-trifluoroethane [76-13-1] $\text{C}_2\text{Cl}_3\text{F}_3$, M 187.4, b $47.6^\circ/760\text{mm}$, d_4^{20} 1.576, n_D^{20} 1.360. Wash it with water, then with weak alkali. Dry it with CaCl_2 or H_2SO_4 and distil it. [Locke et al. *J Am Chem Soc* **56** 1726 1934, DOI: 10.1021/ja01323a023; Beilstein **1** III 157, **1** IV 142.]

Tridecanoic acid [638-53-9] $C_{13}H_{26}O_2$, M 214.4, m 41.8°, 44.5-45.5° (several forms), b 199-200°/24mm, 236°/100mm, $pK_{Est} \sim 5.0$. Recrystallise the acid from acetone or distil it in a vacuum. [Beilstein 2 IV 1117.] The *methyl ester* [1731-88-0] $C_{14}H_{24}O_2$, M 228.4 has m 5.5°, b 131°/4mm. [Beilstein 2 IV 1118.]

7-Tridecanone (dihexyl ketone) [462-18-0] $C_{13}H_{26}O$, M 198.4, m 33°, b 255°/766mm, 264°/atm. Crystallise the ketone from EtOH. [Beilstein 1 H 715.]

Tri-*n*-dodecylamine (Hydrogen ionophore I) [102-87-4] $C_{36}H_{75}N$, M 522.0, m 15.7°, b 220-228°/0.03mm, d_4^{20} 0.833, n_D^{20} 1.4577, $pK_{Est} \sim 11.0$. Distil tridodecylamine under high vacuum and N_2 , and store it in the absence of CO_2 . It can be recrystallised from 95%EtOH/* C_6H_6 at low temperature under vacuum. The *hydrochloride* has m 78-79°. [Ra et al. *J Org Chem* 09 259 1944, DOI: 10.1021/jo01185a009; Beilstein 4 III 413, 4 IV 801.]

Tri-*n*-dodecylammonium nitrate [2305-34-2] $C_{36}H_{75}N \cdot HNO_3$, M 585.0. Crystallise the salt from *n*-hexane/acetone (95:5) and keep it in a desiccator over P_2O_5 under vacuum. [Beilstein 4 IV 801 for tridodecylamine.]

Tri-*n*-dodecylammonium perchlorate [5838-82-4] $C_{36}H_{75}N \cdot HClO_4$, M 622.4. Recrystallise the salt from *n*-hexane or acetone and keep it in a desiccator over P_2O_5 . (Potentially explosive.)

Triethanolamine (2,2',2''-nitriloethanol) [102-71-6] $C_6H_{15}NO_3$, M 149.2, m 20-22°, b 190-103°/5mm, 206-207°/15mm, 335.4°/760mm, d_4^{20} 1.124, n_D^{20} 1.485, pK^{25} 7.92. Shake the amine gently with Linde type 4A molecular sieves for 24 hours, filter and fractionate it under a vacuum, and preferably in the presence of N_2 . Store it in dark stoppered bottles under N_2 as it is *hygroscopic*, and turns brown in air and light. It has a strong ammoniacal odour (like diethanolamine). It is miscible with H_2O , MeOH and Me_2CO and its solubilities at 25° in *n*-heptane, Et_2O and * C_6H_6 are 0.4%, 1.6% and 4.2%, respectively. [See diethanolamine above, Beilstein 4 IV 1524.]

Triethanolamine hydrochloride [637-39-8] $C_6H_{15}NO_3 \cdot HCl$, M 185.7, m 177°, 177-179°, pK^{25} 7.92 (free base). Crystallise the salt from EtOH. Dry it at 80°. [Beilstein 4 IV 1525.]

1,1,2-Triethoxyethane [4819-77-6] $C_8H_{18}NO_3$, M 162.2, b 74°/28mm, 164°/~760mm, 167.2°/760mm, d_4^{20} 0.897, n_D^{20} 1.401. Dry it with Na_2SO_4 , and distil it. [McElvain & Walters *J Am Chem Soc* 64 1963 1942, DOI: 10.1021/ja01260a060; Beilstein 1 H 818, 1 I 418, 1 III 3184, 1 IV 3958.]

Triethylamine [121-44-8] $C_6H_{15}N$, M 101.2, m -115°, b 88.8°/atm, 89.4°/atm, d_4^{20} 0.7280, n_D^{20} 1.4005, pK^{25} 10.82. Dry triethylamine with $CaSO_4$, $LiAlH_4$, Linde type 4A molecular sieves, CaH_2 , KOH, or K_2CO_3 , then distil it, either alone or from BaO, sodium, P_2O_5 or CaH_2 . It has also been distilled from zinc dust, under nitrogen. To remove traces of primary and secondary amines, triethylamine has been refluxed with acetic anhydride, benzoic anhydride, phthalic anhydride, then distilled, refluxed with CaH_2 (ammonia-free) or KOH (or dried with activated alumina), and again distilled. Another purification method involved refluxing for 2 hours with *p*-toluenesulfonyl chloride, then distilling. Grovenstein and Williams [*J Am Chem Soc* 83 412 1961, DOI: 10.1021/ja01463a039] treated triethylamine (500ml) with benzoyl chloride (30ml), filtered off the precipitate, and refluxed the liquid for 1 hour with a further volume of benzoyl chloride (30ml). After cooling, the liquid was filtered, distilled, and allowed to stand for several hours with KOH pellets. It was then refluxed with, and distilled from, stirred molten potassium. Triethylamine has been converted to its *hydrochloride* (see below), crystallised from EtOH (to m 254°), then liberated with strong aqueous NaOH, dried with solid KOH and distilled from sodium under N_2 . It is a strong base and should not be inhaled. [Beilstein 4 H 99, 4 I 348, 4 II 593, 4 III 194, 4 IV 322.]

Triethylammonium bromide [636-70-4] $C_6H_{15}N \cdot HBr$, M 229.1, m 248°. Equimolar portions of triethylamine and aqueous solutions of HBr in acetone are mixed with cooling. The precipitated salt is washed with anhydrous acetone and dried in a vacuum for 1-2 hours. [Odinokov et al. *JCS Faraday Trans 2* 80 899 1984, DOI: 10.1039/F29848000899.] Recrystallise it from $CHCl_3$ or EtOH. [Beilstein 4 IV 322.]

Triethylammonium chloride [554-68-7] $\text{C}_6\text{H}_{15}\text{N} \cdot \text{HCl}$, M 137.7, m 257-260°(dec), 261°(dec). Purify it like the bromide above. [Beilstein 4 IV 327.]

Triethylammonium iodide [4636-73-1] $\text{C}_6\text{H}_{15}\text{N} \cdot \text{HI}$, M 229.1, m 181°. Purify it as for triethylammonium bromide, except the solution for precipitation should be precooled in acetone at -10°, and the precipitate is twice recrystallised from a cooled acetone/hexane mixture at -10°. Store it in the dark. [Beilstein 4 IV 327.]

Triethylammonium trichloroacetate [4113-06-8] $\text{C}_6\text{H}_{15}\text{N} \cdot \text{C}_2\text{Cl}_3\text{O}_2\text{H}$, M 263.6. Equimolar solutions of triethylamine and trichloroacetic acid in *n*-hexane are mixed at 10°. The solid so obtained is recrystallised from CHCl_3 /*benzene. [Hoigné & Gäumann *Helv Chim Acta* 42 437 1959, DOI: 10.1002/hlca.19590420206; Beilstein 4 IV 330.]

Triethylammonium trifluoroacetate [454-49-9] $\text{C}_6\text{H}_{15}\text{N} \cdot \text{C}_2\text{F}_3\text{O}_2\text{H}$, M 196.2. Purify as for the corresponding trichloroacetate but in Et_2O . Evaporation of the Et_2O gives the salt as a colourless viscous liquid at ambient temperature. [Emmons et al. *J Am Chem Soc* 76 3472 1954, DOI: 10.1021/ja01642a031; Beilstein 4 IV 330.]

Triethylene glycol [112-27-6] $\text{C}_6\text{H}_{14}\text{O}_4$, M 150.2, m -7°, b 125-127°/0.1mm, 165°/14mm, 278°/760mm, d_4^{15} 1.1274, n_D^{15} 1.4578. Dry the glycol with CaSO_4 for 1 week, then it is repeatedly and very slowly fractionally distilled under a vacuum. Store it in a vacuum desiccator over P_2O_5 . It is very *hygroscopic*. [Matignon et al. *Bull Chem Soc Fr* 1 [5] 1308 1934, Beilstein 1 IV 2400.]

Triethylene glycol dimethyl ether (triglyme) $\text{C}_8\text{H}_{18}\text{O}_4$, [112-49-2] M 178.2, m -45°, b 20°/0.9mm, 103.5°/10mm, 216°/atm, 225°, d_4^{20} 0.987, n_D^{20} 1.425. Reflux it with, and distil it from sodium hydride or LiAlH_4 . [Beilstein 1 IV 2401.]

Triethylenetetramine (TRIEN, TETA, trientine, $\text{H}_2\text{NCH}_2\text{CH}_2\text{NHCH}_2\text{CH}_2\text{NHCH}_2\text{CH}_2\text{NHCH}_2\text{CH}_2\text{NH}_2$) [112-24-3] $\text{C}_6\text{H}_{18}\text{N}_4$, M 146.2, m 12°, b 157°/20mm, 266-267°/atm, d_4^{20} 0.971, n_D^{20} 1.497, pK_1^{25} 3.32, pK_2^{25} 6.67, pK_3^{25} 9.20, pK_4^{25} 9.92. Dry the amine with sodium, then distil it under a vacuum. Further purification has been *via* the nitrate or the chloride salts. For example, Jonassen and Strickland [*J Am Chem Soc* 80 312 1958, DOI: 10.1021/ja01535a015] separated TRIEN from admixture with TREN (38%) by solution in EtOH, cooling to approximately 5° in an ice-bath and adding concentrated HCl dropwise from a burette, keeping the temperature below 10°, until all of the white crystalline precipitate of TREN.HCl (see below) had formed and was removed. Further addition of HCl then precipitated thick, creamy white TRIEN.HCl (see following entry) which was crystallised several times from hot water by adding an excess of cold EtOH. The crystals were finally washed with Me_2CO , then Et_2O and dried in a vacuum desiccator. The *hydrate* [305808-21-3, $x\text{H}_2\text{O}$] has m 45-48°, b 272°/atm. [Beilstein 4 H 255, 4 II 695, 4 III 542, 4 IV 1242.]

Triethylenetetramine tetrahydrochloride (TRIEN HCl) [4961-10-4] $\text{C}_6\text{H}_{18}\text{N}_4 \cdot 4\text{HCl}$, M 292.1, m 266-270°, 269-270°. Crystallise the salt repeatedly from hot water by precipitation with cold EtOH or EtOH/HCl. Wash it with acetone and absolute EtOH and dry it in a vacuum oven at 80° (see TRIEN above). The *tetrabenzoyl* derivative has m 230° after recrystallisation from boiling EtOH. [Peacock *J Chem Soc* 1518 1936, DOI: 10.1039/JR9360001518; Beilstein 4 H 255, 4 II 695, 4 III 543.]

Triethyl orthoformate (ethyl orthoformate, 1,1,1-triethoxymethane) [122-51-0] $\text{C}_7\text{H}_{16}\text{N}_3$, M 148.2, m -76°, b 60°/30mm, 144-146°/760mm, d_4^{20} 0.891, n_D^{20} 1.392. Fractionate it first at atmospheric pressure, then in a vacuum. If impure, then shake it with aqueous 2% NaOH, dry it with solid KOH and distil it from sodium through a 20cm Vigreux column. Alternatively, wash it with H_2O , dry it over anhydrous K_2CO_3 , filter and fractionate it through a Widmer column. [Sah & Ma *J Am Chem Soc* 54 2964 1932, DOI: 10.1021/ja01346a048; Ohme & Schmitz *Justus Liebigs Ann Chem* 716 207 1968, DOI: 10.1002/jlac.19687160130; Beilstein 2 IV 25.] **IRRITANT** and **FLAMMABLE**.

Triethyloxonium tetrafluoroborate [368-39-8] $\text{C}_6\text{H}_{15}\text{O} \cdot \text{BF}_4$, M 190.0, m 92-93°(dec). Crystallise it from

diethyl ether. It is *very hygroscopic*, and must be handled in a dry box and stored at 0°. [Meerwein *Org Synth Coll Vol* **5** 1096 1973, DOI: 10.15227/orgsyn.046.0120.] Pure material should give a clear and colourless solution in dichloromethane (1 in 50, w/v is a 0.1M solution available commercially). [Beilstein **1** IV 1322.]

Trifluoroacetic acid (TFA) [76-05-1] $\text{C}_2\text{HF}_3\text{O}_2$, **M 114.0**, **m -15.5°**, **b 72.4°/atm**, d_4^{20} **1.494**, n_D^{20} **1.2850**, **pK²⁵ 0.52**. Although an improved preparation of the acid (in 87% yield) involving the oxidation of 2,3-dichloro-1,1,1,4,4,4-hexafluorobut-2-ene (b 65-66°/745mm, [303-04-8], steam volatile, d_4^{25} **1.605**, n_D^{20} **1.3458**) via oxidation with $\text{KMnO}_4/\text{KOH}/95^\circ/8\text{-}10$ hours has been reported [Henne & Trott *J Am Chem Soc* **69** 1820 1947, DOI: 10.1021/ja01199a076], the next purification procedure should be avoided. The purification of trifluoroacetic acid, reported in earlier editions of this work, by refluxing over KMnO_4 for 24 hours and slowly distilling has resulted in very **SERIOUS EXPLOSIONS** on various occasions, but not always. This apparently depends on the source and/or age of the acid. The method is **NOT RECOMMENDED**. It forms an azeotropic mixture of acid and H_2O (80:20) which boils at 103-105°/745mm. Water can be removed by adding trifluoroacetic anhydride (0.05%, to diminish water content) and distilling. [Conway & Novak *J Phys Chem* **81** 1459 1977, DOI: 10.1021/j100530a006]. It can be refluxed and distilled from P_2O_5 , but do not use an excess of P_2O_5 as it produces the anhydride (see below and Schmidt & Staub *Chem Ber* **87** 388 1954, DOI:10.1002/cber.19540870317) which can be separated by fractional distillation because the anhydride distils about 30° lower than the acid at atmospheric pressure. It is further purified by fractional crystallisation by partial freezing and again distilled. It is a strong acid and attacks skin (wear gloves) and eyes (wear safety glasses). It is soluble in H_2O and most organic solvents. **Highly TOXIC vapour — do not inhale it. Work in an efficient fume hood.** [Roepke *Biotechnol Forum Eur* **7** 161 1990, Beilstein **2** II 186, **2** III 426, **2** IV 458.]

Trifluoroacetic anhydride (TFAA) [407-25-0] $\text{C}_4\text{F}_6\text{O}_3$, **M 210.0**, **m -65°**, **b 38-40°/760mm**, d_4^{20} **1.508**. Purification by distilling over KMnO_4 , as for the acid above, is **EXTREMELY DANGEROUS** due to the possibility of **EXPLOSION** (see preceding acid). It is best purified by distilling from P_2O_5 slowly, and collecting the fraction boiling at 39.5°. Store it in a dry atmosphere. It is soluble in most organic solvents, but reacts with protic solvents. **Highly TOXIC vapour and attacks skin, work in an efficient fume hood — see previous entry.** [Tedder *Chem Rev* **55** 787.1955, DOI: 10.1021/cr50005a001; Beilstein **2** II 186, **2** III 427, **2** IV 469.]

1,1,1-Trifluoroacetone [421-50-1] $\text{C}_3\text{H}_3\text{F}_3\text{O}$, **M 112.1**, **b 21°/atm**, **22°/atm**, d_4^{25} **1.252**, n_D^{20} **1.3**. The ketone was obtained by refluxing ethyl 4,4,4-trifluoroacetate (see [372-31-6]) for 2 hours with 10% aqueous H_2SO_4 and collecting the distillate into a receiver containing P_2O_5 , and redistilling it from P_2O_5 . The **2,4-dinitrophenylhydrazone** has **m 139°** [Henne et al. *J Am Chem Soc* **69** 1820 1947, DOI: 10.1021/ja01199a076] synthesis in a few steps [Huguenot & Brigaud *J Org Chem* **71** 7075 2006, DOI: 10.1021/jo0607717]. It has also been used to synthesise 2-trifluoromethyl-7-azaindoles from 2,6-dihalopyridines [Schirok et al. *Synthesis* 251 2007, DOI: 10.1055/s-2006-958935].

2,2,2-Trifluoroethanol [75-89-8] $\text{C}_2\text{H}_3\text{F}_3\text{O}$, **M 100.0**, **m -44°**, **b 72.4°/738mm**, **77-80°/atm**, n_D^{20} **1.30**, d_4^{20} **1.400**, **pK²⁵ 12.8**. Dry it with CaSO_4 and a little NaHCO_3 (to remove traces of acid) and distil it. It is used as a calibration standard for NMR spectroscopy. **Highly TOXIC vapour.** [Beilstein **1** IV 1370.]

Trifluoromethanesulfonic anhydride (triflic anhydride) [358-23-6] $\text{C}_2\text{F}_6\text{O}_5\text{S}_2$, **M 282.1**, **b 82-85°/atm**, **84°/atm**, d_4^{20} **1.71**, n_D^{20} **1.322**. Distil it through a short Vigreux column. It can be freshly prepared from the anhydrous acid (11.5g) and P_2O_5 (11.5g, or half this weight) by setting aside at room temperature for 1 hour, distilling off volatile products then distil it through a short Vigreux column. It is readily hydrolysed by H_2O and decomposes appreciably after a few days to liberate SO_2 and produce a viscous liquid. Store it dry at low temperatures. [Burdon et al. *J Chem Soc* 2574 1957, DOI: 10.1039/JR9570002574; Beard et al. *J Org Chem* **38** 3673 1973, DOI: 10.1021/jo00961a003; Beilstein **3** IV 35.] **Highly TOXIC vapour.**

1,1,1-Trifluoro-2,4-pentanedione (α,α,α -trifluoroacetylacetone, tfacac or facac) [367-57-7] $\text{C}_5\text{H}_5\text{F}_3\text{O}_2$, **M 154.1**, **b 105-107°/atm**, **107°/760mm**, d_4^{25} **1.27**, n_D^{20} **1.3890**. The diketone was prepared from ethyl trifluoroacetate (74.5g) and Na wire (10g, or an equivalent of NaOEt) in Et_2O (5ml) at 0°, by adding Me_2CO (80ml) at a rate to maintain a steady reaction, kept at 0° for 2 hours, and then allowed to warm to room tempera-

ture. Ice-water (5ml) was added, the aqueous layer was acidified with AcOH, and the Cu complex was precipitated with a warm saturated solution of $\text{Cu}(\text{OAc})_2$. The **copper trifluoroacetylacetone** crystallised in long blue needles from EtOH and had **m 189°**; its UV has λ_{max} at 240nm (ϵ 19,500) and 294nm (ϵ 31,000) in EtOH, and the effect of fluorine substitution on the complexing properties has been studied in detail by Calvin and coworkers [Belford et al. *J Inorg Nucl Chem* **2** 11 1956, DOI: 10.1016/0022-1902(56)80100-0].

An ethereal mixture of the complex (partly dissolved) was shaken with 2N H_2SO_4 , then H_2O , the organic layer was separated, dried (Na_2SO_4), filtered and distilled to give a 62% yield of **trifluoroacetylacetone**. Alternatively, the metal can be removed by bubbling H_2S through the ethereal mixture, the CuS was filtered off and the filtrate was distilled. [Reid & Calvin *J Am Chem Soc* **72** 2948 1950, DOI: 10.1021/ja01163a038; Henne et al. *J Am Chem Soc* **69** 1819 1947, DOI: 10.1021/ja01199a076; Haszeldine et al. *J Chem Soc* 609 1951, DOI: 10.1039/JR9510000609; Belford et al. *J Inorg Nucl Chem* **2** 11 1956, DOI:10.1016/0022-1902(56)80100-0].

Its UV has λ_{max} at 284nm (ϵ 10,500) in CHCl_3 ; the FT-IR (film) has ν_{max} at 1604.7, 1422.3, 1369.5, 1282.2, 1154.7, 887.4, 857.4, 795.2, 726.4 cm^{-1} ; and the ^1H NMR (300MHz, CDCl_3 , TMS) has δ at 2.36 (s), 4.60 (s, 3H, CH_3), 8.30 (s, CH); and the ^{13}C NMR (75MHz, CDCl_3 , TMS) has δ at 193.47, 176.71, 176.22, 175.74, 175.25, 127.75, 119.00, 115.25, 111.50, 96.37, 96.35, 24.83. It enolises in aqueous solution, and rates of enolisation were determined by Ried and Calvin (*vide infra*) in H_2O , 0.1N and 0.5N HCl, and in aqueous EtOH of varying concentrations; and are temperature dependent. Thus at equilibrium, at 0.08-0.09M solution of *tfacac* and 25°, 16% aqueous EtOH contains 2.5% of enol+enolate, whereas at 95% it contains 20.7% of enol+enolate. This enolisation should be taken into account when measuring spectra and other physical parameters.

Complexation with Cu, Fe and Al can be used to extract the metals from aqueous medium (acetate buffered) by a 0.10M of the diketone in CHCl_3 [Scribner et al. *Anal Chem* **37** 1136 1965, DOI: 10.1021/ac60228a017], and Be into organic solvents [Scribner et al. *Anal Chem* **38** 1779 1966, DOI: 10.1021/ac60244a041]. The *oxime*, **m 86-87°**, crystallises from H_2O or aqueous EtOH. [Beilstein **1** III 3123, **1** IV 3680.]

Trimethylamine [75-50-3] $\text{C}_3\text{H}_9\text{N}$, **M 59.1**, **m -117°**, **b 3.5°/atm**, **pK²⁵ 9.80**. Dry triethylamine by passing the gas through a tower filled with solid KOH. Water and impurities containing labile hydrogen were removed by treatment with freshly sublimed, ground, P_2O_5 . It has been refluxed with acetic anhydride, and then distilled through a tube packed with HgO and BaO. [Comyns *J Chem Soc* 1557 1955, DOI: 10.1039/JR9550001557.] For more extensive purification, trimethylamine is converted to the hydrochloride, crystallised (see below), and regenerated by treating the hydrochloride with excess aqueous 50% KOH; the gas is passed through a CaSO_4 column into a steel cylinder containing sodium ribbon. After 1-2 days, the cylinder is cooled to -78° and hydrogen and air are removed by pumping. [Day & Felsing *J Am Chem Soc* **72** 1698 1950, DOI: 10.1021/ja01160a077.] Me_3N has been distilled using trap-to-trap methods and degassed by freeze-pump-thaw [Halpern et al. *J Am Chem Soc* **108** 3907 1986, DOI: 10.1021/ja00274a009]. It is commercially available under pressure in tin cylinders. It is available commercially as a ~ 4.2M (31-35 wt%) in EtOH, and as ~25 w% and ~45 w% solutions in H_2O . [Adams & Brown, *Org Synth Coll Vol I* 528 1941, DOI: 10.1522/orgsyn.001.0075; Beilstein **4** H 43, **4** I 322, **4** II 553, **4** III 99, **4** IV 134.]

Trimethylamine hydrochloride [593-81-7] $\text{C}_3\text{H}_9\text{N} \cdot \text{HCl}$, **M 95.7**, **m >280°(dec)**, **283-284°(dec)**. The salt crystallises from CHCl_3 , EtOH or *n*-propanol, and is dried under vacuum. It also crystallises from *benzene/MeOH, MeOH/diethyl ether and is dried under vacuum over paraffin wax and H_2SO_4 . It is kept over P_2O_5 as it is *hygroscopic*. [Beilstein **4** H 262, **4** I 419, **4** IV 138.]

Trimethylamine hydroiodide [20230-89-1] $\text{C}_3\text{H}_9\text{N} \cdot \text{HI}$, **M 186.0**, **m 263°**. It crystallises from MeOH.

Trimethylolpropane (1,1,1-tris-hydroxymethylpropane, 2-ethyl-2-hydroxymethyl-1,3-propanediol) [77-99-6] $\text{C}_6\text{H}_{11}\text{O}_3$, **M 134.2**, **m 57-59°, 60-62°**, **b 159-161°/2mm**. Crystallise it from acetone and ether, and it distils at high vacuum. [Beilstein **1** III 2349, **1** IV 2786.]

2,2,3-Trimethylpentane [564-02-3] C_8H_{18} , **M 114.2**, **b 109.8°/atm**, **d₄²⁰ 0.7161**, **n_D²⁰ 1.40295**, **n_D²⁵ 1.40064**. Purify it by azeotropic distillation with 2-methoxyethanol, which is subsequently washed out with water. The trimethylpentane is then dried and fractionally distilled. [Forziati et al. *J Res Nat Bur Stand* **36** 129 1946, DOI:org/10.6028/jres.036.005; Beilstein **1** IV 439.]

2,2,4-Trimethylpentane (isooctane) [540-84-1] C_8H_{18} , M 114.2, m -107° , b $99.2^\circ/\text{atm}$, d_4^{20} 0.693, n_D^{20} 1.39145, n_D^{25} 1.38898. Distil isooctane from sodium, pass it through a column of silica gel or activated alumina (to remove traces of olefins), and again distilled from sodium. Extract it repeatedly with concentrated H_2SO_4 , then agitate it with aqueous $KMnO_4$, wash it with water, dry ($CaSO_4$) and distil it. Purify it also by azeotropic distillation with EtOH, which is subsequently washed out with water, and the trimethylpentane is dried and fractionally distilled. [Forziati et al. *J Res Nat Bur Stand* **36** 129 1946, DOI: org/10.6028/jres.036.005.] [Beilstein **1** IV 439.]

2,4,4-Trimethylpent-2-ene (β -diisobutylene) [107-40-4] C_8H_{16} , M 112.2, m -106° , b $104^\circ/\text{atm}$, d_4^{20} 0.720, n_D^{20} 1.4160. Fractionate it under N_2 as it is highly flammable. [Beilstein **1** III 848, **1** IV 891.]

Trimethylsulfonium iodide [2181-42-2] C_3H_9S . I, M 204.1, m $211-212.5^\circ(\text{dec})$, $215-220^\circ(\text{dec})$. Crystallise the iodide from EtOH. [Emeleus & Heal *J Chem Soc* 1126 1946, DOI: 10.1039/JR9460001126; Swain & Kaiser *J Am Chem Soc* **80** 4089 1958, DOI: 10.1021/ja01548a067; Borredon et al. *J Org Chem* **55** 501 1990, DOI: 10.1021/jo00289a022; Bouda et al. *Synth Commun* **17** 503 1987, DOI: 10.1080/00397918708056436; Beilstein **1** IV 1280.]

Trimyristin [glyceryl trimyristate, glyceryl tri(tetradecanoate)] [555-45-3] $C_{45}H_{86}O_6$, M 723.2, m 56.5° , $56-57^\circ$, d^{60} 0.885. Crystallise it from diethyl ether. [Beilstein **2** IV 1135.]

Tri-*n*-octylamine [1116-76-3] $C_{24}H_{51}N$, M 353.7, b $164-168^\circ/0.7\text{mm}$, $365-367^\circ/760\text{mm}$, d_4^{20} 0.813, n_D^{20} 1.450, pK^{25} 10.65. It is converted to the amine hydrochloride etherate which is recrystallised four times from diethyl ether at -30° (see below). Neutralisation of this salt regenerates the free amine which distil under high vacuum. [Wilson & Wogman *J Phys Chem* **66** 1552 1962, DOI: 10.1021/j100814a513.] Distil the strong base amine at $<1-2\text{mm}$ pressure. [Beilstein **4** H 196, **4** III 382, **4** IV 754.]

Tri-*n*-octylammonium chloride [1188-95-0] $C_{24}H_{51}N$. HCl, M 384.2, m $78-79^\circ$, pK^{25} 8.35 (in 70% aqueous EtOH). Crystallise it from Et_2O , then *n*-hexane (see above). [Borrows et al. *J Chem Soc* 197 1947, DOI: 10.1039/JR9470000197; Beilstein **4** H 196.]

Tri-*n*-octylammonium perchlorate [2861-99-6] $C_{24}H_{51}N$. $HClO_4$, M 454.2, m $>300^\circ(\text{dec})$. Crystallise the perchlorate from *n*-hexane. (Possibly explosive.) [Beilstein **4** IV 754.]

Tripalmitin [glyceryl tripalmitate, glyceryl tri(hexadecanoate)] [555-44-2] $C_{51}H_{98}O_6$, M 807.4, m 66.4° . Crystallise it from acetone, diethyl ether or EtOH. It exists in an α -form (m 56.0°), a β' -form (m 63.5°) and a β -form (m 65.5°). [Beilstein **2** H 373, **2** I 167, **2** II 340, **2** III 971, **2** IV 1176.]

Tri-*n*-propylamine [102-69-2] $C_9H_{21}N$, M 143.3, m -93.5° , b $40-42^\circ/11\text{mm}$, $155-158^\circ/\text{atm}$, 156.5° , d_4^{20} 0.757, n_D^{20} 1.419, pK^{25} 10.66. Dry the amine with KOH and fractionally distil it. Also reflux it with toluene-*p*-sulfonyl chloride and with KOH, then fractionally distil it. The distillate, after addition of 2% phenyl isocyanate, was redistilled and the residue fractionally distilled from sodium. It is a strong base. [Takahashi et al. *J Org Chem* **52** 2666 1987, DOI: 10.1021/jo00389a007; Beilstein **4** IV 470.]

Tris(2-aminoethyl)amine (TREN) $C_6H_{18}N$, [4097-89-6] M 146.2, b $114^\circ/15\text{mm}$, $263^\circ/744\text{mm}$, d_4^{20} 0.977, n_D^{20} 1.498, pK_1^{25} 8.42, pK_2^{25} 9.44, pK_3^{25} 10.13. For a separation from a mixture containing 62% TRIEN, see entry under triethylenetetramine [112-24-3] above. Also purify it by conversion to the hydrochloride (see below), recrystallise it and regenerate the free base [Xie & Hendrickson *J Am Chem Soc* **109** 6981 1987, DOI: 10.1021/ja00257a013].

Tris(2-aminoethyl)amine trihydrochloride (TREN.HCl) [14350-52-8] $C_6H_{18}N$. HCl, M 255.7, m $300^\circ(\text{dec})$. Crystallise the salt several times by dissolving it in the minimum of hot water and precipitating it with excess of cold EtOH. The precipitate is washed with acetone, then diethyl ether and dried in a vacuum desiccator. [Beilstein **4** H 256, **4** II 695, **4** III 545, **4** IV 1250.]

Tris(dimethylamino)methane (*N,N,N',N',N'',N''*-hexamethylmethanetriamine) [5762-56-1] $C_7H_{19}N_3$, M

145.3, b 42-43°/12mm, n_D^{20} 1.4349, $pK_{Est} \sim 10$. Dry it over KOH and distil it through a Vigreux column at water pump vacuum. Store it in the absence of CO₂. [Bredereck et al. *Chem Ber* **101** 1885 1968, DOI: 10.1002/cber.19681010541; and *Angew Chem Int Ed* **5** 132 1966, DOI: 10.1002/anie.196601321.]

1,1,1-Tris(hydroxymethyl)ethane (2-hydroxymethyl-2-methyl-1,3-propanediol) [77-85-0] C₅H₁₂NO₃, M 120.2, m 193-195°, 200°. Dissolve it in hot tetrahydrofuran, filter and precipitate it with hexane. It has also been crystallised from acetone/water (1:1). Dry it in a vacuum. [Beilstein **1** H 520, **1** IV 2780.]

Tris(hydroxymethyl)methylamine (TRIS, tris(hydroxymethyl)aminomethane, Trizma, trometamol) [77-86-1] C₄H₁₁NO₃, M 121.1, m 172°, b 219-220°/10mm, pK^{25} 8.07. TRIS can ordinarily be obtained in highly pure form suitable sources for use as an acidimetric standard. If only impure material is available, it should be crystallised from 20% EtOH, aqueous MeOH (m 171.1°) or isopropanol (m 172-173°). Dry it in a vacuum desiccator over P₂O₅ or CaCl₂. Its solubility in H₂O at 20° is 80% w/v, and the pH of a 0.1 M in H₂O is 10.36. Alternatively, it is dissolved in twice its weight of water at 55-60°, filtered, concentrated to half its volume and poured slowly, with stirring, into about twice its volume of EtOH. The crystals which separate on cooling to 3-4° are filtered off, washed with a little MeOH, air dried by suction, then finally ground and dried in a vacuum desiccator over P₂O₅. It has also been recrystallised from water, MeOH or aqueous MeOH, and vacuum dried at 80° for 2 days. The *amidosulfate* [NH₂SO₃H] salt has m 103.5°. [Beilstein **4** H 303, **4** III 857, **4** IV 1903.]

Tris(hydroxymethyl)methylammonium hydrochloride (TRIS-HCl) [1185-53-1] C₄H₁₁NO₃·HCl, M 157.6, m 149-150°(dec). Crystallise the salt from 50% EtOH, then from 70% EtOH. TRIS-hydrochloride is also available commercially in a highly pure state. Otherwise, recrystallise it from 50% EtOH, then 70% EtOH, and dry it below 40° to avoid risk of decomposition. [Beilstein **4** H 304.]

N-Tris(hydroxymethyl)methyl-2-aminomethanesulfonic acid (TES) C₆H₅NO₆S [7365-44-8] M 229.3, m 224-226°(dec), 231°(dec), pK^{37} 7.14, pK^{20} 7.50, pK^0 7.92. Crystallise the acid from hot EtOH containing a little water. It is a useful buffer.

Tris(hydroxymethyl)nitromethane [2-(hydroxymethyl)-2-nitro-1,3-propanediol] [126-11-4] C₄H₉NO₅, M 151.1, m 160°(dec), 174-175°(dec, tech. grade), 214°(pure). Crystallise it from CHCl₃/ethyl acetate or ethyl acetate/*benzene. It is an acid and a 0.1M solution in H₂O has pH 4.5. **IRRITANT.** [Beilstein **1** H 520, **1** IV 2777.]

Tris[2-(methylamino)ethyl]amine [65604-89-9] C₉H₂₄N₄, M 188.3, b 77-78°/0.1mm, d_4^{20} 0.896, $pK_{Est(1)} \sim 8.8$, $pK_{Est(2)} \sim 9.4$, $pK_{Est(3)} \sim 10.4$. If this strong base contains carbonate (check IR) it should be shaken with solid KOH, decanted and distilled at high vacuum to give a colourless oil with a strong amine odour. Store it in the dark under N₂ as it absorbs CO₂ in moist air. It is synthesised in two steps. Ethyl chloroformate (33.4g, 310mmol, lachrymatory, see [541-41-3]) is added dropwise to a mixture of TREN (29.2g, 200mmol, see [4097-89-6]) in *C₆H₆ (225ml) and H₂O (100ml), and cooled to 5°. Then a solution of KOH (36.4g, 650mmol) in H₂O (35ml) is added simultaneously with more ethyl chloroformate (33.4g, 310mmol), with stirring, while keeping the reaction mixture below 5° for 2 hours followed, still with stirring, by 8 hours at ~25°. The *C₆H₆ layer is separated, the aqueous layer is extracted with CHCl₃ (3 x 100ml), the combined organic layers are dried (MgSO₄), filtered and the filtrate is evaporated to leave *tris*[2-(ethoxycarbonylamino)ethyl]amine (~85%) as a thick oil which is used directly in the subsequent step. [The crude tri-ester has IR (film) bands with ν_{max} at 3300, 1720, 1530, and 1250 cm⁻¹; and the ¹H NMR (300MHz, CDCl₃, TMS) has δ at 1.27 (t, 9H, ³J_{HH} = 7.1Hz), 2.60 (t, 6H, ³J_{HH} = 5.7 Hz), 3.23 (br s, 6H), 4.10 (q, 6H, ³J_{HH} = 7.1 Hz) and 5.50 (br s, 3H)].

In the second step the preceding crude tri-ester (61.3g, 170mmol) in THF (250ml) is added dropwise to a suspension of LiAlH₄ (30.0g, 790mmol) in THF (700ml), and the reaction mixture is refluxed overnight. Water (50ml), followed by a solution of KOH (50g) in H₂O (50ml) are very carefully added to it (cool if necessary), the solvent is decanted from the inorganic gel, evaporated *in vacuo*, and the residual yellow oil is fractionated in a vacuum to give the desired *amine* in 88% yield. It has ¹H NMR (300MHz, CDCl₃, TMS) with δ at 1.30 (br s, 3H, NH), 2.39 (s, 9H, CH₃), 2.48 (m, 6H, ³J_{HH} = 6.1 Hz, 3CH₂) and 2.52 (m, 6H, ³J_{HH} = 6.1 Hz, 3CH₂); the ¹³C NMR (75MHz, CDCl₃, TMS) has δ at 54.1 (CH₂), 49.6 (CH₂) and 36.3 (CH₃); and HRMS has m/z 189.2082 (calc for M + H is 189.20793). [Schmidt et al. *Z Anorg Allg Chem* **578** 75 1989, DOI: 10.1002/zaac.19895780109.]

Triuret (1,3-dicarbamoylurea, carbonyldiurea) [556-99-0] $\text{C}_3\text{H}_6\text{N}_4\text{O}_3$, **M 146.1, m 233°(dec)**. It crystallises from aqueous ammonia or H_2O (plates **m 232-234°**), and is soluble in liquid NH_3 . It gives mono and dipotassium salts. [Haworth & Mann *J Chem Soc* 603 1943, DOI: 10.1039/JR9430000603; *Beilstein* 3 H 72, 3 I 35, 3 II 60, 3 III 142.]

Undecan-1-ol [112-42-5] $\text{C}_{11}\text{H}_{24}\text{O}$, **M 172.3, m 11°, 16.5°, 146°/30mm, d_4^{25} 0.830, n_D^{20} 1.440**. Purify the alcohol by repeated fractional crystallisation from its melt or by distillation in a vacuum. [*Beilstein* 1 H 427, 1 IV 1835.]

1,11-Undecanedicarboxylic acid (Brassylic acid, 1,13-tridecanedicarboxylic acid) [505-52-2] $\text{C}_{13}\text{H}_{24}\text{O}_4$, **M 244.3, m 103-107°, 113°, 112-114°, pK_{Est} ~5.0**. The dicarboxylic acid was prepared by Mislow and Steinberg from azelaic acid (see above) \rightarrow 1,9-nonadiol (92.4%) \rightarrow 1,9-dibromononane (89.7%) \rightarrow 1,9(bis-diethylmalonyl)nonane \rightarrow 1,9-bis(carboxy)nonane (= 1,11-undecanedioic acid, 80%). Brassylic acid is purified by recrystallisation from $^*\text{C}_6\text{H}_6$, aqueous EtOH, EtOAc or 50% EtOH/ CCl_4 . The *diethyl ester*, prepared in the usual way (76.6%), has **m 20°, b 162-163°/1.3mm, 204°/12mm, n_D^{25} 1.4408**. The *monomethyl ester* [3927-59-1] $\text{C}_{14}\text{H}_{26}\text{O}_4$, **M 258.3, has m 58°**, and the *dimethyl ester* [1472-87-3] $\text{C}_{15}\text{H}_{28}\text{O}_4$, **M 272.3, has m 32°(36°), 32.3°, b 192-194°/11mm, 326-328°/atm**. The *diamide*, prepared from the acid *via* SOCl_2 then 25% NH_4OH , has **m 177°** (white fine needles from EtOH). [Chuit *Helv Chim Acta* 9 264 1926, DOI: 10.1002/hlca.19260090131; Potter & Taylor *J Chem Soc* 3514 1951, within Note pp 3508-3514, DOI: 10.1039/JR9510003508; Mislow & Steinberg *J Am Chem Soc* 77 3807 1955, DOI: 10.1021/ja01619a038; for synthesis from methylene-bis-dihydroresorcinol and conversion to the diamide see Stetter & Dietrichs *Chem Ber* 85 290 1952, DOI: 10.1002/cber.19520850405; *Beilstein* 2 H 731.]

Undecanoic acid (C11, undecylic acid, hendecanoic acid) [112-37-8] $\text{C}_{11}\text{H}_{22}\text{O}_2$, **M 186.3, m 28.5°, b 164°/18mm, 228°/160mm, 248-250°/~760mm, d_4^{20} 0.8907, n_D^{25} 1.4294, pK_{Est} ~5.0**. Purify the acid by repeated fractional crystallisation from its melt or by distillation in a vacuum. [*Beilstein* 2 H 358, 2 IV 1068.]

Undec-10-enoic acid (undecylenic acid) [112-38-9] $\text{C}_{11}\text{H}_{20}\text{O}_2$, **M 184.3, m 24.5°, 25-25.5°, b 131°/1mm, 137°/2mm, 168°/15mm, 275°(dec)/atm, d_4^{20} 0.912, n_D^{25} 1.447, pK_{Est} ~5.0**. Purify the acid by repeated fractional crystallisation from its melt or by distillation preferably in a high vacuum as it decomposes at atmospheric pressure. It was originally isolated from castor oil by destructive distillation and gave **ω -bromo-undecylic acid (m 50.5°, 51°**, by recrystallisation from petroleum ether). The *methyl ester* has **b 124°/10mm, 148°/atm, n_D^{20} 1.43928**. The *Zinc salt* [557-08-4], prepared by dissolving ZnO in dilute undecylenic acid in MeOH and evaporated giving a white powder with **m ~115-116°**, and is sold commercially as a topical fungicide (Fungex). [Das et al. *J Am Oil Chem Soc* 66 938 1989, DOI: 10.1007/BF02682613; Perkin & Cruz *J Am Chem Soc* 49 1070 1927, DOI: 10.1021/ja01403a030; *Beilstein* 2 IV 1612.]

Urea (carbamide) [57-13-6] $\text{CH}_4\text{N}_2\text{O}$, **M 60.1, m 132.7-132.9°, 132-135°, pK^{25} 0.12**. Crystallise urea twice from conductivity water using centrifugal drainage and keeping the temperature below 60°. The crystals are dried under vacuum at 55° for 6 hours. Levy and Margoulas [*J Am Chem Soc* 84 1345 1962, DOI: 10.1021/ja00867a003] prepared a 9M solution in conductivity water (keeping the temperature below 25°) and, after filtering through a medium-porosity glass sinter, added an equal volume of absolute EtOH. The mixture was set aside at -27° for 2-3 days and filtered cold. The precipitate was washed with a small amount of EtOH and dried in air. Crystallisation from 70% EtOH between 40° and -9° has also been used. Ionic impurities such as ammonium isocyanate have been removed by treating the concentrated aqueous solution at 50° with Amberlite MB-1 cation- and anion-exchange resin, and allowing it to crystallise on evaporation. [Benesch et al. *J Biol Chem* 216 663 1955, PMID: 13271343, <http://www.jbc.org/content/216/2/663.citation>.] It can also be crystallised from MeOH or EtOH, and is dried under vacuum at room temperature. [*Beilstein* 3 H 42, 3 I 19, 3 II 35, 3 III 80, 3 IV 94.]

Urea nitrate [124-47-0] $\text{CH}_3\text{N}_3\text{O}_4$, **M 123.1, m 152°(dec), 157-158°, 163°**. Crystallise it from dilute HNO_3 or EtOH (**m 157-158°**), and dry it in a vacuum over P_2O_5 . Its solubility (w/w%) in H_2O is 9.3 (at 0°) and 40 (at 65.3°); and in EtOH is 1.35 (at 0°) and 8.8 (at 65.3°). May **EXPLODE** if shocked or heated. [*Beilstein* 3 H 54,

3 I 25, 3 II 45, 3 III 105, 3 IV 94.]

Urethane (ethyl carbamate, ethyl urethane) [51-79-6] $C_3H_7NO_2$, **M 89.1, m 48-50°, b 182-184°/~760mm, d₄²⁰ 0.986, n_D²⁵ 1.4144.** Urethane is best purified by fractional distillation, but it can be sublimed at ~103°/~50mm. It has also been recrystallised from *benzene. Its solubility at room temperature is 2g/ml in H₂O, 1.25g/ml in EtOH, 1.1g/ml in CHCl₃, 0.67g/ml in Et₂O and 0.03g/ml in olive oil. It is a suspected **human carcinogen**. [Beilstein 3 H 22, 3 IV 40.]

cis-Vaccenic acid (octadec-11-enoic acid) [506-17-2] $C_{18}H_{34}O_2$, **M 282.5, m 14-15°, b 158-163°/0.4mm, d₄²⁰ 0.880, n_D²⁵ 1.4598, pK_{Est} ~ 4.9.** Purify the acid by fractional distillation under high vacuum or crystallisation from its melt in an inert atmosphere away from light. [Beilstein 2 I 198, 2 III 1384, 2 IV 1639.]

trans-Vaccenic acid (octadec-11-enoic acid) [693-72-1] $C_{18}H_{34}O_2$, **M 282.5, m 43-44°, n_D⁵⁰ 1.4472, pK_{Est} ~ 4.9.** Crystallise the acid from acetone (m 45-45.5°) or aqueous MeOH (m 43.5-43.7°). The **methyl ester** $C_{19}H_{36}O_2$ has **b 174-175°/5mm**. [Böeseken et al. *Recl Trav Chim Pays-Bas* 46 619 1927, DOI: 10.1002/recl.19270460902; Ahmad et al. *J Am Chem Soc* 70 3391 1948, DOI: 10.1021/ja01190a051; IR: Rao & Daubert *J Am Chem Soc* 70 1102 1948, DOI: 10.1021/ja01183a069.]

n-Valeraldehyde (pentanal) [110-62-3] $C_5H_{10}O$, **M 86.1, m -92°, b 103°/atm, d₄²⁰ 0.811, n_D²⁵ 1.40233.** Purify pentanal via the bisulfite derivative (see 2-butanone above for the preparation and decomposition of the bisulfite derivative). [Birrell & Trotman-Dickenson *J Chem Soc* 2059 1960, DOI: 10.1039/JR9600002059; Beilstein 1 H 676, 1 IV 3268.] The **2,4-dinitrophenylhydrazone** [2057-84-3] **M 266.3** has **m 103-105°** (from EtOH). [Beilstein 15 III/IV 429.]

n-Valeramide (pentanamide) [626-97-1] $C_5H_{11}NO$, **M 101.1, m 115-116°.** Crystallise the amide from EtOH. It sublimes at 80°. [Philbrook *J Org Chem* 19 623 1954, DOI: 10.1021/jo01369a022; Beilstein 2 H 301, 2 I 131, 2 II 266, 2 III 674, 2 IV 874.]

Valeric acid (n-pentanoic acid) [109-52-4] $C_5H_{10}O_2$, **M 102.1, m -20° to -18°, b 95°/22mm, 186.4°/~760mm, d₄²⁰ 0.938, n_D²⁰ 1.4080, pK₂₅ 4.81.** Water is removed from the acid by distillation using a Vigreux column, until the boiling point reaches 183°. A few crystals of KMnO₄ are added, and after refluxing, the distillation is continued. [Andrews & Keefer *J Am Chem Soc* 83 3708 1961, DOI: 10.1021/ja01478a032; Beilstein 2 H 299, 2 I 130, 2 II 263, 2 III 663, 2 IV 868.] The **ethyl ester** [539-82-2] $C_7H_{14}O_2$, **M 130.2**, has **b 145-146°/atm, d₄²⁰ 0.877, and n_D²⁰ 1.401.** It is used as a substrate for assaying the activities of some esterases, and as an intermediate in the preparation of perfumes. [Beilstein 2 IV 872.]

Valeronitrile (butyl cyanide) [110-59-8] C_5H_9N , **M 83.1, m -96°, b 45-47°/15mm, 142.3°/~760mm, d₄²⁰ 0.799, n_D¹⁵ 1.39913, n_D³⁰ 1.39037.** Wash the nitrile with half its volume of concentrated HCl (twice), then with saturated aqueous NaHCO₃, dry it with MgSO₄ and fractionally distil it from P₂O₅. [Beilstein 2 H 301, 2 I 131, 2 II 267, 2 III 675, 2 IV 875.]

Vinyl acetate [108-05-4] $C_4H_6O_2$, **M 86.1, m -93°, b 72.3°/atm, d₄²⁰ 0.938, n_D²⁰ 1.396.** Inhibitors such as hydroquinone and other impurities are removed by drying with CaCl₂ and fractionally distilling under nitrogen, then refluxing briefly with a small amount of benzoyl peroxide and redistilling it under nitrogen. Store it in the dark at 0°. Add inhibitor (~0.004%) for storage. [Beilstein 2 IV 176.]

Vinyl butoxyethyl ether (ethylene glycol butyl vinyl ether) [4223-11-4] $C_8H_{16}O_2$, **M 144.2, b 70-72°/20mm, d₄²⁰ 0.866, n_D²⁰ 1.4220.** Wash this ether with aqueous 1% NaOH, dry with CaH₂, then reflux with, and distil it from sodium. Stabilise it with 0.5% of 2,6-di-*tert*-butyl-*p*-cresol for storage. [Beilstein 1 IV 2387.] **IRRITANT.**

Vinyl chloroformate [5130-24-5] $C_3H_3ClO_2$, **M 106.5, b 46.5°/80mm, 67-69°/atm, 109-110°/760mm, d₄²⁰ 1.136, n_D²¹ 1.420.** It has been fractionated through a Todd column (Model A with ~60 plates) under atmospheric pressure and the purity can be checked by gas chromatography. Stabilise it with 0.5% of 2,6-di-*tert*-butyl-*p*-

cresol. It has IR with ν_{\max} at 3100 + 2870 (CH₂), 1780 (C=O), 1640 (C=C) and 940 (CH₂ out-of-plane) and 910 (CH₂ wagging) cm⁻¹. [IR: Lee *J Org Chem* **30** 3943 1965, DOI: 10.1021/jo01022a511; Levailant *Ann Chim (Paris)* **6** 504 1936.] It is used for protecting NH₂ groups in peptide synthesis [Olofson et al. *Tetrahedron Lett* 1563 1977, DOI:10.1016/S0040-4039(01)93103-X]. [*Beilstein* **3** III 28.]

Vinyl stearate [111-63-7] C₂₀H₃₈O₂, **M 310.5, m 35°, 35-36°, b 166°/1.5mm, 167°/2mm, 187-188°/4.3mm, d₄⁴⁰ 0.8517, n_D⁴⁰ 1.4423**. Distil the ester in a vacuum under nitrogen, then crystallise it from acetone (3ml/g) or ethyl acetate at 0°. Store it under nitrogen in the dark. When polymerised with a small amount of benzoyl peroxide, a white wax-like polymer was formed. [Swern & Jordan *J Am Chem Soc* **70** 2334 1948, DOI: 10.1021/ja01187a008; Swern & Jordan *Org Synth* **30** 106 1950, DOI: 10.15227/orgsyn.030.0106; *Beilstein* **2** III 1019.]

ALICYCLIC COMPOUNDS

Abietic acid [514-10-3] $\text{C}_{20}\text{H}_{30}\text{O}_2$, **M 302.5**, **m 172-175°**, $[\alpha]_{\text{D}}^{25}$ **-116 (-106°)** (c 1, EtOH), **pK²⁵ 5.27**. Crystallise it by dissolving 100g of acid in 95% EtOH (700ml), adding to H_2O (600ml) and cooling. Filter, dry it in a vacuum (over KOH or CaSO_4) and store it in an O_2 -free atmosphere. It can also be purified *via* the anhydride, tritylabietate and the potassium, piperidine and brucine salts. It has λ_{max} : nm(log ϵ): 2343(4.3), 241(4.4), 2505(4.2), 235(4.34) and 240(4.36) in EtOH. [Harris & Sanderson *Org Synth Coll Vol* 4 1 1963, DOI: 10.15227/orgsyn.032.0001; Lambard & Frey *Bull Soc Chim Fr* 1194 1948, Buchbauer et al. *Monatsh Chem* 116 1345 1985, DOI: 10.1007/BF00811105.] [Beilstein 9 IV 2175.]

S-Abscisic acid (natural; 2-cis-4-trans-5S-abscisic acid, Dormin) [S(+)] 21293-29-8] $\text{C}_{15}\text{H}_{20}\text{O}_4$, **M 264.3**, **m 160-161°, 161-163°** (sublimation), $[\alpha]_{\text{D}}^{20}$ **+411 (1, EtOH)**, $[\alpha]_{287}^{20}$ **+24,000**, $[\alpha]_{245}^{20}$ **-69,000** (c 1-50 $\mu\text{g}/\text{ml}$ in acidified MeOH or EtOH), **pK_{Est} ~3.9**. Crystallise the acid from CCl_4 /petroleum ether, EtOH/hexane and sublime it at 120°. Also purify it by dissolving ~30g in 30ml of EtOAc, adding 100ml of hexane and allow to crystallise overnight (yield 8.4g), **m 156-158°, 161-163°**, $[\alpha]_{\text{D}}^{20}$ **+426** (c 1, 0.005M H_2SO_4 in MeOH). [Cornforth et al. *Nature (London)* 206 715 1965, DOI:10.1038/206715a0; Soukup et al. *Helv Chem Acta* 72 361 1989, DOI: 10.1002/hlca.19890720222.] The **non-natural 2-cis-4-trans-5R-abscisic acid** [R(-)] 14398-53-9] has **m 162-163°**, $[\alpha]_{\text{D}}^{20}$ **-426** (c 1, 0.005N H_2SO_4). **Racemic 2-cis-4-trans-5RS-abscisic acid** [RS(\pm)] 14375-45-2] was purified on a Kieselgel F₂₅₄ plate with toluene/EtOAc/AcOH (50:50:3) and has **m 188-190°** [Cornforth et al. *Aust J Chem* 45 179 1992, DOI: 10.1071/CH9920179]. [Absolute configuration: Ryback *JCS Chem Commun* 1190 1972, DOI: 10.1039/C39720001190.] [Beilstein 17/3 V 13.]

Acetylcyclohexane (cyclohexyl methylketone) [823-76-7] $\text{C}_8\text{H}_{14}\text{O}$, **M 126.2**, **b 64°/11mm, 76.2-77°/25mm, d₄²⁰ 0.9178, n_D²⁰ 1.4519**. Dissolve acetylcyclohexane in Et_2O , shake it with H_2O , dry, evaporate and fractionate it under reduced pressure. [UV: Mariella & Raube *J Am Chem Soc* 74 518 1952, DOI: 10.1021/ja01122a069; enol content: Gero *J Org Chem* 19 1960 1954, DOI: 10.1021/jo01377a013.] The **semicarbazone** has **m 174°** and the **2,4-dinitrophenylhydrazone** has **m 139-140°** [Theus & Schinz *Helv Chem Acta* 39 1290 1956, DOI: 10.1002/hlca.19560390516].

2-Acetylcyclohexanone [874-23-7] $\text{C}_8\text{H}_{12}\text{O}_2$, **M 140.2**, **m -11°, b 62-64°/2.5mm, 95-98°/10mm, 111-112°/18mm, d₄²⁰ 1.08, n_D²⁰ 1.51**. Dissolve it in ligroin (b 30-60°), wash it with saturated aqueous NaHCO_3 , dry over Drierite and fractionate in a vacuum. [Perfett & Levine *J Am Chem Soc* 75 626 1953, DOI: 10.1021/ja01099a031; Manyik et al. *J Am Chem Soc* 75 5030 1953, DOI: 10.1021/ja01116a042; Eistert & Reiss *Chem Ber* 87 108 1954, DOI: 10.1002/cber.19540870116.] It forms a **Cu salt** which crystallises in green leaflets from EtOH, **m 162-163°** [UV: McEntee & Pinder *J Chem Soc* 4419 1957, DOI: 10.1039/JR9570004419]. [Beilstein 7 IV 1997.]

2-Acetylcyclopentanone [1670-46-8] $\text{C}_7\text{H}_{10}\text{O}_2$, **M 126.2**, **b. 72-75°/8mm, 82-86°/12mm, 88°/18mm, d₄²⁰ 1.043, n_D²⁰ 1.490**. Dissolve the ketone in petroleum ether (b 30-60°), wash it with saturated aqueous NaHCO_3 , dry over Drierite and fractionate in a vacuum. It gives a violet colour with ethanolic FeCl_3 and is only slowly hydrolysed by 10% aqueous KOH, but rapidly on boiling to yield 6-oxoheptanoic acid. [Manyik et al. *J Am Chem Soc* 75 5030 1953, DOI: 10.1021/ja01116a042; Acheson *J Chem Soc* 4232 1956, DOI: 10.1039/JR9560004232; UV: Martin & Fernelius *J Am Chem Soc* 81 2342 1959, DOI: 10.1021/ja01519a017.] It gives a grey green **Cu salt** from Et_2O /pentane, **m 237-238°** [House & Wasson *J Am Chem Soc* 79 1488 1957, DOI: 10.1021/ja01563a058]. [Beilstein 7 IV 1993.]

2-Acetyl-5,5-dimethylcyclohexane-1,3-dione (2-acetyldimedone) [1755-15-3] $\text{C}_{10}\text{H}_{14}\text{O}_3$, **M 182.2**, **m 36°, 36-40°, b 132-133°/20mm, 138°/23mm, pK ~4.5**. It can be purified by fractional distillation. *Alternatively*, convert it into the insoluble Cu salt in H_2O and recrystallise it from EtOH, **m ~260°**. The Cu salt is decomposed with N H_2SO_4 , extracted into Et_2O , dried (Na_2SO_4), evaporated and distilled in a vacuum. The residual oil, which solidifies on cooling can be recrystallised from AcOH. It gives a red colour with Fe^{3+} ions. The **oxime** has **m 115°**(dec, from EtOH), and with concentrated NH_3 it forms the **mono-imide** which crystallises from H_2O

in needles with **m** 133°. Its UV in EtOH has $\lambda_{\max}(\epsilon)$ 231(10,620) and 273(10,800)nm. It is used as a protecting group for primary amines. [Dieckmann & Stein *Chem Ber* **37** 3370 1904, DOI: 10.1002/cber.190403703147; Birch *J Chem Soc* 3026 1951, DOI: 10.1039/JR9510003026; Crossley & Renouf *J Chem Soc* **101** 1524 1912, DOI: 10.1039/CT9120101524; Nash et al. *Tetrahedron Lett* **37** 2625 1996, DOI: 10.1016/0040-4039(96)00344-9; Kellam et al. *Tetrahedron Lett* **38** 4849 1997, DOI: 10.1016/S0040-4039(97)01010-1; Beilstein **7** H 860, **7** I 471, **7** IV 2756.]

4-Acetyl-1-methyl-1-cyclohexene [6090-09-1] $\text{C}_9\text{H}_{14}\text{O}$, **M** 138.2, **b** 73-75°/7.5mm, 85-86°/13mm, 94-94.7°/20mm, 204.5-206°/747mm, d_4^{20} 1.0238, n_D^{20} 1.469. Purify it by fractionation under reduced pressure *in vacuo*, and if it is almost pure it can be fractionated at atmospheric pressure, preferably in an inert atmosphere. It forms two *semicarbazones* one of which is more soluble in C_6H_6 , and both can be recrystallised from EtOH; the more soluble has **m** 149°(151°), and the less soluble has **m** 172-175°(191°). The **4-nitrophenylhydrazone** has **m** 166-167° and the **2,4-dinitrophenylhydrazone** has **m** 114-115°. [Pfau & Plattner *Helv Chem Acta* **17** 129, 142 1934, DOI: 10.1002/hlca.19340170118; Adler & Vogt *Justus Liebigs Ann Chem* **564** 109 1949, DOI: 10.1002/jlac.19495640204.]

2-Acetyl-1-methyl-3,5-dioxo-1-methylcyclohexanecarboxylic acid (ADCC-linker) [181486-37-3] $\text{C}_{10}\text{H}_{12}\text{O}_5$, **M** 212.2, **m** 95-99°, $\text{pK}_{\text{est}} \sim 4.5$. It is prepared from 3,5-dioxo-1-methylcyclohexane-1-carboxylic acid methyl ester (obtained from 3,5-dimethoxybenzoic acid [119-52-8] *via* Birch reduction, methylation with MeI and treatment with aqueous HCl) by acetylation to 3-acetoxy-5-oxo-1-methylcyclohex-3,4-ene-1-carboxylic acid methyl ester, rearrangement (heating with DMAP) to 4-acetyl-3,5-dioxo-1-methylcyclohexane carboxylic acid methyl ester (93% yield) followed by hydrolysis (LiOH/THF/MeOH/H₂O) and H⁺ ion-exchange purification to give the ADCC-linker as a crystalline solid. It should be stored at ~5°. After attaching to a solid support through the carboxy function (e.g. to amino-modified polystyrene beads), it is used for linking to primary amines (including α -amino-acid esters) *via* enamine formation of the 4-acetyl group for combinatorial synthesis. The linker is stable to acids such as $\text{CF}_3\text{CO}_2\text{H}$, bases such as piperidine or BTU, and uronium type coupling agents; and by treatment with 2% hydrazine in DMF, the primary amine is released quantitatively from the support. Its ¹H NMR (250MHz, Me₂SO) has δ at 1.25 (s, Me-C1(1)), 2.50 (s, MeCO-C(4)), 2.50-3.00 (m, 2H-C(3), 2H-C(5)), 17.80 (2H, enol OH and COOH). [Bannwarth et al. *Bioorg Med Chem Lett* **6** 1525 1966, DOI: 10.1016/S0960-894X(96)00258-2.]

Adamantane (tricyclo[3.3.1.1^{3,7}]-decane) [281-23-2] $\text{C}_{10}\text{H}_{16}$, **M** 136.2, **m** 269.6-270.8° (sublimes). Crystallise adamantane from acetone or cyclohexane, and sublime it in a vacuum below its melting point [et al. *JCS Faraday Trans 2* **82** 535 1986, DOI: 10.1039/F29868200535]. Adamantane is also purified by dissolving it in *n*-heptane (*ca* 10ml/g of adamantane) on a hot plate, adding activated charcoal (2g/100g of adamantane), and boiling for 30 minutes, filtering the hot solution through a filter paper, concentrating the filtrate until crystallisation just starts, adding one quarter of the original volume of *n*-heptane, and allowing to cool slowly over a period of hours. The supernatant is decanted off and the crystals are dried *in vacuo* at 25°. [Prelog & Seiwert *Chem Ber* **74** 1769 1941, DOI: 10.1002/cber.19410741110; Schleyer et al. *Org Synth Coll Vol* **5** 16 1973, DOI: 10.15227/orgsyn.042.0008; Walter et al. *J Am Chem Soc* **107** 793 1985, DOI: 10.1021/ja00290a010.] [Beilstein **5** III 393, **5** IV 469.]

1-Adamantane acetic acid [4942-47-6] $\text{C}_{12}\text{H}_{18}\text{O}_2$, **M** 194.3, **m** 136°, $\text{pK}_{\text{Est}} \sim 4.8$. Dissolve the acid in hot NaOH, treat with charcoal, filter and acidify. Collect the solid, wash it with H₂O, dry and recrystallise it from MeOH. [Stetter et al. *Chem Ber* **92** 1629 1959, DOI: 10.1002/cber.19590920722.] The **acid chloride** [2094-72-6] has **M** 168.7, **m** 51-54°, and **b** 135-136°/1mm. [Beilstein **9** IV 256.] **LACHRYMATORY**.

1-Adamantane carboxylic acid [828-51-3] $\text{C}_{11}\text{H}_{16}\text{O}_2$, **M** 180.3, **m** 175-176.5°, 177°, $\text{pK}_{\text{Est}} \sim 4.9$. Possible impurities are trimethylacetic acid and C9 and C13 acids. Dissolve 15g of the acid in CCl₄ (300ml) and shake with 110ml of 15N aqueous NH₃ whereby the ammonium salt separates and is collected. Acid impurities form soluble ammonium salts. The salt is washed with cold Me₂CO (20ml) and suspended in H₂O (250mL). This is treated with 12N HCl and extracted with CHCl₃ (100ml). The dried (Na₂SO₄) extract is evaporated and the residue is recrystallised from a mixture of MeOH (30ml) and H₂O (*ca* 10ml) to give the pure acid (10-11g). [Koch & Haaf *Org Synth Coll Vol* **5** 20 1973, DOI: 10.15227/orgsyn.044.0001.] It was also recrystallised from

absolute EtOH and dried under vacuum at 100°. *Alternatively*, the acid (5g) is refluxed for 2 hours with 15ml of MeOH and 2ml of 98% H₂SO₄ (cool when mixing this solution). Pour into 10 volumes of H₂O and extract with the minimum volume of CHCl₃ to give clear separation of phases. The extract is washed with H₂O, dried (CaCl₂) and distilled. The *methyl ester* is collected at 77-79°/1mm, **m 38-39°**. The ester is hydrolysed with the calculated amount of N KOH and refluxed until clear. Acidification with HCl provides the pure acid with 90% recovery. The *amide* [5511-18-3] C₁₁H₁₇NO, **M 179.3**, crystallises from cyclohexane, and has **m 189-190°**. [Stetter et al. *Chem Ber* **92** 1629 1959, DOI: 10.1002/cber.19590920722.] [*Beilstein* **9** IV 253.]

1,3-Adamantane diamine dihydrochloride [26562-81-2] **M 239.2**, **m >310°**, **pK_{Est(1)} ~8.1**, **pK_{Est(2)} ~10.1**. Dissolve it in boiling concentrated HCl (400mg in 15ml) and evaporate to dryness. Dissolve it in absolute EtOH and add dry Et₂O to crystallise the *dihydrochloride*. [Stetter & Wulff *Chem Ber* **93** 1366 1960, DOI: 10.1002/cber.19600930619; *Beilstein* **13** III 27,]

1,3-Adamantane dicarboxylic acid [39269-10-8] C₁₂H₁₆O₄, **M 224.3**, **m 276°**, **276-278°**, **279°**, **pK_{Est(1)} ~4.9**, **pK_{Est(2)} ~5.9**. Dissolve the acid in aqueous NaOH, treat with charcoal, filter and acidify with dilute HCl. It crystallises from MeOH. [Stetter & Wulff *Chem Ber* **93** 1366 1960, DOI: 10.1002/cber.19600930619; *Beilstein* **9** III 4066, **9** IV 2997.]

1-Adamantane methylamine [17768-41-1] C₁₁H₁₉N, **M 165.3**, **b 83-85°/0.3mm**, **d₄²⁰ 0.935**, **pK_{Est} ~10.2**. Dissolve the amine in Et₂O, dry over KOH and distil it. The *N-Tosyl* derivative has **m 134-135°** (from EtOH). [Stetter & Goebel *Chem Ber* **96** 550 1963, DOI: 10.1002/cber.19630960228.]

1-Adamantanol (1-hydroxyadamantane) [768-95-6] C₁₀H₁₆O, **M 152.4**, **m 288.5-290°**. If 2-adamantanol is a suspected impurity, then dissolve the substance (10g) in acetone (100ml) and add Jones's reagent [CrO₃ (10.3g) in H₂O (30ml)], then concentrated H₂SO₄ (8.7ml) is added dropwise (turns green in colour) until excess reagent is present (slight red colour). Stir overnight, decant the acetone solution from the Cr salts and adamantan-2-one, dry (Na₂SO₄) and evaporate to dryness. The residue (*ca* 7g) is chromatographed through Al₂O₃ (250g) and washed with 50% *benzene/petroleum ether (b 40-60°), then 100% Et₂O (to remove any adamantan-2-one present) and the 1-adamantanol is then eluted with 5% MeOH in Et₂O. The eluate is evaporated, and the residue is recrystallised from petroleum ether (b 30-60°) at -70°, **m 287.2-288.5°**. It also crystallises from MeOH and can be sublimed *in vacuo*. It has characteristic IR, with ν_{\max} at 3640, 1114, 1086, 982 and 930 cm⁻¹. [Schleyer & Nicholas *J Am Chem Soc* **83** 182 1961, DOI: 10.1021/ja01462a036.] [*Beilstein* **6** IV 391.] *Alternatively*, if free from the 2-isomer, dissolve it in tetrahydrofuran, and dilute with H₂O to precipitate the alcohol. Collect, dry and sublime it in a vacuum at 130°. [Stetter et al. *Chem Ber* **92** 1629 1959, DOI: 10.1002/cber.19590920722.]

2-Adamantanol (2-hydroxyadamantane) [700-57-2] C₁₀H₁₆O, **M 152.4**, **m 296-299°**, **296.2-297.7°**. Best obtained by reduction of adamantanone (30.0g) in Et₂O (300ml) and LiAlH₄ (7.3g.) in Et₂O (150ml) at ~25° for 2hrs, decomposed in the usual way, and extracted into CH₂Cl₂ to give the adamantan-2-ol (30g). It can be purified by chromatography as for the 1-isomer. It crystallises from cyclohexane and has characteristic IR with ν_{\max} at 3600, 1053, 1029 and 992 cm⁻¹ [Schleyer & Nicholas *J Am Chem Soc* **83** 182 1961, DOI: 10.1021/ja01462a036].

2-Adamantanone [700-58-3] C₁₀H₁₄O, **M 150.2**, **m 256-258°(sublimes)**. Purify 2-admantanone by repeated sublimation *in vacuo*. [Butler et al. *JCS Faraday Trans 2* **82** 535 1986, DOI: 10.1039/F29868200535.]

N-(1-Adamantyl)acetamide [880-52-4] C₁₂H₁₉NO, **M 193.3**, **m 149°**. Wash the amide well with H₂O, dry and recrystallise it from cyclohexane. *It is an irritant*. [Stetter et al. *Chem Ber* **92** 1629 1959, DOI: 10.1002/cber.19590920722.]

1-Adamantylamine (1-adamantanamine, Amantadine) [768-94-5] C₁₀H₁₇N, **M 151.2**, **m 160-190° (sealed tube)**, **180-192°**, **208-210°**, **pK²⁵ 10.58**. Dissolve the amine in Et₂O, dry it over KOH, evaporate and sublime it *in vacuo*. [Stetter et al. *Chem Ber* **93** 226 1960, DOI: 10.1002/cber.19600930133.] It is physiologically active (see next entry).

1-Adamantylamine hydrochloride (Amantadine hydrochloride, Amazolone, Mantadan, Mantadix, Virofral) [665-66-7] $C_{10}H_{17}N$. HCl, M 187.7, m >360° (dec), pK_{25}^{25} 10.58. Dissolve the salt in dry EtOH, add a few drops of dry EtOH saturated with HCl gas, followed by dry Et₂O to crystallise the hydrochloride out. Dry the salt in a vacuum. Its solubility in H₂O is >5%; and it is soluble in EtOH but insoluble Et₂O. [Stetter et al. *Chem Ber* **93** 226 1960, DOI: 10.1002/cber.19600930133.] It is an antiviral, and an anti-Parkinsonian agent [Kornhuber et al. *J Neural Transm* **46** (Suppl) 399 1995].

2-Adamantylamine hydrochloride [10523-68-9] $C_{10}H_{17}N$. HCl, M 187.7, m >300°, pK_{Est} ~10.4. The *free amine* in Et₂O, liberated by the action of alkali in H₂O, is dried over KOH, filtered, evaporated and sublimed at 110°/12torr, m 230-236°. The base is dissolved in EtOH, sufficient ethanolic HCl is added dropwise and crystallised by the addition of Et₂O. Dry it *in vacuo*. [Stetter et al. *Justus Liebigs Ann Chem* **658** 151 1962, DOI: 10.1002/jlac.19626580113].

1-Adamantyl bromide [768-90-1] $C_{10}H_{15}Br$, M 215.1, m 117-119°, 118°, 119.5-120°. If coloured, dissolve it in CCl₄, wash with H₂O, treat with charcoal, dry (CaCl₂), filter and evaporate to dryness. Dissolve the residue in a small volume of MeOH and cool in a CO₂/trichloroethylene bath and collect the crystals. Sublime it at 90-100°/water pump vacuum. [Stetter et al. *Chem Ber* **92** 1629 1959, DOI: 10.1002/cber.19590920722; Schleyer & Nicholas *J Am Chem Soc* **83** 2700 1961, DOI: 10.1021/ja01473a024; *Beilstein* **5** IV 469.] **2-Adamantyl bromide** [7314-85-4] $C_{10}H_{15}Br$, M 215.1, has m 138-140°. Purify as for the 1-isomer.

1-Adamantyl bromomethylketone [5122-82-7] $C_{12}H_{17}BrO$, M 257.2, m 76-79°, 78-79°. Dissolve the ketone in Et₂O, wash it with H₂O, dry (MgSO₄), evaporate and crystallise the residue from small volumes of MeOH. **LACHRYMATORY**. [Stetter & Rauscher *Chem Ber* **93** 2054 1960, DOI: 10.1002/cber.19600930922.] Also recrystallise it from aqueous MeOH and sublime it at 100°/12torr. It crystallises from MeOH at -70°. Do not keep in contact with MeOH for too long. [Stetter et al. *Chem Ber* **92** 1629 1959, DOI: 10.1002/cber.19590920722.]

1-Adamantyl chloride [935-56-8] $C_{10}H_{15}Cl$, M 170.7, m 164.3-165.6°. It has been prepared by refluxing 1-acetamidoadamantane (3g, m 148° from dry cyclohexane, and obtained by hydrolysis of 1-cyanoadamantane with H₂SO₄) in concentrated HCl (100ml) for 2 hours during which time some 1-chloroadamantane sublimed in the condenser. 1-chloroadamantane was collected (on evaporation and addition of H₂O), recrystallised from aqueous MeOH, and sublimed to provide pure 1-adamantyl chloride (3.3g, 98%), m 165°. [Stetter et al. *Chem Ber* **92** 1269 1959, DOI: 10.1002/cber.19590920722; Schleyer & Nicholas *J Am Chem Soc* **83** 2700 1961, DOI: 10.1021/ja01473a024; *Beilstein* **5** IV 469.]

1-Adamantyl chloroformate [5854-52-4] $C_{11}H_{15}ClO_2$, M 214.6, m 46-47°. Crystallise it from petroleum ether (b 30-60°) at -20°. Also purify it as for 1-adamantyl fluoroformate below. Its IR has ν_{max} at 4.2, 5.6 and 8.4 μ (2380, 1786 and 1190 cm⁻¹). [Haas et al. *J Am Chem Soc* **88** 1988 1966, DOI: 10.1021/ja00961a024; cf. Moroder et al. *Hoppe-Seyler's Z Physiol Chem* **357** 1647 1976.]

RS-1-(1-Adamantyl)ethylamine hydrochloride (Rimantadine hydrochloride, Flumadine, Roflual) [1501-84-4] $C_{12}H_{21}N$. HCl, M 215.8, m >300°, 373-375°(sealed tube), pK_{Est} ~10.4. It is prepared by adding the oxime of 1-adamantyl methyl ketone (6 parts, see [1660-04-4] below) in dry THF (200 parts) to a cold suspension of excess of LiAlH₄ in Et₂O, refluxing for 1 hour, decomposing with cold brine, making strongly alkaline and extracting thoroughly with Et₂O. The dried (Na₂CO₃) extract, is filtered, concentrated, and saturated with dry HCl. The hydrochloride (~5.25 parts) is collected, dried and recrystallised as for 2-adamantylamine hydrochloride above. [US patent to du Pont de Nemours 1069563 (1969, Brit amended), *Chem Abstr* **75** 140372w 1971.] It is an antiviral agent [Burkinskaya et al. *J Gen Virol* **60** 49 1982, DOI: 10.1099/0022-1317-60-1-49].

1-Adamantyl fluoride (1-fluoroadamantane) [768-92-3] $C_{10}H_{15}F$, M 154.2, m 210-212°(dec, sealed tube), 259-260°(dec). Dissolve it in Et₂O, dry over Na₂SO₄, evaporate to dryness and sublime the residue at 90-100°/12mm. Recrystallise the sublimate from MeOH, m 259-260°. To remove 1-hydroxyadamantane impurity, dissolve it in cyclohexane, cool for many hours, filter off the hydroxyadamantane, and evaporate to dryness, or

by passage through an Al_2O_5 column in dry cyclohexane. Recrystallise the residue from petroleum ether at -77° and sublime it in vacuum, **m 210-212°dec** (sealed tube). [Bhandari & Pinock *Synthesis* 655 1974, DOI: 10.1055/s-1974-23391; NMR: Fort et al. *J Org Chem* **30** 789 1965, DOI: 10.1021/jo01014a033.]

1-Adamantyl fluoroformate [62087-82-5] $\text{C}_{11}\text{H}_{15}\text{FO}_2$, **M 198.2**, **m 31-32°**. Dissolve it in *n*-hexane (ca 10g in 150 ml) and keep at 0° for 24 hours. Any 1-adamantanol present will separate. Filter and evaporate to dryness. The crystalline residue has **m 31-32°** and is recrystallised from *n*-hexane (90g/500ml), (IR (KBr): ν_{max} 1242, 1824 and 2340 cm^{-1}). There should be no OH str band above 2500 cm^{-1} . [Moroder et al. *Hoppe-Seyler's Z Physiol Chem* **357** 1647 1976, cf. Haas et al. *J Am Chem Soc* **88** 1988 1966, DOI: 10.1021/ja00961a024.]

1-Adamantyl iodide (1-iodoadamantane) [768-93-4] $\text{C}_{10}\text{H}_{15}\text{I}$, **M 262.1**, **m 75.3-76.4°**. The iodide was prepared by heating 1-adamantanol (5g) in 47% hydroiodic acid (75ml) in a sealed glass tube at $100^\circ/1\text{hr}$, then poured into H_2O and extracted with Et_2O , shaken with aqueous NaHSO_3 , aqueous K_2CO_3 , and H_2O , dried (Na_2SO_4), evaporated and recrystallised from MeOH at -70° (to avoid alcoholysis) to give white crystals of the iodide (3.56g) **m 75.3-76.4°**. [Schleyer & Nicholas *J Am Chem Soc* **83** 2700 1961, DOI: 10.1021/ja01473a024; Literature **m** of **151-152.5°** (Landa & Hála *Coll Czech Chem Commun* **24** 93 1959, DOI: 10.1135/cccc19590093) is incorrect.] Also purify by recrystallisation from petroleum ether ($40\text{-}60^\circ\text{C}$) followed by rigorous drying and repeated sublimation. [Beilstein **5** IV 470.]

1-Adamantyl isocyanate [4411-25-0] $\text{C}_{11}\text{H}_{15}\text{NO}$, **M 177.3**, **m 144-145°**. Recrystallise the isocyanate from *n*-hexane and sublime it. **Irritant**. [Stetter & Wulff *Chem Ber* **95** 2302 1962, DOI: 10.1002/cber.19620950932.]

1-Adamantyl isothiocyanate [4411-26-1] $\text{C}_{11}\text{H}_{15}\text{NS}$, **M 193.3**, **m 168-169°**. Dissolve it in Et_2O , wash with H_2O , dry (Na_2SO_4), evaporate and sublime the residue in a vacuum at 140° , then recrystallise it from MeOH. **Irritant**. [Stetter & Wulff *Chem Ber* **95** 2302 1962, DOI: 10.1002/cber.19620950932.]

1-Adamantyl methyl ketone [1660-04-4] $\text{C}_{12}\text{H}_{18}\text{O}$, **M 178.3**, **m 53-55°, 54-55°**. The ketone is prepared by bubbling acetylene through a vigorously stirred solution of 1-adamantylbromide (1g, see [769-90-1]) in 96% H_2SO_4 (40ml) at 5° for 5 hours with evolution of HBr . The mixture is poured onto ice, extracted with Et_2O , the extract is dried (Na_2SO_4), filtered and evaporated to give an ~80% yield of ketone. This was purified *via* the **2,4-dinitrophenylhydrazine derivative** (by passage of a solution through silica gel/ C_6H_6) which gave orange needles from AcOH/EtOH or EtOH with **m 221° (219-222° and 220-222° also reported)**. The **ketone** has an IR (KBr) peak at ν_{max} 1690 cm^{-1} (C=O); its $^1\text{HNMR}$ has τ at 7.96 (s, 3H), 7.90 and 8.30 (15H); and its MS has m/z at 178 (M^+). [Kell & McQuillin *JCS Perkin Trans I* 2100 1972, DOI: 10.1039/P19720002100; Sasaki et al. *Chem Commun (London)* 780 1968, DOI: 10.1039/C19680000780.]

Alternatively, 1-(ethoxycarbonylmethylcarbonyl)adamantane [1-($\text{EtOCOCH}_2\text{CO}$)-Ad], **b 108-115°/0.06mm**, prepared from 1-adamantylcarbonyl chloride and diethyl malonate] (~25g) was hydrolysed and decarboxylated by boiling in AcOH (50ml), H_2O (30ml) and concentrated H_2SO_4 (5.5ml) until evolution of CO_2 ceased, poured into ice-water (~300ml), the oily **ketone** solidified (94-96% yield), and was recrystallised from MeOH or aqueous MeOH. It sublimates at $40^\circ/0.1\text{mm}$. **1-Adamantyl methyl ketone oxime**, **m 182-184°**, (used for the preparation of *Rimantadine hydrochloride* see [1501-84-4] above) is obtained by shaking hydroxylamine hydrochloride (2.5g) and recrystallised NaOAc (4g) in H_2O (10ml) in a glass test tube, then warming to 40° and adding the **ketone** (2.5g), replacing the stopper and shaking vigorously for a few minutes. The crystalline oxime that soon separates is filtered off, washed with H_2O , and gives colourless plates in high yield upon recrystallising from aqueous dioxane. [Stetter & Rauscher *Chem Ber* **93** 2054 1960, DOI: 10.1002/cber.19600930922; see also Hála & Landa *Coll Czech Chem Commun* **25** 2692 1960, DOI: 10.1135/cccc19602692.]

N-(1-Adamantyl)urea [13072-69-0] $\text{C}_{11}\text{H}_{18}\text{N}_2\text{O}$, **M 194.3**, **m >250°(dec), 268-272°(dec)**. Wash the urea with H_2O and dioxane and recrystallise it from EtOH . [Stetter & Wulff *Chem Ber* **95** 2302 1962, DOI: 10.1002/cber.19620950932.]

(-)-Alloaromadendrene [25246-27-9] $C_{15}H_{24}$, M 204.4, b 96°/2mm, 265-267°/atm, $[\alpha]_D^{25}$ -22 (neat), d_4^{20} 0.923, n_D^{23} 1.501. Fractionally distil it from Na. It has IR bands at 6.06 and 11.27 μ due to C=CH₂. [Birch *J Chem Soc* 715 1953, DOI: 10.1039/JR9530000712; cf. Buechi et al. *J Am Chem Soc* 91 6473 1969, DOI: 10.1021/ja01051a051; *Beilstein* 5 IV 1187.]

Cis-(±)-(1-RS,2-SR)-6-Amino-3-cyclohexene-1-carboxylic acid [*cis-(±)-1,2,3,6-tetrahydroanthranilic acid*] [54162-90-2] $C_7H_{11}O_2N$, M 141.5, m 216-218°, $pK_{Est(1)}$ ~3.5, $pK_{Est(2)}$ ~10.2. Purify the free amino-acid by dissolving it in H₂O and passing it through a Dowex 50W (acid form) column and eluting with 1M aqueous NH₄OH. The eluate is evaporated (*in vacuo*) and the residue is dissolved in H₂O. Me₂CO is added to turbidity, cooled at 0° and the colourless crystals of the amino-acid are collected and dried *in vacuo* [Bernáth et al. *Tetrahedron* 41 1353 1958, DOI: 10.1016/S0040-4020(01)96537-8. cf. Mazza & Crapetta *Gazzetta* 57 297 1927]. In earlier work, it was recrystallised from aqueous EtOH and had reported melting points of 265-265° [Kricheldorf *Justus Liebigs Ann Chem* 1387 1975, DOI: 10.1002/jlac.197519750716] and 269-271° [Moriconi & Mazzochi *J Org Chem* 31 1372 1966, DOI: 10.1021/jo01343a015]. The *hydrochloride* [57266-56-5] M 177.6 has m 210-213° and the *methyl ester hydrochloride* [52766-61-2] has m 85-87° (from Et₂O). The *trans-(±)-(1RS,2RS)-amino-acid* [97945-19-2] crystallises from aqueous Me₂CO with m 267-269°. [*Beilstein* 14 II 203.]

α -Amyrin (Urs-12-en-3 β -ol, α -amyrenol, Viminalol) [638-95-9] $C_{30}H_{50}O$, M 426.7, m 186°, 244°/0.8mm, $[\alpha]_D^{25}$ +85 (c 2, CHCl₃). Purify it by acetylation to the acetate followed by hydrolysis and recrystallisation from aqueous MeOH or from EtOH. The *acetate* when crystallised from petroleum ether, *n*-heptane or CHCl₃/MeOH, and sublimed *in vacuo* has m 227° (225-226°) and $[\alpha]_D^{20}$ +76.4 (c 0.6, CHCl₃). The *benzoate* forms prisms from *C₆H₆/Me₂CO with m 195-196° and $[\alpha]_D^{20}$ +94.6 (c 0.1.9, CHCl₃). [Bentley et al. *J Am Chem Soc* 3673 1953, DOI: 10.1039/JR9530003673; IR: Cole & Thornton *J Am Chem Soc* 1332 1957, DOI: 10.1039/JR9570001332; Corey & Cantrall *J Am Chem Soc* 81 1745 1958, DOI: 10.1021/ja01516a056; *Beilstein* 6 III 2889, 6 IV 4191.]

β -Amyrin (Olean-12-en-3 β -ol, β -amyrenol) [508-04-3] $C_{30}H_{50}O$, M 426.7, m 197-197.5°, 204-205°, b 260°/0.8mm, $[\alpha]_D^{23}$ +91 (c 0.9, CHCl₃). Purify it through an Al₂O₃ column and elute with petroleum ether (40-60°) then Et₂O and recrystallise from petroleum ether or EtOH. The *acetate* crystallises from Ac₂O or petroleum ether and has m 242-143° and $[\alpha]_D^{20}$ +82.8 (c 0.81, CHCl₃) [Crow & Michael *Aust J Chem* 8 129 1955, DOI: 10.1071/CH9550129; Barton et al. *J Chem Soc (C)* 1031 1968, DOI: 10.1039/J39680001031; *Beilstein* 6 III 1894, 6 IV 4195.]

1,1'-Azobis(cyclohexane carbonitrile) [2094-98-6] $C_{14}H_{20}N_4$, M 244.3, m 114-114.5°, 114-115°, 114-118°, ϵ_{350nm} 16.0. Purify the nitrile by dissolving it in boiling 95%EOH as rapidly as possible, cool overnight at 0°, filter, wash with a little EtOH and dry it in a vacuum desiccator over CaCl₂. Note that prolonged heating >80° causes decomposition. Recrystallise it from EtOH. It should be regarded as **potentially explosive**. It is a radical initiator. [Overberger et al. *Org Synth Coll Vol* 4 66 1963, DOI: 10.15227/orgsyn.032.0016; *J Am Chem Soc* 71 2661 1949, DOI: 10.1021/ja01176a018; *Beilstein* 16 II 97, 16 IV 328.]

Bicyclohexyl (dodecahydrobiphenyl) [92-51-3] $C_{12}H_{22}$, M 166.3, (*cis-cis*): m 4°, b 238°/atm, d_4^{20} 0.8914, n_D^{20} 1.48325; and (*trans-trans*): m 4.2°, b 95-96°/9mm, 217-218°/atm, d_4^{20} 0.8592, n_D^{20} 1.4766. Shake bicyclohexyl repeatedly with aqueous KMnO₄ and with concentrated H₂SO₄, wash it with water, dry, first with CaCl₂ then with sodium, and distil it. The two isomers can be separated by fractionation through a very efficient column (a spinning band column using a high reflux ratio). [Mackenzie *J Am Chem Soc* 77 2214 1955, DOI: 10.1021/ja01613a057; *Beilstein* 5 IV *cis-cis* 273, *trans-trans* 334.]

Bicyclohexyl of unspecified stereochemistry (probably a mixture of isomers) has been obtained by catalytic reduction of biphenyl (PtO₂/ 5% EtOH + EtOH-HCl/ 70°/3 atm of H₂ pressure/10hr) and had b 220-228°/atm, d_4^{20} 0.8804, n_D^{20} 1.4792 [Brown et al. *J Am Chem Soc* 58 1594 1936, DOI: 10.1021/ja01300a025]. *Bicyclohexyl* also of unspecified stereochemistry (a mixture with high ratio of *trans* isomer) was prepared by Kolbe electrolysis of cyclohexanecarboxylic acid [separation of isomers is possible by g.l.c. using standard commercial packings, e.g. 2.5% SE30, or 20% DEGS, in stainless steel columns (12ft X 0.25 in, 5ft x 0.25 in

for preparation work, and 2 m x 1/8 in analytical column at 90 to 165°)]. *Alternatively, bicyclohexyl* also of unspecified stereochemistry was obtained in 20% yield by reaction cyclohexylmagnesium bromide (0.184 mol) and AgBr (0.186 mol) in dry Et₂O with stirring (0.5 hr) then reflux (1hr), then after isolation it was boiled under reflux over Na and finally fractionated having **b 90-100°/9mm**, IR (film) ν_{\max} 855, 890, 995, 1262 and 1356 cm⁻¹; ¹H NMR (60 MHz; CCl₄) δ 1.8–2.0 (m); M^+ 166.172 (C₁₂H₂₂). [Hawkes et al *JCS Perkin Trans* 2 1709 1976, DOI: 10.1039/P29760001709.] **Note:** There are theoretically *three* bicyclohexyls that are isomeric around the 1,1'-inter-cyclohexyl bond, viz: *cis-cis* (e,e; equatorial-equatorial), *trans-trans* (a,a; axial-axial) and *cis-trans* (e,a; equatorial-axial).

Bicyclo[3.2.1]octane [6221-55-2] C₈H₁₄, **M 110.2, m 141°**. Purify it by zone melting. It has been sublimed under N₂ at 70° and atmospheric pressure (closed vessel), and resublimed over P₂O₅ to give an analytically pure sample **m 137.5-139.5°**. [Von E Doering & Farber *J Am Chem Soc* **71** 1514 1949, DOI: 10.1021/ja01172a526; Cope et al. *J Am Chem Soc* **82** 4299 1960, DOI: 10.1021/ja01501a046; NMR: Stothers et al. *Can J Chem* **55** 841 1977, DOI: 10.1139/v77-117.]

1R-2-endo-Borneol {endo-(1R)-1,7,7-trimethylbicyclo[2.2.1]heptan-2-ol, (+)-borneol} [464-43-7] C₁₀H₁₈O, **M 154.3, m 204.5-205.5°, 208°, b 212°/atm, d₄²⁰ 1.011, [α]_D²⁰ +37.7 (c 5, EtOH), [α]_D²²₅₄₆ +44.4 (c 0.5, toluene)**. It can be steam distilled, the distillate is extracted into Et₂O, the extract dried with Drierite, filtered and evaporated. Recrystallise the residue from petroleum ether. It is almost insoluble in H₂O, soluble in EtOH (176% w/w), very soluble in Et₂O, soluble in petroleum ether (17% w/w), *C₆H₆ (20% w/w), and quite soluble in many other organic solvents. The **1R-2-endo-acetate** [20347-65-3] C₁₂H₂₀O₂, **M 196.3, has m 26.5°, 29°, b 92-93°/8mm, 223-224°/atm, d₄²⁰ 0.9920, n_D²⁰ 1.4634, [α]_D²⁰ +44.7 (neat)**. The enantiomers: **1S-2-endo-borneol** [464-45-9] and **1S-2-endo-acetate** [5655-61-8] have identical properties except that the signs of the optical rotations are negative. [Beilstein **6** III 297.] (**±**)-**Borneol** [6627-72-1] C₁₀H₁₈O, **M 154.3, has m 206-207°, 210-215°**. Crystallise borneol from petroleum ether (b 60-80°) and sublime it *in vacuo*. [Beilstein **6** II 81, **6** IV 281.]

3-Bromoadamantane-1-carboxylic acid [21816-08-0] C₁₁H₁₅BrO₂, **M 259.1, m 145-146°, 146.5°, 147-150°, pK²⁵ 6.28 (50% aqueous EtOH)**. Purify the acid by recrystallising it from cyclohexane and/or subliming at 130°/10mm. It can be converted to the *methyl ester* (diazomethane) with **m 32°** (from petroleum ether at -10°). [Stetter & Mayer *Chem Ber* **95** 667 1962, DOI: 10.1002/cber.19620950314; Stetter & Wulff *Chem Ber* **93** 1366 1960, DOI: 10.1002/cber.19600930619; Bayal & Lantvoev *J Org Chem USSR (Engl Trans)* **9** 291 1973.]

1-Bromoadamantan-4-one (**5-bromo-2-adamantanone, 1-bromotricyclo[3.3.1.1^{3,7}]decan-4-one**) [20098-20-8] C₁₀H₁₃BrO, **M 229.1, m 149-153°, 150-152°, 153°, 151-155°**. Brinker and coworkers [Wagner et al. *Org Lett* **12** 332 2010, DOI: 10.1021/ol902667a] prepared it by dissolving 1-hydroxyadamantan-4-one (1g, 6.02mmol, 20098-14-0) in 48% aqueous HBr (20ml) solution and boiling under reflux for 16 hours, then adding H₂O (15ml) and extracting with Et₂O. The Et₂O extract is washed with brine, dried (MgSO₄) and evaporated to dryness *in vacuo* to give the bromo-ketone (0.8g, 3.49mmol, 58%) which has m 150-152°, and is recrystallised from hexane. It has ¹H NMR (400 MHz CDCl₃) δ : 1.97-2.10 (m 4H, 4 peaks), 2.24 (bs, 1H), 2.49-2.63 (m, 8H, four peaks); ¹³C NMR (150 MHz CDCl₃) δ : 31.2, 37.5, 47.8, 49.0, 59.9 and 214.3; and HRMS (EI) has m/z 228.0144, calc for C₁₀H₁₃BrO (M^+) 228.0150. The bromoketone is then converted into **1-bromo-4-aziadamantane** as follows: Anhydrous NH₃ was bubbled through a solution of 1-bromoandamantan-4-one (1.9g, 8.29mmol) in dry MeOH (30ml) at 0° for 75min while keeping the temperature between 0° and ~10°; then the solution is cooled to -12° and hydroxylamine-*O*-sulfonic acid (1.13g, 9.99mmol) is added in small amounts under stirring during 2.5hrs, and the mixture is kept at +7° overnight. The MeOH and NH₃ of this solution are removed *in vacuo* at 40°, the solid residue is suspended in Me₂CO (50ml) and CrO₃ (1.25g, 12.50mmol) in 20% H₂SO₄ (14ml) is added dropwise over 20min at 0°, then stirred for 1hr at ~25° and then poured onto ice (300g). The aqueous phase is extracted with CH₂Cl₂, the organic layer is dried (MgSO₄), filtered, evaporated to dryness and the residue is subjected to chromatography on SiO₂ and eluting with hexane. The eluent gave **1-bromo-4-aziadamantane** (0.96g, 3.98mmol, 48%) with **m 85-90°**, and IR (KBr) with ν_{\max} at 2934, 2858, 1577, 1451 1079, 1014, 809, 789 and 716 cm⁻¹; UV (cyclohexane) has λ_{\max} nm(ε) at 344(165), 349(141) and 362(226); ¹H NMR (400MHz, CDCl₃) with δ at 0.81 (bs, 2H), 1.77 (dm, 2H), 2.07 (bd, 2H, J = 12.9Hz), 2.24-2.27 (m, 1H),

2.33 (dm, 2H), 2.41 (s, 2H), 2.68 (dm, 2H); ^{13}C NMR (150MHz, CDCl_3) with δ at 31.6, 33.4, 37.9, 46.5, 62.5; HRMS (ESI) found m/z 455.0532 and calcd for $\text{C}_{20}\text{H}_{27}\text{Br}_2\text{N}_2$ (corresponding azine) $\text{M}+\text{H}^+$ 455.0521; and elemental analysis found C 49.46, H 5.11, N 11.50, calcd for $\text{C}_{10}\text{H}_{13}\text{BrN}_2$ C 49.81, H 5.43, N 11.62. [see also Geluk & Schlattmann *Tetrahedron* **24** 5369 1968, DOI: 10.1016/S0040-4020(01)96330-6.]

2-tert-Butyl-1-cyclohexanol [13491-79-7 isomeric mixture] $\text{C}_{10}\text{H}_{20}\text{O}$, M 156.2, m 43-46°, 52-53°, d_4^{25} 0.902. This is a mixture of geometric isomers obtained by the catalytic reduction of *o*-tert-butylphenol with a Ni-kieselguhr catalyst. The liquid residue deposited white rhombs m 52-53° of the isomeric mixture. However, upon oxidation with CrO_3 in $\text{AcOH-H}_2\text{O}$ (10:75) at 30-35° (overnight) all isomers yield the same racemic 2-tert-butyl-1-cyclohexanone described below. [Schmerling *J Am Chem Soc* **69** 1121 1947, DOI: 10.1021/ja01197a040; *Beilstein* **7** IV 82.]

When the urine of rabbits fed with (\pm)-2-tert-butyl-1-cyclohexanone (see next entry) is acidified (+)-*cis*-2-tert-butylcyclohexyl- β -D-glucosiduronic acid, $\text{C}_{16}\text{H}_{28}\text{O}_7 \cdot \text{H}_2\text{O}$, separates and can be obtained pure after two recrystallisations from 10% aqueous EtOH. It has m 167-168°, $[\alpha]_D^{18} +3.4$ (c 2.6, MeOH). By boiling a solution of this *glycosiduronic acid* under reflux in N HCl for 15 minutes, neutralising with NaHCO_3 , steam distilling, extracting the distillate with Et_2O , drying (Na_2SO_4), filtering, evaporating then distilling the residue under high vacuum, *S*-(+)-2-tert-butyl-1-cyclohexanol is obtained. This *S*-(+)-alcohol gave *S*-(+)-2-tert-butyl-1-cyclohexanone upon oxidation with CrO_3 (see below). [Cheo et al. *J Chem Soc (C)* 1988 1966, DOI: 10.1039/J39660001988.]

(\pm)-2-tert-Butyl-1-cyclohexanone [1728-46-7, racemate, also for (\pm) 13495-19-7] $\text{C}_{10}\text{H}_{18}\text{O}$, M 154.2, b 62.5°/4mm, d_4^{25} 0.896, n_D^{20} 1.4579. This ketone is obtained when the CrO_3 oxidation mixture from the preceding entry is diluted with MeOH- H_2O ((1:10) and steam distilled. The distillate is washed to remove any formaldehyde and AcOH present and then distilled under high vacuum. The (\pm)-semicarbazone, m 182-183°, separates slowly after the ketone is added to a mixture of $\text{NH}_2\text{CONHNH}_2 \cdot \text{HCl-NaOAc-H}_2\text{O}$ (1:1.5:10 w/w). [Schmerling *J Am Chem Soc* **69** 1121 1947, DOI: 10.1021/ja01197a040; *Beilstein* **7** IV 82.] *S*-(+)-2-tert-Butyl-1-cyclohexanone [14123-21-9] $\text{C}_{10}\text{H}_{18}\text{O}$, M 154.2, has b 40°/4x10⁻⁵mm, $[\alpha]_D^{22} -45.5$ (c 2.75, CHCl_3). Optical resolution of the *trans*-2-tert-butyl-1-cyclohexyl 3 β -acetoxy- Δ^5 -etienate {see (-)-3 β -acetoxy-5-etienic acid [51424-66-9] in 'Steroids' in Chapter 6} diastereoisomers by chromatographic separation on neutral Al_2O_3 (grade II) is achieved by eluting with $^*\text{C}_6\text{H}_6$ /hexane whereby the (-)-acid(-)-alcohol (m 181-182° from MeOH/ CH_2Cl_2 , $[\alpha]_D^{20} -45$) elutes before the (-)-acid(+)-alcohol [m 171-174° from MeOH/ CH_2Cl_2 , $[\alpha]_D^{20} -9$] which comes through on further elution with $^*\text{C}_6\text{H}_6$. LiAlH_4 in Et_2O cleaves the respective diastereomeric esters to provide *R*-(+)-2-tert-butyl-1-cyclohexanol [*trans*-1*R*,2*S*- 98104-30-4] {m 50-52° $[\alpha]_D^{20} +44.4$ (c 0.76, CHCl_3)} from the ester with m 181-182°, and *S*-(+)-2-tert-butyl-1-cyclohexanol [*trans*-1*S*,2*R*- 13492-07-4] {m 50-52° $[\alpha]_D^{20} +44.2$ (c 0.79, CHCl_3)} from the ester with m 171-174°. These have been used as chiral auxiliaries. [Djerassi et al. *J Am Chem Soc* **86** 78 1964, DOI: 10.1021/ja01055a019; *Beilstein* **6** III 126.]

Oxidation of the enantiomeric cyclohexanols by titration with a standardised solution of CrO_3 [Bowden, Heilbron, Jones and Weedon *J Chem Soc* 39 1946, DOI: 10.1039/JR9460000039] with occasional addition of small portions of anhydrous MgSO_4 , furnished *S*-(+)-2-tert-butyl-1-cyclohexanone [from the (+)-alcohol] and *R*-(+)-2-tert-butyl-1-cyclohexanone [from the (-)-alcohol], which are purified by distilling in a vacuum system at 3×10^{-5} mm using cooled receivers. Distillation can also be carried out in distilling equipment at high vacuum. The ORD for the *S*- enantiomer in MeOH at 22° exhibited Cotton extrema (λ_{max}) at $[\phi] -1650^\circ$ (317nm) and $+1670^\circ$ (275nm) with amplitude (a) equal to -33. The ORD of the *R*- enantiomer is $[\phi] +1690^\circ$ (317nm) and -1720° (275nm) with amplitude (a) equal to +34. These rotatory dispersions in MeOH were unchanged during 64 days at room temperature. Also in *isopropanol* solution containing one drop of concentrated HCl did not affect the amplitude of its Cotton effect at room temperature, however this decreased rapidly on heating at 65°: e.g. $[\alpha]_{317} -1089$ (25°) $\rightarrow [\alpha]_{317} -447$ (6 min at 65°) $\rightarrow [\alpha]_{317} -102$ (14 min at 65°). The IR and the retention times with g.l.c. were identical with those of the racemic ketone. [Djerassi et al. *J Am Chem Soc* **86** 78 1964, DOI: 10.1021/ja01055a019; *Beilstein* **7** IV 82.]

4-tert-Butyl-1-cyclohexanone [98-53-3] $\text{C}_{10}\text{H}_{18}\text{O}$, M 154.2, m 49-50°, 52-52.5, b 90-92°/9mm, 113-116°/20mm. Purify it *via* the *semicarbazone* (crystallised from EtOH with m 203-205°), hydrolyse this with dilute HCl and steam distil it. The distillate is extracted into Et_2O , dried, evaporated and the residue is recryst-

allised from pentane, aqueous EtOH or EtOH [Houlihan *J Org Chem* **27** 3860 1962, DOI: 10.1021/jo01058a024]. The *oxime* recrystallises from 1,2-dichloropropane and has **m 137.5-138.5°**. [Harvill et al. *J Org Chem* **15** 58 1950, DOI: 10.1021/jo01147a011; *Beilstein* **7** IV 82.]

(+)-Calarene [(+)- β -gurjunen, 1,3,3,11-tetramethyltricyclo[5.4.0.0^{2,4}]undecan-7-ene, (1*aR*)-1,1,7*c*,7*ac*-tetramethyl-1*a*,2,3,5,6,7,7*a*,7*b*-octahydro-1*H*-cyclopropa[α]naphthalene, new name 1(10)aristolene)] [17334-55-3] **C₁₅H₂₄**, **M 204.35**, **b 45-47°/0.008-0.01mm**, **120-123°/13mm**, **255-258°/atm**, **d₄²⁰ 0.9340**, **n_D²⁰ 1.55051**, **[α]_D²⁰ +73** (c 2, EtOH), **+81.8** (neat). Purify the Balsam sesquiterpene Calarene by gas chromatography (7% propylene glycol adipate on unglazed tile particles of size 0.2-0.3mm, 400 cm column length and 0.6 cm diameter, at 184°, with N₂ carrier gas at a flow rate of 0.54 ml/sec using a thermal detector). Also purify it by chromatography on alumina (200 times the weight of calarene) and elute with petroleum ether. Its UV has λ_{\max} at 200 and 210 nm (ϵ 9560, 5480) in EtOH. [IR: Šorm *Coll Czech Chem Commun* **18** 512 1953, DOI: 10.1135/cccc19530512; **29** 795 1964, DOI: 10.1135/cccc19640795, Buchi et al. *Tetrahedron Lett* **3** 827 1962, DOI: 10.1016/S0040-4039(00)70529-6; Vrkoc et al. *Tetrahedron Lett* **4** 225 1963, DOI: 10.1016/S0040-4039(01)90611-2; Palmod et al. *Bull Soc Chim Fr* 1050 1963, *Beilstein* **5** II 225, **5** III 1093.]

1*S*,4*R*-(-)-Camphanic acid (1*S*,4*R*-3-oxo-4,7,7-trimethyl-2-oxabicyclo[2.2.1]heptane-1-carboxylic acid) [13429-83-9] **C₁₀H₁₄O₄**, **M 198.2**, **m 190-192°, 198-200°**, **[α]₅₄₈²⁰ -22.5** (c 1, dioxane), **-4.4** (c 8, EtOH), **pK_{Est} ~3.8**. Dissolve the acid [derived from (+)-camphor] in CH₂Cl₂, dry (MgSO₄), filter, evaporate and the residue is sublimed at 120°/0.5mm or 140°/1mm. [Gerlach *Helv Chem Acta* **61** 2773 1978, DOI: 10.1002/hlca.19780610804; *Beilstein* **18/8** V 100.] The enantiomer **1*R*,4*S*-(+)-camphanic acid** [67111-66-9] has identical properties except for the opposite optical rotations. **1*S*,4*R*-(-)-Camphanic acid chloride** [39637-74-6] **C₁₀H₁₃O₃**, **M 216.7**, has **m 65-66.5°, 70.5-71°**, **[α]₅₄₈²² -23** (c 2, CCl₄), **[α]₃₆₄²⁰ -29.2**, **[α]₄₀₅²⁰ -18.0**, **[α]₄₃₆²⁰ -13.5**, **[α]₅₄₆²⁰ -7.8**, **[α]₅₇₈²⁰ -6.0**, (c 0.67, *C₆H₆), is soluble in toluene (50g/100ml at 0°) and crystallises from petroleum ether (b 40-60°). It sublimes at 70°/5mm. Store it dry at 0°, IR (CCl₄) has ν_{\max} at 1805s and 1780m cm⁻¹. [Armarego et al. *JCS Perkin Trans I* 2229 1976, DOI: 10.1039/P19760002229; Gerlach *Helv Chem Acta* **51** 1587 1968, DOI: 10.1002/hlca.19680510712; Gerlach *Helv Chem Acta* **68** 1815 1985, DOI: 10.1002/hlca.19850680702; *Beilstein* **18/8** V 101.]

RS-Camphene (2,2-dimethyl-3-methylenebicyclo[2.2.1]heptane [79-92-5, 565-00-4] **C₁₀H₁₆**, **M 136.2**, **m 51-52°**, **b 40-70°/10mm**, **55-56°/16mm**, **92.4°/100mm**, **55-56°/16mm**, **158.5-159.5°/760**, **d₄⁵⁴ 0.85**, **d₄⁵⁴ 0.8422**, **n_D⁵⁴ 1.45514**. Crystallise it twice from EtOH, then repeatedly melted and frozen at 30mm pressure. [Williams & Smyth *J Am Chem Soc* **84** 1808 1962, DOI: 10.1021/ja00869a007.] Alternatively, it is dissolved in Et₂O, dried over CaCl₂ and Na, filtered, evaporated and the residue is sublimed in a vacuum [NMR: Hana & Koch *Chem Ber* **111** 2527 1978, DOI: 10.1002/cber.19781110707]. [*Beilstein* **5** I 82, **5** II 105, **5** III 380, **5** IV 461.] **1*S*-(-)-Camphene** (1*S*-2,2-dimethyl-3-methylene norbornane) [5794-04-7] **C₁₀H₁₆**, **M 136.2**, has **m 49.2-49.6°, 49-50°, 52°**, **b 52°/17mm**, **79-80°/58mm**, **91.5°/100mm**, **158-160°/atm**, **d₄⁵⁴ 0.8412**, **n_D⁵⁴ 1.4564**, **[α]_D²¹ -119.1** (c 2.3, *C₆H₆), **-117.5** (c 19, toluene), **-113.5** (c 9.7, Et₂O). Purify this norbornane by fractionation through a Stedman column (see Chapter 1) at 100mm in a N₂ atmosphere, crystallise it from EtOH and sublime it in a vacuum below its melting point. It is characterised by its *camphenilone semicarbazone*, **m 217-218.5°**, or *camphor semicarbazone*, **m 236-238°**. [NMR: Hana & Koch *Chem Ber* **111** 2527 1978, DOI: 10.1002/cber.19781110707; Bartlett et al. *Justus Liebigs Ann Chem* **623** 217 1959, DOI: 10.1002/jlac.19596230124; Bain et al. *J Chem Soc* **72** 3124 1950, DOI: 10.1021/ja01163a087; *Beilstein* **5** H 156, **5** IV 461.] The enantiomer **1*R*-(+)-Camphene** [5794-03-6] has similar properties except for the opposite optical rotations. It is a flowering agent and is used in perfumery.

Camphor (1*R*-bornan-2-one, 1,7,7-trimethyl-2-oxabicyclo[2.2.1]heptan-2-one) [*R*-(+)- 464-49-3, *S*-(-)- 464-48-2, *RS*-(\pm)- 76-22-2] **C₁₀H₁₆O**, **M 136.2**, **m 178.8°, 179.97°**(open capillary), **b 204°/atm**, **[α]₅₄₆³⁵ (+) and (-) 59.6** (in EtOH), **[α]_D²⁰ (+) and (-) 44.3** (c 10, EtOH), **[α]₅₇₉¹⁷⁹ (+) and (-) 70.85** (melt). Recrystallise it from EtOH, 50% EtOH/water, MeOH, or petroleum ether or from glacial acetic acid by addition of water. It can be sublimed (50°/14mm) and also fractionally crystallised from its own melt. It is steam volatile. It should be stored in tight containers as it is appreciably volatile at room temperature. The solubility is 0.1% (H₂O), 100% (EtOH), 173% (Et₂O) and 300% (CHCl₃). The *R*-oxime [2792-42-9] **C₁₀H₁₇NO**, **M 167.2**, has **m 119°**

(from Et₂O, CHCl₃, or aqueous EtOH) and $[\alpha]_D^{20}$ -42.4 (c 3, EtOH); the \pm **oxime** has **m 118-119°**. It has a characteristic odour. [Asahina et al. *Chem Ber* **67** 1432 1934, DOI: 10.1002/cber.19340670828; Allen & Rogers *J Chem Soc (B)* 632 1971, DOI: 10.1039/J29710000632; UV, NMR: Fairlie et al. *JCS Perkin Trans 1* 2109 1973, DOI: 10.1039/P19730002109; White & Bishop *J Am Chem Soc* **62** 8 1940, DOI: 10.1021/ja01858a004; *Beilstein* **7** H 135, **7** II 104, **7** IV 213, 215.] It is a topical anti-itching agent, antiseptic and an analgesic material.

Camphoric acid (1,2,2-trimethylcyclopentan-1 α ,3 ϵ -dicarboxylic acid) [*1R,3S*]-(+)- 124-83-4, [*1S,3R*]-(-)- 560-09-8, 5394-83-2 ??] **C₁₀H₁₆O₄**, **M 200.2**, **m 186-188°, 187°, 186.5-189°, $[\alpha]_D^{20}$ (+) and (-) 57 (c 1, EtOH), $[\alpha]_D^{20}$ (+) and (-) 47.7 (c 4, EtOH), pK_1^{25} 4.71, pK_2^{25} 5.83 (for + isomer)**. Purify the acid by reprecipitation from an alkaline solution with HCl, filter it off, and recrystallise it from water several times, rejecting the first crop. It forms leaflets from EtOH, Me₂CO and H₂O, and is insoluble in CHCl₃. Its solubility in H₂O is 0.8% at 25° and 10% at 100°, 50% in EtOH and 5% in ethylene glycol. The (\pm)-**acid** has **m 202-203°**. The (+)-**1-methyl ester** has **m 86°** (from petroleum ether) $[\alpha]_D^{20}$ +45° (c 4, EtOH), and the (+)-**3-methyl ester** has **m 77°** (from petroleum ether) $[\alpha]_D^{17.5}$ +53.9° (c 3, EtOH). [Rupe & Thommen *Helv Chem Acta* **30** 933 1947, DOI: 10.1002/hlca.19470300402; Tiovonon et al. *Acta Chem Scand* **2** 597 1948, DOI: 10.3891/acta.chem.scand.02-0597; Howell & Fisher *J Am Chem Soc* **80** 6316 1958, DOI: 10.1021/ja01556a038; *Beilstein* **9** III 3878, **9** IV 2851.] (\pm)-**Camphoric anhydride** {(\pm)-**1,8,8-trimethyl-3-oxabicyclo[3.2.1]octane-2,4-dione**} [595-30-2, 76-32-4] **C₁₀H₁₄O₃**, **M 182.2**, has a **transition temperature of 135°, m 222-225°**. Crystallise the anhydride from EtOH. If it contains too much of the acid (check the IR), then reflux it in Ac₂O, concentrate and collect the crystals, wash them with petroleum ether and dry them *in vacuo*. [Bunton et al *J Chem Soc* 2918 1963, DOI: 10.1039/JR9630002918; NMR: Baker & Davis *Tetrahedron* **24** 1663 1968, DOI: 10.1016/S0040-4020(01)82473-X; *Beilstein* **18** H 400, 401.]

Camphorquinone (borna-2,3-dione) [*1R*]-(-)- 10334-26-6, [*1S*]-(+)- 2767-84-2] **C₁₀H₁₄O₂**, **M 166.2**, **m 198.7°, 198-199°, 197-201°, $[\alpha]_D^{25}$ (-) and (+) 101.1 (c 2, EtOH, or toluene)**. It can be purified by steam distillation, recrystallisation (yellow prisms) from EtOH, *C₆H₆ or Et₂O/petroleum ether and it can be sublimed in a vacuum. The (\pm)-**quinone** forms needles from EtOH, **m 197-198°, 203°**. [Rupe Buxtorf & Flatt *Helv Chem Acta* **13** 1026 1930, DOI: 10.1002/hlca.19300130523; Asahina et al. *Chem Ber* **67** 1432 1934, DOI: 10.1002/cber.19340670828; *Beilstein* **7** I 325, **7** III 3299, **7** IV 2039.] For biotransformation to the enantiomeric chiral 3-*exo*-hydroxycamphors using red algae see Utsukihara et al. *Tetrahedron Asymmetry* **17** 1179 2006, DOI: 10.1016/j.tetasy.2006.04.007. Forster described the preparation and separation of the four isomeric optically active **camphorquinone dioximes** {*anti* α $[\alpha]_D$ (-) 98.3, *syn* β $[\alpha]_D$ (-) 24.1, *amphi* γ $[\alpha]_D$ (+) 12.6, and *amphi* δ $[\alpha]_D$ (+) 98.3 all in 2% NaOH} obtained by heating an alcoholic solution of isonitrosocamphor with NH₂OH.HCl and NaOAc [Forster *J Chem Soc* **83** 514 1903, DOI: 10.1039/CT9038300514].

RS-Camphorquinone [10373-78-1] **C₁₀H₁₄O₂**, **M 166.2**, **m 199-202°**. Purification is the same as for above enantiomers above. [Huckel & Fechtig *Justus Liebigs Ann Chem* **652** 81 1962, DOI: 10.1002/jlac.19626520112]. A useful preparation is by the oxidation of camphor (5g) with SeO₂ (6g) in Ac₂O (5ml) at 140-150°/3-4hrs, cooling, filtering, washing the precipitated Se with AcOH, and the yellow filtrate is carefully neutralised with aqueous KOH. The crude quinone (5.22g, m 190-195°) is then recrystallised from ligroin and has **m 198°**. Its **semicarbazone** has **m 228-229° (dec)**, and the **p-bromophenylhydrazone** has **m 215-216°**. Use of dioxane as solvent instead of Ac₂O gave lower yields and a product that is more difficult to purify. This quinone provides the simplest route from camphor to camphoric anhydride. [Evans et al. *J Chem Soc* 137 1934, DOI: 10.1039/JR9340000137]. [*Beilstein* **7** I 325, **7** III 3299, **7** IV 2039.]

(*1R,E*)-(+)- and (*1S,E*)-(-) **Camphorquinone 3-oxime** [*1R,E*]-(+)- 31571-14-9, [*1S,E*]-(-)- 251645-83-7] **C₁₀H₁₅NO₂**, **M 181.2**. The **1R-3-seqcis-** enantiomer when purified by recrystallisation from *C₆H₆/petroleum ether (b 90-100°) has **m 156-157°**, and from AcOH it provides needles **m 156°**, $[\alpha]_D^{35}$ (+) 189 (c 1, CHCl₃) and $[\alpha]_D^{35}$ (+) 178.7 (c 1, *C₆H₆), $[\alpha]_D^{25}$ (+) 200 (c 1, EtOH). It has UV: λ_{max} 240nm (ϵ 9270) and on adding dilute aqueous KOH it changes to λ_{max} 298nm (ϵ 12,500); and IR has ν_{max} 3440, 1735 and 1640cm⁻¹; whereas the acetate [from reaction with AcO and recrystallised from *C₆H₆/petroleum ether (b 90-100°)] has UV: λ_{max} 233nm (ϵ 9150) and IR: ν_{max} 1780, 1745 and 1645cm⁻¹. [cf. Hassner et al. *J Org Chem* **28** 304 1963, DOI: 10.1021/jo01037a006.] It forms distinct Na and Cu salts (complexes). The **1S-3-seqcis-** enantiomer when

purified by recrystallisation from EtOH has **m 114-115°**, $[\alpha]_{\text{D}}^{35}$ (-) 169 (c 1, CHCl₃) and $[\alpha]_{\text{D}}^{35}$ (-) 150 (c 1.5, *C₆H₆). [Beilstein 7 H 583, 7 I 327, 7 II 554, 7 III 3301, 7 IV 2040.]

4-Carbethoxy-3-methyl-2-cyclohexen-1-one (Hagemann's ester, ethyl 2-methyl-4-oxo-2-cyclohexene-carboxylate) [487-51-4] C₁₀H₁₄O₃, M 182, b 79-80°/0.2mm, 121-123°/4mm, 142-144°/15mm, 268-272°/atm, n_{D}^{20} 1.488, d_4^{20} 1.038. Dissolve the ester in ether, shake with solid K₂CO₃, aqueous saturated NaHCO₃, brine, dry (MgSO₄) and distil it. The *semicarbazone* has **m 165-167° (169°)**. [Smith & Rouault *J Am Chem Soc* 65 631 1943, DOI: 10.1021/ja01244a040; Beilstein 10 H 631, 10 I 300, 10 III 2899, 10 IV 2666.]

α-Caryophyllene is now known as *α-humulene*, **β-caryophyllene** as *caryophyllene* and **γ-caryophyllene** as *isocaryophyllene* [see below and Aebi, Barton and Lindsey *J Chem Soc* 3124 1953, DOI: 10.1039/JR9530003124; Simonsen & Barton *The Terpenes* Vol III, Cambridge University Press 1952,]. Other CAS Registry numbers used are [54061-81-3 for (1S-(1R*,4Z,9S*))], [61217-74-1 for (1R*4Z-9S*)] and [33993-33-7],

Caryophyllene {previously known as **β-caryophyllene** see above, *trans*-(*-*)-**β-caryophyllene** (*-*)-*trans*-(1R,9S)-8-methylene-4,11,11-trimethyl-bicyclo[7.2.0]undec-4-ene} [87-44-5] C₁₅H₂₄, M 204.4, b 118-119°/9.7mm, 123-125°/10mm, 122°/13.5mm, 262-264°/atm, d_4^{20} 0.9075, n_{D}^{20} 1.4988, $[\alpha]_{\text{D}}^{20}$ -10 (neat), $[\alpha]_{\text{D}}^{21}$ -15 (c 2.6, CHCl₃), also -9.15 has been reported. Purify by fractional distillation *in vacuo* and collecting the fraction boiling at 123-125°/10mm with $[\alpha]_{\text{D}}^{20}$ -9.5 (neat). It can be obtained directly by fractionating clove-bud oil sesquiterpenes. [Barton & Lindsey *J Chem Soc* 2988 1951, DOI: 10.1039/JR9510002988, for absolute configuration see Barton & Nickon *J Chem Soc* 4665 1954, DOI: 10.1039/JR9540004665.] When dry HCl gas is passed through it, crystals of C₁₅H₂₄·2HCl are formed which have **m 69-70°**. Caryophyllene forms a crystalline *nitrosyl chloride* C₁₅H₂₄ClNO [-C(NO)—C(Cl)-] which has **m 159°(dec)** and $[\alpha]_{\text{D}}^{17}$ -98.1; and its *nitrobenzylamine* C₁₅H₂₄ClNO [-C(NO)—C(CH₂Ph)-] has **m 146-148°** after recrystallisation from *C₆H₆. When caryophyllene in petroleum ether is shaken with concentrated aqueous NaNO₂ with cooling, followed by slow addition of glacial acetic acid (volume equal to caryophyllene used) with shaking, the petroleum ether turns blue in colour and blue crystals of *caryophyllene nitrosite* C₁₅H₂₄N₂O₃ separate. Recrystallisation from petroleum ether or cold dilute acetone gives blue crystals with **m 115°**, $[\alpha]_{\text{D}}^{20}$ +103. It exhibits a 20° depression in melting point when mixed with humulene nitrosite (m 114°). [Deussen & Lewinsohn *Justus Liebigs Ann Chem* 356 1 1907, DOI: 10.1002/jlac.19073560102; 359 245 1908, DOI: 10.1002/jlac.19083590112; Chapman *J Chem Soc* 785 1928, DOI: 10.1039/JR9280000785; Beilstein 5 IV 1182.] For the conversion of *caryophyllene* (trans, E) to *isocaryophyllene* (cis, Z) see the latter below.

(*-*)-Caryophyllene oxide {1-S-5c-6t-epoxy-6c,10,10-trimethyl-2-methylene-1r,9t-bicyclo[7.2.0]undecane, (1R,4R,6R,10S)-9-methylene-4,12,12-trimethyl-5-oxatricyclo[8.2.0.0^{4,6}]dodecane} [1139-30-6] C₁₅H₂₄O, M 220.4, m 61-62°, 62-63°, 63.5-64°, 64°, b 114-117°/1.8mm, 141-142°/11mm, d_4^{20} 0.967, n_{D}^{20} 1.4956, $[\alpha]_{\text{D}}^{20}$ -79 (c 2, CHCl₃), $[\alpha]_{\text{D}}^{20}$ -68 (supercooled melt). Purify the oxide by TLC on silica gel with EtOAc/petroleum ether (b 60-80°) (15:85), and recrystallise it from MeOH or *C₆H₆. Also best purified by low-temperature crystallisation from MeOH. [Barton & Lindsey *J Chem Soc* 2988 1951, DOI: 10.1039/JR9510002988; NMR: Warnhoff *Can J Chem* 42 1664 1964, DOI: 10.1139/v64-249; Ramage & Whitehead *J Chem Soc* 4336 1954, DOI: 10.1039/JR9540004336; Beilstein 17 IV 392.]

(+)-Cedrol [octahydro-3,6,8,8-tetramethyl-1-3a,7-methanoazulen-6-ol, 8aS-6c-hydroxy-3c,6t,8,8-tetramethyl(8ar-H)-octahydro-3H,3at,7t-methanoazulene] [77-53-2] C₁₅H₂₆O, M 222.4, m 82-86°, 86-87°, $[\alpha]_{\text{D}}^{28}$ +10.5 (c 5, CHCl₃), $[\alpha]_{\text{D}}^{18}$ +13.1 (c 5.5, EtOH), $[\alpha]_{\text{D}}^{18}$ +14.3 (c 10, dioxane). Purify cedrol (fragrant oil from cedar and cypress) by recrystallisation from aqueous MeOH (colourless needles). It is estimated colorimetrically with H₃PO₄ in EtOH followed by vanillin and HCl [Hayward & Seymour *Anal Chem* 20 572 1948, DOI: 10.1021/ac60018a022]. The 3,5-dinitrobenzoyl derivative has **m 92-93°**. [Stork & Clarke *J Am Chem Soc* 83 3114 1961, DOI: 10.1021/ja01475a030; Beilstein 6 III 424.] The synthetic *racemic* (±)-Cedrol [22567-44-8] also crystallises nicely from aqueous MeOH and has **m 94-96°** and ¹H NMR (360 MHz CCl₄, TMS) with δ at 0.82 (3H, d, J = 6Hz, CH₃COH), 0.97 (3H, s, *endo*-CH₃), 1.18 (3H, s, *exo*-CH₃), 1.26 (3H, s, CH₃COH) [Breitholle & Fallis *J Org Chem* 43 1964 1978, DOI: 10.1021/jo00404a025].

S(+)-Chaulmoogric acid [13(cyclopent-2-enyl-1(*S*)-yl)tridecanoic acid] [29106-32-9] $C_{18}H_{32}O_2$, **M 280.4**, **m 68.5°**, **b 247-248°/20mm**, $[\alpha]_D^{20} +62.1$ (c 4, $CHCl_3$), $pK_{Est} \sim 5.0$. The acid was isolated from the oil of Chaulmoogra seeds *Taraktogenos kurzii* (King) [Power & Gornall *J Chem Soc* **85** 838, 851 1904, DOI: 10.1039/CT9048500838]. Crystallise the acid from petroleum ether or EtOH. Mislow and Steinberg prepared it electrolytically from *S*(+)-2-cyclopentene-1-acetic acid and ethyl hydrogen brassylate, after determining the absolute configuration of cyclopenteneacetic acid by two different methods. The **Me ester** [24828-59-9] $C_{19}H_{34}O_2$, **M 294.4**, prepared by bubbling HCl gas through a methanolic solution of the acid, has **m 22°**, **b 227°/20mm** and $[\alpha]_D^{15} +50$ (c 5, $CHCl_3$). The **amide**, prepared from the acid chloride and NH_3 , has **m 106°**, $[\alpha]_D^{27} +57.3$ (c 4.3, $CHCl_3$) after recrystallisation from EtOH. [Barrowcliff & Power *J Chem Soc* **91** 557 1907, DOI:10.1039/CT9079100557; Mislow & Steinberg *J Am Chem Soc* **77** 3807 1955, DOI: 10.1021/ja01619a038.] **S(+)-Chaulmoogric ethyl ester** (**Moogrol**, **Chaulmestrol**, **ethyl 13-cyclopent-2-enyltridecanoate**) [623-32-5] $C_{20}H_{36}O_2$, **M 308.5**, has **b 230°/20mm**, $d_{20}^{20} 0.90456$, $[\alpha]_D^{20} +50.7$ (neat), prepared as for the methyl ester (previous entry) with EtOH instead of MeOH, is a light yellow liquid with a fruity odour and nasty taste. It is active against *Mycobacterium leprae* and largely used to treat leprosy and various skin disorders [see Power & Gornall *J Chem Soc* **85** 838 1904, DOI: 10.1039/CT9048500838; Levy *Am Rev Respir Dis* **111** 703 1975, PMID: 1093460]. The ethyl ester is used in the treatment of leprosy and sarcoidosis [Levy *Am Rev Respir Dis* **111** 703 1975, PMID: 1093460].

Chlorendic anhydride (**1,4,5,6,7,7-hexachloro-5-norbornene-2,3-dicarboxylic anhydride**) [115-27-5] $C_9H_2Cl_6O_3$, **M 370.9**, **m 234-236°**, **235-237°**, **238°**. Steam distil the anhydride or recrystallise it from H_2O to yield pure diacid. The pure diacid yields the anhydride with Ac_2O . [Prill *J Am Chem Soc* **69** 62 1947, DOI: 10.1021/ja01193a015.]

1-Chloroadamantan-4-one (**5-chloro-2-adamantanone**, **1-chlorotricyclo[3.3.1.1^{3,7}]decan-4-one**) [20098-17-3] $C_{10}H_{13}BrO$, **M 229.1**, **m 197-200°**, **199-200°**. Brinker and coworkers [Wagner et al. *Org Lett* **12** 332 2010, DOI: 10.1021/ol902667a] prepared it by adding $SOCl_2$ (38ml, 0.52mol, CARE lachrymatory) to 4-hydroxyadamantan-1-one (6.16g, 37.1mmol, see [20098-14-0] below) and boiling under reflux for 2 hours, evaporating to dryness *in vacuo*, and dissolving the residue in CH_2Cl_2 . The solution CH_2Cl_2 is washed twice with aqueous 0.1N NaOH, twice with brine, dried (Na_2SO_4), filtered and evaporated to dryness. The residue is recrystallised from hexane or petroleum ether to give the 1-chloroadamantan-4-one (4.2g, 61% yield) with **m 199-200°**, and has IR (KBr) with ν_{max} at 2935s and 2860m (C-H), 1725vs (C=O, + 5 peaks as in adamantanone), 1289m, 1060s, 1024s, 829s, 664m and 473m cm^{-1} ; the 1H NMR (400 MHz $CDCl_3$) has δ at 1.92-2.04 (m 4H, 4 peaks), 2.27-2.31 (m, 2H), 2.32-2.35 (s, 3H), 2.42 (d, 2H, $J = 12.3Hz$), 2.61 (s, 2H); ^{13}C NMR (150 MHz $CDCl_3$) with δ at 30.6, 37.6, 46.5, 47.5, 48.0, 64.5 and 214.6; and HRMS (EI) has m/z 184.0653, calc for $C_{10}H_{13}ClO$ (M^+) 184.0655. [See also Geluk & Schlatmann *Tetrahedron* **24** 5369 1968, DOI: 10.1016/S0040-4020(01)96330-6.] **1-chloro-4-aziadamantane** is prepared in the same way as for **1-bromo-4-aziadamantane** above but in 27% yield, and after recrystallisation from hexane it started to decompose at *ca* 80-90°, and has IR (KBr) with ν_{max} at 2934, 2859, 1577, 1453, 1079, 1020, 828 and 587 cm^{-1} ; UV (pentane) has λ_{max} nm(ϵ) at 348(134), 353(114) and 367(207); 1H NMR (400MHz, $CDCl_3$) with δ at 0.84 (bs, 2H), 1.73 (dm, 2H), 2.01 (bd, 2H, $J = 12.9Hz$), 2.11 (dm, 2H), 2.20 (brs, 2H), 2.28-2.32 (m, 1H), 2.48 (m, 2H, $J = 11.3Hz$); ^{13}C NMR (150MHz, $CDCl_3$) with δ at 30.9, 33.5, 37.1, 45.0, 47.0 and 66.4; HRMS (ESI) found m/z 365.1558 and calcd for $C_{20}H_{27}Cl_2N_2$ (corresponding azine) $M+H^+$ 365.1551; and elemental analysis found C 61.07, H 6.07, Cl 18.17, N 14.13, calculated for $C_{10}H_{13}ClN_2$ C 61.07, H 6.66, Cl 18.03, N 14.24. [see also Geluk & Schlatmann *Tetrahedron* **24** 5369 1968, DOI: 10.1016/S0040-4020(01)96330-6.]

Chlorocyclohexane (cyclohexyl chloride) [542-18-7] $C_6H_{11}Cl$, **M 118.6**, **b 46-48°/26mm**, **142.5°/atm**, $d_4^{20} 1.00$, $n_D^{25} 1.46265$. It has been prepared using the general procedure of Norris and coworkers by refluxing the alcohol with 5 volumes of 'constant boiling HCl' for 2-4 hours [Norris et al. *J Am Chem Soc* **38** 1071 (1077) 1916, DOI: 10.1021/ja02262a014]. It has also been prepared in 76% yield by the ' $CaCl_2$ ' procedure described for chlorocyclopentane (see below [930-28-9]). Wash the chlorocyclohexane several times with dilute $NaHCO_3$, then repeatedly with distilled water. Dry it with $CaCl_2$ and fractionally distil it slowly at atmospheric pressure or better under vacuum. [Perlman et al. *J Org Chem* **1** 288 (294) 1936, DOI: 10.1021/jo01232a008; IR: Roberts & Chambers *J Am Chem Soc* **73** 5030 1951, DOI: 10.1021/ja01155a004; *Beilstein* **5** H 21, **5** I 8, **5** II 11, **5** III 37, **5** IV 48.]

2-Chlorocyclohexanone [822-87-7] $\text{C}_6\text{H}_9\text{ClO}$, M 132.6, m 23°, b 82-83°/10mm, 98-99°/14-15mm, d_4^{25} 1.161, n_D^{20} 1.484. The chlorocyclohexanone is prepared in a well ventilated fume cupboard by bubbling chlorine (215g, ~3moles, use mercury traps) rapidly through a mixture of cyclohexanone (284g, 3moles) and H_2O (900ml) while being stirred in an ice bath at a reaction temperature between 20° and 50°. After about 45 minutes all the chlorocyclohexanone separates as a heavy oil which is combined with the Et_2O (3 x 150ml) extract of the upper aqueous layer, washed with H_2O (150ml), saturated aqueous NaCl (200ml), filtered (by gravity) through anhydrous Na_2SO_4 , the solvent is evaporated, the residue is distilled in a Claisen flask, and the fraction boiling below 110° (300-340g, b 110°/13mm, 92°/4mm) is collected. This is then carefully fractionated through a heated 42-inch Vigreux column with a variable take-off head to give pure 2-chlorocyclohexanone (240-265g, 61-66%, b 90-91°/14-15mm) with 5-13% recovery of cyclohexanone b 52°/14-15mm. [Newman et al. *Org Synth Coll Vol* 3 188 1855, DOI: 10.15227/orgsyn.025.0022; *Beilstein* 7 H 10, 7 I 8, 7 II 11, 7 III 36, 7 IV 32.]

Chlorocyclopentane (cyclopentyl chloride) [930-28-9] $\text{C}_5\text{H}_9\text{Cl}$, M 104.6, b 113-115°/atm, 114°/atm, d_4^{20} 1.005, n_D^{20} 1.4512. It is prepared by mixing cyclopentanol (43g) with concentrated HCl (125ml) and anhydrous CaCl_2 (50g), and stirring under reflux on a steam bath for 10 hours. [As HCl gas evolves continuously, the reaction should be carried out in an sufficient fume cupboard.] After cooling, the upper layer is collected, washed with brine, saturated aqueous NaHCO_3 (CARE, as evolution of CO_2 will occur), brine again and dried over anhydrous CaCl_2 for at least 24 hours. Filter off the solid and fractionate through an efficient column at atmospheric pressure to obtain a ~ 58% yield of cyclopentyl chloride (30g). Bubbling dry HCl gas through the reaction mixture does not appear to increase the yield. See preparation and purification of chlorocyclohexane above. [*Beilstein* 5 IV 18.]

(-)- α -Copaene (1R,2S,6S,7S,8S-8-isopropyl-1,3-dimethyltricyclo[4.4.0.0^{2,7}]dec-3-ene) [3856-25-5] $\text{C}_{15}\text{H}_{24}$, M 204.4, b 119-120°/10mm, 246-251°, d_4^{20} 0.908, n_D^{20} 1.489, $[\alpha]_D^{20}$ -6.3 (c 1.2, CHCl_3). It is a naturally occurring liquid tricyclic sesquiterpene [Simonen & Barton *The Terpenes Vol III*, pp 88-91 1952, University Press, Cambridge]. Purify it by distillation, preferably under vacuum. Other purifications involved a Varian Aerograph A-700 gas chromatograph (He carrier gas with thermal conductivity detection) and Hewlett-Packard Model 810 and 5750 gas chromatographs (N_2 carrier gas, 30 mL/min, flame ionisation detection) which separated the copaene from the isomeric *ylangene* which also occur naturally. The racemate has been synthesised. [Heathcock *J Am Chem Soc* 88 4110 1966, DOI: 10.1021/ja00969a051; Heathcock et al. *J Am Chem Soc* 89 4133 1967, DOI: 10.1021/ja00992a032; Corey & Watt *J Am Chem Soc* 95 2303 1973, DOI: 10.1021/ja00788a034; *Beilstein* 5 IV 1189.]

Cyclobutane [287-23-0] C_4H_8 , M 56.1, m -50°, -80°, b 13°/740mm, 12°/atm, d_4^{20} 0.721, n_D^{20} 1.426. This easily liquefiable gas is dried over Na at melting ice temperature for 4 days and distilled at low temperature through a Podbielniak precision still. A dry sample has been prepared by passage through P_2O_5 and distilled repeatedly until all fractions had similar vapour pressures at 0°. [Cason & Way *J Org Chem* 14 31 1949, DOI: 10.1021/jo01153a006; Heisig *J Am Chem Soc* 63 1698 1941, DOI: 10.1021/ja01851a055; Heisig & Stodola *Org Synth Coll Vol* 3 213 1955, DOI: 10.15227/orgsyn.023.0016.]

Cyclobutane carboxylic acid [3721-95-7] $\text{C}_5\text{H}_8\text{O}_2$, M 100.1, m 3-4°, -5.4°, -7.5°, b 84-84.5°/10mm, 110°/25mm, 135-138°/110mm, 194°/760mm, d_4^{20} 1.061, n_D^{20} 1.453, pK^{25} 4.79. Dissolve the acid in aqueous HCO_3^- then acidify with HCl and extract it into Et_2O , wash with H_2O , dry (Na_2SO_4), concentrate to a small volume, then distil it through a glass helices packed column. The *S-benzylisothiuronium salt* has m 176° (from EtOH), the *anilide* has m 112.5-113°, and the *p-toluide* has m 123°. [Payne & Smith *J Org Chem* 22 1680 1957, DOI: 10.1021/jo01363a041; Kantro & Gunning *J Am Chem Soc* 73 480 1951, DOI: 10.1021/ja01145a520; Stodola & Heisig *Org Synth Coll Vol* 3 213 1955, DOI: 10.15227/orgsyn.023.0016; *Beilstein* 9 H 5, 9 IV 6.] The *acid chloride* [5006-22-4] $\text{C}_5\text{H}_7\text{ClO}$, M 118.6, b 60°/50mm, d^{25} 1.039, n_D^{20} 1.455, is best purified by adding 5% v/v of oxalyl chloride (or more if the OH band in the IR is intense) and distilling it *in vacuo*.

Cyclobutane-1,1-dicarboxylic acid [5445-51-2] $\text{C}_6\text{H}_8\text{O}_4$, M 144.1, m 154-156°, 156.5°, 157°, 158°, 159°(dec), pK_1^{25} 3.13, pK_2^{25} 5.88. Recrystallise the acid from Et_2O , $^*\text{C}_6\text{H}_6$ or $^*\text{C}_6\text{H}_6/\text{EtOAc}$. It forms trans-

parent colourless crystals on evaporating spontaneously a concentrated Et₂O solution. It is freely soluble in H₂O, moderately soluble in Et₂O, CHCl₃, and *C₆H₆ but only sparingly in light petroleum. It forms, Cu (blue needles), Pb (colourless prisms), Ba (slender tiny needles) and Na salts. When heated rapidly it melts at the stated temperatures but if heating is slow it decomposes at lower temperatures with evolution of CO₂ to form an oil which is the *monocarboxylic acid* (**b** 191°/720mm, see preceding acid). [Perkin *J Chem Soc* **51** 1 1887, DOI: 10.1039/CT8875100001; Jeffrey & Vogel *J Chem Soc* 1804 1948, DOI: 10.1039/JR9480001804; German et al. *J Chem Soc* 1624 1935, DOI: 10.1039/JR9350001624.] The *dimethyl ester* has **b** 87°/8mm, **d**₄²⁰ 1.1182, **n**_D²⁰ 1.4415; the *diethyl ester* has **b** 102-105°/10mm, 226.3°/760mm, **d**₄²⁰ 1.0466, **n**_D²⁰ 1.4391 [Cason & Allen *J Org Chem* **14** 1036 1949, DOI: 10.1021/jo01158a011; Heisig & Stodola *Org Synth* **23** 16 1943, DOI: 10.15227/orgsyn.023.0016], and the *diamide* has **m** 275-277°, 278° (from EtOH) [Ingold et al. *J Chem Soc* **121** 1177 1922, DOI: 10.1039/CT9222101177; Dox & Yoder *J Am Chem Soc* **43** 677 1921, DOI: 10.1021/ja01436a040]. [Beilstein **9** H 724, **9** II 514, **9** III 3797, **9** IV 3979.]

cis-Cyclobutane-1,2-dicarboxylic acid (*meso*-isomer) [1461-94-5] C₆H₈O₄, **M** 144.1, **m** 139.5°, 139.5-140°, 139-140°, **pK**₁²⁵ 4.16, **pK**₂²⁵ 6.23 (**pK**₁²⁰ 4.20, **pK**₂²⁰ 6.56). Purify the acid by crystallisation from H₂O or ligroin, or by hydrolysis with H₂O of the *anhydride* (3-oxabicyclo[3.2.0]hepta-2,4-dione) [7687-27-6] C₆H₆O₃, **M** 126.1 [**b** 120-150°/40mm, **m** 77-77.5° (from *C₆H₆, 74-75° from H₂O or ligroin), Georgian et al. *Tetrahedron* **19** 1219 1963, DOI:10.1016/S0040-4020(01)98583-7]. It isomerises to the *trans*-isomer at 190° in the presence of HCl. The *dimethyl ester* [2607-03-6] C₈H₁₂O₄, **M** 172.1, has **b** 114°/20mm, 225°/atm. The *diphenacyl ester* has **m** 113° (from EtOH) and the *p*-bromodiphenacyl ester has **m** 153° (from EtOH/Me₂CO). [Buchman et al. *J Am Chem Soc* **64** 2696 1942, DOI: 10.1021/ja01263a048; Vogel *Justus Liebigs Ann Chem* **615** 1 1958, DOI: 10.1002/jlac.19586150102; Reed *J Chem Soc* 685 1951, DOI: 10.1039/JR9510000685; Ellingboe & Fuson *J Am Chem Soc* **56** 1774 1934, DOI: 10.1021/ja01323a037; **pK**: Bode *Chem Ber* **67** 332 1934, DOI: 10.1002/cber.19340670236; Georgian et al. *Tetrahedron* **19** 1219 1963, DOI: 10.1016/S0040-4020(01)98583-7; Beilstein **9** IV 2788.]

trans-Cyclobutane-1,2-dicarboxylic acid [(±)-1*RS*,2*RS* 58616-94-7; 1124-13-6] C₆H₈O₄, **M** 144.1, **m** 130.5-131°, 131° 130.5-131°, **b** 140°/2mm, **pK**₁²⁵ 3.94, **pK**₂²⁵ 5.55. Crystallise the acid from H₂O (**m** 130.5-131°), *C₆H₆ (**m** 131°), *C₆H₆/dioxane (**m** 130.5-131°), or *C₆H₆/EtOAc. The *cis*-acid isomerises to the *trans*-acid on heating in concentrated HCl at 190°. The *methyl ester* [3396-20-1] C₈H₁₂O₄, **M** 172.2, has **m** 106°, **b** 225°/atm and **d** 1.17. The *diphenacyl ester* has **m** 98° (from EtOH) and the *p*-bromodiphenacyl ester has **m** 158° (from EtOH). [Buchman et al. *J Am Chem Soc* **64** 2696 1942, DOI: 10.1021/ja01263a048; Reed *J Chem Soc* 685 1951, DOI: 10.1039/JR9510000685; Fuson et al. *J Am Chem Soc* **51** 1536 1929, DOI: 10.1021/ja01380a034; Ellingboe & Fuson *J Am Chem Soc* **56** 1774 1934, DOI: 10.1021/ja01323a037; **pK**: Bode *Chem Ber* **67** 332 1934, DOI: 10.1002/cber.19340670236; Beilstein **9** H 725, **9** II 515, **9** III 3799, **9** IV 2788.] The dicarboxylic acid was resolved via the (+)-quinine salt whereby the least soluble (+)-quinine/(-)-acid salt crystallised from H₂O, and from which pure *1*R*,2*R*-(+)-dicarboxylic acid* [17224-72-5] has 116-117°, [**α**]_D²⁹ -150 (**c** 0.8, H₂O) and [**α**]_D²⁵ -158 (MeOH, highest observed), can be isolated (by basifying the salt with NH₃ and extracting with Et₂O) and recrystallised from hydrochloric acid or *C₆H₆ [Goldsworthy *J Chem Soc* **125** 2012 1924, DOI: 10.1039/CT9242502012; Coyner & Hillman *J Am Chem Soc* **71** 324 1949, DOI: 10.1021/ja01169a091]. Its absolute configuration *1*R*,2*R*-(+)* was deduced by comparison of the ORD, [**φ**]_D -223 (MeOH), of sulfur derivatives with those of related compounds [Inouye et al. *Tetrahedron* **23** 3237 1967, DOI: 10.1016/S0040-4020(01)92293-8]. Its *di-N-methylamide* has [**α**]_D -249 (MeOH). Less pure *1*S*,2*S*-(+)-dicarboxylic acid* [58616-97-0], [**α**]_D³⁰ +123 (**c** 1.2, H₂O), was similarly obtained from the more soluble (+)-quinine/(+)-acid salt, and its pure *diethyl ester*, C₁₀H₁₂O₄, has **b** 236°/761mm [**α**]_D³⁰ +77.9 (**c** 1.7, Me₂CO) [Goldsworthy *J Chem Soc* **125** 2012 1924, DOI: 10.1039/CT9242502012]. [Beilstein **9** II 515, **9** III 3799, **9** IV 2788.]

Cyclobutanone [1191-95-3] C₄H₆O, **M** 70.1, **b** 96-97°, 99-100°/atm, **d**₄²⁰ 0.931, **n**_D²⁰ 1.4210, **n**_D⁵² 1.4189. Treat cyclobutanone with dilute aqueous KMnO₄, dry it with molecular sieves and fractionally distil it. Purify it via the semicarbazone, then regenerate the ketone, dry it (CaSO₄), and distil it in a stainless steel spinning-band (or Vigreux column). Alternatively, purify it by preparative gas chromatography using a Carbowax 20-M column at 80° (this treatment also removes acetone). It has FT-IR (NaCl) with **v**_{max} at 3543.3, 2970.3 (CH), 1783.2 (C=O), 1392.5 (OH enol ?), 1208.9, 1136.4, 1080.8, 724.9, and 461.4 cm⁻¹; ¹H NMR (60MHz, CDCl₃,

TMS) with δ at 2.01 (q, 2H, C-3 methylene), 3.09 (t, 4H, C-2 and C-4 methylenes); ^{13}C NMR (15MHz, CDCl_3 , TMS) with δ at 9.74 (C-3), 47.73 (C-2,4) and 208.90 (C-1). The *oxime* has *m* 84-85° (from petroleum ether) and the *semicarbazone* has *m* 212-212.5° (220-221° from MeOH or H_2O , Buchman et al. *J Am Chem Soc* **64** 2701 1942, DOI: 10.1021/ja01263a049). [Salaun et al. *Org Synth* **57** 36 1977, DOI: 10.15227/orgsyn.057.0036, Fitjer & Quabeck *Synthesis* 299 1987, DOI: 10.1055/s-1987-27926; *Beilstein* **7** IV 3.]

Cyclobutylamine [2516-34-9] $\text{C}_4\text{H}_9\text{N}$, *M* 71.1, *b* 82-83°/atm, 83.2-84.2°/760mm, d_4^{20} 0.839, n_D^{20} 1.437, *pK*²⁵ 10.04 (9.34 in 50% aqueous EtOH). It has been purified by steam distillation. The aqueous distillate (e.g. 2L) is acidified with 3N HCl (90ml) and evaporated to dryness in a vacuum. The *hydrochloride* is treated with a few ml of H_2O , cooled in ice and a slush of KOH pellets ground in a little H_2O is added slowly in portions and keeping the solution very cold. The amine separates as an oil from the strongly alkaline solution. The oil is collected, dried over solid KOH and distilled using a vacuum jacketed Vigreux column and protected from CO_2 using a soda lime tube. The fraction boiling at 79-83° is collected, dried over solid KOH for 2 days and redistilled over a few pellets of KOH (*b* 80.5-81.5°). Best distil in a dry N_2 atmosphere. The purity can be checked by GLC using a polyethylene glycol on Teflon column at 72°, 15 psi, flow rate of 102 ml/min of He. The sample can appear homogeneous but because of tailing it is not possible to tell if H_2O is present. The NMR in CCl_4 should show no signals less than 1 ppm from TMS. The *hydrochloride* has a multiplet at ca 1.5-2.6ppm (H 2,2,4,3,3,4,4), a quintet at 3.8 ppm (H 1) and a singlet at 4.75 for NH_2 [Roberts & Mazur *J Am Chem Soc* **73** 2509 1951, DOI: 10.1021/ja01112a048]. The *benzenesulfonamide* has *m* 85-86° (from aqueous MeOH) and the *benzoyl* derivative has *m* 120.6-121.6°. [Roberts & Mazur *J Am Chem Soc* **73** 2509 1951, DOI: 10.1021/ja01150a029; Iffland et al. *J Am Chem Soc* **75** 4044 1953, DOI: 10.1021/ja01112a048; Werner & Casanova Jr *Org Synth Coll Vol* **5** 273 1973, DOI: 10.15227/orgsyn.047.0028; *Beilstein* **12** IV 3.]

(±)-**Cyclobutylol** (α -ethyl-1-hydroxycyclohexaneacetic acid) [512-16-3] $\text{C}_{10}\text{H}_{18}\text{O}_3$, *M* 186.25, *m* 81-81°, *b* 167-170°/16mm, 164°/24mm, d_4^{20} 1.001, n_D^{20} 1.4680. It forms colourless crystals from petroleum ether, and can be distilled *in vacuo* without decomposition. It is obtained by hydrolysis of ethyl α -ethyl-1-hydroxycyclohexaneacetate [b 125-127°/12mm, 135-136°/20mm, Wallach *Justus Liebigs Ann Chem* **360** 26 1908, DOI: 10.1002/jlac.19083600104] which is better performed as follows, otherwise decomposition to butyric acid and cyclohexanone occurs. The ester (100g) is added to a hot solution of KOH (100g) in H_2O (100ml), then MeOH (100ml) is added gradually and the mixture is refluxed for 2-3hrs. The solution is freed from MeOH under a vacuum, diluted with H_2O , extracted with Et_2O , and the aqueous solution is acidified with ice-cold 30% H_2SO_4 , the free hydroxyl-acid is extracted into Et_2O , the extract is dried (Na_2SO_4), filtered, evaporated and the residue is distilled under reduced pressure. It is very soluble in organic solvents, sparingly in petroleum ether and acidic aqueous solution but very soluble in aqueous alkaline solutions. [Kon and Narayanan *J Chem Soc* 1536 1927, DOI: 10.1039/JR9270001536; Kandiah & Linstead *J Chem Soc* 2139 1929, DOI: 10.1039/JR9290002139; Maillard et al. *Bull Soc Chim Fr* 244 1958, *Beilstein* **10** III 38.]

The *sodium salt* (*Trade names: Colepan, Dimene, Hebutol, Bilimix*) [1130-2300] $\text{C}_{10}\text{H}_{17}\text{O}_3\text{Na}$, *M* 208.2, has *m* 299-300°. It is a white crystalline powder which should be stored in sealed containers as it is slightly hygroscopic. It is a choleric agent which inhibits biliary secretions.

Cyclodecanone [1502-06-3] $\text{C}_{10}\text{H}_{18}\text{O}$, *M* 154.2, *m* 21-24°, *b* 100-102°/12mm. Purify the ketone *via* the *semicarbazone* (*m* 205-207°, from EtOH) and distil it through an efficient column. It sublimes in a vacuum. The *oxime* has *m* 80°, from MeOH or by sublimation in a high vacuum. [Cope et al. *Org Synth Coll Vol* **4** 218 1963, DOI: 10.15227/orgsyn.036.0014; Prelog et al. *Helv Chim Acta* **30** 1741 1947, DOI: 10.1002/hlca.19470300637; Ruzicka et al. *Helv Chim Acta* **11** 670 1928, DOI: 10.1002/hlca.19280110178; *Beilstein* **7** III 134, **7** IV 76.]

cis-**Cyclodecene** [935-31-9] $\text{C}_{10}\text{H}_{18}$, *M* 138.3, *m* -3°, -1°, *b* 73°/15mm, 90.3°/33mm, 194-195°/740mm, 197-199°/atm, d_4^{20} 0.8770, n_D^{20} 1.4854. *cis*-Cyclodecene has been prepared by the Pd-catalysed hydrogenation of cyclododecyne, and by dechlorination of 1-chlorocyclodecene using Na and liquid NH_3 . Purify it by fractional distillation preferably after chromatographing three times through silica gel. The IR has ν_{max} (film) at 1656m, 1684m, 1715w (C=C), 2680w, 2915vs (C-H) cm^{-1} . It forms an AgNO_3 complex which crystallises from MeOH, *m* 167-187°. Attempted thermal isomerisation of pure *cis*-isomer (sealed tube, under N_2 with hydroquinone and 2-naphthalenesulfonic acid, at 170-180°/1.5 h) gave a *cis/trans* mixture. [Cope et al. *J Am*

Chem Soc **77** 1628 1955, DOI: 10.1021/ja01611a066; IR: Blomquist et al. *J Am Chem Soc* **74** 3636 1952, DOI: 10.1021/ja01134a051; Prelog et al. *Helv Chim Acta* **35** 1598 1952, DOI: 10.1002/hlca.19520350526; *Beilstein* **5** IV 295.]

trans-Cyclodecene [2198-20-1] $C_{10}H_{18}$, **M 138.3**, **b 68-70°/10mm, 70.5-71.5°/11mm, 78°/18mm, 194°/740mm**, d_4^{20} **0.8681**, n_D^{20} **1.4846**. It has been prepared by shaking cyclodecyl-trimethylammonium iodide with Ag_2O in MeOH (2 h) then filtered, the filtrate was evaporated and the residual oil (free base) was decomposed by heating (in a bath at 100°) at 150mm. The distillate was extracted with Et_2O , the extract was washed with dilute HCl, followed by aqueous Na_2CO_3 solution, H_2O , then was distilled through a short Vigreux column to provide **trans-cyclodecene** in 50% yield. Chromatography through silica gel prior to distillation has been used. Final purification was by distillation over Na *in vacuo*. The IR has ν_{max} (film) at 1667m (C=C), 2675w, 2915vs (C-H) cm^{-1} . [Cope et al. *J Am Chem Soc* **77** 1628 1955, DOI: 10.1021/ja01611a066; IR: Blomqvist et al. *J Am Chem Soc* **74** 3636 1952, DOI: 10.1021/ja01134a051; Prelog et al. *Helv Chim Acta* **35** 1598 1952, DOI: 10.1002/hlca.19520350526; *Beilstein* **5** IV 295.]

cis-cis-trans-1,5,9-Cyclododecatriene (cyclododec-1c,5c,9t-triene) [2765-29-9] $C_{10}H_{18}$, **M 162.3**, **m -9°, -8°, b 117.5°/2mm, 237-239°/atm, 244°/760mm**, d_4^{20} **0.907**, n_D^{20} **1.5129**. Purify the triene by fractional distillation, preferably in a vacuum under N_2 , and it forms an insoluble $AgNO_3$ complex. [IR: Breil et al. *Makromol Chemie* **69** 18 1963, DOI: 10.1002/macp.1963.020690102; *Beilstein* **5** IV 1114.]

Cyclododecylamine [1502-03-0] $C_{12}H_{25}N$, **M 183.3**, **m 27-29°, b 122-124°/7mm, 140-150°/ca 18mm, 280°/atm, pK 9.62 (in 80% methyl cellosolve)**. It can be purified *via* the **HCl salt m 274-275°** (from EtOH) or the **picrate m 232-234°**, and the free base is distilled preferably at water-pump vacuum. It is a strong base and should be stored away from moisture and CO_2 . [Prelog et al. *Helv Chim Acta* **33** 365 1950, DOI: 10.1002/hlca.19500330220; *Beilstein* **12** IV 164.]

1,3-Cycloheptadiene [4054-38-0] C_7H_{10} , **M 94.2**, **b 55°/75mm, 71.5°/150mm, 120-121°/atm**, d_4^{20} **0.868**, n_D^{20} **1.4972**. Purify the diene by dissolving it in Et_2O , washing with 5% HCl, H_2O , drying ($MgSO_4$), evaporating, and the residue is distilled under dry N_2 through a semi-micro column (some foaming occurs) [Cope et al. *J Am Chem Soc* **79** 6287 1957, DOI: 10.1021/ja01580a047; UV: Pesch & Friess *J Am Chem Soc* **72** 5756 1950 DOI: 10.1021/ja01168a512]. [*Beilstein* **5** H 115, **5** III 317, **5** IV 390.]

Cycloheptane [291-64-5] C_7H_{14} , **M 98.2**, **m ~12°, ~13°, b 114.4°, 118°/atm**, d_4^{20} **0.812**, n_D^{20} **1.4588**. Distil it from sodium using a Vigreux column, under nitrogen. It is highly flammable. [Bocian & Strauss *J Am Chem Soc* **99** 2866 1977, DOI: 10.1021/ja00451a004; Ruzicka et al. *Helv Chim Acta* **28** 395 1945, DOI: 10.1002/hlca.660280152; *Beilstein* **5** H 92, **5** IV 92.]

Cycloheptanol [502-41-0] $C_7H_{14}O$, **M 114.2**, **m 2°, b 77-81°/11mm, 83-84/14mm, 185°/atm**, d_4^{20} **0.955**, n_D^{20} **1.471**. Purify it as described for cyclohexanol. The **2,4-dinitrobenzoyl** derivative has **m 79°** and the **allophanate** has **m 184°** (from EtOAc). [Ruzicka et al. *Helv Chim Acta* **28** 395 1945, DOI: 10.1002/hlca.660280152; *Beilstein* **6** H 10.]

Cycloheptanone (suberone) [502-42-1] $C_7H_{12}O$, **M 112.2**, **b 105°/80mm, 172.5°/760mm, 179°/atm**, d_4^{20} **0.949**, n_D^{20} **1.461**. Shake suberone with aqueous $KMnO_4$ to remove material absorbing around 230-240nm, then dry it with Linde type 13X molecular sieves and fractionally distil it through a glass helix packed column. [Blicke et al. *J Am Chem Soc* **74** 2924 1952, DOI: 10.1021/ja01131a503; Dauben et al. *Org Synth Coll Vol* **4** 221, 1963, DOI: 10.15227/orgsyn.034.0019; *Beilstein* **7** H 13, **7** I 9, **7** II 14, **7** III 46, **7** IV 39.]

Cycloheptatriene [544-25-2] C_7H_8 , **M 92.1**, **b 60.5°/122mm, 114-115°/atm**, d_4^{20} **0.895**, n_D^{20} **1.522**. Wash the triene with alkali, then fractionally distil it. Store it under N_2 or Ar as it resinifies in air. [Dryden *J Am Chem Soc* **76** 2841 1954, DOI: 10.1021/ja01639a076; Kohler et al. *J Am Chem Soc* **61** 1057 1939, DOI: 10.1021/ja01874a021; *Beilstein* **5** IV 765.]

Cycloheptylamine [5452-35-7] **M 113.2**, **b 50-52°/11mm, 60°/18mm**, d_4^{20} **0.887**, n_D^{20} **1.472**, **pK_{Est} ~10.5**

(H₂O), pK²⁴ 9.99 (in 50% aqueous methyl cellosolve). It can be purified by conversion to the *hydrochloride* **m 242-246°**, and the free base is distilled under dry N₂ in a vacuum [Cope et al. *J Am Chem Soc* **75** 3212 1953, DOI: 10.1021/ja01109a049; Prelog et al. *Helv Chim Acta* **33** 365 1950, DOI: 10.1002/hlca.19500330220]. [Beilstein **12** IV 115.]

1,3-Cyclohexadiene (1,2-dihydrobenzene) [592-57-4] C₆H₈, **M 80.1**, **m -89°**, **b 83-84°/atm**, **d₄²⁰ 0.840**, **n_D²⁰ 1.471**. Distil the diene from NaBH₄ or Na under N₂ and collect it in a trap cooled in Dry Ice. Stabilise it with ~0.1% hydroquinone. It is highly flammable. [Marvel & Martzell, *J Am Chem Soc* **81** 448 1959, DOI: 10.1021/ja01511a047; for use see Feng et al. *Organometallics* **25** 5456 2006, DOI: 10.1021/om0606385; Beilstein **5** IV 382.] **1,4-Cyclohexadiene (1,4-dihydrobenzene)** [628-41-1] C₆H₈, **M 80.1**, has **b 83-86°/714mm**, **88.3°/741mm**, **86-88°/atm**, **88.7-89°/760mm**, **d₄²⁰ 0.8573**, **n_D²⁰ 1.4725**. Dry the diene over CaCl₂ and distil it in a vacuum under N₂. Stabilise it with ~0.1% hydroquinone. [Hückel & Wörrfel *Chem Ber* **88** 338 1955, DOI: 10.1002/cber.19550880305; Giovannini & Wegmüller *Helv Chim Acta* **42** 1142 1959, DOI: 10.1002/hlca.19590420403; for use see Feng et al. *Organometallics* **25** 5456 2006, DOI: 10.1021/om0606385; Beilstein **5** IV 385.]

Cyclohexane [110-82-7] C₆H₁₂, **M 84.2**, **m 6.47°**, **6.6°**, **4-7°**, **b 6.7°/40mm**, **14.7°/60mm**, **25.5°/100mm**, **42°/200mm**, **60.8°/400mm**, **80.7°/760mm**, **d₄²⁴ 0.77410**, **n_D²⁰ 1.42623**, **n_D²⁵ 1.42354**. It is best to purify it by washing with concentrated H₂SO₄ until the washings are colourless, followed by water, aqueous Na₂CO₃ or 5% NaOH, and again water until neutral. It is then dried with P₂O₅, Linde type 4A molecular sieves, CaCl₂, or MgSO₄, then Na and distilled. Cyclohexane has been refluxed with, and distilled from Na, CaH₂, LiAlH₄ (which also removes peroxides), sodium/potassium alloy, or P₂O₅. Traces of *benzene can be removed by passage through a column of silica gel that has been freshly heated: this gives material suitable for ultraviolet and infrared spectroscopy. If there is much *benzene in the cyclohexane, most of it can be removed by a preliminary treatment with **nitrating acid** (a cold mixture of 30ml concentrated HNO₃ and 70ml of concentrated H₂SO₄) which converts *benzene into nitrobenzene. The impure cyclohexane and the nitrating acid are placed in an ice bath and stirred vigorously for 15 minutes, after which the mixture is allowed to warm to 25° during 1 hour. The cyclohexane is decanted, washed several times with 25% NaOH, then water, dried with CaCl₂, and distilled from sodium. Carbonyl-containing impurities can be removed as described for chloroform. Other purification procedures include passage through columns of activated alumina and repeated crystallisation by partial freezing. Small quantities may be purified by chromatography on a Dowex 710-Chromosorb W gas-liquid chromatographic column. It is very soluble in most organic solvents, but less so in MeOH (57% w/v at 20°), and is difficultly soluble in H₂O (~0.0052% w/w at 24°). **Flammable liquid**. [Sabatier *Ind Eng Chem* **18** 1005 1926, Scheffland & Jacobs *The Handbook of Organic Solvents* (Van Nostrand) p592 1953, Beilstein **5** IV 27.] It is **TOXIC** at relatively low concentrations causing irritation of the eyes, skin (can cause dermatitis), respiratory tract and can cause necrosis and coma at high concentrations. It is highly volatile with *vapour pressures* ca 77mm at 20° and 169mm at 38°.

Rapid purification: Distil, discarding the forerun. Stand distillate over Grade I alumina (5% w/v) or 4A molecular sieves.

Cyclohexane butyric acid [4441-63-8] C₁₀H₁₈O₂, **M 170.3**, **m 31°**, **26.5-28.5°**, **b 136-139°/4mm**, **169°/20mm**, **188.8°/46mm**, **pK²⁵ 4.95**. Distil the acid through a Vigreux column, and the crystalline distillate is recrystallised from petroleum ether at low temperatures. The *S-benzylisothiuronium salt* has **m 154-155°** (from EtOH) [Friediger & Pedersen *Acta Chem Scand* **9** 1425 1955, DOI: 10.3891/acta.chem.scand.09-1425; English & Dayan *J Am Chem Soc* **72** 4187 1950, DOI: 10.1021/ja01165a099]. [Beilstein **9** II 15, **9** IV 69.]

Cyclohexane carboxylic acid (hexahydrobenzoic acid) [98-89-5] C₇H₁₂O₂, **M 128.2**, **m 31-32°**, **b 63-67°/-0.1mm**, **110°/8mm**, **232-233°/atm**, **d₁₅ 1.480**, **n_D²⁰ 1.460**, **pK²⁵ 4.90**. Crystallise the acid from hot H₂O (solubility is 0.2% w/w at 15°), it is soluble in organic solvents. Also distil it at as high a vacuum as possible and warm the condenser as it solidifies on cooling. The *acid chloride* [2719-27-9] **M 146.6**, has **b 184°/atm**, **d₂₅ 1.096**, the *methyl ester* [4630-82-4] C₈H₁₄O₂, **M 142.2**, has **b 183°/atm**, the *carboxamide* [1122-56-1] C₇H₁₃NO, **M 127.2**, has **m 186-188°** (from aqueous EtOH), the *carbonitrile* [766-05-2] C₇H₁₁N, **M 109.2**, has **m 11°**, **b 75-76°/16mm**, and the *S-benzylisothiuronium salt* has **m 165-166°** (from EtOH). [Beilstein **9** H 7-9, **9** I 5, **9** II 6, **9** III 15, **9** IV 16.]

Cyclohexane-1,2-diaminetetraacetic acid H_2O , (CDTA, 1,2-cyclohexanedinitrilotetraacetic acid) [H_2O : 12333-90-4; $x\text{H}_2\text{O}$: 13291-61-7] $\text{C}_{14}\text{H}_{22}\text{N}_2\text{O}_8$, H_2O , M 364.4, m >210°(dec), 213-216°, pK_1 1.34, pK_2 3.20, pK_3 5.75 (6.12), pK_4 9.26 (12.35). Dissolve CDTA in aqueous NaOH as its disodium salt, then precipitate it by adding HCl. The free acid is filtered off and boiled with distilled water to remove traces of HCl [Bond & Jones *Trans Faraday Soc* **55** 1310 1959, DOI: 10.1039/TF9595501310]. Recrystallise it from water and dry it *in vacuo*. [Beilstein **13** III 10.]

cis-Cyclohexane-1,2-dicarboxylic acid (*cis*-hexahydrophthalic acid) [610-09-3] $\text{C}_8\text{H}_{12}\text{O}_4$, M 172.2, m 191-192°, 191-194°, pK_1^{25} 4.25, pK_2^{25} 6.74. It is purified by recrystallisation from EtOH or H_2O . [Smith & Byrne *J Am Chem Soc* **72** 4406 1950, DOI: 10.1021/ja01166a019; Abell *J Org Chem* **22** 769 1957, DOI: 10.1021/jo01358a012; Beilstein **9** III 3812, **9** IV 2801.] **trans-Cyclohexane-1,2-dicarboxylic acid** (*trans*-hexahydrophthalic acid) [2305-32-0] $\text{C}_8\text{H}_{12}\text{O}_4$, M 172.2, has m 227.5-228°, 228-230.5°, pK_1^{25} 4.30, pK_2^{25} 6.06. It is purified by recrystallisation from EtOH or H_2O . It is formed by hydrolysing the anhydride with water. The *dimethyl ester* has m 95-96° (from C_6H_6 /petroleum ether). [Abell *J Org Chem* **22** 769 1957, DOI: 10.1021/jo01358a012; Smith & Byrne *J Am Chem Soc* **72** 4406 1950, DOI: 10.1021/ja01166a019; Beilstein **9** III 3812, **9** IV 2802.] The *1R,2R*-(-)-*trans-cyclohexane-1,2-dicarboxylic acid* [46022-05-3] has m 171-182° and $[\alpha]_D^{20}$ -20 (c 1, Me_2CO).

cis-Cyclohexane-1,2-dicarboxylic anhydride (*cis*-hexahydrophthalic anhydride) [85-42-7, 13149-00-3] $\text{C}_8\text{H}_{10}\text{O}_3$, M 154.2, m 32-34°, b 158°/17mm. It has been obtained by heating the *trans-acid* or *anhydride* at 200°. Crystallise it from $\text{C}_6\text{H}_6/\text{Et}_2\text{O}$ or distil it. [Kohler & Jansen *J Am Chem Soc* **60** 2142 1938, DOI: 10.1021/ja01276a034; Abell *J Org Chem* **22** 769 1957, DOI: 10.1021/jo01358a012; Beilstein **17** H 452, **17** III/IV 5931.]

trans-Cyclohexane-1,2-dicarboxylic anhydride (*trans*-hexahydrophthalic anhydride) [14166-21-3] $\text{C}_8\text{H}_{10}\text{O}_3$, M 154.2, m 140-142°, 145-146°. Crystallise the anhydride from $\text{C}_6\text{H}_6/\text{Et}_2\text{O}$. It has been obtained by heating the *cis-acid* or *anhydride* with HCl at 180° for 3 hours. It is formed from the acid by heating in Ac_2O . It sublimes at 125-135°/0.02mm. [Kohler & Jansen *J Am Chem Soc* **60** 2142 1938, DOI: 10.1021/ja01276a034; Fichter & Simon *Helv Chim Acta* **17** 1218 1934, DOI: 10.1002/hlca.193401701152; Beilstein **17** H 452, **9** IV 2802,]]

(±)-trans-1,2-Cyclohexanediol [1460-57-7] $\text{C}_6\text{H}_{12}\text{O}_2$, M 116.2, m 104°, 105°, 120°/14mm. Crystallise the diol from Me_2CO and dry it at 50° for several days. It can also be recrystallised from CCl_4 or EtOAc and it can be distilled. The *2,4-dinitrobenzoyl* derivative has m 179°. [Winstein & Buckles *J Am Chem Soc* **64** 2780 1942, DOI: 10.1021/ja01264a020.] [Beilstein **6** IV 5194.] **trans-1,2-Cyclohexanediol** [*1R,2R*-(-)- 1072-86-2, *1S,2S*-(+)- 57794-08-8] $\text{C}_6\text{H}_{12}\text{O}_2$, M 116.2, has m 107-109°, 109-110.5°, 111-112°, 113-114°, $[\alpha]_D^{22}$ (-) and (+) 46.5 (c 1, H_2O). The enantiomers have been recrystallised from C_6H_6 or EtOAc. The (±) diol has been resolved *via* the distychnine salt of the hemisulfate [Hayward et al. *JCS Perkin Trans 1* 2413 1976, DOI: 10.1039/P19760002413], or the *l*-menthoxy acetates. {*l*-*trans*-diastereoisomeric salt has m 64°, $[\alpha]_D$ -91.7 (c 1.4, EtOH) from petroleum ether or aqueous EtOH and yields the (-)-*trans*-diol} and {*d*-*trans*-diastereoisomeric salt has m 126-127°, $[\alpha]_D$ -32.7 (c 0.8, EtOH) from petroleum ether or aqueous EtOH and yields the (+)-*trans*-diol}. The **bis-4-nitrobenzoate** has m 126.5° $[\alpha]_D$ (-) and (+) 25.5 (c 1.1, CHCl_3), and the **bis-3,5-dinitrobenzoate** has m 160° $[\alpha]_D$ (-) and (+) 83.0 (c 1.8, CHCl_3) [Wilson & Read *J Chem Soc* 1269 1935, DOI: 10.1039/JR9350001269]. [Beilstein **6** III 4060.]

cis-1,3-Cyclohexanediol (hexahydroresorcinol) [823-18-7] $\text{C}_6\text{H}_{12}\text{O}_2$, M 116.2, m 86°, 87°, 137°/13mm. Crystallise the *cis*-diol from ethyl acetate and acetone or distil it in a vacuum. The *dibenzoyl* derivative has m 65.5° (from MeOH or petroleum ether). [Rigby *J Chem Soc* 1586 1949, DOI: 10.1039/JR9490001586; Furberg & Hassell *Acta Chem Scand* **4** 597 1950, DOI: 10.3891/acta.chem.scand.04-0597 Beilstein **6** III 4077, **6** IV 5208.] **trans-1,3-Cyclohexanediol** (hexahydroresorcinol) [5515-64-0] $\text{C}_6\text{H}_{12}\text{O}_2$, M 116.2, has m 117°, 118-118.5°, 135°/13mm. Crystallise the *trans*-diol from ethyl acetate or Me_2CO . The *dibenzoyl* derivative has m 123.5° (from EtOH or petroleum ether). [Rigby *J Chem Soc* 1586 1949, DOI: 10.1039/JR9490001586; Beilstein **6** III 4077, **6** IV 5208.]

cis-1,4-Cyclohexanediol (hexahydrohydroquinone) [931-71-5] $C_6H_{12}O_2$, M 116.2, m 102.5°, 113-114°. Crystallise the *cis*-diol from acetone (charcoal), then dry and sublime it under vacuum. It also crystallises from Me_2CO or $Me_2CO/*C_6H_6$. The *diacetate* has m 40.6-41.1° (from petroleum ether or 34-36° from EtOH). [Grob & Baumann *Helv Chim Acta* **38** 594 1955, DOI: 10.1002/hlca.19550380306; Owen & Robins *J Chem Soc* 320 1949, DOI: 10.1039/JR9490000320; *Beilstein* **6** III 4080, **6** IV 5209.] **trans-1,4-Cyclohexanediol (hexahydrohydroquinone)** [6995-79-5] $C_6H_{12}O_2$, M 116.2, has m 142.6-143.1°. Crystallise the *trans*-diol from MeOH or Me_2CO . The *diacetate* has m 104.5-105° (from petroleum ether or 102-103° from EtOH). [Grob & Baumann *Helv Chim Acta* **38** 594 1955, DOI: 10.1002/hlca.19550380306; Owen & Robins *J Chem Soc* 320 1949, DOI: 10.1039/JR9490000320; *Beilstein* **6** III 4080, **6** IV 5209.]

Cyclohexane-1,2-dione [765-87-7] $C_6H_8O_2$, M 112.1, m 34-38°, 39°, b 75-79°/16mm, 193-195°/atm, pK_1^{20} 10.30 (40% enol form, stabilised by intramolecular H-bonding). It is prepared by SeO_2 oxidation of cyclohexanone (8 hours at 87°) in 60% yield [Hach et al. *Org Synth Coll Vol* **4** 229 1963, DOI: 10.15227/orgsyn.032.0035]. The distillate solidifies to ice-like crystals m 34°. [*Beilstein* **7** IV 1982.] See below for the dioxime. **Cyclohexane-1,2-dione dioxime (Nioxime)** [492-99-9] $C_6H_{10}N_2O_2$, M 142.2, has m 189-190°, 199°, pK_1^{25} 10.68, pK_2^{25} 11.92. Crystallise Nioxime from alcohol/water and dry it in a vacuum at 40°. Also 2.5g of oxime have been recrystallised from 550ml of H_2O using Fe free Norit. [Hach et al. *Org Synth Coll Vol* **4** 229 1963, DOI: 10.15227/orgsyn.032.0035.] It forms complexes with some metals, e.g. Fe^{2+} , Cu^{2+} , Co^{2+} , Ni^{2+} and Pd^{2+} , and the *urea complex*, $C_6H_{10}N_2O_2 \cdot 2CH_4N_2O$, m 116-117°, crystallises from H_2O . [Banks et al. *Anal Chim Acta* **19** 531 1958, DOI: 10.1016/S0003-2670(00)88214-8; *Beilstein* **7** III 3210, **7** IV 1982.]

Cyclohexane-1,3-dione (dihydroresorcinol) [504-02-9] $C_6H_8O_2$, M 112.1, m 101-105°, 107-108°, pK_1^{25} 5.25. Crystallise the dione from *benzene. Dissolve ~50g of the diol in 140ml of $*C_6H_6$ under N_2 , cool, collect the solid and dry it in a vacuum desiccator overnight. It is unstable and should be stored under N_2 or Ar at ~0°. [Thompson *Org Synth Coll Vol* **3** 278 1955, DOI: 10.15227/orgsyn.027.0021; *Beilstein* **7** IV 1985.]

Cyclohexane-1,4-dione [637-88-7] $C_6H_8O_2$, M 112.1, m 76-77°, 78°, 79.5°, 79-80°, b 130-133°/20mm, d_4^{21} 1.0861, n_D^{102} 1.4576. Crystallise the dione from water, then *benzene. It can also be recrystallised from $CHCl_3$ /petroleum ether or Et_2O . It has been purified by distillation in a vacuum, and the pale yellow distillate which solidified is then recrystallised from CCl_4 (14.3 g/100 ml) and has m 77-79°. The *di-semicarbazone* has m 231°, the *dioxime HCl* has m 150° (from $MeOH/*C_6H_6$) and the *bis-2,4-dinitrophenylhydrazone* has m 240° (from $PhNO_2$). [Nielsen & Carpenter *Org Synth Coll Vol* **5** 288 1973, DOI: 10.15227/orgsyn.045.0025; IR: LeFevre & LeFevre *J Chem Soc* 3549 1956, DOI: 10.1039/JR9560003549.] [*Beilstein* **7** IV 1986.]

1,4-Cyclohexanedione monoethylene acetal (1,4-dioxaspiro[4.5]decan-8-one) [4746-97-8] $C_8H_{12}O_3$, M 156.2, m 70-73°, 73.5-74.5°. Recrystallise it from petroleum ether. It sublimes slowly on attempted distillation. Also purify it by dissolving it in Et_2O and adding petroleum ether (b 60-80°) until turbid, and cooling. [Gardner et al. *J Org Chem Soc* **22** 1206 19 57, DOI: 10.1021/jo01361a021; Britten & Lockwood *JCS Perkin Trans I* 1824 1974, DOI: 10.1039/P19740001824; cf: Revial et al. *J Org Chem* **67** 2252 2002, DOI: 10.1021/jo0110597.] [*Beilstein* **19/4** V 93.]

cis,cis-1,3,5-Cyclohexane tricarboxylic acid [16526-68-4] $C_9H_{12}O_6$, M 216.2, m 216-218°, $pK_{Est(1)} \sim 4.1$, $pK_{Est(2)} \sim 5.4$, $pK_{Est(3)} \sim 6.8$. Purify the acid by recrystallisation from toluene/EtOH or H_2O . It forms a *1.5 hydrate* with m 216-218°, and a *dihydrate* m 110°. Purify it also by conversion to the *triethyl ester* b 217-218°/10mm, 151°/1mm, the distillate solidifies on cooling, m 36-37°, which is hydrolysed by boiling in aqueous HCl. The *trimethyl ester* can be distilled and recrystallised from Et_2O , m 48-49°. [Newman & Lowrie *J Am Chem Soc* **76** 4598 1954, DOI: 10.1021/ja01647a028; Lukes & Galik *Coll Czech Chem Comm* **19** 712 1954, DOI: 10.1135/cccc19540712; *Beilstein* **9** III 4749, **9** IV 3723.]

Cyclohexanol [108-93-0] $C_6H_{12}O$, M 100.2, m 20-22°, 25.2°, b 161.1°/atm, d_4^{25} 0.946g/ml, n_D^{25} 1.466, n_D^{30} 1.437, n_D^{30} 1.462. Reflux it with freshly ignited CaO, or dry it with Na_2CO_3 , then fractionally distil it. Redistil it from a very small amount of sodium, about 0.5-2% depending on the amount of H_2O estimated to be present. It is further purified by fractional crystallisation from the melt in dry air. Peroxides and aldehydes can be re-

moved by prior washing with ferrous sulfate and water, followed by distillation under nitrogen from 2,4-dinitrophenylhydrazine, using a short fractionating column: water distils as the azeotrope. Dry cyclohexanol is *very hygroscopic*, store in a dry atmosphere. The **3,4-dinitrobenzoate** has **m 111-112°** (EtOH or aqueous EtOH). It has **TOXIC** vapours. [Beilstein 6 III 10, 6 IV 20.]

Cyclohexanone [108-94-1] $C_6H_{10}O$, **M 98.2**, **m -32.1°, -47°, b 1.4°/1.0mm, 26.4°/5mm, 38.7°/10mm, 52.5°/20mm, 67.8°/40mm, 77.5°/60mm, 90.4°/100mm, 110.3°/400mm, 132.5°/400mm, 155.7°/atm, d_4^{20} 0.947, n_D^{15} 1.452, n_D^{20} 1.451, pK^{25} -6.8 (aqueous H_2SO_4), pK^{25} 11.3 (enol), 16.6 (keto).** Dry cyclohexanone over $MgSO_4$, $CaSO_4$, Na_2SO_4 or Linde type 13X molecular sieves, then distil it. Cyclohexanol and other oxidisable impurities can be removed by treatment with chromic acid or dilute $KMnO_4$. More thorough purification is possible by conversion to the bisulfite addition compound, or the semicarbazone, followed by decomposition with Na_2CO_3 and steam distillation. [For example, equal weights of the bisulfite adduct (crystallised from water) and Na_2CO_3 are dissolved in hot water and, after steam distillation, the distillate is saturated with NaCl and extracted with Et_2O which is then dried (anhydrous $MgSO_4$ or Na_2SO_4), filtered, and the solvent evaporated prior to further distillation.] Its solubility in H_2O is 15%w/v (at 10°) and 5%w/v (at 30°). Conversely the solubility of H_2O in cyclohexanone is 7.8% at 20°. **FLAMMABLE.** The **semicarbazone** has **m 167°**, the **4-nitrophenylhydrazone** has **m 147°**, the **2,4-dinitrophenylhydrazone** has **m 162°**, and the **benzal derivative** has **m 118°**. [Beilstein 7 III 14, 7 IV 15.] **Cyclohexanone cyanohydrin** [931-97-5] $C_7H_{11}NO$, **M 125.2**, has **m 32-35°, b 132°/19mm**, and should be stored in a tight container as it may slowly liberate HCN [Beilstein 10 H 5]. **Cyclohexanone dimethyl ketal (1,1-dimethoxycyclohexane)** [933-40-4] $C_8H_{16}O_2$, **M 144.2**, has **b 83°/50mm, d_4^{25} 0.948, n_D^{20} 1.4360**, and is purified by fractional distillation after washing with dilute aqueous acid and drying ($MgSO_4$). Should not have a C=O frequency in the IR spectrum. [Beilstein 7 IV 19.] **Cyclohexanone oxime** [100-64-1] $C_6H_{11}NO$, **M 113.2**, has **m 91°, b 100-105°/10-12mm, 206-210°/atm**. Crystallise the oxime from water or petroleum ether (b 60-80°). [Bousquet *Org Synth Coll Vol 2* 313 1943, DOI: 10.15227/orgsyn.011.0054; Beilstein 7 III 32, 7 IV 21.] **Cyclohexanone phenylhydrazone** [946-82-7] $C_{12}H_{15}N_2$, **M 187.3**, has **m 77°, 81°**. Crystallise it from EtOH.

Cyclohexanone-2-carboxylic acid (2-oxocyclohexane carboxylic acid) [18709-01-8] $C_7H_{10}O_3$, **M 142.1, de-carboxylates at 81-82°**. A preparation which is reproducible on a large scale involves adding cyclohexanone (49g, 52ml, 0.5mole) to a stirred suspension of $NaNH_2$ in Et_2O [obtained by dissolving Na (14g, 0.61g atom) in liquid NH_3 (500ml), and then replacing the NH_3 by Et_2O (500ml) carefully]. The mixture is boiled for 30 minutes, kept at ~35° while dry CO_2 is bubbled through for 3 hours, then poured into a slurry of ice and excess of 2M aqueous HCl, extracted with Et_2O (5 x 300ml), and the Et_2O is extracted with excess of saturated aqueous Na_2CO_3 . The ice cold Na_2CO_3 solution is acidified with 2M aqueous HCl and again extracted with Et_2O (5 x 300ml). The combined extracts are dried ($MgSO_4$), filtered and evaporated *in vacuo* to give the carboxylic acid (29g, 41%) which can be recrystallised from Et_2O to form colourless needles. [Christie & Reid *JCS Perkin Trans I* 880 1976, DOI: 10.1039/P19760000880; Gardner et al. *J Chem Soc* 1910 1764, cf. Chiba et al. *Chem Lett* 1387 1978, DOI: 10.1246/cl.1978.1387.] The **methyl ester (methoxycarbonyl cyclohexanone)** [41302-34-5] $C_8H_{12}O_3$, **M 156.2**, has **b 50°/0.2mm, d_4^{20} 1.10**, and is purified by distillation through a short column if a film shows no OH bands in the IR spectrum; however if these bands are present then treat it with diazomethane in dry Et_2O until the yellow colour persists, and then distil under *in vacuo*. [Beilstein 10 IV 2606.]

Cyclohexene (tetrahydrobenzene) [110-83-8] C_6H_{10} , **M 82.2, m -104°, b 83°/atm d_4^{20} 0.810, n_D^{20} 1.4464, n_D^{25} 1.4437**. Free cyclohexene from peroxides by washing with successive portions of dilute acidified ferrous sulfate, or with $NaHSO_3$ solution, then with distilled water, drying with $CaCl_2$ or $CaSO_4$, and distilling under N_2 . Alternative methods for removing peroxides include passage through a column of alumina, refluxing with sodium wire or cupric stearate (then distilling from sodium). The diene is removed by refluxing with maleic anhydride before distilling under vacuum. Treatment with 0.1moles of $MeMgI$ in 40ml of diethyl ether removes traces of oxygenated impurities. Other purification procedures include washing with aqueous NaOH, drying and distilling under N_2 through a spinning band column, redistilling from CaH_2 , storing under sodium wire, and passing through a column of alumina, under N_2 , immediately before use. Store it at <0° under argon. [Coleman & Johnstone *Org Synth Coll Vol I* 183 1941, DOI: 10.15227/orgsyn.005.0033; Corson & Ipatieff *Org Synth Coll Vol 2* 151 1943, DOI: 10.15227/orgsyn.019.0036; Woon et al. *J Am Chem Soc* 108 7990 1986, DOI:

10.1021/ja00285a018; Wong et al. *J Am Chem Soc* **109** 3428 1987 DOI: 10.1021/ja00245a039.] [*Beilstein* **5** IV 218.] It can also be freed from peroxides by distilling over silica gel or alumina [Hershberg & Ruhoff *Org Synth* **17** 25 1937, DOI: 10.15227/orgsyn.017.0025] which may be another way of further purifying it.

(±)-2-Cyclohexen-1-ol (3-hydroxycyclohex-1-ene) [822-67-3] $C_6H_{10}O$, **M 98.1** **b 63-65°/12mm, 65-66°/13mm, 67°/15mm, 74°/25mm, 85°/35mm, 166°/atm, d_4^{20} 0.9865, n_D^{20} 1.4720.** Purify 2-cyclohexen-1-ol by distillation through a short Vigreux column. The **2,4-dinitrobenzoyl** derivative has **m 120.5°**, and the **phenylurethane** has **m 107°**. [Pedersen et al. *Org Synth* **48** 18 1968, DOI: 10.15227/orgsyn.048.0018; Cook *J Chem Soc* 1774 1938, DOI: 10.1039/JR9380001774; Dreiding & Hartman *J Am Chem Soc* **75** 3723 1953, DOI: 10.1021/ja01111a034; *Beilstein* **6** IV 196.]

Cyclohexylamine [108-91-8] $C_6H_{13}N$, **M 99.2**, **m -17°, -17.7°, b 30.5°/15mm, 36.4°/20mm, 45.1°/30mm, 56°/50mm, 72°/100mm, 102.5°/300mm, 118.8°/1500mm, 134.5°/atm, water azeotrope (44.2% amine) has b 96.4°/760mm, d_4^{20} 0.866, d_4^{25} 0.863, n_D^{20} 1.4593, n_D^{25} 1.456, pK^{25} 10.63.** Dry the amine with $CaCl_2$ or $LiAlH_4$, then distil it from BaO, KOH or Na, under N_2 . Also purify it by conversion to the hydrochloride (which is crystallised several times from water), then liberation of the amine with alkali and fractional distillation under N_2 . The **hydrochloride** has **m 205-207°** (dioxane/EtOH). [Lycan et al. *Org Synth Coll Vol* **2** 318 1943, DOI: 10.15227/orgsyn.011.0058; *Beilstein* **12** III 10, **12** IV 8.]

Cyclohexyl bromide [108-85-0] $C_6H_{11}Br$, **M 163.1**, **b 72°/29mm, 163-165°/atm, d_4^{20} 0.902, n_D^{25} 1.4935.** Shake the bromide with 60% aqueous HBr to remove the free alcohol. After removing excess HBr, the sample is dried and fractionally distilled. [IR: Roberts & Chambers *J Am Chem Soc* **73** 5030 1951, DOI: 10.1021/ja01155a004; *Beilstein* **5** III 48, **5** IV 67.]

1-Cyclohexylethylamine [*S*-(+)- 17430-98-7, *R*-(-)- 5913-13-3] $C_8H_{17}N$, **M 127.2**, **b 177-178°/atm, d_4^{20} 0.866, n_D^{20} 1.446, $[\alpha]_D^{20}$ (-) and (+) 3.8(4.0) (neat), $pK_{Est} \sim 10.6$.** Purify it by conversion to the **bitartrate salt** (**m 172°**), then decomposing with strong alkali and extracting into Et_2O , drying (KOH), filtering, evaporating and distilling. The **hydrochloride** has **m 242°** (from $EtOH/Et_2O$), $[\alpha]_D^{15}$ -5.0 (c 10 H_2O , from (+) amine). The **oxalate salt** has **m 132°** (from H_2O). The **(±)-base** has **b 176-178°/760mm**, and its **hydrochloride** has **m 237-238°**. [Reihlen et al. *Justus Liebigs Ann Chem* **534** 247 1938, DOI: 10.1002/jlac.19385340115; Leithe *Chem Ber* **65** 660 1932, DOI: 10.1002/cber.19320650432; *Beilstein* **12** III 95.]

Cyclohexylidene fulvene (6,6-pentamethylene fulvene) [3141-04-6] $C_{11}H_{14}$, **M 146.2**. Purify the fulvene by column chromatography and eluting with *n*-hexane [Abboud et al. *J Am Chem Soc* **109** 1332 1987, DOI: 10.1021/ja00239a008].

Cyclohexyl methacrylate [101-43-9] $C_{10}H_{16}O_2$, **M 168.2**, **b 81-86°/0.1mm, d_4^{20} 0.964, n_D^{20} 1.458.** Purify it as for methyl methacrylate (see [80-62-6]). [Tong & Kenyon *J Am Chem Soc* **68** 1355 1946, DOI: 10.1021/ja01211a078; *Beilstein* **6** III 25, **6** IV 39.]

Cyclononanone [3350-30-9] $C_9H_{16}O$, **M 140.2**, **m 142.0-142.8°, b 220-222°, 100-101.5°/15mm.** Purify it via the **semicarbazone** (**m 179.5-180.5°** from 90% MeOH) and regenerate it by steam distilling a mixture of 13.1g of semicarbazone, 22g of phthalic anhydride and 45ml of H_2O . After collecting 300ml of distillate, the latter is extracted with Et_2O . The dried extract ($MgSO_4$) gives on evaporation 8.4g of ketone **b 100-101.5°/15mm**. The **oxime** has **m 76.5-77.5° (79° from MeOH)** and the **iso-oxime** has **m 138-139°**. [Ruzicka et al. *Helv Chim Acta* **32** 544 1949, DOI: 10.1002/hlca.19490320224.] It has also been repeatedly sublimed at 0.05-0.1mm pressure. [Blomquist et al. *J Am Chem Soc* **74** 3639, 3645 1952, *Beilstein* **7** III 110, **7** IV 62.]

cis,cis-1,3-Cyclooctadiene [3806-59-5] C_8H_{12} , **M 108.2**, **m -5°, -49°, -53 to -51°, b 55°/34mm, 142-144°/760mm, d_4^{20} 0.8690, n_D^{20} 1.48921.** Purify the diene by GLC. Fractional distillation through a Widmer column gives a mobile liquid, and redistil it with a Claisen flask or through a semi-micro column [Gould, et al. *Anal Chem* **20** 361 1948, DOI: 10.1021/ac60016a027]. **NB:** It has a strong characteristic disagreeable odour detectable at low concentrations and causes headaches on prolonged exposure. *Do not breath it internally*. [IR: Cope & Estes *J Am Chem Soc* **72** 1128 1950, DOI: 10.1021/ja01159a016; UV: Cope & Bumgardner *J Am*

Chem Soc **78** 2812 1956 DOI: 10.1021/ja01593a045.] [*Beilstein* **5** IV 401.]

cis-cis-1,5-cyclooctadiene (COD) [111-78-4, 1552-12-1] C_8H_{12} , **M 108.2**, **m** -69.5°, -70°, **b** 51-52°/25mm, 97°/144mm, 150.8°/757mm, 149-150°/atm, d_4^{20} 0.880, n_D^{20} 1.4935. Purify it by GLC. It has been purified via the $AgNO_3$ salt. This is prepared by shaking with a solution of 50% aqueous $AgNO_3$ w/w several times (e.g. 3 x 50 ml and 4 x 50 ml) at 70° for ca 20 minutes to get a good separation of layers. The upper layers are combined and further extracted with $AgNO_3$ at 40° (2 x 20 ml). The upper layer (19 ml) of original hydrocarbon mixture gives colourless needles of the $AgNO_3$ complex on cooling. The adduct is recrystallised from MeOH (and cooling to 0°). The hydrocarbon is recovered by steam distilling the complex. The distillate is extracted with Et_2O , dried ($MgSO_4$), filtered, evaporated and re-distilled. [Jones *J Chem Soc* 312 1954, DOI: 10.1039/JR9540000312.] [*Beilstein* **5** H 116, **5** IV 403.]

Cyclooctanone [502-49-8] $C_8H_{14}O$, **M 126.2**, **m** 32-41°, 42°, 43.8°, **b** 115-115.5°/60mm, 195-197°/atm, d_4^{25} 0.958. Purify the ketone by sublimation after drying an ethereal solution over Linde type 13X molecular sieves, filtering and evaporating. The **semicarbazone** has **m** 168-169° (from dioxane) [Kohler et al. *J Am Chem Soc* **61** 1057 1939, DOI: 10.1021/ja01874a021]. The **oxime** has **m** 36-37° after subliming at high vacuum, or distillation, and has **b** 128-129° /14mm. The **iso-oxime** has **m** 72-73° [Ruzicka et al. *Helv Chim Acta* **32** 544 1949, DOI: 10.1002/hlca.19490320224]. [*Beilstein* **7** III 77, **7** IV 49.]

1,3,5,7-Cyclooctatetraene [629-20-9] C_8H_8 , **M 104.2**, **m** -5° to -3°, **b** 141-141.5°/atm, 142-143°/atm, d_4^{20} 1.537, n_D^{25} 1.5350. Purify the triene by shaking 3ml with 20ml of 10% aqueous $AgNO_3$ for 15 minutes, then filtering off the $AgNO_3$ complex which precipitates. The precipitate is dissolved in water and added to cold concentrated ammonia to regenerate the cyclooctatetraene which is fractionally distilled under vacuum onto molecular sieves and stored at 0°. Add ~0.1% of hydroquinone as stabiliser. It is passed through a dry alumina column before use [Broadley et al. *JCS Dalton Trans* 373 1986, DOI: 10.1039/DT9860000373]. [*Beilstein* **5** I 228, **5** IV 1331.]

cis-Cyclooctene [931-87-3; 931-88-4] C_8H_{14} , **M 110.2**, **m** -16°, **b** 32-34°/12mm, 66.5-67°/60mm, 88°/141mm, 140°/170mm, 143°/760mm, 145-146°/760mm, d_4^{20} 0.84843, n_D^{20} 1.4702. The *cis*-isomer is freed from the *trans*-isomer by fractional distillation through a spinning-band column, followed by preparative gas chromatography on a Dowex 710-Chromosorb W GLC column. It is passed through a short alumina column immediately before use [Collman et al. *J Am Chem Soc* **108** 2588 1986, DOI: 10.1021/ja00270a016]. It has also been distilled in a dry N_2 glove box from powdered fused NaOH through a Vigreux column, then passed through activated neutral alumina before use [Wolf et al. *J Am Chem Soc* **109** 4328 1987, DOI: 10.1021/ja00248a031]. Alternatively, it can be purified via the $AgNO_3$ salt. This salt is obtained from crude cyclooctene (40 ml) by shaking at 70-80° with 50% w/w $AgNO_3$ (2 x 15 ml) to remove cyclooctadienes (aqueous layer). Extraction is repeated at 40° (4 x 20 ml, of 50% $AgNO_3$). Three layers are formed each time. The middle layer contains the $AgNO_3$ adduct of cyclooctene which crystallises on cooling the layer to room temperature. The adduct (complex 2:1) is highly soluble in MeOH (at least 1g/ml) from which it crystallises in large flat needles when cooled at 0°. It is dried under slight vacuum for 1 week in the presence of $CaCl_2$ and paraffin wax soaked in cyclooctene. It has **m** 51° and loses hydrocarbon on exposure to air. *cis*-Cyclooctene can be recovered by steam distillation of the salt, collected, dried ($CaCl_2$) and distilled in a vacuum. [Braude et al. *J Chem Soc* 4711 1957, DOI: 10.1039/JR9570004711; $AgNO_3$: Jones *J Chem Soc* 1808 1954, DOI: 10.1039/JR9540001808; Cope & Estes *J Am Chem Soc* **72** 1128 1950, DOI: 10.1021/ja01159a016; *Beilstein* **5** I 35, **5** IV 263.] **FLAMMABLE LIQUID.**

Cyclopentadecanone (Exaltone) [502-72-7] $C_{15}H_{28}O$, **M 224.4**, **m** 63°, 65°, 65-66°, **b** 120°/0.3mm, 155-157°/5mm. Subliming Exaltone is better than crystallising it from aqueous EtOH for purification. The **semicarbazone** has **m** 186-187°. [Stevens & Erickson *J Am Chem Soc* **64** 144 1942, DOI: 10.1021/ja01253a038; Mathur et al. *J Chem Soc* 3505 1963, DOI: 10.1039/JR9630003505; Bienz & Hesse *Helv Chim Acta* **71** 1704 1988, DOI: 10.1002/hlca.19880710717; *Beilstein* **7** III 203, **7** IV 113.]

Cyclopentadiene [542-92-7] C_5H_6 , **M 66.1**, **b** 41-42°/atm, pK^{25} 15. Dry the diene with $Mg(ClO_4)_2$ and distil it rapidly as it dimerises readily at room temperature. It should be used immediately or stored in a Dry Ice or an

ice-salt bath. **HIGHLY FLAMMABLE.** [Moffett *Org Synth Coll Vol 4* 238 1963, DOI: 10.15227/orgsyn.032.0041.] **Cyclopentadiene Dimer (dicyclopentadiene) (4,7-methano-3a,4,7,7a-tetrahydroindene)** has [77-73-6], $C_{10}H_{12}$, **M 132.3**, **m 33°**, **b 170°/atm**, and **d₄²⁵ 0.986**; add ~0.05% of 2,6-di-*tert*-butyl-4-methylphenol as stabiliser. Cyclopentadiene is prepared when required by *de-polymerising* the technical grade dimer by heating (and distilling) it carefully under a fractionating column [Wilkinson *Org Synth Coll Vol 4* 473 1963, DOI: 10.15227/orgsyn.036.0031], as described by Moffett (above reference), or by adding the dimer at a steady rate onto mineral oil heated at 240-270° when it distils off. [Korach et al. *Org Synth* **42** 50 1962, DOI: 10.15227/orgsyn.042.0050]. [*Beilstein* **5** II 391.]

Cyclopentane [287-92-3] C_5H_{10} , **M 70.1**, **m -94°**, **b 49.3°/atm**, **50°/atm**, **d₄²⁰ 0.745**, **n_D²⁰ 1.40645**, **n_D²⁵ 1.4340**. Free it from cyclopentene by two passages through a column of dried and degassed activated silica gel. It occurs in petroleum and is **HIGHLY FLAMMABLE.** [NMR: Christl *Chem Ber* **108** 2781 1975, DOI: 10.1002/cber.19751080836; Whitesides & Gutowski *J Org Chem* **41** 2882 1976, DOI: 10.1021/jo00879a019; *Beilstein* **5** III 10, **5** IV 4.]

Cyclopentane carbonitrile [4254-02-8] C_6H_9N , **M 95.2**, **m -75.2°**, **-76°**, **b 43-44°/7mm**, **50-62°/10mm**, **67-68°/14mm**, **74.5-75°/30mm**, **d₄²⁰ 0.912**, **n_D²⁰ 1.441**. Dissolve the nitrile in Et₂O, wash it thoroughly with saturated aqueous K₂CO₃, dry (MgSO₄) and distil it through a 10 cm Vigreux column. [McElvain & Starn *J Am Chem Soc* **77** 4571 1955, DOI: 10.1021/ja01622a040; Bailey & Daly *J Am Chem Soc* **81** 5397 1959, DOI: 10.1021/ja01529a036; *Beilstein* **9** IV 14.]

Cyclopentanecarboxaldehyde [872-53-7] $C_6H_{10}O$, **M 98.1**, **b 36°/12mm**, **74-78°/100mm**, **140-141°/atm**, **d₄²⁵ 0.919**, **n_D²⁰ 1.4420-1.4428**. Several preparations of this aldehyde have been described but only two will be briefly mentioned here, and both start from cyclohexene. The *first* requires decomposition of the mercuric sulfate complex where, under N₂, concentrated H₂SO₄ (43.5ml, 80.0g, 0.82mole) in H₂O (3L) and HgSO₄ (740.0g, 2.49moles, CARE **POISONOUS**) are stirred to form the deep-yellow basic salt. To this stirred mixture, under N₂ at 55°, is added all at once cyclohexene (101ml, 82.0g, 1mole, freshly distilled b 82-84°/atm [110-83-8]), and the temperature kept at 55-65° for 1 hour (optimal conditions for decomposition of complex). The colour of the mixture turns from deep-yellow to cream. The equipment is altered for distillation at the end of the hour, and the temperature is raised, stirring and the N₂ flow are continued as the crude *aldehyde* and H₂O distil over during *ca* 2 hours. The layers are separated, the H₂O layer is extracted with Et₂O (3 x 50ml) which is combined with the product, dried (Na₂SO₄), filtered, and distilled rapidly (b 74-78°/100mm) to give the *aldehyde* (46-52g, 46-53%), which if not used immediately should be stored in a brown bottle at 0° for under N₂, after 0.1g of hydroquinone is added as stabiliser. The *aldehyde* readily forms a *trimer*, and does so on prolonged storage. [Grummit et al. *Org Synth Coll Vol 5* 320 1973, DOI: 10.15227/orgsyn.044.0026; English et al. *J Am Chem Soc* **73** 615 1951, DOI: 10.1021/ja01146a036.] When the distillation residue is cooled a solid may be formed which can be distilled above 78°/100mm to give a clear liquid that solidifies, and can be recrystallised from 95% EtOH to give a white solid **m 122-124°**, identical with *cyclopentanecarboxaldehyde trimer* obtained from the *aldehyde* and 85% H₃PO₄ [Brook & Wright *Can J Chem* **29** 308 1951, DOI: 10.1139/v51-036]. In the *second* procedure, Tl(III)(NO₃)₃ · 3H₂O (4.4g, 10mmol, TTN [13453-38-8]) is dissolved in MeOH (50ml), cyclohexene (10mmol) is added, and the mixture is stirred at room temperature or heated until a starch-iodide paper indicates complete reduction of Tl(III) to Tl(I) (usually within a few minutes). The mixture is filtered, and an alcoholic solution containing 10mmol of 2,4-dinitrophenylhydrazine is added. This is evaporated to 1/3 its volume and after addition of H₂O (10ml) the mixture is heated on a steam bath for 10 minutes. On cooling to 0°, the **2,4-dinitrophenylhydrazone** [20956-07-4] crystallises out and is recrystallised from EtOH to **m 195.5-196°** (85% yield). Hydrolysis of the hydrazone in the usual way provides the free *aldehyde*. On a preparative scale the precipitated Tl(I)(NO₃) salt is filtered off, the filtrate is evaporated to a small volume, the mixture of aldehyde and its methyl acetal formed is heated on a steam bath with excess of 5% H₂SO₄ for 30 minutes, and *cyclopentanecarboxaldehyde* is isolated by ether extraction followed by distillation as in the first preparation above. [McKillop et al. *J Am Chem Soc* **95** 3635 1973, DOI: 10.1021/ja00792a028.] It should be stabilised with 0.1% of hydroquinone as before.

Cyclopentane carboxylic acid [3400-45-1] $C_6H_9O_2$, **M 114.1**, **m 3-5°**, **b 87-89°/2-3mm**, **106.5-107°/10mm**, **120-123°/27mm**, **216°/atm**, **d₄²⁵ 1.053**, **n_D²⁰ 1.4530**, **n_D²⁵ 1.4522**, **pK_a²⁵ 4.98**. If it is discoloured, shake it with

saturated aqueous NaCl, extract it with Et₂O, dry the extract (MgSO₄), filter, evaporate and distil the residue preferably under a vacuum. The **lachrymatory acid chloride** [4524-93-0] has **C₆H₉ClO**, **M 132.6**, **m 4°** and **b 161-162°/atm**, **d₄²⁰ 1.091**, **n_D²⁰ 1.4622**. [Beilstein 9 H 6, 9 IV 11.] The **amide** [3217-94-5] **C₆H₁₁NO**, has **m 179°**. Like the methyl ester below, the acid can also be prepared *via* a Favorskii reaction, but by using a little over 2 moles of alkoxide (1mol provides the ester). Thus, to sodium (83g, 3.61mol) in absolute EtOH (2.3L) is added dropwise 2-chlorocyclohexanone (240.5g, 1.82mol, [822-87-7]) during several hours, left overnight, the EtOH is distilled off, replaced by an equal volume of H₂O, acidified to pH 3, and the oily acid that separates is extracted with Et₂O. The extract is washed with H₂O, dried (Na₂SO₄), filtered, and distilled to give the pure *acid* (110g, 53%). [Jackman et al. *J Am Chem Soc* 70 497 1948, DOI: 10.1021/ja01182a019; Mourisson et al. *Bull Soc Chim Fr* 767 1952.] [Beilstein 6 H 6, 6 I 4, 6 II 6, 6 III 11, 6 IV 9.]

Methyl cyclopentanecarboxylate [4630-80-2] **C₇H₁₂O₂**, **M 128.1**, **b 70-73°/48mm**, **38-39°/7mm**, **158.2°/760mm**, **n_D²⁵ 1.4341**, can be obtained by reacting the acid chloride with MeOH and distilling; but it can also be obtained in ~55% yield directly from 2-chlorocyclohexanone and NaOMe (1mol) in dry Et₂O, *via* a Favorskii reaction, and purified by fractional distillation, using a Podbielniak column filled with tantalum wire spirals and a partial reflux head, under reduced pressure [Goheen & Vaughan *Org Synth Coll Vol* 4 594 1963 DOI: 10.15227/orgsyn.039.0037]. The *methyl ester* has also been prepared in 98% yield by direct esterification of cyclopentane carboxylic acid with MeOH in CH₂Cl₂ in the presence of catalytic amounts of H₂SO₄. The acid has FTIR (CCl₄) with ν_{\max} at 2952(s), 2873 (m), 1734 (s), 1436 (w), 1165 (s) and 1140 (s) cm⁻¹; ¹H NMR (75MHz, CDCl₃, TMS) with δ at 1.5-2.0 (m, 8H, 4-methylenes), 2.7 (pentet, 1H, *J* = 9.0Hz), 3.7 (s, 3H, Me); ¹³C NMR (300MHz, CDCl₃, TMS) with δ at 25.7, 29.9, 43.6, 51.5, 177.2, and the GCMS has *m/z* at 128 (M⁺) [Davis et al. *J Org Chem* 58 6843 1993, DOI: 10.1021/jo00076a055].

1,1-Cyclopentanediactic acid (3,3-tetramethyleneglutaric acid) [16713-66-9] **C₉H₁₄O₄**, **M 186.2**, **m 176-177°, 180-181°, pK₁²⁵ 3.80, pK₂²⁵ 6.77**. Purify it by recrystallisation from H₂O and dry it in air, *in vacuo* or over CaCl₂. However, if it is suspect it is better to convert the acid to the **anhydride (8-oxaspiro[4.5]decane-7,8-dione** [5662-95-3] **C₉H₁₂O₃**, **M 168.2**, **m 64-66° (68° also reported)**, **b 186°/15mm**) by refluxing it for 7 hours in excess of Ac₂O, evaporating to dryness, and distilling the residue in a vacuum or recrystallising it from light petroleum. [Kon & Thorpe *J Chem Soc* 115 686 1919, DOI: 10.1039/CT9191500686.] The anhydride (e.g. 30g) is then hydrolysed by refluxing it for 2.5 hours (i.e. until it dissolves) in aqueous KOH (35g, 3mols in 55ml of H₂O), then acidified with concentrated HCl, extracted with Et₂O (3 x 50ml) after saturating the aqueous solution with (NH₄)₂SO₄, drying the extract (Na₂SO₄), filtering, evaporating the Et₂O and recrystallising the *di-acid* from hot H₂O. The **dimethyl ester** [70179-60-3] has **b 141°/17mm** [Vogel *J Chem Soc* 1758 1934, DOI: 10.1039/JR9340001758; Dickens et al. *J Chem Soc* 1496 1922, DOI: 10.1039/CT9222101496], and the **diethyl ester** has **b 153°/14mm** [Kon *J Chem Soc* 513 1922, DOI: 10.1039/JR9340001758]. [Beilstein 9 I 319, and anhydride Beilstein 17/11 V 80.]

Cyclopentane-1,1-dicarboxylic acid [5802-65-3] **C₇H₁₀O₄**, **M 158.1**, **m 183°, 184°, (176-178°)**, **pK₁ 3.23, pK₂ 4.08**. The dicarboxylic acid can be prepared in 71% yield by using the nickel(0) catalyst 2-{2-[2-(dicyclohexyl-phosphino)ethyl]pyridine}-4-oxo-2-nickela-3-oxa-cis-bicyclo[3.3.0]octane which promotes the di-carbonylation of cyclopentene (1.16g, 2.44mmol) in THF (30ml)/pyridine (30ml), in the presence of BeCl (0.2g, 2.50mmol), by CO₂ (10 bar pressure) in an autoclave, and stirring at 60° for 24 hours. The solvent is evaporated off *in vacuo*, the residue is treated with Et₂O/HCl and the solid is collected, dried and recrystallised from pentane/Et₂O (10:1), or H₂O. The pure *dicarboxylic acid* melts at 183° with decarboxylation to *cyclopentanecarboxylic acid* (see preceding acid), and has FTIR (KBr) with ν_{\max} at 3200-2500 (COOH) and 1700 (C=O) cm⁻¹; ¹H NMR (200MHz, THF-*d*₈/TMS) with δ at 1.6 (m, 4H, 2-methylenes), 2.1 (m, 4H, 2-methylenes), 9.6 (br, 2H, 2-COOH) ppm; ¹³C NMR (50MHz, THF-*d*₈/TMS) with δ at 26.3 (t), 35.2 (t), 60.6 (s), 174.2 ppm; and the MS (70 eV) has *m/z* at 158 (M⁺). [Hoberg et al. *Synthesis* 395 1991, DOI: 10.1055/s-1991-26475]. **Dimethyl 1,1-cyclopentanedicarboxylate** [74090-15-6] **M 186.2**, **b 102°/10mm**, can be prepared in 76% yield from the preceding *methyl cyclopentanecarboxylate* (40mmol, [4630-80-2]) in THF (60ml), with (iPr)₂NH (60mmol) and *n*-BuLi (60mmol) followed by MeOCOCl (60mmol), and finally distilling. The *dimethyl ester* has FTIR (CCl₄) with ν_{\max} at 2953(s), 2928 (w), 2876 (w), 1736 (s), 1434 (w), 1266 (s), 1165 (s) and 1159 (m) cm⁻¹; ¹H NMR (300MHz, CDCl₃, TMS) with δ at 1.7 (m, 4H, 2-methylenes), 2.1 (m, 4H, 2-methylenes), 3.7 (s, 6H, 2-Me) ppm; ¹³C NMR (75MHz, CDCl₃, TMS) with δ at 25.3, 34.5, 52.4, 60.2, 173.0 ppm, and the GCMS has *m/z* at 187 (M⁺+1) [Davis et al. *J Org Chem* 58 6843 1993, DOI: 10.1021/

jo00076a055].

IRS,2SR-(meso)-cis-1,2-Cyclopentanedicarboxylic acid [1461-96-7] $C_7H_{10}O_4$, **M 158.2**, **m 139°, 140°, 141°, pK_1^{20} 4.42, pK_2^{20} 6.57**. The *cis*-acid has been prepared by hydrogenation of cyclopent-1-ene-1,2-dicarboxylic acid with Pt as catalyst [Peters *J Chem Soc* 1757 1959, DOI: 10.1039/JR9590001757], heating diethyl 1-cyanocyclopentane-1,2-dicarboxylate (b 135-136°/3.5mm) with aqueous HCl and isolating the anhydride [Fuson & Cole *J Am Chem Soc* **60** 1237 1938, DOI: 10.1021/ja01272a066; Dutta *J Indian Chem Soc* **17** 611, 617 1940], but is best prepared by boiling the *trans*-acid with excess of acetic anhydride for at least 2 hours (which provides the *cis*-anhydride), dissolving this in boiling H_2O or aqueous KOH with cooling (**care**, may be exothermic), and acidifying to pH ~2 in order to precipitate the *cis*-acid. The *cis*-acid crystallises from hot H_2O as colourless needles melting at **140°**, and is rapidly converted to the *cis*-anhydride at **150-160°**. The *cis*-acid is also obtained from the *trans*-acid by heating at 300 ° [Perkin Jnr *J Chem Soc* **51** 240 1887, DOI: 10.1039/CT8875100240]. It is more soluble in H_2O than the (\pm)-*trans*-acid. It had not been possible to prepare *trans*-cyclopentane-1,2-dicarboxylic anhydride because all attempts gave the *cis*-anhydride (*cis*-tetrahydro-1*H*-cyclopent[*c*]furan-1,3(3*aH*)-dione, **m 73.5-74°, b 100-102°/1.5mm**, [5763-49-5]). [Perkin Jnr *J Chem Soc* **65** 572 1894, DOI: 10.1039/CT8946500572; cf. Fuson & Cole *J Am Chem Soc* **60** 1237 1938, DOI: 10.1021/ja01272a066; Goldsworthy & Perkin Jnr *J Chem Soc* **105** 2639 1914, DOI: 10.1039/CT9140502639.]. The nickel(0) catalyst 2-{2-[2-(dicyclohexyl-phosphino)ethyl]pyridine}-4-oxo-2-nickela-3-oxa-*cis*-bicyclo-[3.3.0]octane promotes the carbonylation of cyclopentene in THF by CO at 1bar pressure and -40°, then at ~25°/10 hours followed by treatment with 2N H_2SO_4 and recrystallisation from pentane/ Et_2O (10:1), which provides cyclopentane-1,2-dicarboxylic acid (78% yield) presumed to be mainly the *cis*-isomer. However, its melting point of 124-125° suggests that it must be contaminated with some of the *trans*-isomer [Hoberg et al. *Synthesis* 395 1991, DOI: 10.1055/s-1991-26475]. Its FT-IR (KBr) has ν_{max} at 3300-2500 (COOH) and 1710 (C=O) cm^{-1} ; its 1H NMR (200MHz, THF-*d*₈/TMS) has δ_H at 10.4 (br s, 2H, OH), 2.9 (m, 2H, 2CH), 1.51-2.1 (m, 6H, 3CH₂), 1.69 (m, 4H); its ^{13}C NMR (200MHz, THF-*d*₈/TMS) has δ_C at 176.10 (s, carboxylic C), 47.5 (d, C1 and C2), 29.4 (t, C-3 and C-5) and 24.4 (t, C-4); and its MS (70 eV) has m/z = 158 (M^+), see GC-MS [below, Lu et al. *J Org Chem* **55** 2503, 2507 1990, DOI: 10.1021/jo00295a048]. **Dimethyl cis-cyclopentane-1,2-dicarboxylate**, **b 129°/20mm, n_D^{21} 1.4528** [4841-91-2], is obtained by treating the acid with diazomethane in Et_2O , evaporating and distilling the residue [Owen & Peto *J Chem Soc* 2383, 2386 1955, DOI: 10.1039/JR9550002383]. The *cis*-imide (*cis*-tetrahydrocyclopent[*c*]pyrrole-1,3(2*H*,3*aH*)-dione, [5763-44-0]) has **m 85-87°** (also **90°** reported) after recrystallisation from H_2O or petroleum ether (b 40-60°) [Menon & Simonsen *J Chem Soc* 302 1929, DOI: 10.1039/JR9290000302; Rice et al. *J Org Chem* **24** 7 1959, DOI: 10.1021/jo01083a002], and *cis*-cyclopentane-1,2-dicarboxylic acid dihydrazide, prepared from the dimethyl ester and $NH_2NH_2 \cdot H_2O$ in EtOH and recrystallised from MeOH/ Et_2O , has **m 125-126°** [Müller et al. *J Am Chem Soc* **73** 2487 1951, DOI: 10.1021/ja01150a023]. [Beilstein **9** H 728, **9** III 3807, **9** IV 3793.]

IRS,2RS-(\pm)-trans-1,2-Cyclopentanedicarboxylic acid [1461-97-8] $C_7H_{10}O_4$, **M 158.2**, **m 160-161°, 161.5°, 162°, 162-163°, 163°, 163-165°, 164-165°, pK_1^{20} 4.14, pK_2^{20} 5.99**. Several syntheses of the (\pm)-*trans*-acid have been described. The first preparation involved tetraethyl cyclopentane-1,2,3,4-tetracarboxylate [b 185-190°/1mm, from trimethylene-1,3-dibromide (96g), Na (22g) and diethyl malonate (150g) in absolute EtOH (300ml)] which was refluxed with 2-3 volumes of AcOH, 1 volume of concentrated H_2SO_4 and 1 volume of H_2O for 2 days, and the AcOH and H_2O were distilled off to give *trans*-cyclopentane-1,2-dicarboxylic acid [Perkin Jnr *J Chem Soc* **65** 572 1894, DOI: 10.1039/CT8946500572; Wassermann *Helv Chim Acta* **13** 207 1930, DOI: 10.1002/hlca.19300130212; Perkin Jnr & Prentice *J Chem Soc* **59** 818 1891, DOI: 10.1039/CT8915900818]. Other preparations include heating the *cis*-acid with concentrated HCl at 180° [Perkin Jnr *J Chem Soc* **65** 572, 590 1894, DOI: 10.1039/CT8946500572], refluxing diethyl 1-cyanocyclopentane-1,2-dicarboxylate with concentrated aqueous HCl for 4 days which provided a 92% yield of the *trans*-acid [Fuson et al. *J Org Chem* **10** 121, 126 1945, DOI: 10.1021/jo01178a005; Fuson & Cole *J Am Chem Soc* **60** 1237 1938, DOI: 10.1021/ja01272a066], and electrochemical coupling between 1,3-dibromopropane and dimethyl maleate which gave a 10/1 ratio of *trans/cis* dimethyl esters in 42% yield [Lu et al. *J Org Chem* **55** 2503, 2507 1990, DOI: 10.1021/jo00295a048]. Invariably the crude *trans*-diesters formed are contaminated with some *cis*-isomer and it was found that hydrolysing the ester mixture by refluxing the acidic solution for 4 to 5 days provided only the *trans*-acid in ~95% yields. The *trans*-acid crystallises (charcoal) from hot H_2O , and it is easily soluble in AcOH, and EtOH, but very soluble in Et_2O , $CHCl_3$, *C_6H_6

and petroleum ether. Its FT-IR (Nujol) has ν_{\max} at 3008.3, 2653.8, 1692.6, 1419.2, 1311.1, 1253.5, 1199.3, 935.7 and 689.4 cm^{-1} ; its ^1H NMR (60MHz, CDCl_3) has δ_{H} at 12.2 (br s, 2H, OH), 2.96 (m, 2H, C1 and C2 methines), 1.99 (m, 2H,), 1.69 (m, 4H); its ^{13}C NMR (75MHz, CDCl_3) has δ_{C} at 176.10 (s, carboxylic C), 46.77 (s, C1 and C2), 30.03 (s, C3 and C5) and 25.01 (s, C-4); and its GC-MS has m/z (relative intensity) *trans*-isomer: 157(37, M^+-2), 155(15), 126(67), 113(29), and 67(100), and *cis*-isomer: 157(6, M^+-2), 155(40), 126(32), 113(56), and 67(100) [Lu et al. *J Org Chem* **55** 2503, 2507 1990 DOI: 10.1021/jo00295a048]. **Dimethyl trans-cyclopentane-1,2-dicarboxylate**, **b 118.5-119°/17mm**, n_{D}^{20} **1.4491** [941-75-3], is obtained by treating the acid with diazomethane in Et_2O , evaporating and distilling the residue [Owen & Peto *J Chem Soc* 2383 1955, DOI: 10.1039/JR9550002383]. **trans-Cyclopentane-1,2-dicarbonyl dichloride**, has **b 98-100°/2mm** [Aspinall & Baker *J Chem Soc* 743 1950 DOI: 10.1039/JR9500000743], and **trans-cyclopentane-1,2-dicarboxylic acid dihydrazide**, prepared from the dimethyl ester and $\text{NH}_2\text{NH}_2 \cdot \text{H}_2\text{O}$ in EtOH, and recrystallised from EtOH, has **m 224-225°** [Müller et al. *J Am Chem Soc* **73** 2487 1951, DOI: 10.1021/ja01150a023]. [Beilstein **9** H 728, **9** I 316, **9** III 3807, **9** IV 2793.]

1S,2S-(+)-trans-1,2-Cyclopentanedicarboxylic acid [21917-20-4] $\text{C}_7\text{H}_{10}\text{O}_4$, **M 158.2**, **m 181°**, $[\alpha]_{\text{D}} +87.6$ (c **0.88**, H_2O) and **1R,2R-(-)-trans-1,2-cyclopentanedicarboxylic acid** [17224-73-6] **m 180-181°**, $[\alpha]_{\text{D}} -85.9$ (c **1.2**, H_2O). Optical resolution of the (\pm)-acid (above) is readily achieved via separation of the diastereoisomeric brucine salts (the base being *l*-brucine). Thus pure (\pm)-acid (15g, ~0.1mole) is dissolved in hot H_2O , and while stirring, the finely powdered brucine tetrahydrate (90g, 0.2 moles, [5892-11-5], care **POISONOUS**) that is added gradually dissolves. At the end of the addition, the mixture is stirred for ~10 minutes and filtered hot from a small amount of undissolved brucine. The brucine is washed well with hot H_2O , and the combined filtrate and washings are concentrated on a boiling water bath until crystals commence to form on the surface. When the liquid is cooled and vigorously stirred, copious crystals of the *d*-acid (*l*-base)₂ separate, and the whole mixture sets into a mass of crystals. These are filtered off and recrystallised from hot H_2O to constant specific rotation. An analytical sample of the *d*-acid (*l*-base)₂ is obtained by dissolving it in hot H_2O and allowing it to crystallise slowly in a desiccator over concentrated H_2SO_4 . The decahydrate is obtained on drying the crystalline plates in air but they effloresce in a vacuum desiccator or on heating at 125° for 1 hour to give **anhydrous d-acid (*l*-base)₂** that has $[\alpha]_{\text{D}} -19.9$ (c **1.3**, H_2O). The *d*-acid (*l*-base)₂ is then decomposed by dissolving it in hot H_2O , adding aqueous ammonia, filtering off the solid brucine, washing it well with aqueous ammonia, and the combined hot filtrate and washings are cooled and acidified with hydrochloric acid. On cooling the **(+)-acid** (m 178-180°) separates and is recrystallised from hot H_2O as plates with m 180°. **Diethyl 1S,2S-(+)-trans-1,2-cyclopentanedicarboxylate** **b 170°/100mm**, $[\alpha]_{\text{D}} +70.3$ (c **1.3**, Me_2CO), is prepared by boiling the (+)-acid with 5 times its weight of 10% ethanolic H_2SO_4 for 6 hours, cooling, adding H_2O , the ester is extracted into Et_2O which is washed with dilute Na_2CO_3 , dried (Na_2SO_4), filtered, evaporated and the residue is distilled in a vacuum. The **1S,2S-(+)-trans-dianilide**, **m 245-247°**, $[\alpha]_{\text{D}} +110.1$ (c **0.85**, Me_2CO) is prepared by heating the (+)-acid with SOCl_2 in a sealed tube at 100° for 1 hour, evaporating, the di-acid chloride formed is then dissolved in $^*\text{C}_6\text{H}_6$, treated with excess of aniline, evaporated again, the residue is stirred with dilute aqueous HCl, the precipitate is collected and recrystallised twice from large volumes of MeOH.

1R,2R-(-)-trans-1,2-cyclopentanedicarboxylic acid is obtained by concentrating the mother liquors from the first crystallisation of the *d*-acid (*l*-base)₂ above, which provides copious crystals of the more soluble **diastereoisomeric l-acid (*l*-base)₂**. Treatment of this salt with aqueous ammonia as above followed by recrystallisation from H_2O gives the pure **(-)-acid** with **m 180-181°**. **Diethyl 1R,2R-(-)-trans-1,2-cyclopentanedicarboxylate**, prepared in the same way as its enantiomer above has similar properties except for the optical rotation which is $[\alpha]_{\text{D}} -69.8$ (c **1.7**, Me_2CO). [Goldsworthy & Perkin Jnr *J Chem Soc* **105** 2639 1914, DOI: 10.1039/CT9140502639]. [Beilstein **9** I 316] The absolute configuration of the **(-)-acid** was determined as **1R,2R** by comparing the ORD spectra of the respective 1,2-bis(N-methylthionamide) [1,2-(CSNHMe)₂] with those of the corresponding 1,2-disubstituted cyclopropane, cyclobutane and cyclohexane homologues [Inouye et al. *Tetrahedron* **23** 3237 1967, DOI :10.1016/S0040-4020(01)92293-8]. [Beilstein **9** III 3807.]

1R,2SR-cis-1,2-Cyclopentanediol (meso-cis-cyclopentane-1,2-diol) [5057-98-7] $\text{C}_5\text{H}_{10}\text{O}_2$, **M 102.1**, **m 24-28°**, **30°**, **b 88-92°/2mm**, **108-109°/20mm**, **123.5°/29mm**, n_{D}^{21} **1.4770**. The *cis*-diol can be obtained by treating cyclopentene (21.8g, [142-29-0]) in EtOH (600ml) at -40° with a solution of KMnO_4 (40g) and anhydrous MgSO_4 (30g) in H_2O (800ml) dropwise, with vigorous stirring, during 2 hours. The MnO_2 is filtered off, washed with hot H_2O , the combined aqueous filtrates are evaporated *in vacuo* to 100ml and continuously extr-

acted with Et₂O (liquid-liquid extractor) for 48 hours. The dried (MgSO₄) Et₂O extract is evaporated and the residual oil is distilled to give the *cis*-diol (11.1g, 34%). The *cis-di-p-nitrobenzoate* has **m** 117° (117-118°) (from EtOH) the *cis-di-methanesulfonate* has **m** 82° (needles from MeOH), and the *cis-di-toluene-p-sulfonate* has **m** 92° (prisms from EtOH) are obtained in the usual way in pyridine. The *cis-monoacetate* has **b** 104-106°/20mm, n_D^{17} 1.4576, the *cis-monoacetate-mono-p-nitrobenzoate* has **m** 93-94° (nodules from EtOH), and the *cis-diacetate* has **m** -5°, **b** 80°/2mm. The *cis-dibenzoate* has **m** 47°. [Owen & Smith *J Chem Soc* 4026 1952, DOI: 10.1039/JR9520004026; *Beilstein* 6 I 369, 6 II 742, 6 III 4053, 6 IV 5187.]

1R,2R-(±)-trans-1,2-Cyclopentanediol [*trans-(±)-cyclopentane-1,2-diol*] [5057-99-8, (±) 86703-52-8] C₅H₁₀O₂, **M** 102.1, **m** 50°, 55°, **b** 93°/2mm, 122°/10mm, 136°/22mm. The *cis*-diol can be prepared in a seven step synthesis from *racemic* diethyl tartrate [Cunningham & Küendig *J Org Chem* 53 1823 1988, DOI: 10.1021/jo00243a048], from *trans*-2-tri-methylsilylcyclopentan-1-ol and MeOH [Kono & Ngai *Org Prep Proced Int* 6 19 1974], and by performate oxidation of cyclopentene which is described herein. A mixture of 30% aqueous H₂O₂ (13g) and formic acid (105g) is added to cyclopentene (6.6g, **b** 44-45°/atm [142-29-0]) under a reflux condenser, when much heat is generated, the temperature is allowed to fall to 40° and is so maintained for 4 hours. The solvent is removed *in vacuo*, the residue is dissolved in 10% aqueous NaOH (50ml), boiled under reflux for 45 minutes, the solvent is distilled off, and the residue is twice distilled under reduced pressure to give pure *trans*-pentane-1,2-diol (6g, 60%) which solidified in the distillate. The *trans-dibenzoate* has **m** 63°, and the *trans-(±)-di-p-nitrobenzoate* (**m** 145°, from EtOH), the *trans-(±)-di-methanesulfonate* (**m** 82°, small needles from EtOH), and the *trans-(±)-di-p-toluenesulfonate* (**m** 109°, colourless plates from MeOH) are prepared in the usual way in pyridine. [Owen & Smith *J Chem Soc* 4026 1952, DOI: 10.1039/JR9520004026; Englund *J Prakt Chem* 129 1, 10 1931, DOI: 10.1002/prac.19311290101; *Beilstein* 6 H 739, 6 I 369, 6 II 743, 6 III 4053, 6 IV 5187.]

1R,2R-(-)-trans-1,2-Cyclopentanediol [*trans-(-)-cyclopentane-1,2-diol*] [(-) 930-46-1, (-) 86703-52-8] C₅H₁₀O₂, **M** 102.1, **m** 48-48.5°, $[\alpha]_D^{20}$ -24.88 (**c** 6, EtOH) and **1S,2S-trans-1,2-cyclopentanediol** [*trans-(+)-cyclopentane-1,2-diol*] [(+)/107492-82-0], **m** 47-48°, $[\alpha]_D^{20}$ +24.54 (**c** 5.4, EtOH). The *1R,2R*-enantiomer is obtained optically pure in a seven step synthesis from diethyl D(-)-tartrate by benzylation (TIOEt/PhCH₂Br, 81%), reduction (LAH/Et₂O, 94%) to *2R,3R-(-)-2,3-bis(benzyloxy)-1,4-butanediol*, ditosylation (TsCl/py, 87%), dibromination (with LiBr/DMSO, 94%) to *2R,3R-(-)-2,3-bis(benzyloxy)-1,4-butanedibromide*, and formation of the cyclopentane ring (PhSO₂CH₂SO₂Ph/K₂CO₃/DMF, 92%) to give *1R,2R-(-)-1,2-bis(benzyloxy)-4,4-bis(phenylsulfonyl)cyclopentane*. Removal of the bis-sulfonyl groups (Mg/MeOH, 82%) provides *1R,2R-(-)-1,2-bis(benzyloxy)cyclopentane*, and finally debenylation (H₂/10% Pd/C, 99%) gives pure *1R,2R-(-)-cyclopentane-1,2-diol* in 47% overall yield. The final step only is detailed here. The mixture of *trans-(-)-1,2-bisbenzyloxy*cyclopentane (5.11g, 18.1mmol) in EtOH (35ml) containing 10% Pd/C (1.35g) is degassed three times, H₂ gas is introduced into the flask at a pressure of 6 atmospheres for 5 hours, the catalyst is filtered off, and the filtrate is evaporated to provide the **1R,2R-trans-(-)-diol** (8.2g, 99%) as a white hygroscopic solid **m** 47-48° which should be stored under dry N₂ in a sealed vessel. The enantiomeric **1S,2S-trans-(-)-diol** can be prepared in 40% overall yield from diethyl L(+)-tartrate *via* the same high yielding steps. Its IR (CH₂Cl₂) has ν_{\max} at 3600, 3390br, 2920, 1450, 1280, 1075, 1040 and 970 cm⁻¹; and its ¹H NMR (200MHz, CDCl₃, TMS) with δ at 3.93-4.06 (m, 2H), 2.24 (s, 2H, exchanges with D₂O), 1.86-2.12 (m, 2H), 1.63-1.80 (m, 2H) and 1.42-1.62 (m, 2H). The enantiomers are >96% enantiomerically pure. The **1R,2R-(-)** [113350-86-0] and **1S,2S-(+)** [113303-27-8] **1,2-bis(dimethoxyphosphinoxy)cyclopentane** derivatives, prepared from the respective diols (2NEt₃/2(MeO)₂PCl₂ in 80% and 73% yields respectively), are yellow oils with **b** 120°/0.1mm and $[\alpha]_D^{20}$ -41.39 and +41.78 (**c** 4.8, CH₂Cl₂). The **1R,2R-(-)** [113303-35-8] and **1S,2S-(+)** [113303-28-9] **1,2-bis(difluorophosphinoxy)cyclopentane** derivatives, prepared from the respective diols (PCl₃ in CH₂Cl₂/SbF₃ in pentane, 49% and 48% respectively) are mobile oils with **b** 58°/80mm and $[\alpha]_D^{20}$ -12.22 and +12.96 (**c** 4.8, CH₂Cl₂). [Cunningham & Küendig *J Org Chem* 53 1823 1988, DOI: 10.1021/jo00243a048].

1,2-Cyclopentanedione [3008-40-0] C₅H₆O₂, **M** 98.1, **m** 55-56°, 56°, **b** 67-69°/1.4mm, 87-88°/16mm, 97°/20mm, 105°/20mm, **pK**²⁰ 9.14 (H₂O). The dione has a weak amine-like odour and is soluble in H₂O, EtOH, and Et₂O, but poorly soluble in petroleum ether and CS₂, and is steam volatile. It has been prepared by various routes from cyclopentanone *via* oxidation by SeO₂ (23%) [Goto et al. *Bull Chem Soc, Jpn* 52 2589 1952, DOI: 10.1246/bcsj.52.2589], conversion to the α -methylthio ether then oxidation with CuCl₂/CuO (42%)

[Gregoire et al. *J Org Chem* **51** 1419 1986, DOI: 10.1021/jo00359a008], and autooxidation with O_2/t -BuOK/DME-*t*-BuOH (35.7%). Only the first is described because of its simplicity. *The following procedure should be carried out in a well-ventilated fume cupboard as selenium compounds have foul odours.* To cyclopentanone (1.5L) warmed at 30° is added dropwise, with stirring, a solution of selenium dioxide (361g, 3.25ml) in dioxane (420ml) and H_2O (155ml) during 24 hours, and stirring is continued for a further 18 hours at 35°. The red selenium that is formed is filtered off (Buchner) and extracted with EtOH (500ml) by boiling for 3 hours. The liquids are combined and distilled through a 60cm Vigreux column at 20mm, the lower boiling portion (b 35-90°) is removed, and the remainder is distilled at 10mm until a thick brown residue is left in the distilling flask. This distillate is redistilled, and the fraction with **b 86-88°/16mm** is collected to give pure **1,2-dione** (74g, 23% based on SeO_2 used) which solidifies (**m 56-58°**). It should be stored at -30° under N_2 , in the dark. Cyclopentane-1,2-dione has IR (NaCl) with ν_{max} at 1650 (C=C), 1700 (C=O), 3300 (OH enol) cm^{-1} ; 1H NMR (400MHz, CCl_4 , TMS) with δ at 2.20-2.67 (m, 4H, 2-methylenes), 6.47-6.62 (m, 1H, C=CH), 6.93-7.36 (m, 1H, C=COH); ^{13}C NMR (100MHz, $CDCl_3$, TMS) with δ 22.01, 32.58, 131.36, 153.62, 187.53.

In ethanolic solution the 'dione' is almost entirely in the *mono-enol*-form which slowly tautomerises to a mixture containing ~9% of the *diketo*-form after 36 hours at 50°. The UV in hexane has λ_{max} (log ϵ) at 246nm (4.003) and 300nm (1.900) due mostly to the *diketo*-form, whereas in 0.05N aqueous NaOH it has λ_{max} (log ϵ) at 288nm (3.383) due to the *enol*-form [Hesse & Bücking *Justus Liebigs Ann Chem* **563** 31 1949, DOI: 10.1002/jlac.19495630105]. *Cyclopentane-1,2-dione dioxime* (see above [66,35-29-6]) decomposes at ~210° after recrystallisation from H_2O , and a 0.01M aqueous solution of it is used for the gravimetric determination of Ni. [Voter et al. *Anal Chem* **21** 1320 1949, DOI: 10.1021/ac60035a004.] The *bis-phenylhydrazone* has **m 146°**.

1,3-Cyclopentanedione [3859-41-4] $C_5H_6O_2$, **M 98.1, m 149-150°, 151-152.5°, 151-154°, 151-153°, pKa 4.5.**

Purify the dione by Soxhlet extraction with $CHCl_3$. The $CHCl_3$ is evaporated and the residue is recrystallised from EtOAc and/or sublimed at 120°/4mm. [IR: Boothe et al. *J Am Chem Soc* **75** 1732 1953, DOI: 10.1021/ja01103a505; DePuy & Zaweski *J Am Chem Soc* **81** 4920 1959, DOI: 10.1021/ja01527a040; *Beilstein* 7 IV 1981.]

Cyclopentanol [96-41-3] $C_5H_{10}O$, **M 86.1, m -20°, -19.5°, b 53°/10mm, 56.9-57.4°/34mm, 139.5°/752mm, 140.85°/760mm, d₄²⁰ 0.9478, n_D²⁰ 1.4531.** Cyclopentanol has been prepared by hydration of cyclopentene, or from cyclopentanone (see next entry) by catalytic reduction with H_2 (Ni/MeOH, Ra-Ni/EtOH, PtO_2 , Pt-black in EtOH or AcOH, Cu-chromite), $Al(isoPrO)_3/NaOH$, LAH/ Et_2O or $NaBH_4$ in MeOH or H_2O . The last named has many advantages in ease of use and efficiency. Thus cyclohexanone in H_2O (solubility is 15% at 10°, 8.7% at 20°, 5% 30°) is stirred with excess of $NaBH_4$ (theoretically 0.25 mol/mol ketone) at room temperature until the reaction is complete (addition of a drop of dilute HCl causes effervescence). The pH of the solution is adjusted to ~3-4, NaCl is added to almost saturation and the alcohol is extracted several times with Et_2O , dried (K_2CO_3), filtered, and the filtrate is distilled to give cyclopentanol in ~90% yield. [Chaikin & Brown *J Am Chem Soc* **71** 122 1949, DOI: 10.1021/ja01169a033]. Its FT-IR (neat) has ν_{max} at 3334.9 (br OH), 2959.6 (CH), 1437.2 (OH ?), 1341.3, 1282.9, 1174.3, 1073.7, 994.3, and 837.1 cm^{-1} ; the 1H NMR (60MHz, $CDCl_3$, TMS) has δ at 1.55 (m, 4H, C-3 and C-4 -methylenes), 1.75 (m, 4H, C-2 and C-5 -methylenes), 2.30 (s, 1H, H-1) and 4.30 (brs, 1H, OH); and its ^{13}C NMR (15MHz, $CDCl_3$, TMS) has δ at 23.32 (C-3,4), 35.51 (C-2,5) and 73.86 (C-1). At atmospheric pressure cyclopentanol forms azeotropes with H_2O (96.3°, 42%), tetrachloroethylene (118.8°, 8%), chlorobenzene, *m*-xylene (132.8°, 40%), *p*-xylene (132.8°, 38%), *n*-Bu₂O (139.0°, 75%), among other solvents. It can be characterised as the **4-nitrobenzoate (m 56°)**, **3,5-dinitrobenzoate (m 123°)**, and **phenylcarbamate (m 137-138°)**. [*Beilstein* 6 H 5, 6 I 3, 6 II 3, 6 III 4, 6 IV 5.]

Cyclopentanone (dumasine, adipic ketone) [keto-form 120-92-3, enol-form 59557-02-7] C_5H_8O , **M 84.1, m -58°, -51°, b 23-24°/10mm, 130°/atm, d₄²⁰ 0.947, n_D²⁰ 1.4366, n_D²⁵ 1.4340. pK_a²⁵ 16.7.** This is a flammable liquid (flash point 26-31°) with a pleasant odour, but which is a strong **SKIN** and **EYE IRRITANT**, and should be used in a well ventilated fume cupboard. It is sparingly soluble in H_2O , and forms explosive mixtures with HNO_3 and H_2O_2 . It is volatile with Et_2O and care should be taken that careful fractionation is require when it is extracted with Et_2O . It is prepared by heating, in a distilling flask, a powdered mixture of adipic acid (200g, 1.34moles) and ground $Ba(OH)_2$ (10g) in a metal bath (e.g. Wood's metal) to 285-295° during 1-1.5 hours. [The temperature is best controlled with the thermometer within 5mm above the bottom of the flask and the temperature held as near as 290° as possible to minimising the amount of adipic acid which distils off.] During

the heating period, the *cyclopentanone* distils slowly with small quantities of adipic acid, and after a further 2 hours of heating a very small amount of dry residue remains in the flask. The distillate is separated from some H₂O and adipic acid by salting out with K₂CO₃, washed with H₂O, dried over CaCl₂ or anhydrous K₂CO₃, filtered and fractionated through an efficient column; fraction with b 128-131°/atm of the pure *ketone* (86-92g, 75-80%) is collected. [Thorpe & Kon *Org Synth Coll Vol 1* 192 1932, DOI: 10.15227/orgsyn.005.0037.] If cyclopentanone has been standing for a while, it can be purified by shaking it with aqueous KMnO₄ to remove materials absorbing around 230 to 240nm. Dry it over Linde-type 13X molecular sieves and fractionally distil it. It has also been purified by conversion to the NaHSO₃ adduct which, after crystallising four times from EtOH/water (4:1), is decomposed by adding to an equal weight of Na₂CO₃ in hot H₂O. The free cyclopentanone is steam distilled from the solution. The distillate is saturated with NaCl and extracted with *benzene (do not use H₂O, see above) which is then dried (anhydrous K₂CO₃), filtered and evaporated. The residue is then distilled [Allen et al. *J Chem Soc* 1909 1960, DOI: 10.1039/JR9600001909]. Its FT-IR (NaCl) has ν_{\max} at 2966 (CH), 1746.4 (C=O), 1407.6 (OH enol ?), 1278.2, 1153.0, 959.2, 834.2, 582.2 and 471.7 cm⁻¹; the ¹H NMR (15MHz, CDCl₃, TMS) has δ at 1.97 (t, 4H, C-3 and C-4 -methylenes), 2.17 (t, 4H, C-2 and C-5 -methylenes); and its ¹³C NMR (60MHz, CDCl₃, TMS) has δ at 23.24 (C-3,4), 38.30 (C-2,5) and 220.16 (C-1). The *oxime* [1192-28-5] C₅H₉NO, **M 99.1** forms prisms with **m 56.5°**, **b 120-121°/45mm**, **196°/atm** [Beilstein 7 H 7], the *semicarbazone* [5459-00-7] has **m 224°**, the *2,4-dinitrophenylhydrazone* [2057-87-6] has **m 145.5**, **146.5°**, and the *ethylene ketal* (1,4-dioxaspiro[4.4]nonane [176-32-9] has **m 153°**. The *enol*-form is present to the extent of 2.5 x 10⁻⁷% in the gas phase.

(±)-2-Cyclopentene-1-acetic acid [13668-61-6, 75247-34-6] C₇H₁₀O₂, **M 126.2**, **M 419°**, **b 93-94°/2.5mm**, **95-100°/4mm**, **109-114°/16mm**, **d**₄²⁵ **1.047g/ml**, **n**_D²⁰ **1.4688**, **1.4682**, **pK_{Est} ~4.8**. Prepared from the sequence cyclopentadiene → 3-chlorocyclopentene → ethyl-2-cyclopentene-1-malonate → 2-cyclopentene-1-malonic acid → 2-cyclopentene-1-acetic acid and purified by redistillation under reduced pressure. Optical resolution *via* the brucine salt gave pure diastereoisomeric salt with $[\alpha]_D^{28} +1.54$ (c 7.1, H₂O) that was basified with NH₃ to provide chiral *S*(+)-2-cyclopentene-1-acetic acid [67886-21-9] **b 105°/8mm**, **n**_D²⁵ **1.4673** and $[\alpha]_D^{30} +109.2$ (c 5.9, CHCl₃). This dextro acid was shown to have the *S* (D) configuration, and on standing at ~25° for eight months it became yellow and viscous with $[\alpha]_D^{30} +66.0$ (c 5.7, CHCl₃) and **n**_D²⁵ 1.4673; and when distilled gave a 55% recovery of pure (+)-acid with b 122-123°/30mm, **n**_D²⁵ 1.4678, and $[\alpha]_D^{30} +107.8$ (c 5.9, CHCl₃); and a red glassy residue. The *S*(+)-methyl ester [769820-82-5] has $[\alpha]_D^{25} +45$ (c 0.94, CHCl₃). The *S*(+)-acid was used to prepare **Chaulmoogric acid** [29106-32-9] see above by electrolysis with ethyl hydrogen brassylate. [Mislow & Steinberg *J Am Chem Soc* 77 3807 1955, DOI: 10.1021/ja01619a038; Beilstein 9 III 152.] *R*(-)-2-cyclopentene-1-acetic acid [696-67-3] and the *S*(-)-methyl ester [67920-83-6] have properties similar to those of their enantiomers except with negative optical rotations. The *racemic RS*(±)-methyl ester [20006-85-3] has **b 77-78°/20mm**. [For reduction to the respective 1-hydroxyethylcyclopent-2-enes see Irwin & Jones *J Am Chem Soc* 99 1625 1977, DOI: 10.1021/ja00447a057; for alternative synthesis from 1-chlorocyclopent-2-ene see Chapman et al. *J Am Chem Soc* 100 4878 1978, DOI: 10.1021/ja00483a039; for preparations using phenylseleno-cyclisations see Nicolaou et al. *JCS Chem Commun* 83 1979, DOI: 10.1039/C39790000083].

Cyclopentyl methyl ether (CPME) [5614-37-9] C₆H₁₂N, **M 100.2**, **fp -134.8°**, **b 105.44°/760mm** **105-106.5°/760mm** **106°/760mm**, **d**₄²⁰ **0.8627**, **n**_D²⁰ **1.4206**. This ether has the advantage of having a high resistance to hydroperoxide formation. It is an 'environmentally friendly' solvent and is an alternative to ether solvents like THF, Et₂O and methyl *tert*-butyl ether. Two methods after Williamson's synthesis have been reported. In the *first* Na (370g, 16g. atoms) is dissolved in MeOH (2370g, 74moles) and excess of MeOH is distilled off until NaOMe starts to separate. The solution is cooled to 60° and cyclopentyl chloride (1254g, 12moles, b 114°/atm [930-28-9]) is added slowly and kept at 60° for 100 hours. The mixture is then fractionated to give recovered cyclopentyl chloride (21%), cyclopentyl methyl ether (278g, 29% based on reacted cyclopentyl chloride) and cyclopentene [31% which distils as an azeotrope with MeOH (b 37-38°/atm)]. [Olson et al. *J Am Chem Soc* 69 2451 1947, DOI: 10.1021/ja01202a060.] In the *second*, Na (15g) is dispersed under hot xylene, cooled, the xylene is replaced with anhydrous Et₂O (150ml), and a solution of cyclopentanol (57g, b 141-142°/769mm, see [96-41-3]) in dry Et₂O (~100ml) is added with stirring during 3 hours and allowed to stand for 12 hours. Methyl iodide (103g) is then added during 2 hours to the preceding sodio compound as the Et₂O boils gently, allowed to stand overnight, the Et₂O is distilled off, the crude CPME is then distilled out and redistilled under N₂ at 105°/763mm to give pure *cyclopentyl methyl ether* (21g, 32%). It can be stabilised with 50ppm of

BHT. [Vogel *J Chem Soc* 1809 1948, DOI: 10.1039/JR9480001809; *Beilstein* 6 III 5, 6 IV 6.]

Cyclotetradecane [295-17-0] $C_{14}H_{28}$, **M 196.4**, **m 56°**. Recrystallise it twice from aqueous EtOH, then sublime it *in vacuo* [Drotloff et al. *J Am Chem Soc* 109 7797 1987, DOI: 10.1021/ja00259a031]. It also crystallises from MeOH (**m 54.2°**) and has **m 55°** after sublimation. [Prelog & Polyák *Helv Chim Acta* 40 816 1957, DOI: 10.1002/hlca.19570400332; *Beilstein* 5 III 152, 5 IV 177.]

Cyclotetradecanone [3603-99-4] $C_{14}H_{26}O$, **M 210.4**, **m 53°**, **b 145°/10mm**, **158-160°/12mm**, d_4^{20} 0.926, n_D^{20} 1.480. It is converted to the *semicarbazone*, $C_{15}H_{29}N_3O$, **m 201-202°** (198° and 198-199° also reported) which is recrystallised from EtOH. It is converted to cyclotetradecanone by hydrolysis [Drotloff et al. *J Am Chem Soc* 109 7797 1987, DOI: 10.1021/ja00259a031]. The ketone crystallises from EtOH (**m 52-53°**) and can be fractionated in a vacuum. The *oxime*, **m 113-114°**, crystallises from MeOH and sublimes at high vacuum. [Ruzicka et al. *Helv Chim Acta* 13 1152 1930, DOI: 10.1002/hlca.19300130540]. [*Beilstein* 7 IV 109.]

Decahydronaphthalene (decalin, mixed isomers) [91-17-8] $C_{10}H_{18}$, **M 138.2**, **m -125°**, **b 191.7°/760mm**, d_4^{20} 0.886, n_D^{20} 1.476. Stir decalin with concentrated H_2SO_4 for several hours. Then the organic phase is separated, washed with water, saturated aqueous Na_2CO_3 , again with water, dried with $CaSO_4$ or CaH_2 (and perhaps dried further with Na), filtered and distilled under reduced pressure (**b 63-70°/10mm**). It has also been purified by repeated passage through long columns of silica gel previously activated at 200-250°, followed by distillation from $LiAlH_4$ and storage under N_2 . Type 4A molecular sieves can be used as drying agent. Storage over silica gel removes water and other polar substances. [For the separation of *cis* and *trans* isomers see Seyer & Walker *J Am Chem Soc* 60 2125 1938, DOI: 10.1021/ja01276a028; and Baker & Schuetz *J Am Chem Soc* 69 1250 1947, DOI: 10.1021/ja01198a005; *Beilstein* 5 III 245.]

***cis*-Decahydronaphthalene** (decalin) [493-01-6] $C_{10}H_{18}$, **M 138.2**, **m -43.2°**, **b 67.0°/9mm**, **81-83°/19mm**, **195.7°/atm**, d_4^{20} 0.897, n_D^{20} 1.48113. Purification methods described for the mixed isomers are applicable here. The individual isomers can be separated by very efficient fractional distillation, followed by fractional crystallisation by partial freezing. The *cis*-isomer reacts preferentially with $AlCl_3$ and can be removed from the *trans*-isomer by stirring the mixture with a limited amount of $AlCl_3$ for 48 hours at room temperature, filtering and distilling. *Note*: the boiling points of the *cis* isomer are higher than those of the *trans* isomer. [Seyer & Walker *J Am Chem Soc* 60 2125 1938, DOI: 10.1021/ja01276a028; Baker & Schuetz *J Am Chem Soc* 69 1250 1947, DOI: 10.1021/ja01198a005.] A very pure authentic sample is best obtained by synthesis from *cis*-1,2-bis-chloroethylcyclohexane [Whitesides & Gutowski *J Org Chem* 41 2882 1976, DOI: 10.1021/jo00879a019]. *cis*-Decalin is a flexible molecule (relative to the *trans*-isomer) and its 1H NMR signals are more grouped together, i.e. the axial and equatorial H signals coalesce being time averaged because of the flexibility of the rings. This is not so in the *trans*-isomer where the rings are rigid and axial and equatorial H signals are more spread out. [Musher & Richards *Proc Chem Soc* 205 (p 230) 1958, DOI: 10.1039/PS9580000205; Gerig & Roberts *J Am Chem Soc* 88 2791 1966, DOI: 10.1021/ja00964a031; see also *cis*- and *trans*- decahydronaphthyridines Armarego *J Chem Soc (C)* 377 1967, DOI: 10.1039/J39670000377]. [*Beilstein* 5 IV 310.] ***trans*-Decahydronaphthalene** [493-02-7] has **m -32°, -30.6°**, **b 62.0°/9mm**, **187.3°/atm**, d_4^{20} 0.870, n_D^{20} 1.46968. See purification of *cis*-isomer above. *Note*: the boiling points of the *cis* isomer are higher than those of the *trans* isomer. [Seyer & Walker *J Am Chem Soc* 60 2125 1938, DOI: 10.1021/ja01276a028; Baker & Schuetz *J Am Chem Soc* 69 1250 1949, DOI: 10.1021/ja01198a005; *Beilstein* 5 IV 311.] See *cis*-decahydronaphthalene above for 1H NMR spectra.

(+)-Dehydroabietylamine (abieta-8,11,13-triene-18-ylamine) [1446-61-3] $C_{20}H_{31}N$, **M 285.5**, **m 41°**, **42.5-45°**, **b 192-193°/1mm**, **250°/12mm**, n_D^{40} 1.546, $[\alpha]_D^{20}$ +56.1 (c 2.4, pyridine), $[\alpha]_{546}^{20}$ +51 (c 1, EtOH), **pK_{Est} ~10.3**. The crude base is purified by converting 2g of base in toluene (3.3ml) into the *acetate salt* by heating at 65-70° with 0.46g of AcOH, and the crystals are collected and dried (0.96g from two crops, **m 141-143°**). The *acetate salt* [2026-24-6] $C_{22}H_{35}NO_2$, **M 345.5**, has **m 139-141°** and $[\alpha]_D^{20}$ +50 (c 2.4, pyridine) [*Beilstein* 12 IV 3006]. The acetate salt is then dissolved in warm H_2O , basified with aqueous NaOH and extracted with $*C_6H_6$. The dried extract ($MgSO_4$) is evaporated in vacuum leaving a viscous oil which crystallises and can be distilled. It is a useful base for optical resolutions. [Gottstein & Cheney *J Org Chem*

30 2072 1965, DOI: 10.1021/jo01017a518.] The *picrate* has **m** 234-236° (from aqueous MeOH), and the *formate salt* has **m** 147-148° (from heptane). [Beilstein **12** IV 3005.]

Diamantane (congressane, pentacyclo[7.3.1.1^{4,12}.0^{2,7}.0^{6,11}]tetradecane, decahydro-3,5,1,7-[1,2,3,4]-butanetetraylnaphthalene) [2292-79-7] C₁₄H₂₀, **M** 188.3, **m** 234-235°, 236-237°, 243-245°. Purify diamantane by repeated crystallisation from MeOH or pentane. Alternatively, purify it by dissolving it in CH₂Cl₂, washing with 5% aqueous NaOH and water, and drying (MgSO₄). The solution is filtered, concentrated to a small volume, an equal weight of alumina is added, and the solvent evaporated. The residue is placed on an activated alumina column (ca 4 x weight of diamantane) and eluted with petroleum ether (b 40-60°). Eight sublimations and twenty zone refining experiments gave material **m** 251° of 99.99% purity by differential analysis [Gund et al. *Tetrahedron Lett* 3877 1970, DOI: 10.1016/S0040-4039(01)98613-7; Courtney et al. *JCS Perkin Trans I* 2691 1972, DOI: 10.1039/P19720002691]. [For spectra see Cupas et al. *J Am Chem Soc* **87** 917 1965, DOI: 10.1021/ja01082a042.] It has been isolated from petroleum [Hala et al. *Angew Chem Int Ed* **5** 1045 1966, DOI: 10.1002/anie.196610451].

1,3-Diaminoadamantane [10303-95-4] C₁₀H₁₈N₂, **M** 166.2, **m** 52°, pK_{Est(1)} ~8.6, pK_{Est(2)} ~10.6. Purify it by zone refining. The *dibenzoyl* derivative has **m** 248° (from EtOH), and the *dihydrochloride salt* has **m** 310° (360°) after recrystallisation from aqueous EtOH or EtOH/Et₂O. The *dipicrate* has **m** 290-295° (yellow crystals from EtOH), and the *H₂PtCl₆ salt* does not decompose at 360°. [Prelog & Seiwert *Chem Ber* **74** 1769 1941, DOI: 10.1002/cber.19410741109; Stetter & Wulff *Chem Ber* **93** 1366 1960, DOI: 10.1002/cber.19600930619.] [Beilstein **13** III 27.]

cis-1,2-Diaminocyclohexane (*meso*-chxn) [1436-59-5] C₆H₁₄N₂, **M** 114.2, **b** 92-93°/8mm, 170°/atm, **d**₄²⁵ 0.952, **n**_D²⁰ 1.493, pK₁²⁰ 6.13 (6.41), pK₂²⁰ 9.93 (9.91). It is prepared by reduction of the *cis*-1,2-dioxime with boiling Na/EtOH, or from *cis*-1,2-dihydrazinocarbonylcyclohexane with NaNO₂/HCl [Jaeger & Blumendal *Z Anorg Chem* **175** 161 1928, DOI: 10.1002/zaac.19281750112]. Dry the diamine over solid KOH and distil it in a vacuum. It is a strong base, keep it away from CO₂, and store it in the dark under N₂. The *dihydrochloride* has **m** 307-310°, and the *dipicrate* (from H₂O) has **m** 260°(dec). This *cis*-diamine is a *meso* form because its mirror image is identical to it, and is superimposable over it (see below) [Beilstein **13** II 3, **13** IV 15.] (*±*)-**trans-1,2-Diaminocyclohexane** [(*±*)-chxn] [1121-22-8] C₆H₁₄N₂, **M** 114.2, has **m** 14-15°, **b** 78-81°/15mm, 85-88°/25mm, **d**₄²⁵ 0.951, **n**_D²⁰ 1.489, pK₁²⁰ 6.47 (6.72), pK₂²⁰ 9.94 (9.86). Purify this *racemic base*, and store it as for the *cis*-isomer above since it is a strong base, and becomes yellow on storage. [Beilstein **13** H 1, **13** III 8, **13** IV 1,5.]

cis/trans-1,2-Diaminocyclohexane resolution. [Note that the *trans*- isomer is a racemic mixture, whereas the *cis*- isomer has a *meso* configuration and its mirror image is identical and superimposable over it.] A solution of L-(+)-tartaric acid (150g, 0.99mol) in distilled H₂O (400ml) in a 1L beaker is stirred until clear, and a mixture of *cis*- and *trans*- 1,2-diaminocyclohexane (240ml, 1.94mol, of ~60/40 *trans/cis* mixture which is commercially available and is cheaper than the pure *trans* compound for which the same procedure applies) is added at a rate whereby the temperature rises to 70°. Glacial acetic acid (100ml, 1.75mol) is then added at such a rate that the temperature rises to 90°. A white precipitate immediately separates as the acid is added, and the slurry is stirred vigorously until the temperature cools slowly to 25° (~2 hours), and then is cooled further to ~5° in an ice bath for 2 hours, and is collected by filtration. The filter cake is washed with H₂O (100ml at ~5°), then EtOH (5 x 100ml), and dried by sucking air through it for 1 hour and analysed for enantiomeric purity *via* the bis-*m*-toluoyl amide*. The salt is then dried *in vacuo* at ~40° to give *R,R*-1,2-diamoniumcyclohexane mono-(+)-tartrate as a white solid (160g, 99%) in ≥99% enantiomeric excess (ee). If the optical purity is not as expected then recrystallise the salt in two crops from Me₂CO (1:20 w/v with ~60-70% recovery). The *S,S*-diastereomeric tartrate salt is obtained by using D(-)-tartaric acid. If the 'ee' is <99%, or if the difference between the top and the bottom of the cake is >0.2%, then the cake should be washed with more MeOH. If product of ≥99% 'ee' is required then recrystallisation of the tartrate salt by dissolving it in H₂O (≈ 1:10 w/v) with heating to 90°, followed by cooling to 5° overnight, gives ~60-70% recovery.

* The diammonium salt (25mg) is mixed with 4N NaOH (0.5ml), CH₂Cl₂ (1.5ml) and *m*-toluoyl chloride (50μl) with vigorous mixing. The lower (organic layer) is diluted with 10μl of *iso*-propanol, and 10μl of this is analysed by HPLC using *iso*-propanol/hexane (1:9) mixture at 1ml/minute flow rate. [Larrow et al. *J Org Chem* **59** 1939 1994, DOI: 10.1021/jo00086a062; Gasbøl et al. *Acta Chem Scand* **26** 3605 1972, DOI:

10.3891/acta.chem.scand.26-3605.]

1R,2R(-)-trans-1,2-Diaminocyclohexane [(-)-chxn] [20439-47-8] $C_6H_{14}N_2$, M 114.2, m 41-45°, $[\alpha]_D^{20}$ -25.5 (c 5 M HCl), m 14-15°. Distil or recrystallise the diamine from petroleum ether under N_2 or argon. It has a plain-negative ORD curve [Gillard *Tetrahedron* **21** 503 1965, DOI: 10.1016/S0040-4020(01)82221-3; O'Brien & Toole *J Am Chem Soc* **77** 1368 1955, DOI: 10.1021/ja01610a089]. Store it as above. The **1R,2R-base L-tartrate salt** has [39961-95-0] $C_6H_{14}N_2 \cdot C_4H_6O_6$, M 264.3, m 273° and $[\alpha]_D^{25}$ +12.5 (c 4, H_2O), and can be used to purify and/or optically enrich the free base. [Beilstein **13** III 6, and references below.] **1S,2S(+)-trans-1,2-Diaminocyclohexane [(+)-chxn]** [21436-03-3] has m 42-45°, b 104-110°/40mm, $[\alpha]_D^{20}$ +25.5 (c 5 M HCl). Distil or recrystallise the diamine from petroleum ether under N_2 or argon. Store it as above. It has a plain-positive ORD curve [Gillard *Tetrahedron* **21** 503 1965, DOI: 10.1016/S0040-4020(01)82221-3; O'Brien & Toole *J Am Chem Soc* **77** 1368 1955, DOI: 10.1021/ja01610a089]. The **1S,2S-base D-tartrate salt** has [67333-70-4] $C_6H_{14}N_2 \cdot C_4H_6O_6$, M 264.3, has m 180-184°(dec) and $[\alpha]_D^{25}$ -12.5 (c 4, H_2O), from which the free base can be purified or optically enriched. It is a useful chiral synthon. [Takahashi et al. *Tetrahedron Lett* **30** 7095 1989, DOI: 10.1016/S0040-4039(01)93432-X; Hanessian et al. *J Org Chem* **58** 1991 1993, DOI: 10.1021/jo00060a004; for absolute configuration see Gillard *Tetrahedron* **21** 503 1965, DOI: 10.1016/S0040-4020(01)82221-3; Beilstein **13** III 7.]

trans-1,4-Diaminocyclohexane [2615-25-0] $C_6H_{14}N_2$, M 114.2, m 69-72°, b 197°/760mm, $pK_{est(1)}$ ~9.4, $pK_{est(2)}$ 10.8. Recrystallise the diamine from petroleum ether under N_2 or argon as it should be an even stronger base than the above 1,2-diamine isomers. It distills under N_2 . Store it in the dark under N_2 . [Beilstein **13** I 3, 13 III 11.]

(1R,2SR)-cis-1,2-Diaminocyclopentane (1R,2SR)-cis-1,2-cyclopentanediamine, meso-1,2-cyclopentanediamine, (meso)-cptn [40535-45-3; no configuration 41330-23-8] $C_5H_{12}N_2$, M 100.1, b 62-65°/13.5mm, 99°/60mm. Whereas reduction of *anti*-cyclopentane-1,2-dione dioxime (see below, and from reaction of cyclopentane-1,2-dione and NH_2OH) with Na/EtOH yields predominantly *trans*-(±)-diamine, the reaction of cyclopentane-1,2-dione monooxime [31597-37-0] with NH_2OH gives a 2:1 mixture of *anti*- and *amphi*-dioximes which provide the less water soluble red bis-*anti*-dioximatoNi(II) complex and the more water soluble yellow-brown bis-*amphi*-dioximatoNi(II) complex respectively. When this mixture of Ni(II) complexes, or the *amphi*-complex, is reduced with KBH_4 in 'diglyme', the base finally isolated is pure *cis*-1,2-cyclopentanediamine.

Thus, **bis(amphi-cyclopentane-1,2-dione dioximato)nickel(II)** (79g, 90.5mol, dried over H_2SO_4 in vacuo) is added to a solution of KBH_4 (30g) dissolved in dry diglyme [600ml, bis(2-methoxyethyl) ether, [111-96-6]] in 2L flask. N_2 is bubbled through the solution, then during 1 hour anhydrous $AlCl_3$ (25g) in dry diglyme (100ml) is added dropwise carefully as vigorous evolution of H_2 ensues; the temperature being kept below 35° by cooling in ice-water. When evolution of H_2 ceases, the mixture is heated at 70° for 18 hours. Then a solution of KOH (100g) in H_2O (150ml) is added carefully, and the *cis*-diamine is steam distilled off; 6L of distillate are collected, acidified with 12M HCl (to pH 3), and the dry *cis*-1,2-cyclopentanediamine dihydrochloride (23g, 27% based on the Ni complex) is isolated as for the *trans*-(±)-isomer below. The free base, *cis*-1,2-cyclopentanediamine b 62-65°/13.5mm (83%, based on the dihydrochloride) is stored under N_2 at -20°. [Toftlund & Pedersen *Acta Chim Scand* **26** 4019 1972, DOI: 10.3891/acta.chem.scand.26-4019]. This *cis*-diamine has been fractionated through a vacuum-jacketed distillation column as described by Ray et al. [*Rev Sci Instr* **28** 200 1957, DOI: org/10.1063/1.1746483] and the fraction with b 99°/60mm contained pure *cis*-isomer [Phillips & Royer *Inorg Chem* **4** 616 1965, DOI: 10.1021/ic50027a004].

The 1H NMR in D_2O (TMS as external standard) can be used to distinguish between *trans*-(±)-1,2-diamine and the preceding *cis*-(*meso*)-1,2-diamine because the triplets from carbon atoms 1 + 2 occur at δ 2.72 and δ 2.77 respectively [Phillips & Royer *Inorg Chem* **4** 616 1965, DOI: 10.1021/ic50027a004].

Distinct bis- and tris- *cis*-cptn complexes such as $[Ni(cis-cptn)_2](ClO_4)_2 \cdot 6H_2O$, $[Ni(cis-cptn)_3](ClO_4)_2$ and $[Co(cis-cptn)_3]Br_2 \cdot 3H_2O$ are readily formed [Toftlund & Pedersen *Acta Chem Scand* **26** 4019 1972, DOI: 10.3891/acta.chem.scand.26-4019]. [Beilstein **4** II 3.]

(1R,2RS)-(±)-trans-1,2-Diaminocyclopentane [(1R,2RS)-trans-1,2-cyclopentanediamine, (±)-trans-1,2-cyclopentanediamine, (±)-cptn] [3145-88-8] $C_5H_{14}N_2$, M 100.1, b 65-67°/13.5mm, 103°/60mm, 170°/atm,

pK_{Est 1} ~7.0 , pK_{Est 2} ~10.0. This diamine is a strong base and is best kept as its salts which are not greatly affected by moisture and CO₂. It is prepared by reduction of **anti-cyclopentane-1,2-dione dioxime** [6635-29-6] **M 128.1, m 210° (215° dec; 210-225° dec, darkening at 192-205°** also reported for an analytically pure sample) which is obtained by adding 2-ethoxycarbonylcyclopentanone (31.2g, b 102-104°/12mm, [611-10-9]) to a solution of NaOH (8.8g) in H₂O (160ml) whereby a heavy white precipitate separates. A solution of NaNO₂ (13.8g) in H₂O (32ml) is then added to it and the mixture, under N₂, is shaken mechanically for 44 hours at ~25°. The resulting clear yellow solution is cooled to 0°, and at this temperature 6N H₂SO₄ is added carefully while CO₂, and oxides of nitrogen evolve. The solution is extracted with Et₂O (in a continuous liquid-liquid extractor) for 5.5 hours, the extract is evaporated under N₂ and reduced pressure to a small volume of ~20ml. This suspension of crude cyclopentane-1,2-dione monooximes is treated with a solution of hydroxylamine hydrochloride (13.9g) and NaOH (8.0g) in H₂O (50ml) when heat is evolved and a precipitate separates immediately. The mixture is allowed to stand overnight at 5°, the light tan crude **anti-cyclopentane-1,2-dione dioxime** (16.8g, 66%, decomposes at 230-240°) is filtered off and is dried *in vacuo*, leaving the more soluble **amphi-cyclopentane-1,2-dione dioxime** in the filtrate. The analytically pure **anti-dioxime** (dec at 210-225° without melting, 83% recovery, [6635-29-6]) can be obtained by dissolving the crude material (1g) in 2% aqueous NaOH (25ml) at ~25° (slightly turbid), stirring with Norit, filtering, neutralising the filtrate with 1 N HCl (to phenolphthalein), filtering off the solid, washing it twice with H₂O, once with Me₂CO and it drying *in vacuo*. [Cope et al. *J Am Chem Soc* **73** 1199 1951, DOI: 10.1021/ja01147a096; Lloyd & Marshall *J Chem Soc* 2597 1956, DOI: 10.1039/JR9560002597.]

Alternatively, to a stirred mixture of granulated Na (38g) and *C₆H₆ (900ml, dried over Na) is added diethyl adipate (225ml, 1.11mole, [141-28-6]) rapidly followed by absolute EtOH (5ml), and is boiled under reflux with stirring for 18 hours then cooled to 0°. The white solid that separated is filtered off, washed with a little *C₆H₆ and dried *in vacuo* over shredded paraffin wax to give **sodium cyclopentanone-2-carboxylate** (180g). This salt (1mol) is partly dissolved in ice-water (600ml), N₂ is bubbled through for 15 minutes to remove O₂; and a mixture of a solution of NaOH (4g, 0.1mol) and NaNO₂ (70g, 1.03mol) in H₂O (160ml) is added dropwise to it. The mixture is then heated under N₂ with vigorous stirring at 40° for 30 hours. After cooling to 0°, 6M H₂SO₄ (200ml, 1.2mol) is added in one hour; and when evolution of CO₂ has ceased the solution is neutralised with 12M ammonia. A solution of NH₂OH HCl (70g, 1.0mol) in H₂O (160ml), neutralised with K₂CO₃ (70g, 0.5mol), is immediately added. After 1 hour, the precipitation of crude **anti-cyclopentane-1,2-dione dioxime** as light tanned needle-like crystals is complete; it is filtered off, washed with a little H₂O and dried *in vacuo* over H₂SO₄ for 48 hours (yield 90g, 62% calculated on diethyl adipate used). The filtrate contains the **amphi-cyclopentane-1,2-dione dioxime** which is isolated as the yellow-brown **bis(amphi-cyclopentane-1,2-dione dioximato)nickel(II) complex** (30g) by slowly adding Ni(II)SO₄ 7H₂O (20g) in H₂O (60ml) to this filtrate, and washing the complex with H₂O and drying in air. [Toftlund & Pedersen *Acta Chim Scand* **26** 4019 1972, DOI: 10.3891/acta.chem.scand.26-4019.] **Bis(amphi-cyclopentane-1,2-dione dioximato)nickel(II) complex** is obtained analytically pure, as a red precipitate in 70% yield, from **anti-cyclopentane-1,2-dione dioxime** (12.8g, 0.1mol) in 4M ammonia (15ml), by adding slowly Ni(II)Cl₂ 6H₂O (12g, 0.05mol) in H₂O (40ml) to it, filtering off the solid, washing it with H₂O and drying it in air. [Toftlund & Pedersen *Acta Chim Scand* **26** 4019 1972, DOI: 10.3891/acta.chem.scand.26-4019.]

Reduction to (±)-diamine: The **anti-cyclopentane-1,2-dione dioxime** (10g) in absolute EtOH (2L) is reduced by addition of freshly cut small pieces of Na metal (180g) over a period of several hours. The solution is refluxed until all the metal has dissolved. The EtOH is distilled off *in vacuo*, the residue is basified strongly with 10N NaOH, and the base is steam distilled. The distillate is made strongly alkaline with aqueous NaOH, extracted with trichloroethylene, the solvent is evaporated and **trans-1,2-cyclopentanediimine** then distils at **65°/13.5mm, n_D²⁵ 1.4850-1.4858** in 29% yield. When the fractional distillation is carried out using a column (90cm x 1 cm) packed with glass helices and the reflux ratio is kept at >40:1, the first 25% of distillate contains a significant amount of lower boiling **cis-diamine, 99°/60mm**; the remaining 75% of material then boils at a constant temperature, **103°/60mm**, and is the **trans-(±)-diamine**. **Alternatively**, the steam distillate is acidified with concentrated HCl, the solution is evaporated to dryness *in vacuo*, and the residue is recrystallised from EtOH/Et₂O to provide **trans-(±)-1,2-cyclopentanediimine dihydrochloride** in 77% yield, which **decomposes at 287-290°** (darkening gradually >230°), and forms a **dihydrate** on exposure to air. The **trans-(±)-dipicrate** has **m 233-233.5° (dec)** after recrystallisation from aqueous EtOH or H₂O, and the **trans-(±)-diacetyl derivative** has **m 226.5-227.5° (dec)** after recrystallisation from EtOH by addition of Et₂O. [Cope et al. *J Am Chem Soc* **73** 1199 1951, DOI: 10.1021/ja01147a096; Phillips & Royer *Inorg Chem* **4** 616 1965, DOI: 10.1021/ic50027a004;

Lloyd & Marshall *J Chem Soc* 2597 1956, DOI: 10.1039/JR9560002597; Jaeger & Blumendal *Z Anorg Allgem Chem* **175** 161 1928, DOI: 10.1002/zaac.19281750112.]

Alternative reduction to (±)-diamine: The *anti*-cyclopentane-1,2-dione dioxime (20g, 156mmol) in absolute EtOH (1.5L) is reduced by heating to 40° (internal temperature), and stirring under a reflux condenser, with Mg powder (8g) and Hg(II)Cl₂ (0.1g) while Na metal (150g, 6.5mol, in 5g pieces) is added *via* the top of the condenser and keeping the temperature below 50° (internal temperature). The mixture is finally heated to boiling in order to dissolve all the Na. The (±)-diamine is isolated by steam distillation (external heating being necessary so as to maintain a small volume in the flask) until the pH of the distillate is below 8 (*ca* 4L). This is acidified (pH 3) with 12M HCl, evaporated almost to dryness *in vacuo* (at ~25°), excess HCl is removed by washing with EtOH/Et₂O (2:1v/v, 20ml), and the hygroscopic salt is dried over KOH in a desiccator to give the (±)-diamine dihydrochloride (5g, 18% based on dioxime). The free base is obtained by adding the dihydrochloride (5g, 29mmol) in small portions with stirring to a 1:1v/v mixture of MeOH/Et₂O (50ml) into which is dissolved Na (1.7g, 74mmol, cooling). After stirring for 1 hour, the precipitated NaCl is filtered off, extracted with MeOH/Et₂O (1:1v/v, 5ml), the combined extract and filtrate are evaporated to ~3ml and this residue is distilled in a N₂ atmosphere to give (±)-*trans*-cyclopentane-1,2-diamine, **b 65-67°/13.5mm** (2.5g, 89%, based on the dihydrochloride) which is stored under N₂ at -20°. Distinct *trans*-(±)-cptn complexes of Ni, Co, Rh and Pt have been prepared; some are enantiomorphous and others are structural isomers [Phillips & Royer *Inorg Chem* **4** 616 1965, DOI: 10.1021/ic50027a004; Jaeger & Blumendal *Z Anorg Allgem Chem* **175** 161 1928, DOI: 10.1002/zaac.19281750112; Toftlund & Pedersen *Acta Chim Scand* **26** 4019 1972, DOI: 10.3891/acta.chem.scand.26-4019]. The ¹H NMR in D₂O (TMS as external standard) can be used to distinguish between *trans*-(±)-1,2-diamine and the preceding *cis*-(*meso*)-1,2-diamine because the triplets from carbon atoms 1 + 2 occur at δ 2.72 and δ 2.77 respectively [Phillips & Royer *Inorg Chem* **4** 616 1965, DOI: 10.1021/ic50027a004].

(1R,2R)-(-)-trans-1,2-Diaminocyclopentane dihydrochloride [(-)-cptn HCl] [1030390-38-5] C₆H₁₄N₂. 2HCl, M 173.1, dec on melting, [α]_D²⁵ -23.5 (c 2, 1M HCl) and (1S,2S)-(+)-trans-1,2-Diaminocyclopentane dihydrochloride [(+)-cptn HCl] [477873-22-6] M 173.1, dec on melting, [α]_D²⁵ +23 (c 2, 1M HCl). **Optical resolution: (±)-Cptn (2g, 20mmol) is resolved by adding it to a solution of D-(+)-tartaric acid (7g, 47mmol) in MeOH (1ml) and H₂O (7ml), and at 40° MeOH (12ml) is added dropwise. The mixture is then kept at 3° for 18 hours, the crystalline solid is filtered off, washed with MeOH (10ml) to give the less soluble hydrogen tartrate salt [5.2g, 65% based on (±)-cptn]. After two recrystallisations, the (-)-cptn—(+)-tartrate salt [2.5g, 31.4% based on (±)-cptn] has **m 134-135°** (143-144° also reported), [α]_D²⁵ +10.1 (c 2, H₂O). When this salt (2.4g, 6mmol) in H₂O (8ml) is treated with a solution of KCl (0.9g, 12mmol) in H₂O (4ml), the highly insoluble potassium hydrogen tartrate separates and is filtered off after standing for 1 hour. Addition of solid KOH (8g, with cooling) allowed solid (-)-cptn hydrate to separate, but the mixture is extracted thoroughly with Et₂O (5 x 10ml), the extract is dried (KOH or MgSO₄), filtered, and evaporated to give (-)-*trans*-cptn (0.55g, 92% based on tartrate salt). When carried out on a larger scale the (-)-*base* can be distilled, **b 166°/atm, d₂₅ 0.9463**, and has [α]_D -54.7 (neat), [α]_D -42 (*C₆H₆), [α]_D -39 (H₂O), and a plain-negative ORD curve. (-)-*trans*-Cptn sulfate has [α]_D -12 (H₂O). [Toftlund & Pedersen *Acta Chim Scand* **26** 4019 1972, DOI: 10.3891/acta.chem.scand.26-4019; Jaeger & Blumendal *Z Anorg Allgem Chem* **175** 161 1928, DOI: 10.1002/zaac.19281750112; Phillips & Royer *Inorg Chem* **4** 616 1965, DOI: 10.1021/ic50027a004]. (+)-Cptn—(+)-tartrate can be obtained from the mother liquors of the above resolution in order to isolate the (+)-base; but it is advisable to collect all the mother liquors, liberate the free base with KOH, isolate it and repeat the process using L-(-)-tartaric acid to form the less soluble diastereoisomeric (+)-cptn—(-)-tartrate salt. Pure (+)-*trans*-cptn can then be isolated as above and will have the same physical properties except for the optical rotations that will be of opposite sign and has a plain-positive ORD curve [Dunlop et al. *J Chem Soc* 3160 1964, Gillard *Tetrahedron* **21** 503 1965, DOI:10.1016/S0040-4020(01)82221-3; O'Brien & Toole *J Am Chem Soc* **77** 1368 1955, DOI: 10.1021/ja01610a089]. [Beilstein **4** II 3, 4 III 5]**

Complexes of optically active cptn with Rh, Co, Ni and Pt have been prepared, and in some cases enantiomeric and geometrical isomers have been identified. One has been used to obtain pure enantiomeric diamine, for example, the complex [Co₂(±-cptn)₇ (H₂O)₂]⁶⁺ has been resolved into three bands on Amberlite CG-50 carboxylic acid resin 400-400 by elution with aqueous HCl (pH 2, flow rate 5ml/hour) and collecting 40ml fractions. Bands I, II and III are probably geometrical isomers because when the isolated complexes are decomposed with concentrated aqueous NaOH, extracted with *C₆H₆, the extract dried and evaporated, the free

base (+)-*cptn* with $[\alpha]_D^{25} +42 \pm 1$ (*C₆H₆) is obtained (compare with above) [Phillips & Royer *Inorg Chem* **4** 616 1965, DOI: 10.1021/ic50027a004].

The optical properties (ORD, CD) of several *cptn*-transition metal complexes have been thoroughly investigated and used to determine and confirm the absolute configuration of *cptn* [Dunlop et al. *J Chem Soc* 3160 1964, DOI: 10.1039/JR9640003160; Gillard *Tetrahedron* **21** 503 1965, DOI: 10.1016/S0040-4020(01)82221-3]. The crystal structure of (-)₅₈₇-[Co(+*trans-cptn*)₃]Cl₃·4H₂O confirms the *1S,2S*-configuration for the dextro-enantiomer [Ito et al. *Acta Cryst B* **27** 2187 1971, DOI: 10.1107/S0567740871005545].

***trans*-1,2-Dibromocyclopentane** [10230-26-9] C₅H₈Br₂, M 227.9, b 72.5°/15mm, 74°/17mm, d₄²⁰ 1.857, n_D²⁵ 1.5460. It is prepared by addition of bromine to cyclopentene at -20° followed by fractional distillation. The ¹H NMR (60MHz, CCl₄) has a complex multiplet centered at ~2.5 ppm (3 x CH₂), and two triplets centered at 4.58ppm (*J* = 7.4Hz, for 2 x CHBr) from TMS. [Altona et al. *Recl Trav Chim, Pays-Bas* **85** 983 1966, DOI: 10.1002/recl.19660851003; cf. Abell et al. *J Am Chem Soc* **82** 3610 1960, DOI: 10.1021/ja01499a034.]

***cis*-3,4-Dichlorocyclobutene (3*RS,4SR*)** [2957-95-1] C₄H₄Cl₂, M 123.0, b 70-71°/55mm, 74-76°/55mm, d₄²⁰ 1.297, n_D²⁰ 1.499. Distil the cyclobutene at 55mm through a 36-in platinum spinning band column, a fore-run b 58-62°/55mm is mainly 1,4-dichlorobutadiene. When the temperature reaches 70° the reflux ratio is reduced to 10:1 and the product is collected quickly. It is usually necessary to apply heat frequently with a sun lamp to prevent any dichlorobutadiene from clogging the exit in the early part of the distillation [Pettit & Henery *Org Synth* **50** 36 1970, DOI: 10.15227/orgsyn.050.0036; Warrenner et al. *Aust J Chem* **26** 389 1973, DOI: 10.1071/CH9730389]. (±)-((3*RS,4RS*)-3,4-Dichlorocyclobutene (3*RS,4SR*) [41326-64-1], m -4°, b 28°/20mm, is the more stable isomer formed by catalytic (AlCl₃) isomerisation of the *cis*-isomer and purified by gas preparative chromatography on a TCP-on-Fluoropak column at 50-75° and dried by distillation through P₂O₅. [Prepn and IR, Raman and NMR spectra of *cis*- and *trans*- isomers: Craig et al. *Spectrochimica Acta Part A*, **47A** 881 1991, DOI: 10.1016/0584-8539(91)80275-N.]

Dicyclohexylamine (Cy₂NH₂) [101-83-7] C₁₂H₂₃N, M 181.3, m -1°, -2°, b 83°/1mm, 99.3°/4mm, 113.5°/9mm, 117-120°/10mm, 135.4°/30mm, 154.3°/50mm, 199°/200mm, 255.8°/atm, 256°/atm, d₄²⁰ 0.912, n_D²⁰ 1.4845, pK²⁵ 11.25. It is a strong base, has a fishy odour, is soluble in H₂O, and organic solvents and forms adducts with some of them. It can be purified by fractional distillation at atmospheric pressure, but if the distillate is coloured (e.g. green) then distillation under reduced pressure gives a clear colourless liquid [Vogel *J Chem Soc* 1825 1948, DOI: 10.1039/JR9480001825]. The *hydrochloride* [4693-92-9] crystallises from H₂O or EtOH and has m 334-335° (sealed tube), and the *acetyl derivative* [1563-91-3] has m 103° (from Et₂O). [Diwoy & Adkins *J Am Chem Soc* **53** 1868 1931, DOI: 10.1021/ja01356a035; *Beilstein* **12** H 6, **12** I 114, **12** II 7, **12** III 19, **12** IV 22.] **SKIN and EYE IRRITANT.**

1,3-Dicyclohexylcarbodiimide (DCC) [538-75-0] C₁₃H₂₂N₂, M 206.3, m 34-35°, b 95-97°/0.2mm, 122-124°/6mm, 155°/11mm. It is sampled as a liquid after melting in warm H₂O. It is sensitive to air, and it is a *potent skin irritant*. It can be distilled in a vacuum, and is best stored in a tightly stoppered bottle in a freezer. It dissolves readily in CH₂Cl₂ and pyridine where the reaction product with H₂O, after condensation, is dicyclohexyl urea which is insoluble and can be filtered off. *Alternatively*, dissolve it in CH₂Cl₂, add powdered anhydrous MgSO₄, shake for 4 hours, filter, evaporate and distil the residue at 0.6mm pressure and oil bath temperature of 145°. Possible SKIN ALLERGEN, handle with gloves, and do not breathe it in. [Bodansky et al. *Biochemical Preparations* **10**, 122 1963, Schmidt & Seefelder *Justus Liebigs Ann Chem* **571** 83 1951, DOI: 10.1002/jlac.19515710110; Schmidt et al. *Justus Liebigs Ann Chem* **612** 11 1958, DOI: 10.1002/jlac.19586120103; *Beilstein* **12** IV 72.]

***N,N*-Dicyclohexylmethylamine (Cy₂NMe, *N*-methyldicyclohexylamine)** [7560-83-0] C₁₃H₂₅N, M 195.3, b 131-133°/13mm, 135-137°/15mm, 150°/50mm, 265°/atm, d₄²⁵ 0.912, n_D²⁵ 1.4900, pK 8.2. When dicyclohexylamine (90.5g, 0.5mol, see 101-83-7) and 55% aqueous formic acid (121.1ml, 2.5mol) are heated at ~100° and 35% aqueous formaldehyde (97.5g, 11.25mol) is added dropwise over an hour, CO₂ evolution occurs. After gas evolution is complete, ~15 hours, concentrated HCl (1mol) is added and the mixture is evaporated *in vacuo*. The free base is liberated from the residue with strong aqueous NaOH and the oil is extracted with Et₂O, dried (solid KOH), filtered, the filtrate is evaporated *in vacuo* and the residue is fract-

ionally distilled to give Cy_2NMe in 51% yield. It was also obtained by alkylating Cy_2NH with dimethylsulfate. [Hünig & Kiessel *J Prakt Chem* **5** 224 1958, DOI: 10.1002/prac.19580050504.] The **hydrochloride** has **m 154-154°** (from EtOAc), and the **methiodide** has **m 228-229°** (from EtOH/Et₂O). [*Beilstein* **12** I 115, **12** II 8, **12** III 21, **12** IV 23.]

3,4-Diethoxy-3-cyclobutene-1,2-dione (diethyl squarate) [5321-87-8] $\text{C}_8\text{H}_{10}\text{O}_4$, **M 170.2**, **b 95°/0.1mm**, **89-91°/0.4mm**, **88-92°/0.4mm**, **d** $^{20}_4$ **1.162**, **n** $^{25}_D$ **1.5000**. Dissolve the ester in Et₂O, wash it with Na₂CO₃, H₂O and dry (Na₂SO₄) it, filter, evaporate and distil it using a Kügelrohr, or purify it by chromatography. Use a Kieselgel column and elute with 20% Et₂O/petroleum ether (b 40-60°), then with Et₂O/petroleum ether (1:1), evaporate and distil the residue *in vacuo*. [Dehmloew & Schell *Chem Ber* **113** 1 1980, DOI: 10.1002/cber.19801130103; Perri & Moore *J Am Chem Soc* **112** 1897 1990, DOI: 10.1021/ja00161a039; IR: Cohen & Cohen *J Am Chem Soc* **88** 1533 1966, DOI: 10.1021/ja00959a040] **It can cause severe dermatitis**. [Foland et al. *J Am Chem Soc* **111** 975 1989, DOI: 10.1021/ja00185a030; Perri et al. *Org Synth* **69** 220 1990, DOI: 10.15227/orgsyn.069.0220].

N,N-Diethylcyclohexylamine [91-65-6] $\text{C}_{10}\text{H}_{21}\text{N}$, **M 155.3**, **b 85-87°/15mm**, **193°/760mm**, **d** $^{25}_{25}$ **0.850**, **n** $^{25}_D$ **1.4562**, **pK²⁵ 10.72**. Dry the amine with BaO and fractionally distil it. It is a strong base, store away from CO₂. The **picrate** has **m 98-99°** (from aqueous EtOH) and the **methiodide** has **m 224°** (from Me₂CO/petroleum ether) [Cadogan *J Chem Soc* 1079 1957, DOI: 10.1039/JR9570001079.] [Bain & Pollard *J Am Chem Soc* **61** 2704 1939, DOI: 10.1021/ja01265a039; *Beilstein* **12** H 6, **12** III 14, **12** IV 19.]

Diethyl cyclopropane-1,1-dicarboxylate [1559-02-0] $\text{C}_9\text{H}_{14}\text{O}_4$, **M 186.2**, **b 94-96°/10mm**, **d** $^{25}_4$ **1.055**, **n** $^{20}_D$ **1.433**. If it is free from OH bands in the IR, then fractionally distil the ester and redistil the middle fraction. Otherwise shake it with aqueous NaHCO₃, dry it (MgSO₄), filter and distil as before or re-esterify it. [As synthon see Danishefsky *Acc Chem Res* **12** 66 1979, DOI: 10.1021/ar50134a004; *Beilstein* **9** I 314, **9** II 512, **9** III 3595, **9** IV 2786.]

Dimedone (5,5-dimethylcyclohexane-1,3-dione, Methone) [126-81-8] $\text{C}_8\text{H}_{12}\text{O}_2$, **M 140.2**, **m 148-149°**, **pK²⁵ 5.27**. Crystallise dimedone from acetone (*ca* 8ml/g), water or aqueous EtOH. Dry it in air. [Shriner & Todd *Org Synth Coll Vol* **2** 200 1943, DOI: 10.15227/orgsyn.015.0014; *Beilstein* **7** H 559, **7** IV 1999.]

cis- and trans-1,4-Dimethylcyclohexane (hexahydro-p-xylene) [cis/trans 589-90-2] C_8H_{16} , **M 112.2**, **m -91.6°, -87.44°**, **b 120°/atm**, **120°/atm**, **d** $^{20}_4$ **0.788**, **n** $^{25}_D$ **1.427**. Free it from olefins by shaking with concentrated H₂SO₄, washing with water, drying and fractionally distilling it. It should be possible to separate **trans-isomer** [2207-04-7] **b 119-119.5°/atm (119.35°/760mm)**, **n** $^{20}_D$ **1.4209**, from the **cis-isomer** [624-29-3] **b 124-125°/atm (125.3°/779.4mm)**, **n** $^{18}_D$ **1.4291**, by using an efficient distillation still by virtue of the difference in boiling points. [*Beilstein* **5** IV 123 (*trans*), 122 (*cis*).] [synthesis from 1,4-bishydroxymethylcyclohexanes: Haggis & Owen *J Chem Soc* 408 1953, DOI: 10.1039/JR9530000408; separation by fractionation: Forziati et al. *J Res Nat Bur Stand* **36** 129 1946, DOI: org/10.6028/jres.036.005; thermodynamic data: Huffman et al. *J Am Chem Soc* **71** 584 1949, DOI: 10.1021/ja01170a058; *Beilstein* **5** H 38, **5** III 102, **5** IV 122.]

1,2-Dimethylcyclohexene [1674-10-8] C_8H_{14} , **M 110.2**, **b 135-136°/760mm**, **d** $^{25}_4$ **0.826**, **n** $^{25}_D$ **1.4587**. Pass it through a column of basic alumina and distil it. If removal of 2-methylmethylenecyclohexane or 2,3-dimethylcyclohexene is required, then fractionation through a centre-rod column operating at ~50 theoretical plates is required. [Hammond & Nevitt *J Am Chem Soc* **76** 4121 1954, DOI: 10.1021/ja01645a020; *Beilstein* **5** H 72, **5** I 36, **5** II 46, **5** III 213, **5** IV 268.]

Ethyl chrysanthemate (ethyl ±2,2-dimethyl-3(c and t)-[2-methylpropenyl]cyclopropane carboxylate) [97-41-6] $\text{C}_{12}\text{H}_{20}\text{O}_2$, **M 196.3**, **b 98-102°/11mm**, **117-121°/20mm**. Purify the ester by vacuum distillation. The free **trans-acid** has **m 54°** (from EtOAc), and the free **cis-acid** has **m 113-116°** (from EtOAc). The **4-nitro-phenyl ester** has **m 44-45°** (from petroleum ether) [Campbell & Harper *J Chem Soc* 283 1945, DOI: 10.1039/JR9450000283; IR: Allen et al. *J Org Chem* **22** 1291 1957, DOI: 10.1021/jo01362a002]. [*Beilstein* **9** II 45.]

Ethylcyclohexane [1678-91-7] C_8H_{16} , **M 112.2**, **m** -111°, **b** 131.2°/742mm, 130-132°/atm, d_4^{20} 0.7839, n_D^{20} 1.43304, n_D^{25} 1.43073. Purify it by azeotropic distillation with 2-ethoxyethanol; then the alcohol is washed out with water and, after drying, the ethylcyclohexane is redistilled. The dried material has been repeatedly fractionated over Na. [Groves & Baker *J Chem Soc* 1144 1939, DOI: 10.1039/JR9390001144; *Beilstein* 5 H 35, 5 III 90, 5 IV 110.]

Ethyl cyclohexanecarboxylate [3289-28-9] $C_9H_{16}O_2$, **M 156.2**, **b** 76-77°/10mm, 92-93°/34mm, 196-196.2°/760mm, d_4^{20} 0.955, n_D^{20} 1.441. Wash the ester with N sodium hydroxide solution, then water, dry with Na_2SO_4 and distil it. The **amide** has **m** 185-186°. [Adkins & Cramer *J Am Chem Soc* 52 4349 1930, DOI: 10.1021/ja01374a023; Newman & Walborsky *J Am Chem Soc* 72 4296 1950, DOI: 10.1021/ja01165a527; *Beilstein* 9 III 17, 9 IV 18.]

Ethyl 2-oxocyclohexanecarboxylate (2-ethoxycarbonylcyclohexanone, 2-carbethoxycyclohexanone, ethyl 2-ketohexahydrobenzoate) [1655-07-8] $C_9H_{14}O_3$, **M 170.2**, **b** 86-88°/3.2mm, 106°/11mm, d_4^{25} 1.064, n_D^{20} 1.47940, **pK²⁵ 10.94 (12.87)**. The ester is obtained by the decarbonylation of *ethyl 2-ketocyclohexylglyoxalate* (which in turn is prepared from cyclohexanone and diethyl oxalate in the presence of NaOEt). The *ethyl glyoxalate* (~250-265g, boiling at 105° to 165°/10-15mm) is mixed with Fe powder (1-3mg) and finely ground soft glass 0.5-1.0g in a Claisen flask (~500ml), and heated in a 40mm vacuum (bath temperature at 165-175°, not higher, to avoid unreacted ester from distilling) while CO [TOXIC] evolves and decarbonylated ester distils between 125-140°, requiring 1.5 to 2 hours for pyrolysis. The desired ester obtained (200-210g, 85.8%) has n_D^{25} 1.476 to 1.479. [Snyder et al. *Org Synth Coll Vol* 2 531 1943, DOI: 10.15227/orgsyn.011.0042.] The ester has FT-IR (neat) with ν_{max} at 2939.5 (CH), 1716.2 (C=O), 1658.3 (C=O ester), 1442.5 (OH enol ?), 1365.2, 1299.3, 1219.0, 1082.9 and 832.9 cm^{-1} ; 1H NMR (60MHz, $CDCl_3$, TMS) with δ at 1.30 (t, 3H, ester CH_3), 1.65 (m, ~4H, 2-methylenes), 2.25 (m, 4H 2-methylenes), 3.35 (t, 1H, partly enolised), 4.20 (q, 2H, ester CH_2) and ~12.0 (s, enolic OH); and ^{13}C NMR (15MHz, $CDCl_3$, TMS) with δ at 14.17, 14.33, 21.97, 22.42, 23.31, 27.13, 29.10, 29.98, 41.55, 57.22, 60.12, 61.04, 97.72, 169.93, 171.93, 172.71, 206.19. [*Beilstein* 10 IV 2606.] The *enol* content of *ethyl 2-oxocyclohexanecarboxylate* is higher than that of the *ethyl 2-oxocyclopentanecarboxylate* below ([611-10-9]) and varies with polarity of solvent, viz: 67% aqueous MeOH (22.2%), MeOH (45.1%), EtOH (61.7%), $CHCl_3$ (54.5%), $*C_6H_6$ (74.8%) [Kabachnik et al. *Tetrahedron* 1 317 1957, DOI:10.1016/0040-4020(57)88007-7; Gero *J Org Chem* 19 1960 1954, DOI: 10.1021/jo01377a013, see also Schreck *J Am Chem Soc* 71 1881 1949, DOI: 10.1021/ja01173a523; Buu-Hoi & Cagniant *Bull Soc Chim Fr* 10 [5] 251 1943, Lewin *Izv Akad S.S.S.R Ser fiz* 11 413 1947, *Chem Abstr* 42 3261 1948]. [*Beilstein* 10 H 601, 10 II 420, 10 III 2813, 10 IV 2606.]

Ethyl 2-oxocyclopentanecarboxylate (2-ethoxycarbonylcyclopentanone, 2-carbethoxycyclopentanone, Dieckmann Ester) [611-10-9, \pm 53229-92-8] $C_8H_{12}O_3$, **M 156.2**, **b** 79-81°/3mm, 86-87°/6mm, 102-104°/11mm, 108-111°/15mm. The ester is prepared in a dry N_2 atmosphere by adding ethyl adipate (202g, 1mole), during 2 hours, to a stirred (important to use a Hershberg stirrer, Hershberg *Ind Eng Chem, Anal Ed* 8 313 1936 DOI: 10.1021/ac50102a041) suspension of Na metal (23g, 1g.atom) in dry toluene (250ml). The reaction starts immediately and the temperature is maintained at 100-115° (oil bath) during the 2 hours and for 5 hours longer, while dry toluene is added (~750ml to 1L) so as to keep the mixture fluid and efficiently stirred, and to avoid 'caking'. The mixture is cooled to 0° and slowly added with stirring to 10% aqueous KOH (below 1°), cold H_2O being added to keep the potassium salt in solution; the toluene layer is separated, washed with cold H_2O (2 x 150ml), cold 10% aqueous KOH (adding cold H_2O to dissolve separated potassium salt), the yellow toluene solution is finally washed with cold H_2O (2 x 150ml). The combined aqueous layers, after extraction with Et_2O (250ml), are run slowly, with stirring at 0°, into cold 10% AcOH. The oily ester which separates is extracted into Et_2O (400ml), the aqueous layer is extracted with Et_2O (4 x 250ml). The combined Et_2O extracts are washed with 7% aqueous Na_2CO_3 , dried (Na_2SO_4), filtered, evaporated and the residue is distilled (**b** 79-81°/4mm) to give the *keto-ester* (100-115g, 64-74%) free from ethyl adipate. An alternative synthesis in $*C_6H_6$ with slight modification provided 79-82% yield of ester. [Pinkney *Org Synth Coll Vol* 2 116 1943, DOI: 10.15227/orgsyn.017.0030; Cornubert & Borrel *Bull Soc Chim Fr* 47 301 1930.] The *enol* content of *ethyl 2-oxocyclopentanecarboxylate* is low and varies little with polarity of solvent, viz: MeOH (4.0%), EtOH (3.9%), $CHCl_3$ (3.7%), $*C_6H_6$ (8.2%) [Kabachnik et al. *Tetrahedron* 1 317 1957, DOI: 10.1016/0040-4020(57)88007-7; Lewin *Izv Akad S.S.S.R Ser fiz* 11 413 1947, *Chem Abstr* 42 3261 1948]. The

ester has FT-IR (neat) with ν_{\max} at 2979.9 (CH), 1725.6 (C=O), 1678.8 (C=O ester), 1454.5 (OH enol ?), 1370.0, 1255.0, 1190.7, 1111.9 and 1026.3 cm^{-1} ; ^1H NMR (60MHz, CDCl_3 , TMS) with δ at 1.29 (t, $J = \sim 7\text{Hz}$, 3H, ester Me), 1.89 (m, 1H, C-3 H_{eq}), 2.24 (m, 1H, C-3 H_{ax}), 2.31 (m, 4H, C-4,5 methylenes), 3.15 (t, 1H, $J = \sim 8\text{Hz}$, C-1 H), 4.20 (q, $J = \sim 7\text{Hz}$, ester CH_2); ^{13}C NMR (15MHz, CDCl_3 , TMS) with δ at 14.18, 20.98, 27.43, 38.04, 54.77, 61.26, 169.37 and 212.25.

***N*-Phenyl 2-oxocyclopentanecarboxamide** [4874-65-1] **M 203.2, m 104°** is obtained by boiling aniline (25mmol) with ethyl 2-oxocyclopentanecarboxylate (100mmol) and pyridine (0.5ml) for 2 minutes, cooling and the solid that separates is filtered off, washed with cold EtOH, then with 4% aqueous NaOH to free it from any anil formed. The solution is filtered and acidified with dilute AcOH, the precipitate is filtered off, dried and the anilide is recrystallised from EtOH. [Barany & Pianka *J Chem Soc* 1420 1947, DOI: 10.1039/JR9470001418.] It is soluble hot EtOH and in Me_2CO . A 0.3% solution of the anilide in EtOH is used for the gravimetric determination of Be, Hg V and U [Chaudhuri & Das *Anal Chim Acta* 57 193 1971, DOI: 10.1016/S0003-2670(01)80144-6]. [Beilstein 10 IV 2602.]

1-Ethynyl-1-cyclohexanol [78-27-3] **$\text{C}_8\text{H}_{12}\text{O}$, M 124.2, m 30-33°, 32-33°, b 74°/12mm, 76-78°/17mm, 171-172°/694mm, 180°/atm, d_4^{25} 0.9734, n_D^{25} 1.4801.** Dissolve it in Et_2O , wash it with H_2O , dilute NaHCO_3 , H_2O again, dry (Na_2SO_4), filter, evaporate and distil the residue. The IR (CCl_4) has ν_{\max} at 3448 (OH), 2941 (CH), 1449-1123 and 956 cm^{-1} , and the ^1H NMR (CCl_4) has δ at 3.2 (OH), 2.5 ($=\text{CH}$), 1.70 (m 10H, CH_2) [Hasbrouck & Anderson *J Org Chem* 38 2103 1972, DOI: 10.1021/jo00951a037]. [Beilstein 6 II 100, 6 IV 348] **TOXIC.**

Eucaliptol (1,8-cineol, 1,8-epoxy-*p*-menthane, 1,3,3-trimethyl-2-oxabicyclo[2.2.2]-octane, cajeputol) [470-82-6] **$\text{C}_{10}\text{H}_{18}\text{O}$, M 154.2, m 1.3°, 1.5°, b 39-39.3°/4mm, 176-176.4°/760mm, d_4^{20} 0.9232, n_D^{20} 1.4575.** It is a major component of eucalyptus oil with a camphor-like odour [Berry *Australasian J Pharm* 203 1929, Wallach *Justus Liebigs Ann Chem* 291 342 1896, DOI: 10.1002/jlac.18962910306; Birch et al. *Tetrahedron Lett* no 3 1 1959, DOI: 10.1016/S0040-4039(01)99411-0]. Purify 1,8-cineol by dilution with an equal volume of petroleum ether, then saturate it with dry HBr. The precipitate is filtered off, washed with small volumes of petroleum ether, then cineole is regenerated by stirring the crystals with H_2O . It can also be purified via its *o*-cresol or resorcinol addition compounds. Store it over Na until required. Purify it also by fractional distillation. It is insoluble in H_2O but soluble in organic solvents. [IR: Kome et al. *Nippon Kagaku Zasshi* [*J Chem Soc Japan* (Pure Chem Sect)] 80 66 1959, *Chem Abstr* 603 1961, *Beilstein* 17 II 32, 17/I V 273.]

(+)- α -Fenchol (1R-1,3,3-trimethylnorbornan-2-ol) [1632-73-1] **$\text{C}_{10}\text{H}_{18}\text{O}$, M 154.3, m 40-43°, 47-47.5°, b 201-202°/atm, $[\alpha]_D^{20} +12.5$ (c 10, EtOH).** It is prepared by reduction of (-)-fenchone and is purified by recrystallisation from $^*\text{C}_6\text{H}_6$ /petroleum ether, or distillation, or both. The **2-carboxybenzoyl (monophthalate) derivative** has **m 146.5-147.5°** $[\alpha]_D^{20} -20.4$ (EtOH), and the **2-phenylurethane** has **m 81°**. [Beckmann & Metzger *Chem Ber* 89 2738 1956, DOI: 10.1002/cber.19560891210]. [Beilstein 6 III 288, 6 IV 278.]

(+)- Fenchone (1S-1,3,3-trimethylnorbornan-2-one) [4695-62-9] **$\text{C}_{10}\text{H}_{16}\text{O}$, M 152.2, m 5-7°, 6.1°, b 63-65°/13mm, 66°/15mm, 122°/10mm, d_4^{20} 0.9434, n_D^{20} 1.4636, $[\alpha]_D^{20} +66.9$ (neat, or in c 1.5, EtOH), $[\alpha]_{546}^{20} +60.4$ (neat).** The oily liquid is purified by distillation in a vacuum and is very soluble in EtOH and Et_2O . [Boyle et al. *J Chem Soc D* 395 1971, DOI: 10.1039/C29710000395; Hüchel *Justus Liebigs Ann Chem* 549 186 1941, DOI: 10.1002/jlac.19415490106, (\pm)-isomer: Braun & Jacob *Chem Ber* 66 1461 1933, DOI: 10.1002/cber.19330661005.] It forms two *oximes*, the **cis-oxime** has **m 167°** (crystallises from petroleum ether) $[\alpha]_D^{20} +46.5$ (c 2, EtOH), the ***O*-benzoyloxime** has **m 81°**, $[\alpha]_D^{18} +49$ (EtOH), and the **oxime-HCl** has **m 136°(dec)**. The **trans-oxime** has **m 123°** (from petroleum ether) $[\alpha]_D^{18} +148$ (c 2, EtOH) and the ***O*-benzoyloxime** has **m 125°** $[\alpha]_D^{20} +128.5$ (c 2, EtOH) [Hüchel *Justus Liebigs Ann Chem* 549 186 1941, DOI: 10.1002/jlac.19415490106; Hüchel & Sachs *Justus Liebigs Ann Chem* 498 166 1932, DOI: 10.1002/jlac.19324980109]. [Beilstein 7 III 212, 7 IV 212.] **(-)- Fenchone (1R-1,3,3-trimethylnorbornan-2-one)** [7787-20-4] has **m 5.2°, b 67.2°/10mm, 191-195°/atm, d_4^{20} 0.9484, n_D^{20} 1.4630, $[\alpha]_D^{20} -66.8$ (neat).** Purification is as for the (+)-enantiomer above and should have the same physical properties except for opposite optical rotations. UV has λ_{\max} 285nm (ϵ 12.29). [Braun & Jacob *Chem Ber* 66 1461 1933, DOI: 10.1002/cber.19330661005; UV: Ohloff et al. *Chem Ber* 90 106 1957, DOI: 10.1002/cber.19570900121.]

5S-(-)-Gabaculine [5S-(-)-5-aminocyclohexa-1,3-diene-1-carboxylic acid, 5-amino-2,3-dihydrobenzoic acid] [*S*- 59556-29-5; 87980-11-8 unspecified stereochemistry] $C_7H_9NO_2$, **M 139.2**, amorphous white powder, $pK_{Est(1)} \sim 3.75$, $pK_{Est(2)} \sim 8.86$, $[\alpha]_D - 454$ (c 1, H_2O). The 5-*S*-aminoacid has been isolated from the culture filtrates of *Streptomyces toyocaensis* subsp 1039, and purified by chromatography through a cellulose column and eluted with *n*-BuOH/ H_2O (100:7); the physiologically active fractions are concentrated to a small volume, the precipitate is collected, dissolved in a little H_2O and lyophilised. The white powder is homogeneous on silica gel TLC plates (60 F-254, Merck, eluting with *n*-BuOH/880 NH_3/H_2O ::6:2:1) with R_F 0.65, and on cellulose plates (F, Merck, eluting with *n*-BuOH/AcOH/ H_2O ::3:1:1) with R_F 0.53. It has UV (H_2O) with λ_{max} at 275nm (ϵ 8600); the IR has ν_{max} at $1650cm^{-1}$ (C=C-C=O); 1H NMR has δ at 6.82 (2-H, d, A of ABX), 6.47 (3-H, q, X of ABX), 6.06 (4-H, q, B of ABX) J_{cis} for BX = 9.5Hz, 4.11 (5-H, d, coupled with 4-H and 6-H), and 2.77 (geminal 6-H, 6-H, q), assignments are deduced by decoupling technique, i.e. irradiating 2-H, 5-H and 6-H respectively, and MS showed abundant peaks as those of benzoic acid except for the molecular ion and base peak at m/e 94 ($M^+ - COOH$). [Kobayashi et al. *Tetrahedron Lett* 537 1976, DOI: 10.1016/S0040-4039(00)77904-4; *FEBS Lett* 76 207 1977, DOI: 10.1016/0014-5793(77)80153-1.] It is a conformationally restricted analogue of γ -aminobutyric acid (GABA), an irreversible and potent inhibitor of γ -aminobutyrate (GABA) aminotransferase [EC 2.6.1.19] at sub μM concentrations, a GABA reuptake inhibitor, and as a drug it increases GABA levels in the brain. [Allan, Johnston and Twitchin *Neuroscience Lett* 4 51 1977, DOI: 10.1016/0304-3940(77)90124-0.] It inhibits the formation of chlorophyll in plants [Gardner & Gordon *Plant Physiol* 77 540 1985, <http://www.jstor.org.virtual.anu.edu.au/stable/4269178>].

(\pm)-Gabaculine [59556-18-2] has been synthesised using mild reagents and reaction conditions as well as judicious protection and deprotection of the NH_2 and CO_2H groups to avoid aromatisation of the cyclohexane ring. Final purification is achieved by dissolving the final product in the minimum volume of H_2O applying onto a column of Bio-Rad AG 11 A8 ion-retardation resin and eluting with H_2O . Alternatively, chromatography through an ion-exchange column (SP Sephadex C-25) and eluting with aqueous NH_4OH gave pure amino-acid. The eluent is visualised with UV light and the fractions which gave positive ninhydrin tests are collected, lyophilised and should have all the above spectroscopic (IR, UV NMR, MS), TLC and amino acid analysis properties. The (\pm)-amino-acid can be recrystallised by dissolving it in the minimum volume of H_2O , adding 1 to 2 drops of H_2O and placing in a closed chamber over Me_2CO (whereby H_2O distils into the Me_2CO) and after 24 hours crystals appear, and are recrystallised in the same manner to give pure crystals with **m 196-197°** (identical with that from other syntheses). Store it below 5°. Its ^{13}C NMR (in D_2O + Na_2HPO_4 , and proton decoupled) has δ 176.0, 133.0, 128.8, 128.3, 128.1, 54.4 and 29.9, and HRMS has m/z 139.0635 (calc for $C_7H_9NO_2$ is 139.0635). **(\pm)-Gabaculine hydrochloride** [59556-17-1] $C_7H_9NO_2 \cdot HCl$, **M 175.6**, has **m 203° (dec), 203-306° (dec), 198-200° (dec)**, (*note*: not very different from the free amino-acid), can be purified by recrystallising from dilute HCl (in needles **m 198-200°**), by dissolving the amino-acid in ice cold MeOH (saturated with dry HCl gas), the solvent is removed under high vacuum, and the white solid is recrystallised from Me_2CO containing a little MeOH [**m 195-199° (dec)**]. Alternatively, the (\pm)-hydrochloride is prepared in MeOH by adding dry HCl and recrystallising by dissolving in the least volume of H_2O , adding 1 to 2 drops of H_2O , evaporating over Me_2CO in a closed chamber and collecting the crystals and drying them *in vacuo* [**m 197-199°**]. Store it below 5°. All salts were undepressed with authentic material. [Kobayashi et al. *Tetrahedron Lett* 537 1976, DOI: 10.1016/S0040-4039(00)77904-4; Singer & Sharpless *J Org Chem* 43 1448 1978, DOI: 10.1021/jo00401a034; Trost & Keinen *J Org Chem* 44 3451 1979, DOI: 10.1021/jo01334a001.] *Note* that the synthetic (\pm)-amino-acid has half the inhibitory GABA-aminotransferase activity compared with the natural (-)-amino-acid [Kobayashi et al.].

Gibberillic acid A₃ (gibberillin A₃, GA₃) [77-06-5] $C_{19}H_{22}O_6$, **M 346.4**, **m 233-235° (dec)**, $[\alpha]_{546}^{20} +92$ (c 1, MeOH), $[\alpha]_D^{20} +93$ (c 0.5, MeOH), **pK 4.0**. It crystallises from EtOAc, EtOAc/petroleum ether, MeOH/petroleum ether or Me_2CO /petroleum ether. The *methyl ester* $C_{20}H_{24}O_6$, **M 360.4**, forms needles from $*C_6H_6/MeOH$ with **m 209-210°** and $[\alpha]_D^{20} +75$ (c 0.5, MeOH). It belongs to a class of plant growth substances. [Cross *J Chem Soc* 3022 1960, DOI: 10.1039/JR9600003022; Beilstein 18 III/IV 6533, 18 V/9 6533.]

(-)-Guaiol {3,8-dimethyl-5-(α -hydroxyisopropyl)- Δ^9 -octahydroazulene, (3*R*,6*S*,10*S*)-6,10, α , α -tetramethyl-bicyclo[5.3.0]deca-1(7)-ene-3-methanol} [489-86-1] $C_{15}H_{26}O$, **M 222.4**, **m 91°, 91-93°, b 148°**

10mm, 165°/17mm, 288°/atm (slight dec), d_{20}^{100} 0.9074, n_D^{100} 1.4716, $[\alpha]_D^{20}$ -30 (c 4, EtOH), $[\alpha]_D^{20}$ -36.7 (c 9, CHCl₃). It occurs in Guaiacum wood oil, and wood oils from *Zygophyllaceae* spp. Purify it by distilling under high vacuum and/or recrystallise from EtOH or Me₂CO by cooling to *ca* -10°. The **3,5-dinitrobenzoate** has **m 137-137.5°** (from EtOH or Me₂CO). [Plattner & Lemay *Helv Chim Acta* **23** 897 1940, DOI: 10.1002/hlca.194002301115]. The synthetic **racemate** [33496-08-1], purified by gas chromatography from the (±)-7-epimer, has **m 55-60°**, IR has ν_{\max} (film) at 3.00, 6.90, 7.38, 7.67, 7.88, 8.04, 8.18, 8.30, 8.52, 8.70, 8.80, 10.05, 10.33, 10.81, 11.00, 11.38 and 12.20 μm ; and ¹H NMR (CDCl₃, TMS) has δ at 1.18 (methyls), 0.98 (CH₃ d, *J* = 7.5 Hz), 0.96 (CH₃ d, *J*=7 Hz) [Marshall & Greene *J Org Chem* **37** 982 1972, DOI: 10.1021/jo00972a013]. The **methyl ether C₁₆H₂₈O**, has **b 142°/9mm, d_4^{25} 0.9332, $n_D^{18.5}$ 1.4896, $[\alpha]_D^{20}$ -31.8 (c 4, EtOH).** [Ruzicka & Haagen-Smit *Helv Chim Acta* **14** 1122 1931, DOI: 10.1002/hlca.19310140521; Absolute Configuration: Eisenbraun et al. *J Am Chem Soc* **82** 3648 1960, DOI: 10.1021/ja01499a045; *Beilstein* **6** III 412.]

1,2,3,4,5,6-Hexachlorocyclohexane [α -319-84-6 α -BHC, γ -58-89-9] C₆H₆Cl₆, M 290.8, m 158° (α -), 312° (β -), 112.5° (γ -isomer). Crystallise it from EtOH. Purify it also by zone melting. **CANCER AGENT, TOXIC.** [α : *Beilstein* **1** H 23, γ : *Beilstein* **5** I 8, many isomers : *Beilstein* **5** III 41, **5** IV 55.]

1,2,3,4,5,5-Hexachlorocyclopenta-1,3-diene [77-47-4] C₅Cl₆, M 272.8, b 80°/1mm, 83-84°/3mm, 234°/atm, d_4^{25} 1.702, n_D^{25} 1.5628. Dry the diene with MgSO₄, filter, and distil it under vacuum in a nitrogen atmosphere. **Irritates skin and eyes, HIGHLY TOXIC.** [McBee et al. *J Am Chem Soc* **77** 4375 1955, DOI: 10.1021/ja01621a056; UV spectra: Idol et al. *J Org Chem* **20** 1743 1955, DOI: 10.1021/jo01364a023; *Beilstein* **5** III 308, **5** IV 381.]

Hexahydromandelic acid [*R*-(-)- 53585-93-6, *S*-(+)- 61475-31-8] C₈H₁₄O₃, M 158.2, m 127-129°, 128-129°, 129.7°, $[\alpha]_D^{20}$ (-) and (+) 25.5 (c 1, AcOH) and $[\alpha]_D^{20}$ (-) and (+) 13.6 (c 7.6, EtOH). It forms hexagonal clusters on recrystallisation from CCl₄ or Et₂O. [Wood & Comley *J Chem Soc* 2630 1924, DOI: 10.1039/CT9242502630; Lettré et al. *Chem Ber* **69** 1594 1936, DOI: 10.1002/cber.19360690708]. The **racemate** has **m 137.2-137.6° (134-135° also reported)** [Smith et al. *J Am Chem Soc* **71** 3772 1949, DOI: 10.1021/ja01179a056]. [*Beilstein R-* **10** II 5, **10** III 25; *S-* **10** II 6.]

Hexamethyldewarbenzene (HMDB, 1,2,3,4,5,6-hexamethyl-bicyclo[2.2.0]hexa-2,5-diene) [7641-77-2] C₁₂H₁₈, M 162.3, m 7.5°, b 60°/20mm, ~152°/760mm, d_4^{20} 0.8125, n_D^{20} 1.4480. HMDB is obtained in ~80% yield when 2-butyne (dimethylacetylene, flammable gas b 27°/atm, [503-17-3]) in *C₆H₆ or CH₂Cl₂ containing anhydrous AlCl₃ is stirred at 35° for 5-7 hours. The brown reaction mixture is poured onto crushed ice, washed with dilute aqueous NaOH, and HMDB is isolated from the organic phase by fractional vacuum distillation. **Hexamethylbenzene (m 165-166°, [87-85-4])** is a by-product (~12-18%) together with a mixture (~2%) of *syn*- (m 127°) and *anti*- (m 196°) octamethyltricyclo[4.2.0.0^{2,5}]octa-3,7-diene, and octamethylcyclooctatetraene (m 113°). Although longer reaction times increase the conversion of 2-butyne, they lower the yields of HMDB in favour of the other by-products. It is fairly thermally stable with half-life time conversions to hexamethylbenzene of 105 hours/120°, 5.5 hours/140°, and 2.1 hours/150°. It is best to store it at low temperature and away from light as radiation from a low pressure UV lamp converts it mainly to hexamethylbenzene with ~20-25% of the valence isomer hexamethylprismane (m 91°). The UV spectrum exhibits tail end absorption from 220 to 250nm; the FT-IR (neat) has ν_{\max} at 2950.0, 1683.5, 1439.5, 1368.8, 1275.4, 1222.2, 1064.8, 736.2 and 490.7 cm⁻¹; the ¹H NMR (60MHz, CDCl₃, TMS) has δ at 1.08 (s, 6H, 1,4-(CH₃)₂) and 1.69 (s, 12H, 2,3,5,6-(CH₃)₄), and the ¹³C NMR (15MHz, CDCl₃, TMS) has δ at 10.06 (1,4-Me carbons) and 11.22 (2,3,5,6-Me carbons), 55.68 (2 saturated carbons) and 144.09 (4 olefinic carbons). [Schäfer & Hellmann *Angew Chem Int Ed* **6** 518 1967, DOI: 10.1002/anie.196705181.] Purify it also by passing it neat through alumina or in *C₆H₆ or CH₂Cl₂ solution. [Traylor & Miksztal *J Am Chem Soc* **109** 2770 1987, DOI: 10.1021/ja00243a033].

α -Humulene [α -caryophyllene (now an obsolete name, see above), *trans-trans-trans*-2,6,6,9-tetramethyl-1,4,8-cycloundecatriene = 1,1,4,8-tetramethylcycloundeca- *trans-trans-trans*-3,7,10-triene] [6753-98-6] C₁₅H₂₄, M 204.4, b 106-107°/5mm, 123°/10mm, 166-168°/atm, d_4^{20} 0.967, d_4^{25} 0.8883, n_D^{20} 1.4956, n_D^{25}

1.5017, n_D³⁰ 1.5004. It is a large component of the *mono-cyclic* sesquiterpenes in the essential oils of hops [Chapman *J Chem Soc* 54, 780 1895, DOI: 10.1039/CT8956700054; DOI: 10.1039/CT8956700780 and 785 1928, DOI: 10.1039/JR9280000785], is present in clove oil, Egyptian Hashish [Simonsen & Todd *J Chem Soc* 188 1942, DOI: 10.1039/JR9420000188], and subsequently found in many other sources [Duessen *J prakt Chem* **83** 483 1911, DOI: 10.1002/prac.19110830134]. Originally isolated by steam distillation of hops and the oil fractionated preferably *in vacuo*. Later purification was further carried out by column chromatography on Al₂O₃ or silica gel (hexane or petroleum ether) to remove polar by-products [Kitigawa et al. *J Am Chem Soc* **99** 3864 1977, DOI: 10.1021/ja00453a069]. It is best to purify it first by fractionation using a Podbielniak still and at a reflux ratio of ~50, or a 1.25 m Dixon ring-packed still under a reflux ratio of ~35. The sesquiterpene should be optically inactive (most purified samples from natural sources have a very small optical rotation due to small amounts of bicyclic impurities, e.g. caryophyllenes — see above). Pure optically inactive material is best obtained *via* its AgNO₃ adduct. After distillation, it is diluted with light petroleum (b 40°-60°) and stirred with an equal volume of 50% aqueous AgNO₃ (w/w). Colourless needles of the **adduct** separate almost immediately with evolution of heat. The solid is filtered after standing, washed with light petroleum and dried *in vacuo*. Upon recrystallisation from aqueous EtOH, the **adduct** has **m 175-175.5°(dec, capillary tube), 179°(copper block)** and analyses for **C₁₅H₂₄· 2 AgNO₃**. It is stable on storage in the dark at 5°. X-ray crystallography confirmed the humulene structure and demonstrated that the two Ag⁺ ions are each complexed to the 7-8 and 10-11 double bonds [McPhail & Sims *J Chem Soc (B)* 112 1966, DOI: 10.1039/J29660000112].

Regeneration of the hydrocarbon is achieved by refluxing the adduct (e.g. 21.4g) with H₂O (e.g. 300 mL) under an oil trap (to avoid loss of hydrocarbon through the top of the condenser), cool, extract with Et₂O, dry over CaCl₂, or Na₂SO₄, filter, evaporate and fractionate through a still containing 1 m of Bower-Cooke packing (reflux ratio ~10) collecting the fraction with b 123°/10mm. *Alternatively*, regenerate by dissolving the adduct (e.g. 5g) in excess saturated NH₃ which liberates the hydrocarbon that is extracted with Et₂O as previously to yield pure α-caryophyllene (2g). The UV shows end absorption with ε 3050 at λ = 220 nm. FTIR (film cm⁻¹) ν_{max}: 2956, 2925, 2866, 2853, 1684, 1446, 1384, 1362, 1210, 1176, 966 and 822; ¹H NMR (CDCl₃, 500 MHz) δ: 5.57 (dt, *J* = 15.9, 7.4 Hz, 1H), 5.14 (d, *J* = 15.9 Hz, 1H), 4.93 (br t, *J* = 6.3Hz, 1H), 4.85 (br t, *J* = 7.5Hz, 1H), 2.49 (d, *J* = 7.4Hz, 2H), 2.05-2.11 (m, 4H), 1.89 (d, *J* = 7.4Hz, 2H), 1.62 (s, 3H), 1.41 (s, 3H), 1.04 (s, 6H); ¹³C NMR (CDCl₃, 100 MHz) δ: 141.0, 139.2, 133.2, 127.7, 125.8, 125.0, 42.0, 40.4, 39.7, 37.3, 23.3, 17.9 and 15.1. This AgNO₃ procedure can be used to obtain pure hydrocarbon from ‘commercial caryophyllene’ which contains ~13% of the hydrocarbon.

The white **nitrosochloride**, **C₁₅H₂₄ClNO**, [-C(NO)—C(Cl)-] (2.56g), prepared from α-caryophyllene (1.99g) and ethyl nitrite and concentrated HCl in MeOH-AcOH, crystallises from CHCl₃-EtOH with **m 172°(dec)** [also reported: 173-174°, 176°, 177°]. The **nitrolpiperidide**, **C₁₅H₂₄NONC₅H₁₀**, is prepared from the **nitrosochloride** and piperidine at ~100° and crystallises from EtOH, to form colourless prisms with **m 148-149°** [also reported: 153°, 154-155°]. The **nitrolbenzylamine**, **C₁₅H₂₄NONHCH₂C₆H₅**, is similarly prepared from the **nitrosochloride** and benzylamine, has **m 136-137°** [also reported: 136°]. [Hildebrand & Sutherland *Aust J Chem* **14** 272 1961, DOI: 10.1071/CH9610272; Chapman *J Chem Soc* 785 1928, DOI: 10.1039/JR9280000785; Clemo & Harris *J Chem Soc* 22 1951, DOI: 10.1039/JR9510000022; *synthesis* Kitigawa et al. *J Am Chem Soc* **99** 3864 1977, DOI: 10.1021/ja00453a069; *biomimetic synthesis* Corey et al. *Tetrahedron Lett* **34** 3675 1993, DOI:10.1016/S0040-4039(00)79198-2; for *X-ray structure of AgNO₃ adduct* see McPhail & Sim *J Chem Soc (B)* 112 1966, DOI: 10.1039/J29660000112; *Beilstein* **5** III 1070.]

The **nitrosate**, **C₁₅H₂₄N₂O₄**, [-C(NO)—C(ONO₂)-] prepared from amyl nitrite and AcOH, then HNO₃-AcOH, crystallises from *C₆H₆, and has **m 161° to m 165°** depending on heating rate [reported also: 163°]. The **nitrosite**, **C₁₅H₂₄N₂O₃**, [-C(NO)—C(ONO)-] separates as blue crystals when equal volumes of α-humulene and light petroleum are shaken with concentrated aqueous NaNO₂ with cooling, followed by slow addition of glacial acetic acid (volume equal to humulene used) with shaking. The organic green-blue layer is separated, and the blue crystals of the nitrosite are collected and crystallised from EtOH to give blue needles of the **dinitrosite compound** (dimer ?) with **m 120-121°**. Boiling a solution of the blue nitrosite in EtOH for several hours slowly caused loss of colour which on cooling deposited almost colourless needles with a blue tinge of the isomeric (tautomeric) **bis-oxime** with **m 165-168°**. [Deussen *J Prakt Chem* **83** 483 1911, DOI: 10.1002/prac.19110830134; Chapman *J Chem Soc* 785 1928, DOI: 10.1039/JR9280000785.]

Humulene trioxide, **C₁₅H₂₄O₃**, MW 252.3, prepared by oxidation of humulene (3g) in CHCl₃ (220 ml) for 3 days at room temperature with perbenzoic acid (1 ml of CHCl₃ containing 64mg of the acid), followed by shaking with cold 5% aqueous NaOH, H₂O to neutrality, drying, evaporating, and distilling to give a viscous oil

which formed crystals on trituration with EtOH-hexane. Recrystallisation from the same solvent gives pure **trioxide** with **m 122.5°** [Sorm et al. *Coll Czech Chem Commun* **14** 699 1949, DOI: 10.1135/ccccc19490693]. Alternatively, a sample of humulene, extracted with other terpenes in the turpentine oil from *Araucaria cunninghamii*, and purified by gas chromatography on 'Apiezon M on Embecel' at 170° with N₂ as carried gas, and the fraction from relevant peak converted into the AgNO₃ complex from which pure *humulene* was recovered, was oxidised with monoperphthalic acid to *humulene trioxide*, **m 121-122°** (hexane-EtOH) [Gallagher & Sutherland *Aust J Chem* **13** 367 1960, DOI: 10.1071/CH9600367; *Beilstein* **19** IV 4748]. Catalytic reduction of α -humulene in glacial acetic acid using Adam's Pt results in the absorption of 3 mols of H₂ (at 19°/747.6mm) to produce **Humulane (hexahydrohumulene, 1,1,4,8-tetramethyl cycloundecane) C₁₅H₃₀, M 210.5**, which has **b 120-122°/13mm, d₄²⁰ 0.8637, n_D²⁰ 1.4723** [Sorm et al. *Coll Czech Chem Commun* **14** 669 1949, DOI: 10.1135/ccccc19490693].

β -Humulene (trans,trans-8-methylene-1,4,4-trimethylcycloundeca-1,5-diene) [116-04-1] C₁₅H₂₄, M 204.4, d₄²⁰ 0.8905, n_D²⁰ 1.5014. The *mono-cyclic* sesquiterpenic fractions from hops, oil of cloves or other sources (see preceding entry) contain a high proportion of α -humulene which cannot be cleanly separated by fractional distillation from the smaller amounts of the isomeric β -humulene. The oil from the leaves of *Lindera strychnifolia* (F.) WILL gave very pure humulene. Column chromatography on alkaline Al₂O₃ (grade I, according to Brockmann) and eluting with light petroleum afforded better separation of the isomers than fractional distillation. The α -isomer elutes first followed by a mixture of α - and β -isomers followed by enriched β -isomer. The fractionation can be followed by evaporating aliquots and examining the IR spectra (film) of the residues. Unlike the α -isomer, the β -isomer has a medium-intense deformation vibration band characteristic of an exocyclic double bond at ν_{\max} 888-889 cm⁻¹. A band at 968 cm⁻¹ characteristic of *trans* — CH=CH— double bonds is present in both isomers. The exocyclic methylene double bond is further confirmed by bands at 1645 and 3075 cm⁻¹. Mixtures of isomers have the two sets of bands of varying intensities according to their ratios. Repeated chromatography of fractions can improve separation. However, pure α -humulene isolated *via* its AgNO₃ adduct and regenerated, did not possess an exocyclic double but repeated chromatography on Al₂O₃ as above caused isomerisation and enriched it in the β -isomer with the exocyclic methylene group. Catalytic reduction of the isomers provided the same *humulane* (see preceding entry). [Benesová, Herout and Sorm et al. *Coll Czech Chem Commun* **26** 1832 1961, DOI: 10.1135/ccccc19611832, Sorm et al. *Coll Czech Chem Commun* **14** 669 1949, DOI: 10.1135/ccccc19490693.] [*Beilstein* **5** III 1070.] **γ -Humulene (isohumulene, 5-methylene-1,8,8-trimethylcycloundeca-cis-1-, trans-6-diene) [26259-79-0] C₁₅H₂₄, M 204.4, d₄²⁰ 0.9, is a second humulene with an exocyclic methylene group. [*Beilstein* **5** III 1070.]**

Humulon [α -lupulic acid, 6R-3,5,6-trihydroxy-4,6-bis(3-methyl-2-butenyl)-2-(3-methyl-1-oxobutyl)-2,4-cyclo-hexadien-1-one [26472-41-3] C₂₁H₃₀O₅, M 362.5, m 65-66.5°, [α]_D²⁰ -232 (*C₆H₆), [α]_D²⁶ -212 (c 6.5, 95% EtOH), -124 (abs EtOH), -162 (dioxane). Originally, it was isolated from the mother liquors of the petroleum ether extract of hop cones, after crystallisation of *lupulone* (see below) as its *o*-phenylenediamine complex. After evaporation of the filtrate, the residue was recrystallised three times from *C₆H₆-petroleum ether (1:2) to constant optical rotation. Also recrystallise *humulon* from Et₂O. It dissolves slightly in hot H₂O but precipitates on cooling. It has λ_{\max} nm (ϵ) at 237 (13,760) and 282 (8,330) in EtOH. It is an antimicrobial agent in commercial hops. **Note:** variations in the specific rotations may well be caused by slight variation in keto-enol tautomeric ratios. [Wöllmer *Chem Ber* **49** 780 1916, DOI: 10.1002/cber.19160490185; Carson *J Am Chem Soc* **73** 4652 1951, DOI: 10.1021/ja01154a046; *Beilstein* **8** II 537, **8** III 4034, **8** IV 3410.]

(\pm)-1-Hydroxyadamantan-4-one (Kemantone, 5-hydroxy-2-adamantanone, 1-hydroxytricycl[3.3.1.1^{3,7}]-decan-4-one) [20098-14-0] C₁₀H₁₄O₂, M 166.2, m 319-322°. The hydroxyketone crystallises from CCl₄ and sublimes unchanged. It can be obtained by stirring 2-hydroxyadamantane (6.1g, see [700-57-2] above) in 70% H₂SO₄ (50ml) at 90° for 2 hours followed by neutralising with 50% NaOH to give a mixture of adamantane (44.5%) and adamantanone (17.5%), and from the aqueous solution a mixture of 1,4-dihydroxyadamantane and 1-hydroxyadamantan-4-one that are combined and oxidised with CrO₃ in 0.5N H₂SO₄/Me₂CO to give 1-hydroxyadamantan-4-one (0.86g, m 315-321°, 98% purity by GLC,), which after recrystallisation from CCl₄ (recovery 76%) has **m 319-322°, m/z** found 166.0994, C₁₀H₁₄O₂ requires M = 166.0994; ¹H NMR has δ at 2.62 (2H, protons α to the C=O), 2.35 (1H, tertiary proton at C-7), 1.98 (10H, secondary protons), 1.89 (1H, -OH); and IR (CH₂Cl₂) has ν_{\max} at 3590 (m, free OH), 2920 (s) and 2855 (m, CH), 1720 (vs, C=O), 1110 (s), 1090

(m), 1066 (m), 1055 (s), 971 (m) and 926 (s) cm^{-1} . **1-Hydroxyadamantan-4-one oxime**, obtained in the usual way, crystallises from EtOAc in colourless crystals with **m** 157-160°; IR (KBr) with ν_{max} at 3300 (vs, OH), 2925 (s) and 2855 (m) (CH), 1660 (w, C=N), 1115 (s), 1095 (s), 1067 (m), 925 (s) and 848 (m) cm^{-1} . Reduction with $\text{LiAlH}_4/\text{THF}/65^\circ/20\text{hrs}$ provided the amine which was converted to the **4-aminoadamantan-1-ol hydrochloride** [*trans*: 62075-23-4] $\text{C}_{10}\text{H}_{17}\text{NO} \cdot \text{HCl}$, **M** 203.7, **m** 356-358°, on treatment with HCl and recrystallising twice from MeOH (and addition of Et_2O) [Geluk & Schlatmann *Tetrahedron* **24** 5369 1968, DOI: 10.1016/S0040-4020(01)96330-6.]. **Alternatively**, adamantanone (12.0g, 80mmol) is stirred into 100% HNO_3 (100ml) in an ice-water bath where the solid dissolves immediately, the temperature rises to 13-15°, and the colour changes to orange-brown. The mixture is allowed to stand for 70 hours, then is heated at 60° for 90min, the acid is distilled off under vacuum on a steam bath and the yellow oil solidified as the HNO_3 -adduct. Water (40ml) and 96% H_2SO_4 (15ml) is added and the resultant clear solution is heated on a steam bath for 1 hour when all the nitrous fumes evolved (FUME HOOD). The cooled solution is extracted with 2:1 petroleum ether (b 40-60)/ Et_2O (50ml x 2) to remove unreacted adamantone (0.07g), the acid layer is neutralised with 30% aqueous NaOH and while warm is extracted with CH_2Cl_2 (50ml x 2). The extract is washed with brine (25ml) and evaporated *in vacuo*, the crude residue (10.75g) is dissolved in CH_2Cl_2 (15-20ml) and petroleum ether (b 40-60) is added until no further precipitation occurs. The hydroxyadamantanone is collected and dried (**m** 315-318°, 9.65g, 77% yield, and has 3% of adamantane-2,6-dione by GLC). Recrystallisation from CCl_4 gave pure product with **m** 319-312°. [Geluk *Synthesis* 374 1972, DOI: 10.1055/s-1972-21884; Noble et al. *J Org Chem.* **48** 1099, 1983, DOI: 10.1021/jo00155a034.] For a rapid conversion of adamantanone to 5-hydroxyadamantan-4-one in 60% yield on a multigram scale see Srivastava & le Noble *Synth Commun* 14 65 1984, DOI:10.1080/00397918408060865]. The (\pm)-form is a T-cell suppressor, and a lymphocyte and antibody stimulant in mice.

1-Acetoxyadamantan-4-one [63382-10-5] $\text{C}_{12}\text{H}_{16}\text{O}_3$, **M** 208.2, **m** 60°, is obtained by $\text{CrO}_3/\text{Ac}_2\text{O}$ oxidation of adamantanone for 10 days at 20°, and purified through an Al_2O_3 column (neutral, activity III) and eluting with petroleum ether. It has IR (melt) with ν_{max} at 2870, 1730, 1235 and 1060 cm^{-1} ; ^1H NMR (CDCl_3) has δ at 2.46 (2H α to C=O), 2.40 (7H), 2.00 (4H + 3H of OAc) ppm; and the ^{13}C NMR (C_6D_6) has δ at 212.9 (C=O) 169.4 (OAc), 77.5 (a), 47.2 (b), 41.3 (c), 40.0 (d), 38.0 (e), 30.2 (f), 22.0 (OAc) ppm [Morat & Rassat *Tetrahedron Lett* **20** 4409 1979, DOI: 10.1016/S0040-4039(01)86603-X].

1-Hydroxymethyladamantane (1-adamantanemethanol) [770-71-8] $\text{C}_{11}\text{H}_{18}\text{O}$, **M** 166.3, **m** 115°, 114-117°. Dissolve the adamantane in Et_2O , wash it with aqueous 0.1N NaOH and H_2O , dry over CaCl_2 , evaporate and recrystallise the residue from aqueous MeOH. [Stetter et al. *Chem Ber* **92** 1629 1959, DOI: 10.1002/cber.19590920722; *Beilstein* **6** IV 400.]

N-Hydroxy-5-norbornene-2,3-dicarboxylic acid imide [21715-90-2] $\text{C}_9\text{H}_9\text{NO}_3$, **M** 179.2, **m** 165-166°, 166-169°, **pK_{Est}** ~6. Dissolve the imide in CHCl_3 , filter, evaporate and recrystallise from EtOAc. The IR (nujol) has ν_{max} at 1695, 1710 and 1770 (C=O), and 3100 (OH) cm^{-1} . The **O-acetyl** derivative has **m** 113-114° (from EtOH) with IR bands at ν_{max} 1730, 1770 and 1815 cm^{-1} only, and the **O-benzoyl** derivative has **m** 143-144° (from propan-2-ol or $\ast\text{C}_6\text{H}_6$). [Bauer & Miarka *J Org Chem* **24** 1293 1959, DOI: 10.1021/jo01091a035; Fujino et al. *Chem Pharm Bull Jpn* **22** 1857 1974, DOI:org/10.1248/cpb.22.1857]. [*Beilstein* **21/10** V 188.]

α -Ionone (*trans*-+) [4-(2,6,6-trimethyl-2-cyclohexenyl)-3-buten-2-one] [127-41-3] $\text{C}_{13}\text{H}_{20}\text{O}$, **M** 192.3, **b** 86-87°/1.9mm, 131°/13mm, 259-263°/atm, d_4^{20} 0.929, n_D^{20} 1.5497, $[\alpha]_D^{20}$ +401 (neat) +415 (EtOH). Purify α -ionone through a spinning band fractionating column. The **semicarbazone** has **m** 157-157.5° (from EtOH) and $[\alpha]_D^{20}$ +433 (c 4, $\ast\text{C}_6\text{H}_6$). [Naves *Helv Chim Acta* **30** 769 1947, DOI: 10.1002/hlca.19470300310; CD: Ohloff et al. *Helv Chim Acta* **56** 1874 1973, DOI: 10.1002/hlca.19730560609; Buchecker et al. *Helv Chim Acta* **56** 2548 1973, DOI: 10.1002/hlca.19730560739; *Beilstein* **7** H 168, **7** III 640, **7** IV 363.]

β -Ionone [4-(2,6,6-trimethyl-1-cyclohexenyl)-3-buten-2-one] [79-77-6] $\text{C}_{13}\text{H}_{20}\text{O}$, **M** 192.3, **b** 150-151°/24mm, d_4^{20} 0.945, n_D^{20} 1.5211, $\epsilon_{296\text{nm}}$ 10,700. **Note:** this isomer is not chiral. Convert β -ionone to the **semicarbazone** (**m** 149°) by adding semicarbazide hydrochloride (50g) and potassium acetate (44g) in water (150ml) to a solution of β -ionone (85g) in EtOH. (More EtOH is added to redissolve any β -ionone that precipitates.) The semicarbazone crystallises on cooling in an ice-bath and is recrystallised from EtOH or 75% MeOH to constant **m** (148-149°). The semicarbazone (5g) is shaken at room temperature for several days with

petroleum ether (20ml) and M H₂SO₄ (48ml); then the petroleum ether layer is washed with water and dilute aqueous NaHCO₃, dried and the solvent is evaporated. The β -ionone is distilled under vacuum. (The customary steam distillation of β -ionone semicarbazone did not increase the purity.) [Young et al. *J Am Chem Soc* **66** 855 1944, DOI: 10.1021/ja01234a001]. [Beilstein **7** H 167, **7** I 109, **7** II 140, **7** III 634, **7** IV 361.]

ψ -Ionone [*pseudo-ionone*; **6,10-dimethyl-3,5,9-undecatrien-2-one**] [141-10-6] C₁₃H₂₀O, M 192.3, b 114-116°/2mm, 124-126°/4mm, 143-145°/12mm, 167-168°/23mm, d₄²⁰ 0.8984, n_D²⁰ 1.53346. It is the precursor of the above ionones. It is purified by mixing the ionone with an aqueous solution containing sodium bisulfite (2-3 moles, not more) and refluxing for a few hours whereby it forms a water soluble monosulfonic acid complex, the impurities do not react and can be extracted with Et₂O. The solution is then treated with aqueous sodium hydroxide and the ionone extracted into Et₂O or *iso*-Pr₂O, dried and finally distilled under a vacuum. The ¹³C NMR (22.63MHz, CDCl₃, proton-decoupled, ¹³C signal of internal TMS is the reference) has δ at 17.7 (1-Me_{cis}), 25.7 (1-Me_{trans}), 132.2 (1-C), 132.3 (2-C), 26.4 (3-C), 40.5 (4-C), 150.8 (5-C), 17.5 (5-Me), 123.8 (6-C), 139.4 (7-C), 128.5 (8-C), 198.3 (9-C=O) and 27.4 (9-Me), showing all-*trans* double bonds and can be used as a check of purity [Englert *Helv Chim Acta* **58** 2367 1975, DOI: 10.1002/hlca.19750580817]. The **2,4-dinitrophenylhydrazone** has m 146-147° (from EtOH). [Russell & Kenyon *Org Synth Coll Vol* **3** 747 1955, DOI: 10.15227/orgsyn.023.0078; Wilkinson et al. US Pat 3161684 A Dec 1964, for synthesis see Ohnisi et al. *Synthesis* 651 1980, DOI: 10.1055/s-1980-29157; see also Vetter et al. *Carotenoids* ed Isler, Birkhauser Verlag, Basel, p 189 1971.]

(\pm)-Irone (6-methylionone, \pm -*trans*-(α)-4*t*-[2,5,6,6-tetramethylcyclohex-2-yl]but-3*en*-2-one) [79-69-6] C₁₄H₂₂O, M 206.3, b 85-86°/0.05mm, 109°/0.7mm, d₄²⁰ 0.9340, n_D²⁰ 1.4998. If large amounts are available, then fractionate through a Podbielniak column or an efficient spinning band column, but small amounts are distilled using a K gelrohr apparatus. The **4-phenylsemicarbazone** has m 174-175° (165-165.5°). [IR: Seidel & Ruzicka *Helv Chim Acta* **35** 1826 1952, DOI: 10.1002/hlca.19520350609; Naves *Helv Chim Acta* **31** 1280 1948, DOI: 10.1002/hlca.19480310513; Lecomte & Naves *J Chim Phys* **53** 462 1956, Beilstein **7** IV 378.]

dl-Isoborneol [124-76-5] C₁₀H₁₈O, M 154.3, m 212° (sealed tube). Crystallise isoborneol from EtOH or petroleum ether (b 60-80°). It sublimes in a vacuum. The **4-nitrobenzoyl** derivative has m 153°. [Yager & Morgan *J Am Chem Soc* **57** 2071 1935, DOI: 10.1021/ja01314a015; Beilstein **6** II 80, **6** III 299, **6** IV 281.]

Isocaryophyllene (γ -caryophyllene, *cis*-caryophyllene, **1*R*,9*S*-8-methylene-4,11,11-trimethylbicyclo [7.2.0] undec-4-ene**) C₁₅H₂₄, [118-65-0] M 204.4, b 93°/3mm, 122-124°/12mm, 131-133°/16mm, 130-131°/24mm, 271-273°/atm, d₄²⁰ 0.8959, n_D²⁰ 1.496, [α]₅₄₆²⁰ -31, [α]_D²⁰ -27 (neat). Purify it by vacuum distillation or GLC using a nitrile-silicone column. It can be characterised by treating an ethereal solution with perphthalic acid to give **isocaryophyllene oxide** m 80-81° [from petroleum ether (b 40-60°)] and [α]_D²⁰ -4 (c 1.0, CHCl₃). Unlike the less stable *caryophyllene*, it does not form a *nitrosite* readily, reacts more slowly towards perphthalic acid oxidation and has the characteristic IR bands at 3100, 1630 and 880 cm⁻¹ of the exocyclic methylene group (>C=CH₂). [Aebi et al. *J Chem Soc* 3124 1953, DOI: 10.1039/JR9530003124; Barton et al. *J Chem Soc* 2210 1952, DOI: 10.1039/JR9520002210; Corey et al. *J Am Chem Soc* **86** 485 1964, DOI: 10.1021/ja01057a040; Ramage & Simonsen *J Chem Soc* 741 1936, DOI: 10.1039/JR9360000741; Kumar et al. *Synthesis* 461 1976 DOI: 10.1055/s-1976-24082]. [Beilstein **5** II 355, **5** III 1085.] *Isocaryophyllene* can be isolated by steam distilling the mother liquors remaining after the preparation of *caryophyllene nitrosite* (see above and Deussen & Lewinsohn *Justus Liebigs Ann Chem* **356** 1 1907, DOI: 10.1002/jlac.19073560102), or by boiling this *nitrosite* with EtOH [Deussen *Justus Liebigs Ann Chem* **359** 245 1908, DOI: 10.1002/jlac.19083590112; Ramage & Whitehead *J Chem Soc* 4336 1954, DOI: 10.1039/JR9540004336]. On a large scale for the perfumery industry, *caryophyllene* (100 g) has been converted into *isocaryophyllene* (71.5 g) by mixing with sulfur (4 g), or selenium, and heating at 225° under N₂ for 8 hours [Rachlin German Patent DE 2044018 (1971 to I.F.F.), *Chem Abs* **75** 49364 1971].

(-)- β -Isolongifolene (1-*R*-(*-*)-2,2,7,7-tetramethyltricyclo[6.2.1.0^{1,6}]undec-5-ene) [1135-66-6] C₁₅H₂₄, M 204.4, b 82-83°/0.4mm, 144-146°/30mm, 255-256°/atm, d₄²⁰ 0.930, n_D²⁰ 1.4992, [α]₅₄₆²⁰ -166, [α]_D²⁰ -38 (c 1, EtOH). Reflux it over, and distil it from Na. [Zeiss & Arakawa *J Am Chem Soc* **76** 1653 1954, DOI: 10.1021/ja01635a056; IR: Reinaecker & Graafe *Angew Chem Int Ed* **97** 348 1985, DOI: 10.1002/

ange.19850970435; UV and NMR: Dev *Tetrahedron* **9** 1 1960, DOI: 10.1016/0040-4020(60)80047-6; Ranganathan et al. *Tetrahedron* **26** 621 1970, DOI: 10.1016/S0040-4020(01)97855-X; *Beilstein* **5** IV 1191.]

Isophorone (3,5,5-trimethyl-2-cyclohexen-1-one) [78-59-1] $\text{C}_9\text{H}_{14}\text{O}$ **M 138.2**, **m -8°**, **b 94°/16mm**, **213-214°/atm**, **d_4^{20} 0.921**, **n_D^{20} 1.4778**. Wash isophorone with aqueous 5% Na_2CO_3 and then distil it under reduced pressure immediately before use. *Alternatively*, it can be purified *via* the semicarbazone. Its solubility in H_2O is 1.20%w/v (20°) and 1.45%w/v (25°). **TOXIC**, do not breathe the vapours. [Erskine & Waight *J Chem Soc* 3425 1960, DOI: 10.1039/JR9600003425; *Beilstein* **7** IV 165.]

Isopinocampheol (pinan-3-ol, 2,6,6-trimethylbicyclo[3.1.1]heptan-3-ol) [1S,2S,3S,5R-(+)- 27779-29-9, 1R,2R,3R,5S-(-) 25465-65-0] $\text{C}_{10}\text{H}_{18}\text{O}$, **M 154.25**, **m 52-55°, 55-56°, 55-57°**, **b 103°/11mm**, **n_D^{20} 1.4832**, **$[\alpha]_D^{20}$ (+) and (-) 43**, **$[\alpha]_D^{20}$ (+) and (-) 36 (c 20, EtOH)**. Dissolve it in Et_2O , dry it over MgSO_4 , filter, evaporate, then recrystallise it from petroleum ether. Also recrystallise it from aqueous EtOH and distil it in a vacuum. [Kergomard & Geneix *Bull Soc Chim Fr* 394 1958, Zweifel & Brown *J Am Chem Soc* **86** 393 1964, DOI: 10.1021/ja01057a021. The **3,4-dinitrobenzoyl** derivative has **m 100-101°**, the **phenylcarbamoyl** derivative has **m 137-138°** and the **acid-phthalate** has **m 125-126°**. [*Beilstein* **6** III 282, 283, **6** IV 277.]

Isopropenylcyclobutane [3019-22-5] C_7H_{12} , **M 96.2**, **b 98.7°/760mm**, **d_4^{20} 0.7743**, **n_D^{20} 1.438**. Purify the cyclobutane by preparative chromatography (silicon oil column), or fractional distillation. Dry it over molecular sieves. Its IR (film) has ν_{max} at 1640 (C=C), 887 and 1773 (C-H) cm^{-1} . [Chiurdohlu & Van Walle *Bull Soc Chim Belg* **66** 612 1957, *Beilstein* **5** IV 255.]

(1S,2S,4R)-(+)-Limonene-1,2-diol [1S,2S,4R-(+)-4-isopropenyl-1-methylcyclohexan-1,2-diol, (1S,2S,4R)-(+)-p-menth-8-en-1,2-diol, (+)-1-hydroxyneodihydrocarveol, neolimonene glycol] [38630-75-0] $\text{C}_{10}\text{H}_{18}\text{O}_2$, **M 170.3**, **m 68-72°, 70-70.5°, 72-73°**, **$[\alpha]_D^{25}$ +55 (c 1, Me_2CO , also +45 and +53.7 were reported)**. It has been prepared from commercial grade (+)-limonene 1,2-oxide [500g, $[\alpha]_D^{27}$ +63.3 (neat)] by stirring at 0° with 6% H_2SO_4 (2.5L) for 5 hours, the solution is filtered, and the crude **diol hydrate** is dissolved in hot CHCl_3 separated from H_2O , concentrated, and crystallised from CHCl_3 or petroleum ether to give the **anhydrous diol** (375g, 67%) **m 70-70.5°**. The **tri-hydrate** crystallises from H_2O in plates with **m 60°**. On hydrogenation in EtOAc over Pt_2O , it provides **(+)-1-hydroxyneocarvomenthol**, **m 88°**, **$[\alpha]_D^{25}$ +48 (Me_2CO)** after recrystallisation from $^*\text{C}_6\text{H}_6$ -petroleum ether (b 70-110°). [Royals & Leffingwell *J Am Chem Soc* **31** 1937 1966, DOI: 10.1021/jo01344a062; Newhall *J Org Chem* **29** 185 1964, DOI: 10.1021/jo01024a042; *Beilstein* **6** H 753, **6** II 758, **6** III 4137, **6** IV 5294.]

Lupulon (β -lupulic acid, bitter acid, [3,5-dihydroxy-2,6,6-tris(3-methyl-2-butenyl)-4-(3-methyl-1-oxobutyl)-2,4-cyclohexadien-1-one] [468-28-0] $\text{C}_{26}\text{H}_{38}\text{O}_4$, **M 414.6**, **m 93°, 92-94°**, **$\text{pK}_{\text{Est}(1)} \sim 4.2$, $\text{pK}_{\text{Est}(2)} \sim 9.7$** . It was originally extracted from hop cones with petroleum ether (b 30-60°) which was concentrated, from which crystalline **Lupulon** separated and was recrystallised first from petroleum ether. Finally it was recrystallised from 90% MeOH. It can also be recrystallised from hexane or petroleum ether at low temperature. It has been purified by chromatography through Kieselgel. It is optically inactive and forms an *o*-phenylenediamine complex. It has antibiotic activity. [Wieland et al. *Chem Ber* **102** 2012 1925, DOI: 10.1002/cber.19250580912; Riedl *Chem Ber* **85** 692 1952, DOI: 10.1002/cber.19520850706; Carson *J Am Chem Soc* **73** 4652 1951, DOI: 10.1021/ja01154a046; *Beilstein* **7** II 856, **7** III 4752, **7** IV 2866.]

***l*-(-)-Menthone (2S,5R-2-isopropyl-5-methylcyclohexan-1-one) [14073-97-3]** $\text{C}_{10}\text{H}_{18}\text{O}$, **M 154.3**, **b 98-100°/14mm**, **205-208°/atm**, **207-210°/760mm**, **d_4^{25} 0.8930**, **n_D^{20} 1.4505**, **$[\alpha]_D^{20}$ -28 (neat)**. It is obtained by adding pure *l*-(-)-menthol (4.5g, see [2216-51-5]) in four portions to chromic acid [prepared from $\text{Na}_2\text{Cr}_2\text{O}_7 \cdot 2\text{H}_2\text{O}$ (60g, CARE CARCINOGENIC) in H_2O (300ml) and concentrated H_2SO_4 (27ml) with cooling] when the temperature rises, but should be kept at 55° (warm if necessary). A black spongy mass is first formed which becomes soft and melts to a dark brown oil on the surface as the temperature rises. The temperature drops when the oxidation is complete, the mixture is cooled and extracted with Et_2O (100ml), the extract is separated, washed with 5% aqueous NaOH (100ml, CARE) several times (\sim x 3) until the colour is light yellow, then H_2O

and dried (Na_2SO_4 or Mg_2SO_4). The ethereal solution is evaporated and the residual oil is redistilled at atmospheric pressure or in a vacuum through a short column to give pure *l*-menthone (38g, 91%). [Brown & Garg *J Am Chem Soc* **83** 2952 1961, DOI: 10.1021/ja01474a037; Hussey & Baker *J Org Chem* **25** 1434 1960, DOI: 10.1021/jo01078a602.] This menthone is the most frequently naturally occurring of the four optically active isomers. It has been used for the optical resolution of diols *via* acetalisation [Harada & Oku *Synlett* 95 1994, DOI: 10.1055/s-1994-22753], and the menthone spirocyclic 1,3-dioxane-4,6-diones have been used for asymmetric [2 + 2]cycloaddition and Diels-Alder reactions [Sato et al. *Tetrahedron* **47** 7271 1991, DOI: 10.1016/S0040-4020(01)89729-5]. The *oxime* has **m** 59° (from petroleum ether), the *semicarbazone* has **m** 189-191° (from MeOH, EtOH or aqueous EtOH), the *phenylhydrazone* has **m** 53° (from aqueous EtOH), and the *2,4-dinitrophenylhydrazone* has **m** 146° (from EtOH, aqueous EtOH or EtOH/EtOAc). [Beilstein **7** H 38, **7** I 34, **7** II 39, **7** III 152, **7** IV 87.] It is used in flavourings and perfume.

1*R*-(*-*)-Menthyl chloride (1*S*,2*R*,4*R*-2-chloro-1-isopropyl-4-methylcyclohexane) [16052-42-9] $\text{C}_{10}\text{H}_{19}\text{Cl}$, **M** 174.7, **m** -20.1° to -16.5°, **b** 88.5°/12.5mm, 101-105°/21mm, d_4^{20} 0.936, n_D^{20} 1.463, $[\alpha]_D^{20}$ -52.4 (neat). Dissolve menthyl chloride in petroleum ether (b 40-60°), wash it with H_2O , concentrated H_2SO_4 until no discoloration of the organic layer occurs (care with concentrated H_2SO_4 in a separating funnel), again with H_2O and dry it (MgSO_4). Evaporate the organic layer, and distil the residual oil through a Claisen head with a Vigreux neck of ca 40 cm length. [Smith & Wright *J Org Chem* **17** 1116 1952, DOI: 10.1021/jo50008a010; Barton et al. *J Chem Soc* 453 1952, DOI: 10.1039/JR9520000453; Beilstein **5** III 134, **5** IV 152]

1-Methyladamantane [768-91-2] $\text{C}_{11}\text{H}_{18}$, **M** 150.2, **m** 103°, 104°. Purify it by zone melting, chromatography through an Al_2O_3 column and eluting with pentane, and sublime it repeatedly at 90-95°/12mm. [Stetter et al. *Chem Ber* **92** 1629 1959, DOI: 10.1002/cber.19590920722; Schleyer & Nicholas *Tetrahedron Lett* **2** 305 1961, DOI: 10.1016/S0040-4039(01)84067-3; Beilstein **5** IV 479,] **2-Methyladamantane** [700-56-1] $\text{C}_{11}\text{H}_{18}$, **M** 150.2, has **m** 144-146°. Purify it by zone melting, chromatography through an Al_2O_3 column and eluting with pentane. Recrystallise it from EtOH and sublime it repeatedly at 90-95°/12mm. [Schleyer & Nicholas *J Am Chem Soc* **83** 182 1961, DOI: 10.1021/ja01462a036; Molle et al. *Can J Chem* **65** 2428 1987, DOI: 10.1139/v87-405.]

Methylcyclohexane (hexahydrotoluene) [108-87-2] C_7H_{14} , **M** 98.2, **m** -126°, **b** 100.9°/atm, d_4^{25} 0.7650, n_D^{20} 1.4231, n_D^{25} 1.42058. Passage through a column of activated silica gel gives material transparent down to 220nm. It can also be purified by passage through a column of activated basic alumina, or by azeotropic distillation with MeOH, followed by washing out the MeOH with H_2O , drying and distilling. Methylcyclohexane can be dried with CaSO_4 , CaH_2 or sodium. It has also been purified by shaking with a mixture of concentrated H_2SO_4 and HNO_3 in the cold, washing with H_2O , drying (CaSO_4) and fractionally distilling from potassium. Percolation through a column of Celite impregnated with 2,4-dinitrophenylhydrazine (DNPH), phosphoric acid and H_2O (prepared by grinding 0.5g DNPH with 6ml 85% H_3PO_4 , then mixing with 4ml of distilled H_2O and 10g of Celite) removes carbonyl-containing impurities. **TOXIC**, do not breathe vapour. [Cowan et al. *J Chem Soc* 1862 1939, DOI: 10.1039/JR9390001862; Beilstein **5** III 65, **5** IV 94.]

cis- and trans- 2-Methylcyclohexanol [583-59-5] $\text{C}_7\text{H}_{14}\text{O}$, **M** 114.2, **b** 65°/20mm, 167.6°/760mm, d_4^{20} 0.922, n_D^{20} 1.46085. Dry 2-methylcyclohexanol with Na_2SO_4 and fractionate it under vacuum. *Note:* The *cis-isomer* [7443-70-1] has **m** 6-8°, **b** 165°/760mm, d_4^{25} 0.936, n_D^{20} 1.465 [Beilstein **6** II 17], and the *trans-isomer* [7443-52-9] has **m** -21°, **b** 167.2-167.6°/760mm, d_4^{25} 0.924, n_D^{20} 1.461 [Beilstein **6** H 11]. [Eliel & Haber *J Org Chem* **23** 2041 1958, DOI: 10.1021/jo01106a638; Beilstein **6** III 61, **6** IV 100.]

cis- and trans-3-Methylcyclohexanol (mixture) [591-23-1] $\text{C}_7\text{H}_{14}\text{O}$, **M** 114.2, **b** 69°/16mm, 172°/760mm, d_4^{20} 0.930, n_D^{20} 1.45757, $n_D^{25.5}$ 1.45444. Dry 3-methylcyclohexanol with Na_2SO_4 and fractionate it under vacuum. *Note:* The *cis-isomer* has **b** 173°/760mm, and the *trans-isomer* has **b** 168-169°/760mm. [Eliel & Haber *J Org Chem* **23** 2041 1958, DOI: 10.1021/jo01106a638; Beilstein **6** IV 102.]

4-Methylcyclohexanone [589-92-4] $\text{C}_7\text{H}_{12}\text{O}$, **M** 112.2, **m** -40.6°, **b** 68°/23mm, 165.5°/743mm, 169-171°/atm, d_4^{20} 0.914, n_D^{20} 1.44506. Dry the ketone with CaSO_4 , then fractionally distil it. The *semicarbazone* has **m** 197°, 203.5°(dec) (from MeOH or EtOH). [White & Bishop *J Am Chem Soc* **62** 8 1940, DOI: 10.1021/

10.1021/ja01858a004; Vogel & Oommen *J Chem Soc* 768 1930, DOI: 10.1039/JR9300000768; *Beilstein* 7 III 63, 7 IV 44.]

1-Methylcyclohex-1-ene (3,4,5,6-tetrahydrotoluene) [591-49-1] C_7H_{12} , M 96.2, m -120.4° , b $107.4-108^\circ/\text{atm}$, $110-111^\circ/760\text{mm}$, d_4^{20} 0.813, n_D^{20} 1.451. Free it from hydroperoxides by passing through a column containing basic alumina or refluxing with cupric stearate, filter and fractionally distil it from sodium. Vogel *J Chem Soc* 1323 1938, DOI: 10.1039/JR9380001323; Cope et al. *J Am Chem Soc* 79 4729 1957, DOI: 10.1021/ja01574a038; *Beilstein* 5 III 197, 5 VI 245.]

Methylcyclopentane [96-37-7] C_6H_{12} , M 84.2, m -142° , b $64.32^\circ/400\text{mm}$, $71.8^\circ/\text{atm}$, d_4^{20} 0.749, n_D^{20} 1.40970, n_D^{25} 1.40700. Purification procedures include passage through columns of silica gel (prepared by heating in nitrogen to 350° prior to use) and activated basic alumina, distillation from sodium-potassium alloy, and azeotropic distillation with MeOH, followed by washing out the methanol with water, drying and distilling. It can be stored with CaH_2 or sodium. [Vogel *J Chem Soc* 1323 1938, DOI: 10.1039/JR9380001323; *Beilstein* 5 III 55, 5 IV 84.]

Methylnorbornene-2,3-dicarboxylic anhydride (5-methylnorborn-5-ene-2-endo-3-endo-dicarboxylic anhydride) [25134-21-8] $C_{10}H_{10}O_3$, M 178.2, m $88.5-89^\circ$. Purify the anhydride by thin layer chromatography on Al_2O_3 (previously boiled in EtOAc) and eluted with hexane/* C_6H_6 (1:2), then recrystallise it from * C_6H_6 /hexane. The *free acid* has m $118.5-119.5^\circ$. [Mironov et al. *Tetrahedron* 19 1939 1963, DOI: 10.1016/S0040-4020(01)97855-X; *Beilstein* 17/11 V 199.]

2,5-Norbornadiene (bicyclo[2.2.1]hepta-2,5-diene, NBD) [121-46-0] C_7H_8 , M 92.1, b $89^\circ/\text{atm}$, d_4^{20} 0.854, n_D^{20} 1.470. Purify the diene by distillation from activated alumina [Landis & Halpern *J Am Chem Soc* 109 1746 1987, DOI: 10.1021/ja00240a025; for an application in prostaglandin synthesis see Bindra et al. *J Am Chem Soc* 95 7522 1973, DOI: 10.1021/ja00803a061]. [*Beilstein* 5 IV 879; see ‘Catalysts-Part 2’ in Chapter 5.]

cis-endo-5-Norbornene-2,3-dicarboxylic anhydride (carbic anhydride, 3 α ,4,7,7, α , α -tetrahydro-4 α ,7 α -methanoisobenzofuran-1,3-dione) [129-64-6] $C_9H_8O_3$, M 164.2, m 164.1° , $164-165^\circ$, $165-167^\circ$, d_4^{20} 1.417. It forms crystals from petroleum ether, hexane or cyclohexane. It is hydrolysed by H_2O to form the acid [Diels & Alder *Justus Liebigs Ann Chem* 460 98 1928, DOI: 10.1002/jlac.19284600106; Maitte *Bull Soc Chim Fr* 499 1959]. The *exo-exo-isomer* has m $142-143^\circ$ (from * C_6H_6 /petroleum ether) [Alder & Stein *Justus Liebigs Ann Chem* 504 216 1933, DOI: 10.1002/jlac.19335040115]. [*Beilstein* 17 II 461, 17/11 V 192.]

(\pm)-endo-2-Norbornylamine hydrochloride (\pm endo[2.2.1]hept-2-ylamine HCl) [14370-45-7] $C_7H_{13}N$. HCl, M 147.7, m $\sim 295^\circ(\text{dec})$, $pK_{\text{Est}} \sim 9.0(\text{free base})$. Recrystallise the salt from MeOH/EtOAc or EtOH/ Et_2O . The *free base* has m $75-80^\circ$, b $156-157^\circ/\text{atm}$ and the *picrate* has m $179-180^\circ$ (from aqueous MeOH). [*Beilstein* 12 III 160.]

Norbornylene (bicyclo[2.2.1]hept-2-ene) [498-66-8] C_7H_{10} , M 94.2, m $44-46^\circ$, b $96^\circ/\text{atm}$. Reflux it over Na, and distil it [Gilliom & Grubbs *J Am Chem Soc* 108 733 1986, DOI: 10.1021/ja00264a027]. It has also been purified by sublimation *in vacuo* onto an ice-cold finger. [Woon et al. *J Am Chem Soc* 108 7990 1986, DOI: 10.1021/ja00285a018; *Beilstein* 5 IV 394.]

(\pm)-exo-2-Norbornylformate [41498-71-9] $C_8H_{12}O_2$, M 140.2, b $65-67^\circ/16\text{mm}$, $80-81^\circ/25\text{mm}$, d_4^{20} 1.048, n_D^{20} 1.4620. Shake with $NaHCO_3$ and distil it *in vacuo* (*exo*-borneol has m $124-126^\circ$). Alternatively, mix the ester with formic acetic anhydride overnight and fractionate, particularly if the sample contains much *exo*-borneol as evidenced by strong OH bands in the IR spectrum. [*Beilstein* 6 III 219.]

Norcamphor (bicyclo[2.2.1]heptan-2-one, \pm norbornan-2-one) [497-38-1] $C_7H_{10}O$, M 110.2, m $94-95^\circ$, $95.5-96.5^\circ$, b $89-94^\circ/60\text{mm}$. Crystallise it from water and sublime it *in vacuo*. It has λ_{max} at 287nm (EtOH). The *semicarbazone* has m $196-196.5^\circ$ (from EtOH/ H_2O). The *2,4-dinitrophenylhydrazone* has m $137-138^\circ$ (from EtOH). [Wildman & Hemminger *J Org Chem* 17 1641 1952, DOI: 10.1021/jo50012a011; Wood &

Roberts *J Org Chem* **23** 1124 1957, Bixler & Niemann *J Org Chem* **23** 742 1958, DOI: 10.1021/jo01099a600; Beilstein **7** III 243, **7** IV 139.]

1,2,3,4,5-Pentamethylcyclopentadiene (Cp') [4045-44-7] $C_{10}H_{16}$, **M 136.2**, **b 55-60°/13mm**, **58°/13mm**, **58.3°/13.5mm**, **d₄²⁵ 0.870**, **n_D²⁰ 1.4740**. Of the many syntheses of this useful ligand, the following is the most economical on materials and can be scaled up. Strictly anhydrous and anaerobic conditions should be used and reagents should be dried appropriately, degassed, and precaution against fire should be exercised. Three steps are involved: **Step 1:** Under argon, lithium (58g, 8.36moles, with 0.02%Na, of ~3.2mm diameter cut into the flask in ~5mm lengths) is covered with Et₂O (1600ml, freshly distilled from ~3:1w/w K/Na benzophenone), and 2-bromo-2-butene (120g, 0.88mole, 90.4ml of a molecular sieves 4Å dried commercial *cis*- and *trans*- mixture or prepared according to Bordwell & Landis *J Am Chem Soc* **79** 1593 1957, DOI: 10.1021/ja01564a020) is added slowly with stirring (10ml aliquots slowly at first until the reaction begins; with evolution of bubbles and cloudiness due to separation of LiBr), and at such a rate as to maintain gentle reflux of the Et₂O. **Step 2:** After addition is complete (~1.5 hours), still under argon, a mixture of ethyl acetate (166g, 1.88mole, 184ml, dried over molecular sieves 4Å) and 2-bromo-2-butene (430g, 3.18mole, 324ml as above) are then added dropwise with stirring, while carefully maintaining gentle reflux over a period of 4-5 hours. When this addition is completed, a further volume of dry EtOAc (50g, 55.4ml) and a further portion of 2-bromo-2-butene (~10-20g, 7.5-15ml) are added, while stirring is continued, until refluxing of Et₂O ceases. The mixture is allowed to cool over 4 hours, saturated aqueous NH₄Cl solution is added dropwise to hydrolyse unreacted Li, the Et₂O layer is collected, the aqueous layer is extracted with Et₂O (3 x 200ml), and the combined Et₂O solutions are evaporated to ~350ml. **Step 3:** The Et₂O concentrate is added, with stirring, to a slurry of *p*-toluenesulfonic acid monohydrate (26g) in Et₂O (500ml) under a reflux condenser, at such a rate that the solvent refluxes gently. The mixture is then stirred for 5 minutes after refluxing ceases, and poured into saturated aqueous NaHCO₃ (1200ml) containing Na₂CO₃ (19g). The yellow aqueous phase is removed, extracted with Et₂O (3 x 200ml), and the combined Et₂O solutions are dried (Na₂SO₄), filtered, evaporated to 250-300ml (rotavap), then trap-to-trap distilled *in vacuo* (bath temperature at 35-40°) to give a yellow liquid (85% pure by GC) which is fractionally distilled in a vacuum under N₂ using a 50cm vigreux column. The fraction with **b 65-70°/20mm** collected (142g, 53% yield based on EtOAc used) as pale-yellow liquid is **1,2,3,4,5-pentamethylcyclopentadiene** (92% pure by GC). An additional fraction (15g, 5%) with **b 70-75°/20mm** is 85% pure by GC. The liquids are colourless to pale-yellow in colour with a sweet olefinic odour, and are pure enough for use as ligands. They should be stored in a freezer under N₂ or argon. In comparison with the unsubstituted *cyclopentadiene* it is a stronger donor of electron density, exerting considerably enhanced thermal stability, and the metal complexes that it forms are generally more soluble and easier to crystallise. The *pentamethylpentadiene* has FT-IR (neat) with ν_{max} at 2960 (vs), 2915 (vs), 2855 (vs), 2735 (w), 1660 (m), 1640 (w), 1390 (s), 1355 (m), 1150 (w), 1105 (mw), 1048 (w), 840 (mw) and 668 (w) cm⁻¹; ¹H NMR (60MHz, CCl₄, TMS) with δ at 2.4 (m, 1H), 1.75 (br s, 13H, 2,3,4,5-Me), 0.95 (d, ³J_{H-H} = 8Hz, 3H, 1-Me); ¹³C NMR (15MHz, CDCl₃, TMS) with δ at 14.18, 20.98, 27.43, 38.04, 54.77, 61.26, 169.37 and 212.25. [Manriquez et al. *Inorg Synth* **XXI** 181 1982, Feitler et al. *Inorg Chem* **15** 466 1976, DOI: 10.1021/ic50156a046; Threlkel et al. *Org Synth* **65** 42 1987, DOI: 10.15227/orgsyn.065.0042.]

Perfluorocyclobutane (octafluorocyclobutane) [115-25-3] C_4F_8 , **M 200.0**, **m -40°**, **b -5°/atm**, **d₄⁻²⁰ 1.654**, **d₀ 1.72**. Purify octafluorocyclobutane by trap-to-trap distillation, retaining the middle portion. [Danus *Ind Eng Chem* **47** 144 1955, Claasen *J Chem Phys* **18** 543 1950, DOI: org.virtual.anu.edu.au/10.1063/1.1747681; Beilstein **5** III 8, **5** IV 8.]

Perfluorocyclohexane (dodecafluorocyclohexane) [355-68-0] C_6F_{12} , **M 300.1**, **m 51°** (sublimes), sublimes on melting at **52°**, **m 58.2°** (sealed tube), **d₄²⁵ 1.720**, **n_D³⁰ 1.269**. Extract it repeatedly with MeOH, then pass it through a column of silica gel (previously activated by heating at 250°). Sublime it *in vacuo*. [Haszeldine & Smith *J Chem Soc* 2689 1950, DOI: 10.1039/JR9500002689; IR: Thompson & Temple *J Chem Soc* 1432 1948, DOI: 10.1039/JR9480001432; Beilstein **5** III 37, **5** IV 48.]

Perfluoro-1,3-dimethylcyclohexane [335-27-3] C_8F_{16} , **M 400.1**, **m -55°**, **b 101-102°/atm**, **d₄²⁰ 1.829**, **n_D²⁰ 1.300**. Fractionally distil it, then 35ml are sealed with about 7g KOH pellets in a borosilicate glass ampoule and

heated at 135° for 48 hours. The ampoule is cooled, opened, and the liquid is resealed with fresh KOH in another ampoule and heated as before. This process is repeated until no further decomposition is observed. The substance is then washed with distilled water, dried (CaSO₄) and distilled. [Grafstein *Anal Chem* **26** 523 1954, DOI: 10.1021/ac60087a029, *Beilstein* **5** III 378.] **IRRITANT, avoid breathing its vapours.**

Perfluoro(methylcyclohexane) [355-02-2] C₇F₁₄, **M 350.1, b 76.3°/atm, d₄²⁵ 1.7878.** Reflux it for 24 hours with saturated acid KMnO₄ (to oxidise and remove hydrocarbons), then neutralise, steam distil, dry with P₂O₅ and pass slowly through a column of dry silica gel. [Glew & Reeves *J Phys Chem* **60** 615 1956, DOI: 10.1021/j150539a026.] It can also be purified by percolation through a 1 metre neutral activated alumina column, and the impurities are checked by ¹H NMR. [*Beilstein* **5** IV 102.] **IRRITANT, avoid breathing its vapours.**

R(-)-α-Phellandrene (p-menta-1,5-diene) [4221-98-1] C₁₀H₁₆, **M 136.2, b 61°/11mm, 171-174°/atm, 175-176°/760mm, d₄²⁰ 0.838, n_D²⁰ 1.471, [α]_D²⁰ -230 (c 10, Et₂O), -153 to -183 (neat).** Purify it by gas chromatography using an Apiezon column. Also purify it by steam distillation (with 0.5% hydroquinone), then re-distil it through a 50 plate bubble cap column and collecting the fraction with **b 72-72.5°/22mm** [Pines & Eschinazi *J Am Chem Soc* **77** 6314 1955, DOI: 10.1021/ja01628a072]. UV: λ_{max} at 263nm (ε 3,345) in octane. [Read & Storey *J Chem Soc* 2770 1930, DOI: 10.1039/JR9300002770; *Beilstein* **5** III 341, **5** IV 436.]

Picrotoxin (cocculin) [124-87-8] C₃₀H₃₄O₁₃, [C₁₅H₁₆O₆, C₁₅H₁₈O₇], **M 602.6, m 203°, 203-204°, [α]_D²⁰ -40 (c 1, EtOH), [α]_D¹⁶ -29.3 (c 4, EtOH).** It is the toxic and bitter principle in the seeds of *Anamirta cocculus*, and is *very poisonous* (use CARE when working with it). Crystallise picrotoxin from H₂O or Me₂CO/H₂O. It is sparingly soluble in Et₂O and CHCl₃. Its solubility (w/v) in H₂O is 0.3% (~20°) and 20% (100°); in 95% EtOH it is 7.4% (~20°) and 33% (78°). It is freely soluble in aqueous NaOH and 880 NH₃. The *monoacetate* has **m 244-245°** (*C₆H₆). [Meyer & Bruger *Chem Ber* **31** 2958 1898, DOI: 10.1002/cber.18980310366; Johns et al. *J Chem Soc* 4715 1956, DOI: 10.1039/JR9560004715; *Beilstein* **19** III/IV 5245.] **HIGHLY TOXIC**, stimulates the CNS and respiration [For toxicity see Setnikar et al. *J Pharmacol Exp Ther* **128** 176 1960, PMID: 14445192; and for the crystal and molecular structure see Dupont et al. *Acta Crystallographica B* **32** S. 2987 1976, DOI: 10.1107/S0567740876009424].

(±)-α-Pinene (2,6,6-trimethylbicyclo[3.1.1]hept-2-ene) [80-56-8] C₁₀H₁₆, **M 136.2, b 52.5°/20mm, 155-156°/760mm, d₄²⁰ 0.8592, n_D²⁰ 1.4664.** Racemic synthetic material can be purified like the chiral forms below. It forms a *hydrochloride*, C₁₀H₁₇Cl, with **m 132°**. Upon per-acid oxidation it forms the epoxide **(±)-α-pinene oxide** [1686-14-2] C₁₀H₁₆O, **M 152.3**, which distils at **102-103°/50mm** and has **d₄²⁵ 0.964, n_D²⁰ 1.469** [*Beilstein* **5** H 152]. It occurs naturally in optically active forms (see below) in turpentine oil (~60% of α-pinene and ~30% of β-pinene) where the pinene from most European sources are *laevo-rotatory* whereas those from North America are *dextro-rotatory*. [Simonsen *The Terpenes Vol II* pp 105-191 1949, Cambridge University Press.]

1R,5S-α-Pinene [7785-70-8] C₁₀H₁₆, **M 136.2, b 61°/30mm, 156.2°/760mm, d₄²⁰ 0.858, n_D¹⁵ 1.4634, n_D²⁰ 1.4658, [α]_D²⁰ +51 (neat).** It is isomerised by heat, acids and certain solvents. It should be distilled under reduced pressure under N₂ and stored in the dark. It has been purified *via* the nitrosochloride [Waterman et al. *Recl Trav Chim Pays-Bas* **48** 1191 1929, DOI: 10.1002/recl.19290481202]. For purification of optically active forms see Lynn [*J Am Chem Soc* **91** 361 1919, DOI: 10.1021/ja01460a010]. It forms a *hydrochloride*, C₁₀H₁₇Cl, with **m 132°, [α]_D²⁰ +33.5 (EtOH)**. Small quantities (0.5ml) have been purified by GLC using helium as carrier gas and a column at 90° packed with 20 wt% of polypropylene sebacate on a Chromosorb support. Larger quantities are fractionally distilled under reduced pressure through a column packed with stainless steel gauze spirals. The material can be dried over CaH₂ or sodium, and stored in a refrigerator: CaSO₄ and silica gel are not satisfactory because they induce spontaneous isomerisation. [Bates et al. *J Chem Soc* 1521 1962, DOI: 10.1039/JR9560004715; *Beilstein* **5** III 366, **5** IV 452.]

1S,5S-α-Pinene [7785-26-4] C₁₀H₁₆, **M 136.2, b 155-156°/760mm, d₄²⁰ 0.858, n_D²⁰ 1.4634, [α]_D²⁰ -47.2 and [α]_D²⁰ -51.28 (neat).** Purify as for 1R,5S-α-Pinene above. It forms a *hydrochloride*, C₁₀H₁₇Cl, with **m 132°, [α]_D²⁰ -33.5 (EtOH)**. [*Beilstein* **5** III 366, **5** IV 455.]

1*S*,5*S*- β -Pinene (1*S*,5*S*-6,6-dimethyl-2-methylenebicyclo[3.1.1]heptane) [18172-67-3] $C_{10}H_{16}$, **M 136.2**, **m -61°**, **b 162-163°/atm**, **165-167°/atm**, **d₄²⁰ 0.858**, **n_D¹⁵ 1.4872**, **n_D²⁰ 1.478**, **[α]_D²⁰ -47.2** and **[α]_D²⁰ -22.4** (**neat**). Purification is as for the previous isomer, i.e. by fractional distillation. *Note* that it differs from the isomeric α -pinene (above) in that the 2-Me group is replaced by a 2-methylene group, and it isomerises irreversibly to α -pinene when shaken with Pt black that had been saturated with hydrogen [Richter & Wolff *Chem Ber* **59** 1733 1926, DOI: 10.1002/cber.19260590816]. [For synthesis see Harwood & Julia *Synthesis* 456 1980, DOI: 10.1055/s-1980-29049 and Crowley *Proc Chem Soc (London)* 237 (p 245) 1962, DOI: 10.1039/PS9620000237.] [*Beilstein* **5** IV 457.]

R(+)-Pulegone (**R-2-isopropylidene-5-methylcyclohexanone**) [89-82-7] $C_{10}H_{16}O$, **M 152.2**, **b 69.5°/5mm**, **84°/6mm**, **103°/17mm**, **151-153°/1100mm**, **224°/atm**, **d₄²⁰ 0.936**, **n_D²⁰ 1.4894**, **[α]_D²⁰ +23.5** (**neat**), **[α]_D²⁰ +24.2** (**neat**). Purify pulegone *via* the *semicarbazone* which has **m 174°** (from MeOH) and **[α]_D²⁰ +68.2** (c 1, $CHCl_3$). Fractionally distil it *in vacuo*. [Short & Read *J Chem Soc* 1306 1939, DOI: 10.1039/JR9390001306]. [Erskine & Waight *J Chem Soc* 3425 1960, DOI: 10.1039/JR9600003425; cf Ort *Org Synth* **65** 203 1987, DOI: 10.15227/orgsyn.065.0203; *Beilstein* **7** III 334, **7** IV 188.] The enantiomer **S(-)-pulegone** [3391-90-0] $C_{10}H_{16}O$, **M 152.2**, has **b 104-108°/20mm**, **[α]_D²³ -22.5** (**neat**). [*Beilstein* **7** IV 188.]

1*R*,3*R*,4*R*,5*R*-Quinic acid (1,3,4,5-tetrahydroxycyclohexane carboxylic acid) [77-95-2] $C_7H_{12}O_6$, **M 192.3**, **m 172°(dec)**, **[α]_D²⁰ -51** (c 20, H_2O), **[α]_D²³ -45** (c 5, H_2O), **pK²⁵ 3.58**. Quinic acid crystallises from H_2O with **m 174°**, and from EtOH with **m 168-169°**. [McComsey & Maryanoff *J Org Chem* **59** 2652 1994, DOI: 10.1021/jo00088a065; pK: Timberlake *J Chem Soc* 2795 1959, DOI: 10.1039/JR9590002795; Anet & Reynolds *Aust J Chem* **8** 280 1955, DOI: 10.1071/CH9550280; *Beilstein* **10** III 2407, **10** IV 2257.]

Reductic acid (1,2-dihydroxycyclopent-1,2-en-3-one) [80-72-8] $C_5H_6O_3$, **M 114.1**, **m 213°**, **pK₁²⁰ 4.80**, **pK₂²⁰ 12.9**. Crystallise reductic acid from EtOH, EtOAc (**m 213-213.5°**) or EtOH/EtOAc. It has been sublimed at 0.5mm. It is soluble in MeOH, EtOH and H_2O , much less soluble in Et₂O, EtOAc and Me_2CO , but almost insoluble in petroleum ether and C_6H_6 . The *osazone* has **m 245°(dec)** (from BuOH). [Hess et al. *Justus Liebigs Ann Chem* **563** 31 1949, DOI: 10.1002/jlac.19555920204; **592** 137 1955, DOI: 10.1002/jlac.19495630105; **736** 134 1970, DOI: 10.1002/jlac.19707360114; for structure see Semmingen *Acta Chem Scand B* **31** 81 1977, DOI: 10.3891/acta.chem.scand.31b-0086; *Beilstein* **8** III 1942, **8** IV 1714.] It is an antioxidant, similar to ascorbic acid, and its **5-methyl derivative** (**m 71°**) has stronger reducing properties.

Squaric acid (3,4-dihydroxy-3-cyclobutene-1,2-dione) [2892-51-5] $C_4H_2O_4$, **M 114.1**, **m 293°(dec)**, **294°(dec)**, **>300°**, **pK₁²⁰ 1.50**, **pK₂²⁰ 2.93**. Purify squaric acid by recrystallisation from H_2O — this is quite simple because the acid is ~ 7% soluble in boiling H_2O and only 2% at room temperature. It is not soluble in Me_2CO or Et₂O; hence it can be rinsed with these solvents and dried in air or a vacuum. It is not hygroscopic and gives an intense purple colour with $FeCl_3$. It has IR with ν_{max} at 1820 (C=O) and 1640 (C=C) cm^{-1} , and UV with λ_{max} at 269.5nm (ϵ 37,000 $M^{-1}cm^{-1}$). [Cohn et al. *J Am Chem Soc* **81** 3480 1959, DOI: 10.1021/ja01522a083; Park et al. *J Am Chem Soc* **84** 2919 1962, DOI: 10.1021/ja00874a015] See also **pK_a** values of 0.59 \pm 0.09 and 3.48 \pm 0.023 [Schwartz & Howard *J Phys Chem* **74** 4374 1970, DOI: 10.1021/j100719a013]. [Schmidt & Ried *Synthesis* 869 1978, DOI: 10.1055/s-1978-24920; *Beilstein* **8** IV 2701.] The *diethyl ester* [5231-87-8] $C_6H_{10}O_4$, **M 170.2**, has **b 95°/0.1mm**.

Terpin hydrate [2451-01-6 *cis*-hydrate, 565-50-4 and 565-48-0 stereoisomers] $C_{10}H_{20}O_2 \cdot H_2O$ **M 190.3**, **m 105.5°** (*cis anhydrous*), **116-117°** (*cis hydrate*), **156-158°**, **157.5°** (*trans*). Crystallise terpin from H_2O or EtOH. The anhydrous *cis*-isomer distils at 258°/760mm, but hydrates on exposure to moist air. *Anhydrous* terpin is also obtained by recrystallisation from absolute EtOH. [Sword *J Chem Soc* **127** 1632 1925, DOI: 10.1039/CT9252701632; Lombard & Ambrose *Bull Soc Chim Fr* 230 1961, *Beilstein* **5** IV 435.]

1,1,2,2-Tetrafluorocyclobutane [374-12-9] $C_4H_4F_4$, **M 128.1**, **b 50-50.7°/atm**, **d₄²⁰ 1.275**, **n_D²⁰ 1.3046**.

Purify 1,1,2,2-tetrafluorocyclobutane by distillation or by preparative gas chromatography using a 2m x 6mm(i.d.) column packed with β , β' -oxydipropionitrile on Chromosorb P at 33°. [Conlin & Fey *JCS Faraday Trans 1* **76** 322 1980, DOI: 10.1039/F19807600322; Coffmann et al. *J Am Chem Soc* **71** 490 1949, DOI: 10.1021/ja01170a033; *Beilstein* **5** III 8, **5** IV 8.]

2,2,4,4-Tetramethylcyclobutan-1,3-dione [933-52-8] $\text{C}_8\text{H}_{12}\text{O}_2$, **M 140.2**, **m 114.5-114.9°**, **114-116°**, **b 95-159°/750mm**. Crystallise the dione from $^*\text{C}_6\text{H}_6$ and dry it *in vacuo* over P_2O_5 in an Abderhalden pistol. [*Beilstein* **7** III 3234, **7** IV 2004.]

3,3,5,5-Tetramethylcyclohexanone [14376-79-5] $\text{C}_{10}\text{H}_{18}\text{O}$, **M 154.3**, **m 11-12°**, **13.2°**, **b 59-61°/5mm**, **80-82°/13mm**, **196°/760mm**, **203.8-204.8°/760mm**, **d** $^{20}_4$ **0.8954**, **n** $^{20}_D$ **1.4515**. Purify the ketone first through a 24 inch column packed with Raschig rings, then a 40cm Vigreux column under reduced pressure (**b 69-69.3°/7mm**). The *oxime* has **m 144-145°** (from 60% EtOH), and the *semicarbazone* has **m 196-197°**, **197-198°** (214.5°, 217-218°) [Karasch & Tawney *J Am Chem Soc* **63** 2308 1941, DOI: 10.1021/ja01854a005; UV: Sandris & Ourisson *Bull Soc Chim Fr* 958 1956]. [*Beilstein* **7** III 163, **7** IV 89.]

(1R)-(-)-Thiocamphor (**1R-bornane-2-thione**, **1R-(-)-1,7,7-trimethylbicyclo[2.2.1]heptane-2-thione**) [53402-10-1] $\text{C}_{10}\text{H}_{16}\text{S}$, **M 168.3**, **m 136-138°**, **146°**, **[α] $^{22}_D$ -22°** (c **3**, EtOAc). It forms red prisms from EtOH and sublimes under vacuum. It possesses a sulfurous odour and is volatile like camphor. [Sen *J Indian Chem Soc* **12** 647 1935, Sen *J Indian Chem Soc* **18** 76 1941.] The *racemate* crystallises from $^*\text{C}_6\text{H}_6$ and has **m 145°** [**138.6-139°** by White & Bishop *J Am Chem Soc* **62** 8 1940, DOI: 10.1021/ja01858a004]. [*Beilstein* **7** III 419.]

1r,2t,4t-Trimethylcyclohexane [2234-75-7, isomeric mixture 2234-57-5] C_9H_{18} , **M 126.2**, **b 145.7-146.7°/760mm**, **d** $^{20}_4$ **0.786**, **n** $^{20}_D$ **1.4330**. Wash the trimethylcyclohexane with concentrated H_2SO_4 (removes aromatic hydrocarbons), then with H_2O , dry it (type 4A molecular sieves), and fractionally distil it through a glass helices packed column with partial take-off and reflux ratio between 50 and 75. **Flammable liquid**. [cf. Henne et al. *J Am Chem Soc* **63** 3474 1941, DOI: 10.1021/ja01857a062; Rossini *Anal Chem* **20** 110 1948, DOI: 10.1021/ac60014a006; *Beilstein* **5** H 42, **5** I 17, **5** II 24, **5** III 121, **5** IV 138.]

R-(-)-2,2,6-Trimethyl-1,4-cyclohexanedione [60046-49-3] **M 154.2**, **m 88-90°**, **91-92°**, **[α] $^{20}_D$ -270°** (c **0.4%**, MeOH), **[α] $^{20}_D$ -275** (c **1**, CHCl_3). It is obtained from yeast fermentation and is purified by recrystallisation from diisopropyl ether. [ORD: Leuenberger et al. *Helv Chim Acta* **59** 1832 1976, DOI: 10.1002/hlca.19760590541.] The *racemate* has **m 65-67°**, and the **4-(4-phenyl)semicarbazone** has **m 218-220°** (from $\text{CH}_2\text{Cl}_2/\text{MeOH}$) [Isler et al. *Helv Chim Acta* **39** 2041 1956, DOI: 10.1002/hlca.19560390717; *Beilstein* **7** IV 2032.]

cis,cis-1 α ,3 α ,5 α -Trimethylcyclohexane-1,3,5-tricarboxylic acid (Kemp's acid) [79410-20-1] $\text{C}_{12}\text{H}_{18}\text{O}_6$, **M 258.3**, **m 241-243°**, **pK₁ 3.30**, **pK₂ 5.85**, **pK₃ 7.3** (H_2O); **pK₁ 4.7**, **pK₂ 7.6**, **pK₃ 8.8** (50% $\text{H}_2\text{O}/\text{MeOH}$). Recrystallise the tricarboxylic acid from Me_2CO after re-precipitating it several times with mineral acid from aqueous alkaline solution. The *trimethyl ester* has **m 78-81°**. [See Kemp *J Org Chem* **46** 5140 1981, DOI: 10.1021/jo00338a014; Jeong et al. *J Am Chem Soc* **113** 201 1991, DOI: 10.1021/ja00001a029; Stack et al. *J Am Chem Soc* **114** 7007 1992, DOI: 10.1021/ja00044a009.]

(\pm)-2,2,6-Trimethylcyclohexanone [2408-37-9] $\text{C}_9\text{H}_{16}\text{O}$, **M 140.2**, **b 69-71.5°/20mm**, **177-178.5°/758mm** **d** $^{20}_4$ **0.904**, **n** $^{20}_D$ **1.4470**. Purify it *via* the *semicarbazone* (**m 218°**, from MeOH or EtOH), decompose this in the usual way (see Chapter 2 or purification of methyl ethyl ketone [78-93-0]) and fractionally distil the liquid ketone through a Vigreux column at $\sim 760\text{mm}$. [Chakravarti *J Chem Soc* 1565 1947, DOI: 10.1039/JR9470001565; Milas et al. *J Am Chem Soc* **70** 1829 1948, DOI: 10.1021/ja01185a053; *Beilstein* **7** I 24, **7** II 32, **7** III 123, **7** IV 69.] **(\pm)-3,3,5-Trimethylcyclohexanone** [873-94-9] $\text{C}_9\text{H}_{16}\text{O}$, **M 140.2**, has **m -10°**, **b 188-192°/758mm** **d** $^{25}_4$ **0.889**, **n** $^{20}_D$ **1.445**. Purify it *via* the *semicarbazone* (**m 218°**, from MeOH or EtOH), decompose this in the usual way (see Chapter 2 or purification of methyl ethyl ketone [78-93-0]) and fractionally distil the liquid ketone through a Vigreux column at $\sim 760\text{mm}$. [Chakravarti *J Chem Soc* 1565 1947, DOI: 10.1039/JR9470001565; Milas et al. *J Am Chem Soc* **70** 1829 1948, DOI: 10.1021/ja01185a053; *Beilstein* **7** I 24, **7** II 32, **7** III 7, **7** IV 69.]

Xanthatin (3-methylene-7-methyl-6-[3-oxo-1-buten-1-yl]cyclohept-5-ene-[10,11-*b*]furan-2-one, (-)-2-[(*IR*)-7*t*-hydroxy-5*c*-methyl-4-(3-oxobut-1-en- ξ -yl)cyclohept-3-en-*r*-yl]-acrylic acid lactone [26791-73-1] $C_{15}H_{18}O_3$, M 246.3, m 114.5-115°, 114.5-115.2°, $[\alpha]_D^{20}$ -20 (c 2, $CHCl_3$, or EtOH). Xanthatin was first isolated from *Xanthium pensylvanicum* [Little, Foote and Johnstone *Arch Biochem* **27** 247 1950, PMID: 14771893], found in several *Xanthium* species, and its structure was firmly established by Geissman and co-workers [Geissman et al.* *J Am Chem Soc* **76** 685 1954, DOI: 10.1021/ja01632a014; Deuel & Geissman* *J Am Chem Soc* **79** 3778 1957, DOI: 10.1021/ja01571a044]. It is a potent antibacterial against *Staphylococcus aureus*, including MRSA [Sato et al. *J Pharm Pharmacol* **49** 1042 1997, PMID: 9364417], and its total synthesis has been achieved by Shishido and coworkers [Hiromasa et al. *Tetrahedron Lett* **49** 3504 2008, DOI: 10.1016/j.tetlet.2008.03.081]. It can be purified by recrystallisation from MeOH, aqueous MeOH, absolute EtOH or aqueous EtOH. It is soluble in Et_2O , Me_2CO and EtOH, slightly soluble in H_2O at neutral pH, but almost insoluble in aqueous N NaOH and N HCl. Its UV has $\lambda_{max}(\epsilon)$ at 213 (22800) and 275nm (7300) in H_2O ; and IR (film) with ν_{max} at 1766 (C=O), 1660, 1609, 1590 (C=O, C=C) and 812 ($R_2C=CCHR$) cm^{-1} . The **2,4-dinitrophenylhydrazone** has m 240°(dec) (twice recrystallised from $CHCl_3/MeOH$ as garnet-red crystals). *Beilstein* **17** III/IV 6221, **17/1** V 305.]

*Geissman and coworkers extracted the dried, ground young leaves of the Cocklebur *X Pennsylvanicum* (from the daisy family of the sunflower group of plants) (50g) with Me_2CO (800ml) at $\sim 25^\circ$ with shaking for 2 hours, filtered with suction, and the filtrate was evaporated to dryness. Water (5L) was added to the dry residue and set aside for 24 hours, then filtered and the filtrate was extracted with Et_2O (1L x 3) and the combined extracts were placed in the freezer to freeze out any H_2O present. After filtration, the filtrate was evaporated down to 15ml and this Et_2O concentrate gave crystals on cooling. These were filtered off washed with cold Et_2O and twice recrystallised from the minimum volume of dilute EtOH to give 0.5-0.6g of **xanthinin** [580-49-4, 64047-79-6] (representing 1-1.2% of dry leaves). **Xanthinin** crystallises as colourless plates from Et_2O , EtOH or aqueous EtOH with m 121-122°, is odourless, insoluble in cold but not hot H_2O , soluble in pyridine, EtOH, Me_2CO , and $*C_6H_6$, slightly soluble in Et_2O but insoluble in petroleum ether. It analyses for $C_{17}H_{22}O_5$, and its UV has $\lambda_{max}(\epsilon)$ at 215 (9037) and 220nm (5974) in 95% EtOH, and 215 (12193) and 220nm (8823) in H_2O ; has IR (film) with ν_{max} at 1765 (C=O), 1720 (C=O, C=C) and 814 ($R_2C=CCHR$) cm^{-1} ; and has $[\alpha]_D^{25}$ -53.0 (c 0.119g in 5ml $CHCl_3$). Its **p-bromophenylhydrazone** forms yellow crystals from EtOH with m 138-138.5°.

Xanthatin (**desacetylxanthinin**) was best obtained by deacetylation of **xanthinin** (e.g. 1.24g) with NaOAc (2.3g) solution in EtOH (20ml) by refluxing (40 minutes) while some EtOH (10ml) was allowed to distil off. The solution was made alkaline (to litmus) with HCl, H_2O was added until the precipitated NaCl had just dissolved, and cooled whereby **xanthatin** crystallised out (0.96g, 96%). It had m 114.5-115.5° upon recrystallisation from absolute EtOH.

AROMATIC COMPOUNDS

Acenaphthene [83-32-9] $C_{12}H_{10}$, **M 154.2, m 94.0°, 95.0°, b 279°/atm.** It is a coal tar component with an odour of naphthalene and is used as a vermicide and insecticide. Crystallise acenaphthene from EtOH. It has also been purified by chromatography from CCl_4 on alumina with *benzene as eluent [McLaughlin & Zainal *J Chem Soc* 2485 1960, DOI: 10.1039/JR9600002485]. Acenaphthalene is almost insoluble in H_2O , but is soluble at ~ 25° (w/v) in EtOH (3.2%), MeOH (1.7%), PrOH (4.0%), $CHCl_3$ (40.0%), * C_6H_6 (20.0%) and AcOH (3.2%) and forms 1:2 complexes with bile acids in EtOH or dioxane. [Beilstein 5 IV 1834.]

Acenaphthenequinone [82-86-0] $C_{12}H_6O_2$, **M 182.2, m 249-252°(dec), 260-261°.** Extract it with, then recrystallise it twice from * C_6H_6 . Dry it *in vacuo*. [LeFevre et al. *J Chem Soc* 974 1963, DOI: 10.1039/JR9630000974; Beilstein 7 IV 2498.] It has been prepared by the oxidation of acenaphthene with $Na_2Cr_2O_7/Ce(OAc)_3$ in AcOH [Allen & VanAllan *Org Synth* 28.4.2003, DOI: 10.1002/0471264180.os024.01].

RS-1-Acenaphthenol [6306-07-6] $C_{12}H_{10}O$, **M 170.2, m 144.5-145.5°, 146°, 148°.** If highly coloured (yellow), dissolve it in boiling *benzene (14g in 200ml), add charcoal (0.5g), filter it through a heated funnel, concentrate to 100ml and cool to give almost colourless needles. *Benzene vapour is **TOXIC**; use an efficient fume cupboard. The **acetate** has **b 166-168°/5mm** (bath temperature 180-185°). [Cason *Org Synth Coll Vol* 3 3 1955, DOI: 10.15227/orgsyn.021.0001.] It can also be recrystallised from EtOH [Fieser & Cason *J Am Chem Soc* 62 432 1940, DOI: 10.1021/ja01859a053]. It forms a brick-red crystalline **complex** with 2,4,5,7-tetranitrofluoren-9-one which is recrystallised from AcOH and is dried in a vacuum over KOH and P_2O_5 at room temperature, **m 170-172°** [Newman & Lutz *J Am Chem Soc* 78 2469 1956, DOI: 10.1021/ja01592a034]. [Beilstein 6 IV 4623.]

Acenaphthylene [208-96-8] $C_{12}H_8$, **M 152.2, m 92-93°, b 280°/~760mm.** Dissolve acenaphthylene in warm redistilled MeOH, filter through a sintered glass funnel and cool to -78° to precipitate the material as yellow plates [Dainton et al. *Trans Faraday Soc* 56 1784 1960, DOI: 10.1039/TF9605601784]. Alternatively it can be sublimed *in vacuo*. [Beilstein 5 H 625, 5 IV 2138.]

4-Acetamidobenzaldehyde (4'-formylacetanilide) [122-85-0] $C_9H_9NO_2$, **M 163.2, m 155°, 156°, 160°.** Recrystallise it from water. The **4-nitrophenylhydrazone**, **m 264-265°**, crystallises as orange needles from EtOH [Hodgson & Beard *J Chem Soc* 21 1927, DOI: 10.1039/JR9270000020; Beilstein 14 H 38, 14 II 25, 14 III 75, 14 IV 71.]

p-Acetamidobenzenesulfonyl chloride (N-acetylsulfanilyl chloride) [121-60-8] $C_8H_8ClNO_3S$, **M 233.7, m 149°(dec).** Crystallise the chloride from dry toluene, $CHCl_3$, or ethylene dichloride. Although difficultly soluble in * C_6H_6 , it does however form thick, light tan, crystals from it. It is insoluble in Et_2O . It is useful for making sulfonamides. **IRRITANT**, do not breath it in, and use protective gloves and eye protection. [Smiles & Stewart *Org Synth Coll Col* I 8 1941, DOI: 10.15227/orgsyn.005.0003; Beilstein 14 IV 2703.]

α-Acetamidocinnamic acid [5469-45-4] $C_{11}H_{11}NO_3$, **M 205.2, m 185-186° (2H₂O), 190-191°(anhydrous), 193-195°, pK_{Est} ~3.2.** It crystallises from H_2O as the *dihydrate*, and on drying at 100° it forms the *anhydrous* compound which is *hygroscopic*. Alkaline hydrolysis yields NH_3 and phenylpyruvic acid. [Erlenmeyer & Früstück *Justus Liebigs Ann Chem* 284 47 1895, DOI: 10.1002/jlac.18952840105; Beilstein 14 IV 1769.]

2-Acetamidofluorene (N-[2-fluorenyl]acetamide) [53-96-3] $C_{15}H_{13}NO$, **M 223.3, m 194°, 196-198°.** Recrystallise it from toluene (1.3mg in 100ml). Its solubility in H_2O is 1.3mg/l at 25°, its UV has λ_{max} at nm(log ϵ): 288(4.43), 313(4.13). [Sawicki *J Org Chem* 21 271 1956, DOI: 10.1021/jo01109a002.] It can also be recrystallised from 50% AcOH. [Diels et al. *Chem Ber* 35 3284 1902, DOI: 10.1002/cber.190203503136]. 9- ^{14}C and ω - ^{14}C 2-acetamidofluorene were recrystallised from aqueous EtOH and had **m** 194-195° and 194° respectively. **Potent CARCINOGEN and handle with care.** It is used as a positive control for measuring carcinogenic activity. [Miller et al. *Cancer Res* 9 504 1949, <http://cancerres.aacrjournals.org/content/9/8/504>; for ^{14}C labeling see Ray & Geiser *Cancer Res* 10 616 1950, <http://cancerres.aacrjournals.org/content/10/10/616>;

Sandin et al. *J Am Chem Soc* **74** 5073 1952, DOI: 10.1021/ja01140a023; *Beilstein* **12** H 3287, **12** IV 3373.]

2-Acetamidophenol [614-80-2] $\text{C}_8\text{H}_9\text{NO}$, **M 151.2**, **m. 209°**, **pK_{Est} ~9.4**. Recrystallise it from water, EtOH or aqueous EtOH. [*Beilstein* **13** H 370, **13** I 113, **13** II 171, **13** III 778.] **3-Acetamidophenol (Metacetamol)** [621-42-1] $\text{C}_8\text{H}_9\text{NO}$, **M 151.2**, has **m 148-149°**, **pK₂₅ ~9.59**. Recrystallise the phenol from water. The **3,5-dinitrobenzamide complex** gives orange-yellow crystals from hot H_2O and has **m 212°**. [*Beilstein* **13** IV 977.]

4-Acetamidophenol (Paracetamol, Acetaminophen, 4'-hydroxyacetanilide) [103-90-2] $\text{C}_8\text{H}_9\text{NO}$, **M 151.2**, has **m 169-170.5°**, **pK_{Est} ~10.0**. Recrystallise *Paracetamol* from water or EtOH. It is a highly used *analgesic* and is a febrifuge. The **3,5-dinitrobenzamide complex** gives orange crystals from hot H_2O and has **m 171.5°**. [*Beilstein* **13** II 243, **13** III 1056, **13** IV 1091.]

p-Acetamidophenylacetic acid (Actarit) [18699-02-0] $\text{C}_{10}\text{H}_{11}\text{NO}_3$, **M 193.2**, **m 167°**, **168-170°**, **173-175°**, **174-175°**, **pK₂₅ 3.49**. Crystallise the acid from MeOH/ Me_2CO , aqueous EtOH or H_2O . The *amide* has **m 231°** (from 50% aqueous EtOH). [Gabriel *Chem Ber* **15** 834 1882, DOI: 10.1002/cber.188201501180; Cerecedo & Sherwin *J Biol Chem* **42** 217 1924; Tramontano et al. *J Am Chem Soc* **110** 2282 1988, DOI: 10.1021/ja00215a045; *Beilstein* **14** II 281.]

Acetanilide [103-84-4] $\text{C}_8\text{H}_9\text{NO}$, **M 135.2**, **m 114°**, **pK₂₅ 0.5**. Recrystallise acetanilide from water, aqueous EtOH, *benzene or toluene. [*Beilstein* **12** IV 373.]

Acetoacetanilide [102-01-2] $\text{C}_{10}\text{H}_{11}\text{NO}_2$, **M 177.2**, **m 85°**, **86°**, **pK₂₅ 10.68**. Crystallise the anilide from H_2O , aqueous EtOH or petroleum ether (b 60-80°). [Williams & Krynitsky *Org Synth Coll Vol* **3** 10 1955, DOI: 10.15227/orgsyn.021.0004.]

4-Acetophenetidide (Phenacetin, p-methoxyacetanilide) [62-44-2] $\text{C}_{10}\text{H}_{13}\text{NO}_2$, **M 179.2**, **m 136°**. Crystallise it from H_2O or EtOH, and its solubility in H_2O is 0.08% (at ~10°) and 1.2% (at ~100°), and in EtOH it is 6.7% (at ~10°) and 36% (at ~100°). *Alternatively*, it can be purified by solution in cold dilute alkali and reprecipitating by addition of acid to neutralisation point. Dry it in air. It is an analgesic and a febrifuge. [*Beilstein* **13** H 461, **13** IV 1092.]

Acetophenone [98-86-2] $\text{C}_8\text{H}_8\text{O}$, **M 120.2**, **m 19.6°**, **20.5°**, **b 54°/2.5mm**, **202°/760mm**, **d₄²⁵ 1.0238**, **n_D²⁵ 1.5322**, **pK₂₆ -7.6(basic)**, **pK₂₅ 19.2(acidic)**. Dry it by fractional distillation or by standing with anhydrous CaSO_4 or CaCl_2 for several days, followed by fractional distillation under reduced pressure (from P_2O_5 , optional), and careful, slow and repeated partial crystallisations from the liquid at 0° excluding light and moisture. It can also be crystallised at low temperatures from isopentane. Distillation can be followed by purification using gas-liquid chromatography [Earls & Jones *JCS Faraday Trans 1* **71** 2186 1975, DOI: 10.1039/F19757102186.] [*Beilstein* **7** H 271, **7** I 146, **7** II 208, **7** III 936, **7** IV 619.] It has hypnotic properties. § A commercial polystyrene supported version is available — scavenger resin (for diol substrates).

Aceto-o-toluidide (2-methylacetanilide) [120-66-1] $\text{C}_9\text{H}_{11}\text{NO}$, **M 149.2**, **m 110°**, **112°**, **b 176°/14mm**, **296°/760mm**. Crystallise the toluidide from hot H_2O (solubility 1g/210ml), EtOH or aqueous EtOH. Its UV has λ_{max} at 230 and 280nm (EtOH). [*Beilstein* **12** H 792, **12** I 376, **12** II 439, **12** III 1853, **12** IV 1755.] **Aceto-m-toluidide (3-methylacetanilide)** [537-92-8] has **m 65.5°**, **b 182-183°/14mm**, **303°/760mm**, **303°/760mm**. Crystallise the toluidide from H_2O , EtOH, aqueous EtOH or Et_2O /petroleum ether (**m 66°**). Its UV has λ_{max} at 245nm (EtOH). [*Beilstein* **12** H 860, **12** I 400, **12** II 468, **12** III 1962, **12** IV 1823.] **Aceto-p-toluidide (4-methylacetanilide)** [103-89-9] has **m 146°**, **b 307°/760mm**. Crystallise it from aqueous EtOH. [*Beilstein* **12** H 920, **12** I 420, **12** II 501, **12** III 2051, **12** IV 1902.]

R(-)- α -Acetoxyphenylacetic (acetyl mandelic) acid [51019-43-3] $\text{C}_{10}\text{H}_{10}\text{O}_4$, **M 194.2**, **m 96-98°**, **97-99°**, **[α]_D²⁰ -153.7 (c 2.06, Me_2CO)**, **[α]₅₄₆²⁰ -194 (c 2.4, Me_2CO)**, **pK_{Est} ~2.9**. It crystallises from H_2O with 1mol of solvent which is removed on drying, or from other solvents as for the *S*-isomer below. [Angus & Owen *J Chem Soc* 227 1943, DOI: 10.1039/JR9430000227; Parker *Chem Rev* **91** 1441 1991, DOI: 10.1021/cr00007a009; *Beilstein* **10** III 453.] **S-(+)- α -Acetoxyphenylacetic (acetyl mandelic) acid** [7322-88-

5] has **m** 80-81°, **95-97.5°**, $[\alpha]_D^{27} +158$ (c 1.78, Me₂CO), $[\alpha]_{546}^{20} +186$ (c 2, Me₂CO). Recrystallise it from *benzene/hexane or toluene. It has characteristic NMR and IR spectra. [Pracejus *Justus Liebigs Ann Chem* **622** 10 1959, DOI: 10.1002/jlac.19596220104; Breitholle & Stammer *J Org Chem* **39** 1311 1974, DOI: 10.1021/jo00923a033; *Beilstein* **10** IV 567.]

1-Acetylanthracene [7396-21-6] C₁₆H₁₂O, **M** 220.3, **m** 109.4-109.8°, **110.5-111°** (uncorrected), **115.5-116°** (corrected), **pK_{Est}** 19.0. Recrystallise from EtOAc. Most likely impurities are the 2-isomer, to a lesser extent the 9-isomer and to a yet lesser extent traces of diacetylanthracenes. These can be checked by TLC on silica gel G (Merck) and one dimensional multiple development with *C₆H₆/CHCl₃ and visualising under a UV lamp. On a larger scale chromatography through a column of activated Al₂O₃ (e.g. Spence, Type H) and the mono-acetylanthracenes are eluted with *C₆H₆. Further elution with *C₆H₆/CHCl₃ affords diacetylanthracenes. The fractions are checked by TLC. [Gore & Thadani *J Chem Soc (C)* 1729 1966, DOI: 10.1039/J39660001729; 1498 1967, DOI: 10.1039/J39670001498.] Also obtained in almost equal proportion with the 9-isomer by acylation with Ac₂O/I₂ in CH₂Cl₂ (7 hours reflux, 43% yield) and purified by TLC. The UV has λ_{max} (EtOH) 2410, 2550, 3640, 3850nm and 3490shnm (ε 61,600, 68,900, 5,500, 6,050 and 4,300 resp.) [Gore & Hoskins *J Chem Soc* 5737 1965, DOI: 10.1039/JR9650005737], and the IR has ν_{max} (CCl₄) at 1686cm⁻¹ (C=O). [Gore & Thadani *J Chem Soc (C)* 1729 1966, DOI: 10.1039/J39660001729]. The **oxime** has **m** 162-163° (from EtOH) [Bergmann & Katz *J Chem Soc* 3216 1958, DOI: 10.1039/JR9580003214], and the **2,4-dinitrophenylhydrazone** has **m** 260° (red crystals from pyridine/EtOH). [For crystallography see Langer & Becker *Z Krist* **206** 155 1993.] 1-Acetylanthracene dimerises in CH₂Cl₂ upon irradiation with light at 400nm [Becker et al. *J Photochem Photobiol* **97** 25 1996, DOI: 10.1016/1010-6030(96)04318-3].

2-Acetylanthracene [10210-32-9] C₁₆H₁₂O, **M** 220.3, **m** 190-192°, **195.5-196°** (uncorrected), **205.5-206°** (corrected), **pK_{Est}** 19.2. This is the thermodynamically stable isomer in the Friedel-Crafts reaction and is obtained, together with a smaller amount of 1-acetylanthracene (m 107.5-109°) by the AlCl₃-catalysed rearrangement of 9-acetylanthracene (see below) in nitrobenzene [Hawkins *J Chem Soc* 3858 1957, DOI: 10.1039/JR9570003858]. Similarly the 1- and 9-isomers are rearranged to the 2-isomer completely when heated with polyphosphoric acid (PPA) at 80-120°. [Mala'bi et al. *Letters in Organic Chemistry* **6** 237 2009, DOI: 10.2174/157017809787893118.] Purify 2-acetylanthracene by recrystallisation from boiling EtOAc (charcoal) then light petroleum (b 80-100°). It also crystallises from EtOH (**m** 183-185°), *C₆H₆ (yellow plates, **m** 189-189.5°) or light petroleum (**m** 190-192°). Important impurities are the 1-isomer and to a lesser extent the 9-isomer and diacetylanthracenes. For purification from these see 1-acetylanthracene above. the IR has ν_{max} (CHCl₃) at 1676cm⁻¹ (C=O). [Gore & Thadani *J Chem Soc (C)* 1729 1966, DOI: 10.1039/J39660001729]. The **oxime** has **m** 247-248 (from EtOH) [Bergmann & Katz *J Chem Soc* 3214 p3216 1958, DOI: 10.1039/JR9580003214], the **phenylhydrazone** has **m** 262° (from EtOH), and the **2,4-dinitrophenylhydrazone** has **m** 297° (bright red crystals from pyridine) [Gore & Thadani *J Chem Soc (C)* 1729 1966, DOI: 10.1039/J39660001729]. The **semicarbazone** has **m** 316° (from EtOH or AcOH). [*Beilstein* **7** H 450, **7** III 2538, **7** IV 1743.]

9-Acetylanthracene [784-04-3] C₁₆H₁₂O, **M** 220.3, **m** 75-76°, **80°**, **76.5-77.3°** (uncorrected), **79.6-80.4°** (corrected), **pK_{Est}** 18.9. The 9-position (*meso*) in anthracene is the most reactive position in the Friedel-Crafts reaction and 9-acylation is kinetically favoured, thus forming 9-acetylanthracene under the mildest conditions [Gore & Thadani *J Chem Soc (C)* 1729 1966, DOI: 10.1039/J39660001729]. Dry 9-acetylanthracene *in vacuo*, then recrystallise it from EtOH (**m** 80°), from EtOH then EtOAc (charcoal, **m** 74-76°), or cyclohexane (**m** 82°). Its UV has λ_{max} (EtOH) 2530, 3300, 3480, 3630 and 3800nm (ε 128,000, 2,650, 5,950, 7,900 and 6,800 resp.) [Gore & Hoskins *J Chem Soc* 5737 p 5744 1965, DOI: 10.1039/JR9650005737], and the IR has ν_{max} (CCl₄) at 1705cm⁻¹ (C=O). [Gore & Thadani *J Chem Soc (C)* 1729 1966, DOI: 10.1039/J39660001729]. The **oxime** has **m** 185-186° (from EtOH) [Bergmann & Katz *J Chem Soc*, 3214 p 3216 1958, DOI: 10.1039/JR9580003214], and the **2,4-dinitrophenylhydrazone** has **m** 258° (dark red crystals from pyridine/EtOH), prepared under forcing conditions because of steric hindrance: i.e. by refluxing the ketone (1g) and 2,4-dinitrophenylhydrazine (1g), conc HCl (5ml), H₂O (5ml) and EtOH (40ml), cool, (seed if necessary), collect the crystals, wash thoroughly with hot 10% aqueous HCl, then H₂O, dry *in vacuo* and recrystallise [Pearson & Greer *J Am Chem Soc* **77** 1294 1955, DOI: 10.1021/ja01610a068]. The **semicarbazone** forms yellow crystals **m** 265° (from EtOH). [Merritt Jr & Braun *Org Synth Coll Vol* **4** 8 1943, DOI: 10.15227/orgsyn.030.0001; *Beilstein* **7** III 2539, **7** IV 1743.]

N-Acetylanthranilic acid (2-acetamidobenzoic acid) [89-52-1] $C_9H_9NO_3$, M 179.1, m 182-184°, 185-186°, 184-187°, 190°(dec), pK^{20} 3.61. Wash the acid with distilled H_2O and recrystallise it from aqueous AcOH, dry it and recrystallise again from EtOAc. Also recrystallise it from water or EtOH. Its UV has λ_{max} at 221, 252 and 305nm (EtOH). The **amide** crystallises from aqueous EtOH and has m 186-187° and λ_{max} 218, 252 and 301nm. [Chattaway *J Chem Soc* 2495 1931, DOI: 10.1039/JR9310002495; Walker *J Am Chem Soc* 77 6698 1955, DOI: 10.1021/ja01629a120; *Beilstein* 14 H 337, 14 I 540, 14 II 219, 14 III 922, 14 IV 1041.]

2-Acetylbenzoic acid [577-56-0] $C_9H_8O_3$, M 164.2, m 115-116°, 115-117°, 116-118°, pK^{20} 4.14, pK^{25} 4.10. It crystallises from $*C_6H_6$ and H_2O (15g/100ml). The **oxime** has m 156-157°, and the **2,4-dinitrophenylhydrazone** has m 185-186°(needles from EtOH). [Yale *J Am Chem Soc* 69 1547 1947, DOI: 10.1021/ja01198a519; Panetta & Miller *Synthesis* 43 1977, DOI: 10.1055/s-1977-24269; *Beilstein* 10 H 690, 10 I 330, 10 II 479, 10 III 3025, 10 IV 2766.] **3-Acetylbenzoic acid** [586-42-5] has m 161-172°, 169-171°, 172°, pK^{25} 3.827, (5.21 in 50% aqueous EtOH). It was synthesised from *m*-nitroacetophenone via *m*-amino- to the *m*-cyano- [6136-68-1, see below] intermediate which was hydrolysed (reflux with concentrated HCl/6hrs) to the acid, and recrystallised from conductivity H_2O or aqueous Me_2CO , and dried in a vacuum desiccator over silica gel. It is slightly soluble in H_2O , EtOH or Et_2O but very soluble in $CHCl_3$ and $*C_6H_6$. [Bray et al. *J Chem Soc* 265 1957, DOI: 10.1039/JR9570000265; *Beilstein* 10 H 694, 10 IV 2769.] **4-Acetylbenzoic acid** [586-89-0] has m 207.5-209.5°, 208.6-209.4°, 208-210°, pK^{25} 3.70, (5.10 in 50% aqueous EtOH). Dissolve the acid in 5% aqueous NaOH, extract it with Et_2O , and acidify the aqueous solution. Collect the precipitate, and recrystallise it from boiling H_2O (100 parts) using decolorising charcoal [Pearson et al. *J Org Chem* 24 504 1959, DOI: 10.1021/jo01086a015; Bray et al. *J Chem Soc* 265 1957, DOI: 10.1039/JR9570000265; Detweiler & Amstutz *J Am Chem Soc* 72 2882 1950, DOI: 10.1021/ja01163a021; pK_a : Bordwell & Cooper *J Am Chem Soc* 74 1058 1952, DOI: 10.1021/ja01124a057]. [*Beilstein* 10 IV 2769.]

3-Acetylbenzonitrile (*m*-cyanoacetophenone) [6136-68-1] C_9H_7NO , M 145.2, m 98-99°, 98-100°. It was prepared by a Sandmeyer reaction on the corresponding 3-aminoacetophenone below. Recrystallise the nitrile from hot EtOH (m 98°) and using decolourising charcoal. UV (hexane) has λ_{max} (log ϵ) 3420 (2.66) and 3518 (3.70) nm. [Prepn. and UV spectra: Pestemer et al. *Monatsh Chem* 68 326 1936, DOI: 10.1007/BF01518872]. [*Beilstein* 10 H 694, 10 III 3028.] **4-Acetylbenzonitrile (*p*-cyanoacetophenone)** [1443-80-7] has m 57-58°, 56-59°. Recrystallise the nitrile (prepared from the corresponding 4-aminoacetophenone below using a Sandmeyer reaction) from EtOH (charcoal). After two sublimations (bath temperature 130°) colourless needles, m 56.8°, are obtained. UV (hexane) has λ_{max} (log ϵ) 3382 (3.18), 3472 (3.28), 3504 (3.28) and 4078 (4.47) nm. [Prepn. and UV spectra: Pestemer et al. *Monatsh Chem* 68 326 1936, DOI: 10.1007/BF01518872; Wagner et al. *J Am Chem Soc* 108 7727 1986, DOI: 10.1021/ja00284a041]. [*Beilstein* 10 H 695, 10 III 3030.]

Acetyl-5-bromosalicylic acid [1503-53-3] $C_9H_7BrO_4$, M 259.1, m (156°), 168°, 168-169°, pK_{Est} ~3.0. Crystallise the acid from EtOH. [Robertson *J Chem Soc* 81 1475 1902, DOI: 10.1039/CT9028101475; *Beilstein* 10 H 108, 10 II 64.]

2-Acetylfluorene [781-73-7] $C_{15}H_{12}O$, M 208.3, m 128-129°, 130-131°, 132°. Crystallise 2-acetylfluorene from EtOH (solubility is 60g/800ml) or Me_2CO (solubility is 60g/400ml). The **oxime** [110827-07-1] has m 192-193.5° and the **2,4-dinitrophenylhydrazone** [109682-26-0] has m 261-262°. [Ray & Rieveschl *Org Synth Coll Vol* 3 23 1973, DOI: 10.15227/orgsyn.028.0003.]

5(3)-Acetyl-2(6)-methoxybenzaldehyde [531-99-7] $C_{10}H_{10}O_3$, M 178.2, m 141-143°, 143-144°, 144°. Extract a solution of the aldehyde in $*C_6H_6$ with 20% aqueous sodium bisulfite, and the bisulfite adduct in the aqueous solution is decomposed by acidifying and heating, whereby the aldehyde separates. It is collected, washed with H_2O , and dried in a vacuum. It is recrystallised from EtOH and then from Et_2O . It sublimes *in vacuo*. The **2,4-dinitrophenylhydrazone** (prepared by adding a cold saturated solution of 2,4-diphenylhydrazine in 95% EtOH to 5mg of aldehyde in 1ml of EtOH followed by a drop of concentrated HCl, whereby the orange derivative separates immediately) has m 258-161° after two crystallisations from EtOH and dried. The derivative is insoluble in most organic solvents but can be recrystallised from nitrobenzene. [Gray & Bonner *J Am Chem Soc* 70 1249 1948, DOI: 10.1021/ja01183a114; Angyal et al. *J Chem Soc* 2141 1950, DOI: 10.1039/JR9500002141; *Beilstein* 8 III 2339, 8 IV 1984.]

4-Acetyl-N-methylaniline (4-methylaminoacetophenone) [17687-47-7] $C_9H_{11}NO$, M 149.2, m 102-106°, 103-107°. This herbicide crystallises from H_2O . The **4-acetyl-N,N-dimethylaniline** derivative forms colourless plates also from H_2O with m 58-59°. [Klingel *Chem Ber* **18** 2687 1885, DOI: 10.1002/cber.188501802177; Staudinger & Kon *Justus Liebigs Ann Chem* **384** 111 1911, DOI: 10.1002/jlac.19113840103; *Beilstein* **14** H 47, **14 I** 366.]

1-Acetylnaphthalene (1-acetonaphthenone) [941-98-0] $C_{12}H_{10}NO$, M 170.1, m 10.5°, b 93-95°/0.1mm, 167°/12mm, 302°/760mm, d_4^{20} 1.12, pK^{25} -6.22 (H_0 scale, aqueous H_2SO_4). If the NMR spectrum indicates the presence of impurities, probably 2-acetylnaphthalene, convert the substance to its **picrate** by dissolving in *benzene or EtOH and adding excess of saturated picric acid in these solvents until separation of picrates is complete. Recrystallise the picrate till the melting point is 118°. Decompose the picrate with dilute NaOH and extract with Et_2O . Dry the extract (Na_2SO_4), filter, evaporate and distil the residue. The **2,4-dinitrophenylhydrazone** crystallises from EtOH and has m 259°. [Stobbe et al. *Justus Liebigs Ann Chem* **380** 93 1911, DOI: 10.1002/jlac.19113800114; Williams & Osborne *J Am Chem Soc* **61** 3438 1939, DOI: 10.1021/ja01267a061; *Beilstein* **7** IV 1292.] **2-Acetylnaphthalene (2-acetonaphthenone, β -acetonaphthone, 2-acetonaphthalene, methyl-2-naphthylketone)** [93-08-3] has m 52-53°, 55°, 55.8°, b 164-166°/8mm, 171-173°/17mm, 301-303°/760mm, pK^{25} -6.16 (H_0 scale, aqueous H_2SO_4). Separate it from the 1-isomer by fractional crystallisation of the **picrate** in EtOH (see entry for the 1-isomer above) to m 82°. Decomposition of the picrate with dilute NaOH and extraction with Et_2O , then evaporation, gives purer 2-acetylnaphthalene. If this residue solidifies, it can be recrystallised from petroleum ether, EtOH or acetic acid; otherwise it should be distilled in a vacuum and the solid distillate recrystallised [Gorman & Rodgers *J Am Chem Soc* **108** 5074 1986, DOI: 10.1021/ja00277a005; Levanon et al. *J Phys Chem* **91** 14 1987, DOI: 10.1021/j100285a006]. Purity should be checked by high field NMR spectroscopy. Its **oxime** has m 145°(dec), and the **semicarbazone** has m 235°. [Stobbe et al. *Justus Liebigs Ann Chem* **380** 93 1911, DOI: 10.1002/jlac.19113800114; Raffauf *J Am Chem Soc* **72** 753 1950, DOI: 10.1021/ja01158a028; Hunsberger *J Am Chem Soc* **72** 5626 1950, DOI: 10.1021/ja01168a074; Immediata & Day *J Org Chem* **05** 512 1940, DOI: 10.1021/jo01211a005; *Beilstein* **7** IV 1294.]

1-Acetyl-2-phenylhydrazine [114-83-0] $C_8H_{10}N_2O$, M 150.2, m 128.5°, pK^{25} 1.3. Crystallise the hydrazine from aqueous EtOH. [*Beilstein* **15** H 241.]

Acetylsalicylic acid (Aspirin) [50-78-2] $C_9H_8O_4$, M 180.2, m 133.5-135°, 134-136° (and various), pK^{25} 3.38, (pK^{17} 3.56). Crystallise aspirin twice from toluene, wash it with cyclohexane and dry it at 60° under vacuum for several hours [Davis & Hetzer *J Res Nat Bur Stand* **60** 569 1958, DOI:org/10.6028/jres.060.057]. It has been recrystallised from isopropanol and from diethyl ether/petroleum ether (b 40-60°). It crystallises from EtOH (m 143-144°), * C_6H_6 (m 143°), hexane (m 115° and 128°), octane (m 121°), and has m 110° after sublimation. It has pK^{26} 3.69(H_2O), 4.15(20% aqueous EtOH), 4.47(30% aqueous EtOH) and 4.94(40% aqueous EtOH). It is a commonly used analgesic. [*Beilstein* **10** H 67, **10** II 41, **10** III 102, **10** IV 138.] The anhydride, **O-acetylsalicylic anhydride** [1466-82-6] $C_{18}H_{14}O_7$, M 342.3, has m 80-83° (* C_6H_6 -petroleum ether). [*Beilstein* **10** IV 165.]

O-Acetylsalicyloyl chloride [5538-51-2] $C_9H_7ClO_3$, M 198.6, m 45°, 46-49°, 48-52°, b 107-110°/0.1mm, 115°/5mm, 135°/12mm, n_D^{20} 1.536. Check first the IR to see if an OH frequency is present. If so, some free acid is present. Then reflux with acetyl chloride for 2-3 hours and fractionate at high vacuum. The distillate should crystallise on cooling. It can be recrystallised from hexane or * C_6H_6 (m 60°, sintering at 52°). [Riegel & Wittcoff *J Am Chem Soc* **64** 1486 1942, DOI: 10.1021/ja01213a038; *Beilstein* **10** H 86, **10** I 43, **10** II 55, **10** III 151, **10** IV 169.]

O-Acetylsalicylsalicylic acid (Salsalate acetate) [530-75-6] $C_{16}H_{12}O_6$, M 300.3, m 152°, 159°. Crystallise the acetate from dilute AcOH or EtOH (m 161-162°), MeOH (m 165-168°), and * C_6H_6 /EtOH (m 163-165°). Its solubilities in boiling Et_2O , * C_6H_6 and EtOH are 1.4%, 2.2% and 33%, respectively. It has analgesic properties. [Baker et al. *J Chem Soc* 201 1951 DOI: 10.1039/JR9510000201; Garrett et al. *J Am Pharm Soc* **48** 684 1959, *Beilstein* **10** I 41, **10** II 54, **10** IV 165.]

N-(4)-Acetylsulfanilamide (Sulfacetamide) [144-80-9] $\text{C}_8\text{H}_{10}\text{N}_2\text{O}_3\text{S}$, **M 214.2**, **m 182-184°**, **216°**, **pK₁ 1.8 (amino, basic)**, **pK₂ 5.4 (sulfonamide NH, acidic)**. Crystallise the amide from aqueous EtOH (colourless or pale yellow prisms). Its solubility (w/v) at 20° in H_2O is 0.67%, in EtOH it is 6.7% and in Me_2CO it is 14.3 %. Its UV has λ_{max} at 258nm (ϵ 17,700) in H_2O ; 217nm (ϵ 18,900) in 0.1M HCl; 270nm (ϵ 8,900) in 1M HCl and 257nm (ϵ 17,300) in 0.1M NaOH. The **sodium salt** [127-56-0 **anhydrous**] **M 236.2**, $\text{C}_8\text{H}_9\text{N}_2\text{NaO}_3\text{S}$, [6029-17-2 **monohydrate**] **M 253.3**, $\text{C}_8\text{H}_{11}\text{N}_2\text{NaO}_4\text{S}$, crystallises in very small prisms from aqueous EtOH with **m 257°**. It is sparingly soluble in 96% EtOH, and mostly insoluble in other organic solvents, but is 67% (w/v) soluble in H_2O . [Beilstein 14 IV 2662.]

Acetyl p-toluenesulfonate [26908-82-7] $\text{C}_8\text{H}_{10}\text{O}_5\text{S}$, **M 230.2**, **m 54-56°**, **b 186-188°/20mm**. The most likely impurity is p-toluenesulfonic acid (could be up to 10%). This can be removed by dissolving it in dry Et_2O and cooling until the anhydride crystallises out. It decomposes on heating; below ~130° it gives the disulfonic anhydride and above ~130° polymers are formed, but it can be distilled in a vacuum if it is free of acid. It is used for cleaving ethers [Prep, IR, NMR: Mazur & Karger *J Org Chem* 36 528, DOI: 10.1021/ja01213a038; Mazur & Karger *J Org Chem* 36 532 1971, DOI: 10.1021/jo00803a010]. [Beilstein 11 III 255.]

Alizarin (1,2-dihydroxyanthraquinone, Mordant Red 11) [72-48-0] $\text{C}_{14}\text{H}_8\text{O}_4$, **M 240.2**, **m 279-283°**, **290°**, **d₄²⁰ 0.884**, **CI 58000**, **pK₁²⁵ 7.45**, **pK₂²⁵ 11.80**. Alizarin crystallises from glacial acetic acid or 95% EtOH. It sublimes at 110°/2mm as orange needles (m 290°). It is an indicator with λ_{max} at 452nm (pH 5.8) and 520nm (pH 7.2,), and colour changes at 5.5-6.8, yellow to red (acidic) and 10.1 to 12.1, red to violet (alkaline); and stains biological materials. [Beilstein 8 IV 3256.]

Alizarin-3-methyliminodiacetic acid (Alizarin Complexone) (2H₂O) [3952-78-1] $\text{C}_{19}\text{H}_{15}\text{NO}_8$, **M 385.3**, **m 189°(dec)**, **pK_{Est(1)}~4.9**, **pK_{Est(2)}~7.5**. It is purified by suspending it in 0.1M NaOH (1g in 50ml), filtering the solution and extracting alizarin with 5 successive portions of CH_2Cl_2 . Then add HCl dropwise to precipitate the reagent, stirring the solution in an ice bath. Filter the precipitate onto a glass filter, wash it with cold water and dry it in a vacuum desiccator over KOH [Ingman *Talanta* 20 135 1973, DOI: 10.1016/0039-9140(73)80243-7; Beilstein 14 IV 931].

Alizarin Yellow R [5-(4-nitrophenylazosalicylic acid), Mordant Orange I] [2243-76-7] $\text{C}_{13}\text{H}_9\text{N}_3\text{O}_5$, **M 287.2**, **m 253-254°(dec)**, **>300°**, **CI 14030**, **pK²⁵ 11.17**. The free acid is precipitated by adding HCl to an aqueous solution of the Na salt. After 2 recrystallisations from aqueous AcOH, it has **m 255°(dec)**; [m 253-254°(dec) was reported by Hewitt & Fox *J Chem Soc* 79 49 1901, DOI: 10.1039/CT9017900049]. The free acid recrystallises from dilute AcOH as orange brown needles. The Na salt changes colour from yellow to red when the pH is increased from 10.2 to 12.0. [Woodland et al. *J Am Chem Soc* 75 5835 1953, DOI: 10.1021/ja01119a018; Beilstein 16 IV 372.]

Allyl phenyl sulfide [5296-64-0] $\text{C}_9\text{H}_{10}\text{S}$, **M 150.2**, **b 59-60°/1.5mm**, **79-80°/3mm**, **114-114.3°/23.5mm**, **225-226°/740mm**, **215-218°/750mm**, **d₄²⁰ 1.0275**, **n_D²⁰ 1.5760**. Dissolve the sulfide in Et_2O , wash with alkali, H_2O , dry over CaCl_2 , evaporate and fractionally distil it, preferably under vacuum. It should **not** give a precipitate with an alcoholic solution of $\text{Pb}(\text{OAc})_2$. [Hurd & Greengard *J Am Chem Soc* 52 3356 1930, DOI: 10.1021/ja01371a054; Tarbell & McCall *J Am Chem Soc* 74 48 1952, DOI: 10.1021/ja01121a013; Beilstein 6 IV 1479.]

Amberlite IRA-904 Anion exchange resin (Rohm and Haas) [9050-98-0]. Wash with 1M HCl, CH_3OH (1:10) and then rinse it with distilled water until the washings are neutral to litmus paper. Finally extract successively for 24 hours in a Soxhlet apparatus with MeOH, *benzene and cyclohexane [Shue & Yen *Anal Chem* 53 2081 1981, DOI: 10.1021/ac00236a030]. It is a strong basic resin also used for base catalysis [Fieser 1 511].

p-Aminoacetanilide [122-80-5] $\text{C}_8\text{H}_{10}\text{N}_2\text{O}$, **M 150.2**, **m 162-163°**, **163°**, **165-167°**, **166-167°**, **pK¹⁵ 4.46**, **pK⁴⁰ 3.94**. Crystallise the anilide from water. It has an unstable crystalline form with **m 141°**. It has IR with ν_{max} (CCl_4) at 1681cm^{-1} . [Atkinson et al. *J Chem Soc* 2023 1954, DOI: 10.1039/JR9540002023; Beilstein 13 H 94, 13 I 28, 13 II 50, 13 III 166, 13 IV 137.] The *ortho*-isomer, **o-aminoacetanilide** [34801-09-7] has **m 133-**

137°, $pK_{\text{Est}} \sim 4.0$, and the *meta*-isomer, *m*-aminoacetanilide [102-28-3] has **m 86-88°**, $pK_{\text{Est}} \sim 4.2$. [Beilstein 13 H 86.]

***o*-Aminoacetophenone (2-acetylaniline)** [551-93-9] $\text{C}_8\text{H}_9\text{NO}$, **M 135.2**, **m 20°**, **b 85-90°/0.5mm**, **103-105°/1mm**, **113°/6mm**, **124°/10mm**, **130-131°/14mm**, **250-255°/atm**, $d_4^{25} 1.123$, $n_D^{25.1} 1.6632$, $pK^{25} 2.22$. It can be purified by conversion into the *hydrochloride* [25384-14-9] $\text{C}_8\text{H}_9\text{NO} \cdot \text{HCl}$, **M 171.6**, **m 264-265° (285-290°**, also reported) which can be recrystallised from EtOH or H_2O , then the *free base* is liberated with ammonia, collected, dried and distilled. It is steam volatile. *Alternatively*, it can be purified *via* the *picrate*, **m 127°**, which recrystallises from H_2O , and from which the free base can be recovered. Its UV (Et_2O) has λ_{max} at 2764 (log ϵ 3.80) and 3965 (log ϵ 3.87)nm [Prepn. and UV spectra: Pestemer et al. *Monatsh Chem* **68** 326 1936, DOI: 10.1007/BF01518872]. The *N*-acetyl derivative has **m 74-75°**. The *semicarbazone*, **m 290°**, crystallises from EtOH (plates). The *oxime* [4964-49-2] crystallises from H_2O in needles with **m 111-112°**. Parks et al. have identified this ketone in the flavor of stale milk [*Nature* **202** 185 1964, DOI:10.1038/202185b0]. Also found in the mandibular secretion of ants. [Synth: Leonard & Boyd *J Org Chem* **11** 405 1946, DOI: 10.1021/jo01174a018; Beilstein **14** H 42, **14** II 29, **14** III 80, **14** IV 91.]

***m*-Aminoacetophenone (3-acetylaniline)** [99-03-6] $\text{C}_8\text{H}_9\text{NO}$, **M 135.2**, **m 98-99°**, **99.5°**, **b 189-290°/760mm**, $pK^{25} 3.59$. Recrystallise it from EtOH, aqueous EtOH (**m 99.5°**) or C_6H_6 /petroleum ether/EtOH (**m 98-99°**). Its UV (Et_2O) has λ_{max} at 2965 (log ϵ 3.40)nm and (hexane) 3704 (log ϵ 3.36)nm [Prepn. and UV spectra: Pestemer et al. *Monatsh Chem* **68** 326 1936, DOI: 10.1007/BF01518872]. The *hydrochloride* has **m 173°** (from EtOH); the *oxime* has **m 129-130°** (from H_2O), **132°** (from Et_2O / petroleum ether); the *semicarbazone* has **m 196°(dec)** (from H_2O); the *thiosemicarbazone* has **m 152-154° (202-204°** also reported) (from aqueous EtOH or MeOH), and the *benzenesulfonamide* has **m 163-164°** (yellow crystals from MeOH). The *N*-acetyl derivative has **m 127-128°**. [Synth: Leonard & Boyd *J Org Chem* **11** 405 1946, DOI: 10.1021/jo01174a018; Beilstein **14** III 88, **14** IV 96.]

***p*-Aminoacetophenone (4-acetylaniline)** [99-92-3] $\text{C}_8\text{H}_9\text{NO}$, **M 135.2**, **m 104-106°**, **105-107°**, **b 293°/atm**, $pK^{25} 2.19$. Recrystallise it from CHCl_3 , C_6H_6 or H_2O . It is soluble in hot H_2O . Its UV (Et_2O) has λ_{max} at 3356nm (log ϵ 4.40) [Prepn. and UV spectra: Pestemer et al. *Monatsh Chem* **68** 326 1936, DOI: 10.1007/BF01518872]. Vandenbelt et al. *Anal Chem* **26** 726 1954, DOI: 10.1021/ac60088a031.] The **2,4-dinitrophenylhydrazone** has **m 266-267°** (from CHCl_3 or EtOH) with λ_{max} at 403nm (log ϵ 4.42) [Johnson *J Am Chem Soc* **75** 2720 1953, DOI: 10.1021/ja01107a051] and the *semicarbazone* has **m 193-194°(dec)**(from MeOH). The *hydrochloride* has **m 98°(dec)**(from H_2O). [Beilstein **14** IV 100.]

***o*-Aminoacetophenone hydrochloride (phenacylamine hydrochloride, 2-aminoacetophenone HCl)** [5468-37-1] $\text{C}_8\text{H}_9\text{NO} \cdot \text{HCl}$, **M 171.6**, **m 188°(dec)**, **194°(dec)**, $pK^{25} 5.34$. Crystallise the salt from Me_2CO /EtOH, EtOH/ Et_2O , 2-propanol or 2-propanol and a little HCl (slowly after a few days). The *oxime* of the free base has **m 140°**, and the *picrate* of the free base has **m 182°** (from EtOH). [Castro *J Am Chem Soc* **108** 4179 1986, DOI: 10.1021/ja00274a055; Baumgarten & Petersen *Org Synth Coll Vol* **5** 909 1973, DOI: 10.15227/orgsyn.041.0082; cf. Beilstein **14** H 49, **14** III 105.]

1-Aminoanthraquinone-2-carboxylic acid [82-24-6] $\text{C}_{15}\text{H}_9\text{NO}_4$, **M 276.2**, **m 295-296°**. Crystallise the acid from nitrobenzene (red needles). It is used for the detection of Al, Mg Cd, Zn, Mn, Cu, Hg, Fe, Co, Ni and Pb. The *methyl ester* gives red needles from AcOH, **m 228°**. The *phenyl ester*, **m 198°**, crystallises as golden-red needles from AcOH. [Locher & Fierz *Helv Chim Acta* **10** 642 1927, DOI: 10.1002/hlca.19270100183; Beilstein **14** II 419, **14** III 168.]

***p*-Aminoazobenzene (*p*-phenylazoaniline, Aniline yellow)** [60-09-3] $\text{C}_{12}\text{H}_{11}\text{N}_3$, **M 197.2**, **CI 11000**, **m 126°**, **128°**, **b >360°**, $pK^{25} \sim 2.82$. Crystallise this dye from EtOH, CCl_4 , petroleum ether/ C_6H_6 , or a MeOH/ H_2O mixture (brown-yellow needles with a blue lustre). It is soluble in most organic solvents. [Beilstein **16** IV 445.]

***o*-Aminoazotoluene (Fast Garnet GBC base, 4'-amino-2,3'-dimethylazobenzene, Solvent yellow 3)** [97-56-3] $\text{C}_{14}\text{H}_{15}\text{N}_3$, **M 225.3**, **m 101-103°**, **101.4-102.6°**, **CI 11160**, $pK^{26} 2.29$ (50% aqueous EtOH). Recrystallise the dye twice from EtOH, once from C_6H_6 , then dry it in an Abderhalden drying apparatus. [Cilento *J Am*

Chem Soc **74** 968 1952, DOI: 10.1021/ja01124a029; Sawicki *J Org Chem* **21** 605 1956, DOI: 10.1021/jo01112a002; *Beilstein* **16** H 334, **16** I 322, **16** II 178, **16** III 386, **16** IV 525.] **CARCINOGENIC**.

2-Aminobenzaldehyde [529-23-7] C_7H_7NO , **M 121.1**, **m 39-40°**, **80-82°/2mm**, **pK²⁰ 1.36**. Distil it in steam and recrystallise it from H_2O or EtOH/ Et_2O . The *semicarbazone* has **m 247°**. [*Beilstein* **14** H 21, **14** I 356, **14** II 14, **14** III 47, **14** IV 42.] The derivative **2-aminobenzaldehyde phenylhydrazone** (**Nitrin**) [63363-93-9] $C_{13}H_{13}N_3$, **M 211.3**, has **m 227-229°**. Crystallise it from acetone. [Knöpfer *Monatsh Chem* **31** 87 1910, DOI: 10.1007/BF01520385; *Beilstein* **14** H 21, **14** II 14, **14** III 47.] **3-Aminobenzaldehyde** [29159-23-7] has **m 28-30°**, **pK_{Est} ~2.0**. The aldehyde crystallises as light yellow plates from ethyl acetate. The UV has λ_{max} at 227 and 327.5nm in cyclohexane. The *acetyl* derivative has **m 122°** (from EtOH) and the *oxime* has **m 195°** (yellow-brown plates from EtOH). [*Beilstein* **14** H 28, **14** I 359, **14** II 21, **14** III 53, **14** IV 46.]

4-Aminobenzamide hydrochloride [59855-11-7] $C_7H_8N_2O$. HCl, **M 172.6**, **m 284-285°**, **pK_{Est} ~1.7**. Recrystallise the salt from EtOH. The *free base* [2835-68-9] **M 136.2**, has **m 182.9°** and crystallises with $0.25H_2O$ (**m 178-179°**). [Rupe & Vogler *Helv Chim Acta* **8** 832 1925, DOI: 10.1002/hlca.192500801115; *Beilstein* **14** H 425, **14** III 1061.]

p-Aminobenzeneazodimethylaniline (*N,N*-dimethyl 4,4'-azodianiline) [539-17-3] $C_{14}H_{16}N_4$, **M 240.3**, **m 186-187°**, **190°(dec)**, **191-191.3°**, **191.5-192°**. Crystallise the azo-dye from aqueous EtOH. It has UV (95% EtOH) has λ_{max} at 410nm (log ϵ 4.48) + 440nm (log ϵ 4.46), and (25% aqueous EtOH) at 418s nm (log ϵ 4.36) + 458nm (log ϵ 4.48). [Sawicki *J Org Chem* **22** 915 1957, DOI: 10.1021/jo01359a016; 1084 1957, DOI: 10.1021/jo01360a024; Brode & Piper *J Am Chem Soc* **63** 1502 1941, DOI: 10.1021/ja01851a005; *Beilstein* **16** H 335, **16** I 319, **16** III 375, **16** IV 516.]

o-Aminobenzoic acid (**2-AA**, **anthranilic acid**) [118-92-3] $C_7H_7NO_2$, **M 137.1**, **m 144.5-144.8°**, **144.9-145.4°**, **144.6-145.6°**, **145°**, **pK₁²⁵ 2.94**, **pK₂²⁵ 4.72**. Crystallise anthranilic acid from water (charcoal). It has also been recrystallised from 50% aqueous acetic acid. It sublimes in a vacuum. Sugihara and Newman [*J Org Chem* **21** 1445 1956, DOI: 10.1021/ja01851a005] found that by adding a small amount (~0.05%) of detergent [Aerosol-OTB (sodium dioctylsulfosuccinate, American Cyanamide Co.), Triton-X-100 (alkyl aryl polyethyleneglycol ether, Rohm & Haas Co.) or Oronite D-60 (sodium alkylarylsulfonate, Oronite Chemical Co.)] to the saturated aqueous (or solvent) solution, purer crystals of the acid are obtained. The crystals are filtered off, washed several times with cold H_2O on the funnel until the filtrate does not foam, indicating removal of detergent from the crystals. These are dried to constant weight *in vacuo* over $CaCl_2$ then in a drying pistol (using P_2O_5) for 24hrs at 40 mm. This procedure should produce crystals with sharper melting points. [*Beilstein* **14** IV 1004.] The amide, **anthranilamide** [88-68-6] $C_7H_7NO_2$, **M 137.1**, **m 111-113°**, is similar to the acid in being used for non-selective fluorescent labeling of glycans containing a free reducing end. Its fluorescence has λ_{ex} at 330nm and λ_{em} 420nm. [Bigge et al. *Anal Biochem* **230** 229 1995, DOI:10.1006/abio.1995.1468; *Beilstein* **14** IV 1010.] **m-Aminobenzoic acid** [99-05-8] has **m 174°**, **178-180°**, **pK₁²⁵ 3.29**, **pK₁²⁵ 5.10**. Crystallise the acid from water. Soluble in hot EtOH and in Et_2O . Darkens in the presence of light. [Nielsen et al. *J Chem Soc* 371 1962, DOI: 10.1039/JR9620000361; *Beilstein* **14** IV 1092.] The **hydrochloride** [15151-51-6] $C_7H_7NO_2$. HCl, **M 173.6**, **m 260-280°(dec)** is prepared in EtOH/HCl, washed with Et_2O and dried *in vacuo*. [*Beilstein* **14** III 992.] **p-Aminobenzoic acid** (**Vitamin H₁**, **PABA**) [150-13-0] has **m 187-188°**, **pK₁²⁵ 2.45**, **pK₁²⁵ 4.85**. Purify *p*-aminobenzoic acid by dissolving it in 4-5% aqueous HCl at 50-60°, decolorising with charcoal and carefully precipitating it with 30% Na_2CO_3 to pH 3.5-4 in the presence of ascorbic acid. It can be recrystallised from water, EtOH or EtOH/water mixtures. Its solubility (w/v) in H_2O is 0.6% (25°) and 1.1% (100°), in EtOH it is 12.5% (~25°), in Et_2O it is 2.0% (~25°), and it is slightly soluble in C_6H_6 but is almost insoluble in petroleum ether. It is an *anti-rickettsial*. [*Beilstein* **14** IV 1126.] The **dimethylammonium salt** [6018-84-4] $C_{17}H_{18}N_2O_2$, **M 282.3**, is freely soluble in H_2O and the crystals from Me_2CO have **m 170-173°**. **Potassium p-aminobenzoate** [138-84-1] $C_7H_7KNO_2$, **M 175.2**, **decomp at high temperatures**, is prepared by dissolving 0.1 mole in MeOH, adding the calculated volume of KOMe, partially evaporating the solvent and recrystallising the salt from MeOH. Dry it in an oven at 105° and store it *in vacuo* over $CaCl_2$. (Meyers et al. *J Am Chem Soc* **89** 3565 1967, DOI: 10.1021/ja00990a038.) Its solubility in cold H_2O is ~20% w/v, and more so in hot H_2O . It crystallises also from EtOH. It is *antifibrotic* and is a lesser gastric irritant than the acid or the **sodium salt** [206557-08-6] $C_7H_7NNaO_2 \cdot xH_2O$, **M 159.1** (anhydr). The

hydrazide [5351-17-7] $C_7H_9N_3O$, M 151.2, has m 225-227°.

o-Aminobenzonitrile (*o*-cyanoaniline) [1885-29-6] $C_7H_6N_2$, M 118.1, m 45-48°, b 267-268°/atm, pK^{20} 1.80, pK^{25} 0.77, 0.95. It has been prepared from *o*-chloronitrobenzene, via *o*-cyanonitrobenzene (with Cu bronze + CN^-) followed by reduction with $SnCl_2/HCl$. It is soluble in most organic solvents, and is extracted into Et_2O , dried ($CaCl_2$), evaporated, and the residual solid can be distilled or dried in *vacuo* over H_2SO_4 to give crystals with m 49.5°. Recrystallisation from CS_2 gives beautiful monoclinic prisms with m 50-51° which can also be crystallised from H_2O . It is a useful synthon for preparing quinazolines and other heterocyclic compounds. The *N*-acetyl derivative has m 107.5-108.5° (from AcOH) and the *N*-tosyl derivative has m 170-171° (from *iso*-PrOH). The *picrate* provides red crystals from H_2O with m 109-110°, or $*C_6H_6$ with m 108-109°. [Synth: Bogert & Hand *J Am Chem Soc* **24** 1031 1902, DOI: 10.1021/ja02025a001; Cooper & Partridge *J Chem Soc* 3429 1954, DOI: 10.1039/JR9540003429; *Beilstein* **14** H 322, **14** I 532, **14** II 210, **14** IV 1013.] *p*-Aminobenzonitrile (*p*-cyanoaniline) [873-74-5] has m 86-86.5°, 85-87°, pK^{25} 1.74. It crystallises from water, 5% aqueous EtOH or EtOH and is dried over P_2O_5 , or dried in *vacuo* for 6 hours at 40°. [Moore et al. *J Am Chem Soc* **108** 2257 1986, DOI: 10.1021/ja00269a022; Edidin et al. *J Am Chem Soc* **109** 3945 1987, DOI: 10.1021/ja00247a019; *Beilstein* **14** IV 1158.]

2-Aminobenzophenone (2-benzoylaniline) [2835-77-0] $C_{13}H_{11}NO$, M 197.2, m 103-107°, 105-106°, 110°, d 1.32, pK^{25} 0.33. Dissolve the pale yellow ketone in aqueous acetic acid, filter and precipitate it with ammonia. This process is repeated several times, then the amine is recrystallised from aqueous EtOH (pale yellow crystals). It is soluble in *iso*-PrOH, $CHCl_3$ and Et_2O , slightly soluble in EtOH, insoluble in H_2O but soluble in acidic solution. A $CHCl_3$ solution is used to extract Pd with which it complexes (has λ_{max} at 475nm in $CHCl_3$ /benzyl alcohol). The *hydrochloride* [40318-20-5] crystallises as needles m 192-193°(dec). The *E*-oxime [15185-37-2] crystallises from $*C_6H_6$ with m 127° whereas the *Z*-oxime [4844-60-4] has m 156°. It also gives a 2,4-dinitrophenylhydrazone which crystallises in two forms from EtOH with m 235-236° and m 261-262°. [Scheifele & DeTar *Org Synth Coll Vol* **4** 34 1963, DOI:10.15227/orgsyn.032.0008; *Beilstein* **14** IV 243.] 3-Aminobenzophenone (3-benzoylaniline) [2835-78-1] has m 81-84°, 87°, pK^{25} 0.33. Purify as for the above 2-, or 4-isomer below, and recrystallise from H_2O (needles). The *hydrochloride* provides needles from dilute HCl with m 187° and the *oxime* has m 156°. [*Beilstein* **14** H 81.] 4-Aminobenzophenone (4-benzoylaniline) [1137-41-3] has m 123-124°, b 246°/13mm, pK^{10} 2.36, pK^{25} 2.17, pK^{40} 2.02. This is prepared from 1 aniline/ 2 $PhCOCl$ / $ZnCl_2$ /200-210° to give a high yield of the 4-benzoylaminobenzophenone which is hydrolysed by EtOH/ H_2O /NaOH to the *free* base as feathery crystals (90-95% yield). These are purified by recrystallisation from CCl_4 or aqueous EtOH. Alternatively, dissolve the aniline in aqueous AcOH, filter and precipitate it with ammonia. This process is repeated several times, then the amine is recrystallised from aqueous EtOH (leaflets). The *oxime* crystallises in two forms from aqueous EtOH (presumably *E* and *Z*) which have m 126° and m 168°; and the 2,4-dinitrophenylhydrazone has m 189-191° (from EtOH). [Clarke & Esselen Jr *J Am Chem Soc* **33** 1135 1911, DOI: 10.1021/ja02220a016; *Beilstein* **14** IV 248.]

2-Aminobiphenyl (*o*-aminobiphenyl) [90-41-5] $C_{12}H_{11}N$, M 169.2, m 47-48°, 49°, 49.3°, 49-50°, b 114°/2mm, 135°/5.5mm, 160°/11mm, 170°/15mm, 182°/30mm, 299.0°/atm, pK^{18} 3.85, pK^{20} 3.39, pK^{20} 3.03, 3.34 (50% aqueous EtOH). It is prepared by reduction of 2-nitrobiphenyl [see 86-00-0] with $SnCl_2/HCl$ [Scarborough & Waters *J Chem Soc* 89 1927, DOI: 10.1039/JR9270000089], or catalytically in batches of 150g by H_2 at 75 atmospheres pressure in the presence of Raney Ni, which begins at ~80° and is not allowed to rise above 100°, in 90% yield [Cookson & Mann *J Chem Soc* 2888 1949, DOI: 10.1039/JR9490002888]. The white solid amine crystallises from aqueous EtOH (charcoal). The *picrate* has m 164-165° (from EtOH or H_2O), the *N*-acetyl derivative has m 121° (corrected, from aqueous EtOH, or petroleum ether b 60-80°), the *N*-phenylsulfonate derivative has m 292° (from MeOH), and the *N*-tosylate has m 194.1-195.6° (from MeOH). [Dewar et al. *Org Synth Coll Vol* **5** 727 1973, **46** 65 1966, DOI:10.15227/orgsyn.046.0065; application: Morgan & Walls *J Chem Soc* 2225 1932, DOI: 10.1039/JR9320002225; *Beilstein* **12** H 1317, **12** I 546, **12** II 747, **12** III 3124, **12** IV 3223.] *p*-Aminobiphenyl [92-67-1] has m 53°, b 191°/16mm, pK^{18} 4.38. Crystallise it from water or EtOH. [*Beilstein* **12** IV 3241.] **CARCINOGENIC.**

5-Amino-2-bromobenzoic acid (3-Amino-6-bromobenzoic acid) [2840-02-0] $C_7H_6BrNO_2$, M 216.0, m 171-179°(bath temp 180°), 180°, 200-201°, $pK_{Est(1)}$ ~1.7, $pK_{Est(2)}$ ~4.4. Crystallise the acid from H_2O or $*C_6H_6$ (m

128°). The *acetyl* derivative crystallises from H₂O (as *monohydrate*) or absolute EtOH with **m 196-197°** (*anhydrous*). [Koopal *Recl Trav Chim Pays-Bas* **34** 115 1915, DOI: 10.1002/recl.19150340402; Bamberger *Chem Ber* **57** 2082 1924, DOI: 10.1002/cber.19240571121; *Beilstein* **14** H 413, **14** II 245.]

2-Amino-5-bromotoluene (4-bromo-2-methylaniline) [583-75-5] C₇H₈BrN, **M 186.1, m 59°, 59.5°, 240°/760mm, pK²⁵ 3.58**. Steam distil the aniline and recrystallise it from EtOH. It has UV with λ_{max} at 292.5nm (H₂O). [*Beilstein* **12** H 838, **12** I 389, **12** II 456, **12** IV 1804.]

2-Amino-5-chlorobenzoic acid (5-chloroanthranilic acid) [635-21-1] C₇H₆ClNO₂, **M 171.6, m 204-205°, 204-206°, 210-212°, pK₁²⁵ 1.69, pK₂²⁵ 4.35**. Crystallise the acid from water, aqueous EtOH or chloroform. The *ethyl ester* crystallises from EtOH with **m 82°**, and the *ethyl ester hydrochloride* crystallises from Me₂CO with **m 148°**; and the *amide* crystallises from H₂O (**m 169-171°**) or aqueous EtOH (**m 172°**). [Endicott et al. *J Am Chem Soc* **68** 1303 1946, DOI: 10.1021/ja01211a057; Chapman *J Chem Soc* 890 1947, DOI: 10.1039/JR9470000890; McKee et al. *J Am Chem Soc* **69** 940 1947, DOI: 10.1021/ja01196a063; *Beilstein* **14** H 365, **14** II 231, **14** III 962, **14** IV 1075.] **3-Amino-4-chlorobenzoic acid** [2840-28-0] has **m 212-215°, 214-216°, 216-217°(corrected), pK_{Est(1)} ~2.7, pK_{Est(2)} ~2.9**. Crystallise the acid from water. The *methyl ester* has **m 82-83°** (from aqueous MeOH), the *acetyl derivative* has **m 263-264°** (from H₂O), and the *amide* crystallises from EtOH or H₂O (**m 264.5-265.5°**, corrected). [*Beilstein* **14** H 412, **14** IV 1115.]

4-Amino-4'-chlorobiphenyl [135-68-2] C₁₂H₁₀ClN, **M 203.6, m 132-133°, 134°, pK_{Est} ~4.0**. Crystallise the amine from petroleum ether, EtOH or aqueous EtOH. The *acetyl* derivative has **m 245°** from EtOH. [Dewar & James *J Chem Soc* 4265 1958, DOI: 10.1039/JR9580004265; Gelmo *Chem Ber* **39** 4175 1906, DOI: 10.1002/cber.190603904117; *Beilstein* **12** H 1319, **12** II 757, **12** IV 3269.]

2-Amino-4,6-dichlorophenol [527-62-8] C₆H₅Cl₂NO, **M 178.0, m 95-96°, pK_{Est(1)} ~3.1, pK_{Est(2)} ~6.8**. Also available commercially as the hydrochloride in the form of a dark brown powder. It can be purified by dissolving 50g of crude hydrochloride in boiling H₂O (500 ml), treating it twice with animal charcoal under reflux, filtering off the charcoal followed by adding NaHCO₃ solution to neutrality (litmus paper). The dark solid is collected and dried first with CaCl₂, then P₂O₅. The dried base (6g) sublimes under high vacuum (0.06 mm during 4 hours, at bath temperature 70-80°) to give pure snow white aminodichlorophenol (5.75g). Crystallise the phenol from CS₂ or *benzene. The *hydrochloride*, crystallises on adding concentrated HCl to a solution in H₂O. It has **m 280-285°** after recrystallisation from EtOH. [Meyer *Helv Chim Acta* **41** 1890 1958, DOI: 10.1002/hlca.19580410643; *Beilstein* **13** II 185, **13** III 856, **13** IV 889.]

4-Amino-N,N-diethylaniline hydrochloride [16713-15-8] C₁₀H₁₆N₂. HCl, **M 200.7, m 233.5°, pK²² 6.61**. Recrystallise the salt from EtOH. The *free base* [93-05-0] C₁₀H₁₆N₂, **M 164.2** distils at **105-108°/3mm, 115-116°/5mm, 139-140°/10mm, 260-262°/~760mm, d₄²⁵ 1.01, n_D²⁰ 1.571**, solidifies on cooling, and the solid has **m 19-21°**. [Bent et al. *J Am Chem Soc* **73** 3100 1951, DOI: 10.1021/ja01151a037; *Beilstein* **13** H 75, **13** I 22, **13** II 40, **13** III 113, **13** IV 109.]

4-Amino-3,5-diiodobenzoic acid [2122-61-4] C₇H₅I₂NO₂, **M 388.9, m ~350°, pK_{Est(1)} 0.4, pK_{Est(2)} ~1.6**. Purify the iodo-acid by dissolving it in dilute NaOH and precipitating with dilute HCl. *Alternatively*, dissolve it in aqueous NH₃ and acidify it with AcOH. Dry it in air. The solubility of the *Na salt* in H₂O is 2.56% at 25°. [Klemme & Hunter *J Org Chem* **05** 508 1940, DOI: 10.1021/jo01211a004; *Beilstein* **14** H 439, **14** III 1161, **14** IV 1284.]

2-Aminodiphenylamine (N-phenyl-o-phenylenediamine) [534-85-0] C₁₂H₁₂N₂, **M 184.2, m 77-80°, 79-80°, pK_{Est(1)} ~3.8 (NH₂), pK_{Est(2)} <~0**. Crystallise the amine from H₂O or EtOH. The *hydrochloride*, prepared by shaking an ethereal solution of the amine with 10% hydrochloric acid, has **m 115.5-117°(dec)**. [Lane & Williams *J Chem Soc* 1468 1955, DOI: 10.1039/JR9550001468; *Beilstein* **13** IV 43.]

4-Aminodiphenylamine [101-54-2] C₁₂H₁₂N₂, **M 184.2, m 68-72°, b 155°/0.026mm, pK²⁵ 5.20, pK_{Est(2)} <~0, pK²⁰ 5.24 (50% EtOH)**. It crystallises from EtOH with **m 66°**, and from ligroin with **m 75°**. It can be distilled at high vacuum. The *hydrochloride* crystallises from MeOH (**m 240-241°**). [Allen et al. *J Chem Soc*

234 1954, DOI: 10.1039/JR9540000234; *Beilstein* **13** H 76, **13** I 23, **13** II 40, **13** III 113, **13** IV 113.]

2-Amino-1,2-diphenylethanol (1,2-diphenylethanolamine, four stereoisomers) [530-36-9] $C_{14}H_{15}NO$, **M 213.3**, **m 165°**, **pK_{Est(1)} ~7.5**. Recrystallise the ethanol from EtOH. The *1R,2S*(-)- [23190-16-1] and the *1S,2R*(+)-[23364-17-2] *enantiomers* also crystallise from EtOH and have **m 142-145°**, **[α]_D²⁵ (-) and (+) 7 (c 0.6, EtOH)**. The *enantiomers 1R,2R*(+)-**2-amino-1,2-diphenylethanol** [88082-66-0] and *1S,2S*(-)-**2-amino-1,2-diphenylethanol** [23190-66-0] have **m 116-119°** and **[α]_D²⁵ (+) 124 and (-) 124 (c 0.87, EtOH)**. [Masters et al. *J Org Chem* **56** 5666 1991, DOI: 10.1021/jo00019a038; Masters & Hegedus *J Org Chem* **58** 4547 1993, DOI: 10.1021/jo00069a012; *Beilstein* **13** IV 2150.]

2-Aminodiphenylmethane (2-benzylaniline) [28059-64-5] $C_{13}H_{13}N$, **M 183.2**, **m 52°, 51-54°, b 172°/12mm and 190°/22mm**, **pK_{Est(1)} ~4.2**. Crystallise 2-benzylaniline from ether and distil it in a vacuum. **3-aminodiphenylmethane (3-benzylaniline)** [61424-26-8] has **m 48-52°**, and **4-aminodiphenylmethane (4-benzylaniline)** [1135-12-2] has **m 38-38°** and **pK_a 4.78**. [*Beilstein* **12** IV 3279.]

1-Aminofluorene (1-fluorenamine) [6344-63-0] $C_{13}H_{11}N$, **M 181.2**, **m 124.0-124.6°, 125-126°, 125-126°, pK₂₅ 3.87, pK₄₀ 3.67, and pK₂₅ 3.57 (70% EtOH)**. Prepared by acid hydrolysis of 1-fluorenylurethane (starting from fluorine-1-carboxylic acid), extracted into Et₂O, and first crystallised from dilute EtOH, then purified by recrystallising from *C₆H₆/petroleum ether. It forms a **2,4,7-trinitrofluorenone complex**, $C_{13}H_{11}N \cdot C_{13}H_5N_3O_7$, which crystallises from *C₆H₆ as beautiful black needles (**m 211-211.8°**, corrected). [Grantham et al. *J Org Chem* **26** 1008 1961, DOI: 10.1021/jo01063a006; Bergmann & Orchin *J Am Chem Soc* **71** 1111 1949, DOI: 10.1021/ja01171a502; *Beilstein* **12** H 1331, **12** IV 337.] **2-Aminofluorene (2-fluorenamine)** [153-78-6] has **m 127.8-128.8°, 132-133°, pK₂₅ 4.64, pK₄₀ 4.42**. Wash the amine well with H₂O, recrystallise it from Et₂O or 50% aqueous EtOH (25g with 400ml), and dry it in a vacuum. Store it in the dark. [Bavin *Org Synth Coll Vol* **5** 30 1973, DOI: 10.15227/orgsyn.040.0005, Grantham et al. *J Org Chem* **26** 1008 1961, DOI: 10.1021/jo01063a006; *Beilstein* **12** H 1331, **12** IV 337.]

9-Aminofluorene (9-fluorenamine) [525-03-1] $C_{13}H_{11}N$, **M 181.2**, **m 64-65°, pK₂₅ 7.56 (70% EtOH)**. Purify it by converting it to the hydrochloride with HCl, then basify it with NH₃ and recrystallise it from petroleum ether (**m 62-63°**) or from hexane. It was originally prepared by reduction of 9-fluorenone oxime with Zn and AcOH, isolated as the hydrochloride then basified as stated here. **Note**: the high basicity of the 9-amino group (compare with above fluorenamines) is probably due to its being attached to a *benzylic-type* carbon atom. The **hydrochloride** [5978-75-6] **M 217.7**, has **m ~255°**(dec, from EtOH). [Ingold & Wilson *J Chem Soc* 1493 1933, DOI: 10.1039/JR9330001493; Mathieu *Bull Soc Chim Fr* 1526 1971, Grantham et al. *J Org Chem* **26** 1008 1961, DOI: 10.1021/jo01063a006; *Beilstein* **12** H 1331, **12** I 553, **12** II 780, **12** III 3297, **12** IV 3390.]

1-Amino-4-hydroxyanthraquinone [116-85-8] $C_{14}H_9NO_3$, **M 239.2**, **m 207-208°, pK_{Est(1)} ~2.6 (NH₂), pK_{Est(2)} ~9.0 (OH)**. Purify it by TLC on SiO₂ gel plates (0.75mm thick) using toluene/acetone (9:1) as eluent. The main band is scraped off and extracted with MeOH. The solvent is evaporated, and the dye is dried in a drying pistol [Land et al. *JCS Faraday Trans 1* **72** 2091 1976, DOI: 10.1039/F19767202091]. It has also been recrystallised from aqueous EtOH. [*Beilstein* **14** H 268, **14** I 503, **14** II 168, **14** III 652, **14** IV 891.]

3-Amino-4-hydroxytoluene (2-amino-*p*-cresol) [95-84-1] C_7H_9NO , **M 123.2**, **m 135°, 137-138°, pK_{Est(1)} ~4.7(NH₂), pK_{Est(2)} ~9.6 (OH)**. Recrystallise the cresol from H₂O, Et₂O or toluene. It sublimes *in vacuo* as plates or needles. The **hydrochloride** has **m 222-224°** (dec, from aqueous EtOH). [*Beilstein* **13** H 601, **13** I 227, **13** II 338, **13** III 1576, **13** IV 1712.] **4-Amino-3(5)-hydroxytoluene (2-amino-*m*-cresol)** [2835-98-5] has **m 159°, 159-162°, pK_{Est(1)} ~5.4 (NH₂), pK_{Est(2)} ~10.2 (OH)**. Crystallise it from H₂O, 50% EtOH, or toluene. [*Beilstein* **13** H 590, **13** III 1552.] **2-Amino-5-hydroxytoluene (4-amino-*m*-cresol)** [2835-99-6] C_7H_9NO , **M 123.2**, **m 162°**(dec), **177-179°**(dec), **pK_{Est(1)} ~5.4 (NH₂), pK_{Est(2)} ~10.4 (OH)**. Crystallise it from 50% EtOH. [*Beilstein* **13** H 593, **13** IV 1698.]

5-Aminoindane [24425-40-9] $C_9H_{11}N$, **M 133.2**, **m 37-38°, b 131°/15mm, 146-147°/25mm, 247-249°/745mm, pK₁₆ 5.31**. Distil the indane, and then recrystallise it from petroleum ether. [*Beilstein* **12** I 511, **12** III 2798.]

1R,2SR-(±)-cis-1-Amino-2-hydroxyindane (cis-1-amino-2-indanol) [7480-35-5] $C_9H_{11}NO$, **M 149.2**, **m 132-133°**, **134-135°**, $pK_{Est} \sim 8.5$. It is obtained by hydrolysis of (±)-cis-indano[1,2-d]-2-oxazolidone [in turn formed by solvolysis of ethyl *N*-(trans-2-iodo-1-indan)carbamate, m 159.5-160° in 87% yield by refluxing in dry diglyme for 14 hours]. The cis-2-oxazolidone (360mg) is dissolved in 0.85N methanolic KOH (20ml), diluted with H₂O (5ml) and refluxed under N₂ for 18 hours, evaporated to dryness *in vacuo*, the residue is extracted with Et₂O, filtered, dried (Mg₂SO₄), concentrated to a small volume and cooled to -12°, and the **amino-2-indanol** is collected as white plates (2 crops, m 134-135° and 133-134°) in 79% yield. [Hassner et al. *J Org Chem* **32** 540 1967, DOI: 10.1021/jo01278a006.] Similarly, hydrolysis of 2-phenyl-3a,8a-dihydro-8H-indeno[1,2-d]oxazole with diluted H₂SO₄ (16 hours reflux), cool, basify with 10N NaOH, extract with CHCl₃, dry (Mg₂SO₄), and evaporation gives an 88% yield of the (±)-**amino-alcohol (m 132-133°)**. It can also be prepared from (±)-hydrindan-1,2-epoxide as described in the following entry. Its ¹HNMR (400MHz, CDCl₃, TMS) has δ at 2.0-2.5 (br s, 3H), 2.95 (dd, *J* = 3 and 15Hz, 1H), 3.1 (dd, *J* = 7 and 15Hz, 1H), 4.28 (d, *J* = 7Hz, 1H), 4.4 (m, 1H), 7.2-7.35 (m, 4H). [Thompson et al. *J Med Chem* **35** 1685 1992, DOI: 10.1021/jm00088a003.] Large scale preparations of the (±)-cis-1-amino-2-indanol from indene *via* the (±)trans- acylamido-indanol by hydrolysis with strong acids (HCl, H₂SO₄ or MeSO₃H) which invert the OH stereochemistry have been patented [Gao et al. USP appl 5,599,985 1997; *Chem Abstr* **123** 55421b 1995, Gao et al. USP 5,616,808 1997, *Chem Abstr* **126** 250997h 1997]. [Beilstein **13** II 398a, 399d, **13** III 1841, **13** IV 2033.]

1S,2R-(-)-cis-1-Amino-2-hydroxyindane [(-)-cis-1-amino-2-indanol] [126456-43-7] $C_9H_{11}NO$, **M 149.2**, **m 117-117.5°**, **118-121°**, $[\alpha]_D^{20} -61^\circ$ (c 0.5, CHCl₃), $[\alpha]_D^{25} -62$ (c 1.0, MeOH). This enantiomer has been prepared in several ways in high state of optical purity. Oxidation of indene [95-13-6] using Jacobsen's salen catalyst *R,R*-MnCl/NaOCl/CH₂Cl₂ gave *1R,2S*-indane-1,2-epoxide in 84%ee, which on treatment with NH₃/MeOH then PhCOCl/NaOH followed by 80% H₂SO₄ provided 2-phenyl-3a*S*,8a*R*-dihydro-8H-indeno[1,2-d]oxazole [see (±) above] by a C-1 chirality transfer process which gave the desired *1S,2R*-(-)-aminoindanol on hydrolysis. Alternatively, oxidation of indene using Jacobsen's salen catalyst *S,S*-MnCl/PhCl/NaOCl and P₃NO {4-(3-phenylpropyl)pyridine-*N*-oxide see [34122-28-6]} gave the enantiomeric *1S,2R*-indane-1,2-epoxide which was reacted with Oleum/MeCN then hydrolysed with H₂O (by C-2 chirality transfer *via* a Ritter reaction) to give the same desired *1S,2R*-(-)-aminoindanol 78-80% yield with high optical purity. [Senanayake *Aldrichimica Acta* **31** 3 1998, Senanayake et al. *Tetrahedron Lett* **36** 3993 1995, DOI: 10.1016/0040-4039(95)00696-A; Senanayake et al. *Tetrahedron Lett* **37** 3271 1996 DOI: 10.1016/0040-4039(96)00565-5.] In a different approach *N*-(L-phenylalanyl)-(±)-amino-2-indanol was prepared [84% yield, from (±)-aminoindanol and BOC-L-phenylalanine *via* the 1-hydroxybenzotriazole/dimethyl 3-{3-(dimethylamino)-propyl}carbodiimide procedure], and the diastereoisomeric mixture was separated by low pressure chromatography on silica gel, eluting with 8% MeOH in CHCl₃ to give the faster moving L-Phe-*1S,2R*-1-aminoindan-2-ol diastereoisomer (40% yield). A mixture of this diastereomer (30g, 0.1mol), EtOH (1L) and 20% aqueous NaOH (265ml) was refluxed for 16 hours, concentrated to remove EtOH, diluted with H₂O (100ml) and brine (100ml), extracted with CHCl₃ (3 x 600ml), the combined extracts were dried (Mg₂SO₄), filtered and evaporated to give **1S,2R-1-aminoindan-2-ol** (14g, 93%) as a white solid **m 114-115°**. Recrystallisation from Et₂O/hexanes gave analytically pure white rods **m 117-117.5°**, with little loss of material, and with $[\alpha]_D^{25} -62$ (c 1.0, MeOH), with ¹HNMR (400MHz, CDCl₃, TMS) which had δ at 2.0-2.5 (br s, 3H), 2.95 (dd, *J* = 3 and 15Hz, 1H), 3.1 (dd, *J* = 7 and 15Hz, 1H), 4.28 (d, *J* = 7Hz, 1H), 4.4 (m, 1H), 7.2-7.35 (m, 4H), same as the racemate. [Thompson et al. *J Med Chem* **35** 1685 1992, DOI: 10.1021/jm00088a003.] Both enantiomers of this *cis*-aminoalcohol have also been obtained *via* enzymatic transformations [Didier et al. *Tetrahedron* **47** 4941 1991, DOI: 10.1016/S0040-4020(01)80959-5]. The oxazolines, oxazolidin-2-ones and the 2-*N*-tosylamides derived from the two enantiomers have been explored extensively as *chiral auxiliaries* [Senanayake *Aldrichimica Acta* **31** 3 1998]. For the preparation of CBS-oxazaboroline catalysis (Corey-Bakshi-Shibata) see Chapter 6, Catalysis-Part 1. **NOTE:** Both enantiomers are commercially available from Sepracore in gram to hundred kilogram quantities [Hong et al. *Tetrahedron Lett* **35** 6631 1994, DOI: 10.1016/S0040-4039(00)73453-8, see also Stinson *Chemical and Engineering News* May 16 p.6 1994.]. [Beilstein **13** II 398a, 399d, **13** III 1841, **13** IV 2033.]

1R,2S-(+)-cis-1-Amino-2-hydroxyindane [(+)-cis-1-amino-2-indanol] [136030-00-7] $C_9H_{11}NO$, **M 149.2**, **m 118-121°**, $[\alpha]_D^{20} +63$ (c 0.2, CHCl₃). This compound was prepared using the same syntheses as for its enantiomer (preceding entry) and purified in the same way. It can be also obtained from the slower moving chromatographic band of the L-phenylalaninyl diastereoisomer (see preceding entry).

2-Amino-5-iodotoluene [13194-68-8] C_7H_8IN , M 233.0, m 87°, $pK_{Est} \sim 3.6$. Crystallise it from 50% EtOH. [Beilstein 12 IV 1807.]

1-Amino-4-methylaminoanthraquinone [1220-94-6] $C_{15}H_{12}N_2O_2$, M 252.3, $pK_{Est(1)} \sim 1$, $pK_{Est(2)} < 4$. Purify the quinone by TLC on silica gel plates using toluene/acetone (3:1) as eluent. The main band is scraped off and extracted with MeOH. The solvent is evaporated, and the residue is dried in a drying pistol [Land et al. *JCS Faraday Trans 1* 72 2091 1976, DOI: 10.1039/F19767202091]. [Beilstein 14 H 198, 14 I 462, 14 III 440, 14 IV 458.]

4-Aminomethylbenzenesulfonamide hydrochloride (Mafenide HCl) [138-37-4] $C_7H_{10}N_2O_2S \cdot HCl$, M 222.3, m 256°, 265-267° (also reported), $pK_1^{20} 8.18$ (NH_2), $pK_2^{20} 10.23$ ($SONH_2$). Crystallise the salt from dilute HCl or 95% EtOH and dry it in a vacuum at 100°. [Miller et al. *J Am Chem Soc* 62 2099 1940, DOI: 10.1021/ja01865a053; Bergeim & Barker *J Am Chem Soc* 66 1459 1944, DOI: 10.1021/ja01237a012.] The **free base** [138-39-6] $C_7H_{10}N_2O_2S$, M 186.2, has m 151-152° (from EtOH), the **acetate salt (Mafatate)** [13009-99-9] $C_7H_{10}N_2O_2S \cdot C_2H_4O_2$, M 246.2, also has m 151-152° (from AcOH), and the **propionate salt (Sulfomyl)** [12001-72-8] $C_7H_{10}N_2O_2S \cdot C_3H_6O_2$, M 260.3, has m 158°. The **sulfate salt** has m 254-255° (from aqueous EtOH). [Beilstein 14 III 2223, 14 IV 2799.] All these compounds are effective antibacterials.

3-Amino-2-naphthoic acid [5959-52-4] $C_{11}H_9NO_2$, M 187.2, m 212-215°(dec), 214°(dec), $pK_{Est(1)} \sim 1.5$, $pK_{Est(2)} \sim 4.0$. Crystallise the naphthoic acid from aqueous EtOH. Ethanolic and ethereal solutions are yellow in colour with green fluorescence. The amino acid is used for the determination of Cu, Ni and Co. The **sodium salt** is difficultly soluble in H_2O , but forms leaflets from it, and is also sparingly soluble in EtOH. The **ethyl ester** $C_{13}H_{13}NO_2$ crystallises from EtOH in yellow needles from aqueous EtOH and has m 115-115.5°. [Allen & Bell *Org Synth Coll Vol* 3 78 1955, DOI: 10.15227/orgsyn.022.0019; Beilstein 14 III 1341.]

4-Amino-5-naphthol-2,7-disulfonic acid [90-20-0] $C_{10}H_9NO_7S_2$, M 319.3, $pK_1^{20} 3.54$, $pK_2^{20} 8.55$; $pK_1^{25} 3.63$, $pK_2^{25} 8.83$. A slightly alkaline solution of Na_2CO_3 (ca 22g) in water (litmus) is added to a solution of 100g of the dry acid in 750ml of hot distilled water, followed by 5g of activated charcoal and 5g of Celite. The suspension is stirred for 10 minutes and filtered by suction. The acid is then precipitated by adding ca 40ml of conc HCl (solution is blue to Congo Red), then filter it by suction through a shark skin filter circular sheet (or hardened filter paper) and wash it with 100ml of distilled water. The purification process is repeated. The acid is dried overnight in an oven at 60° and stored in a dark bottle. The **diethylamine salt** has m 306-307°(dec, from aqueous EtOH). [Post & Moore *Anal Chem* 31 1872 1959, DOI: 10.1021/ac60155a060]. [Beilstein 14 H 840, 14 I 758, 14 II 502, 14 III 2292, 14 IV 2824.]

1-Amino-2-naphthol hydrochloride [1198-27-2] $C_{10}H_9NO \cdot HCl$, M 195.7, m 250°(dec), $pK_{Est(1)} \sim 3.7$ (NH_2), $pK_{Est(2)} \sim 9.9$ (OH). Crystallise the salt from the minimum volume of hot water containing a few drops of stannous chloride in an equal weight of hydrochloric acid (to reduce atmospheric oxidation). Filter the solution and add half its volume of concentrated HCl and set aside. The salt crystallises almost quantitatively. Dry it in a vacuum in the dark (m 260°). The salt is more stable than the **free base** which has m 150° (darkening at 130°) and its **O-methyl ether** has m 49° and b 159-159°/9mm. [Desai et al. *J Chem Soc* 324 1938, DOI: 10.1039/JR9380000321; Beilstein 13 H 666, 13 I 268, 13 II 408, 13 III 1875, 13 IV 2080.]

1-Amino-2-naphthol-4-sulfonic acid [116-63-2] $C_{10}H_9NO_4S$, M 239.3, m 290°(dec), 295°(dec), $pK_{Est(1)} < 0$, $pK_{Est(2)} \sim 2.8$ (NH_2), $pK_{Est(3)} \sim 8.8$. Purify it by warming 15g of the acid, 150g of $NaHSO_3$ and 5g of Na_2SO_3 (anhydrous) with 1L of water to ca 90°, shaking until most of the solid had dissolved, then filtering hot. The precipitate obtained by adding 10ml of concentrated HCl to the cooled filtrate is collected, washed with 95% EtOH until the washings are colourless, and is dried under vacuum over $CaCl_2$. It is stored in a dark-coloured bottle, in the cold [Chanley et al. *J Am Chem Soc* 74 4347 1952, DOI: 10.1021/ja01137a030]. The **Na salt**, $C_{10}H_8NNaO_4S$, forms needles from hot H_2O ; a solution which has blue fluorescence. [Fieser 2 42; Hiers & Hager, DOI: 10.15227/orgsyn.011.0012; Beilstein 14 IV 2825.]

2-Amino-4-nitrobenzoic acid (4-nitroanthranilic acid) [619-17-0] $C_7H_6N_2O_4$, M 182.1, m 257°(dec), 269°(dec), $pK_1^{25} 0.65$, $pK_2^{25} 3.70$. Recrystallise it from water, EtOH (m 271°) or aqueous EtOH (m 269°).

The *acetyl* derivative has **m 217°** (from EtOH), **m 222°** (from aqueous EtOH). [Chapman & Stephen *J Chem Soc* 1791 1925, DOI: 10.1039/CT9252701791; *Beilstein* **14** II 234, **14** III 975, **14** IV 1087.]

5-Amino-2-nitrobenzoic acid [13280-60-9] $C_7H_6N_2O_4$, **M 182.1**, **m 235°(dec)**, **236-238°**, **pK_{Est(1)} ~1.1**, **pK_{Est(2)} ~1.2**. Crystallise the acid from water. [*Beilstein* **14** III 1021.]

1-Amino-4-nitronaphthalene [776-34-1] $C_{10}H_8N_2O_2$, **M 188.2**, **m 190-193°**, **195°**, **196-197°**, **pK²⁰ 0.54**. It crystallises from EtOH, ethyl acetate or aqueous NH_3 as light yellow crystals. The *acetyl* derivative also forms yellow crystals, **m 192.5-193.5°**, from Me_2CO . [*Beilstein* **12** I 530, **12** II 704, **12** III 2971, **12** IV 3114.]

2-Amino-4-nitrophenol (2-hydroxy-5-nitroaniline) [99-57-0] $C_6H_6N_2O_3$, **M 154.1**, **m 80-90° (hydrate)**, **142-143° (anhydrous)**, **146°**, **pK_{Est(1)} ~3.9 (NH₂)**, **pK_{Est(2)} ~9.2**. Crystallise the phenol from water and dry it in a vacuum. It is slightly soluble in MeOH or AcOH, but is more soluble in hot $*C_6H_6$, EtOH and dilute AcOH. With Ac_2O at room temperature it forms the *N-monoacetyl derivative* with **m 279-280°** (crystallising in clusters of needles from a large volume of boiling EtOH). On long boiling with excess of Ac_2O it forms, with difficulty, the *O,N-diacetyl derivative* with **m 187-188°** (crystallising in needles or prisms from boiling EtOH in which it is relatively soluble) [Hewitt & King *J Chem Soc* **129** 817 1926, DOI: 10.1039/JR9262900817]. [Hartman & Silloway *Org Synth Coll Vol* **3** 82 1955, DOI: 10.15227/orgsyn.025.0005; *Beilstein* **13** IV 896.] **2-Amino-5-nitrophenol (2-hydroxy-4-nitroaniline)** [121-88-0] has **m 198-202°(dec)**, **207-208°**, **pK_{Est(1)} ~3.8**, **pK_{Est(2)} ~9.3**. It is best obtained by treating 6-nitrobenzoxazolone with NH_3 in an autoclave, but can be achieved more simply by treatment with hydrazine [Bower & Stephens *J Chem Soc* 325 1951, DOI: 10.1039/JR9510000325]. Recrystallisation of the phenol from water gives orange or dark brown needles **m 207-208°**. Its UV has λ_{max} at 390nm (ϵ 13,000) (EtOH) [Pedersen *J Org Chem* **23** 255 1958, DOI: 10.1021/jo01096a028]. With cold Ac_2O it forms the *N-monoacetyl derivative* with **m 171-172°** (crystallising from large volume of EtOH). On boiling with Ac_2O it forms the *O,N-diacetyl derivative* with **m 193-194°** (crystallising as felt fine needles from boiling EtOH), and the acetylation mother liquors provide a *triacetyl derivative* with **m 138-139°** (crystallising as fern-like needles from boiling absolute EtOH) [Hewitt & King *J Chem Soc* **129** 817 1926, DOI: 10.1039/JR9262900817]. [*Beilstein* **13** H 390, **13** I 121, **13** II 194, **13** III 887.13 IV 903.]

RS-(±)-3-Amino-3-(4-nitrophenyl)propionic acid [35005-61-9] $C_9H_{10}N_2O_4$, **M 210.2**, **m 215°(dec)**, **220-226° (dec)**, **226°(dec)**, **browning at 215°**, **pK_{Est(1)} ~3.0**, **pK_{Est(2)} ~9.5**. The acid crystallises from 50% aqueous EtOH. The *hydrochloride* has **m 218-220°(dec)**. The *N-benzoyl derivative* has **m 204-205°** (from MeOH) and the *N-benzoyl-hydrazide* has **m 226-227°** (from MeOH or EtOH). [Posner *Justus Liebigs Ann Chem* **389** 44 1912, DOI: 10.1002/jlac.19123890102; *Beilstein* **14** I 603, **14** IV 1548.]

2-Aminophenol [95-55-6] C_6H_7NO , **M 109.1**, **m 170-175°**, **175-176°**, **pK₁²⁵ 4.65**, **pK₂²⁵ 9.75**. Purify it by dissolving it in hot water, decolorising with activated charcoal, filtering and cooling to induce crystallisation. Maintain an atmosphere of N_2 over the hot phenol solution to prevent its darkening by oxidation [Charles & Freiser *J Am Chem Soc* **74** 1385 1952, DOI: 10.1021/ja01126a007]. It can also be crystallised from EtOH using the same precautions. Its solubility (w/v) in H_2O is 2.0%, in EtOH it is 4.3%, and is very soluble in Et_2O but much less so in $*C_6H_6$. It sublimes *in vacuo*. Store in the dark, preferably under N_2 . [*Beilstein* **13** IV 805.] Crystals of the *hydrochloride* become grey in colour on standing in air. **3-Aminophenol** [591-27-5] has **m 122-123°**, **b 164°/11mm**, **pK₁²⁵ 4.25**, **pK₂²⁵ 9.90**. Crystallise it from hot water or toluene. Its solubility (w/v) in H_2O is 2.5% at $\sim 10^\circ$, it is very soluble in hot H_2O , alcohols and Et_2O , but much less so in $*C_6H_6$. Store it in the dark, preferably under N_2 . [*Beilstein* **13** IV 952.] **4-Aminophenol** [123-30-8] has **m 186°**, **189.6-190.2°**, **190° (under N_2)**, **b 130.2°/0.3mm**, **150°/3mm**, **167°/8mm**, **284°/atm (dec)**, **pK₁²⁵ 5.38**, **pK₂²⁵ 10.4**. It is light and oxygen sensitive and turns pink on standing. Crystallise it from EtOH, then water, excluding oxygen. It sublimes at $110^\circ/0.3mm$. It has been purified by chromatography on alumina with a 1:4 (v/v) mixture of absolute EtOH/ $*benzene$ as eluent. Its solubility (w/v) in H_2O is 0.4% (15°), 0.65% (25°), 1.5% (50°), 4.7% (80°) and 9% ($\sim 100^\circ$). It is almost insoluble in $CHCl_3$ and $*C_6H_6$, is soluble in absolute EtOH (4.5% at 0°) and ethyl methyl ketone (9.3% at 58.5°). [*Beilstein* **13** IV 1014.] **4-Aminophenol hydrochloride** [51-78-5] $C_6H_7NO \cdot HCl$, **M 145.6**, **m 306°(dec)**. Purify the salt by treating an aqueous solution with saturated $Na_2S_2O_3$, filtering under N_2 , then recrystallising it from 50% EtOH twice and once from absolute EtOH [Livingston & Ke

J Am Chem Soc **72** 909 1950, DOI: 10.1021/ja01158a068]. [*Beilstein* **13** III 993.]

4-Aminophenylacetic acid [1197-55-3] $C_8H_9NO_2$, **M 151.2**, **m 196-197°**(corrected), **199-200°**(dec), **201°**(dec), **pK₁²⁰ 3.60**, **pK₂²⁰ 5.26**. It has been prepared by the Willgerott reaction from 4-aminoacetophenone, sulfur and morpholine; and acidifying with AcOH. [Schwenk & Papa *J Org Chem* **11** 798 1946, DOI: 10.1021/jo01176a023; King & McMillan *J Am Chem Soc* **68** 2335 1946, DOI: 10.1021/ja01215a058]. Also recrystallise the acid from hot water (60-70ml/g, leaflets). The **hydrochloride**, is quite soluble in H₂O, soluble in EtOH, but has been recrystallised from H₂O and then from EtOH, and melts in the range **215 to 240°** depending on the heating rate. The **ethyl ester** forms platelets from H₂O (**m 49.5°, 51°**) or EtOH, and the **N-benzoyl derivative**, **m 205-206°**, forms needles from EtOH. [Albert & Goldacre *Nature* **149** 245 1942, DOI: 10.1038/149245a0; *Beilstein* **14** H 456, **14** I 589, **14** II 280, **14** III 1182.]

IRS,2RS-2-Amino-1-phenylbutan-1-ol [α -(α -aminopropyl)benzyl alcohol] [(\pm)-*threo* 5897-76-7] $C_{10}H_{15}NO$, **M 165.1**, **m 79-80°**, **pK_{Est} ~9.7**. Crystallise the **free base** of the *threo* isomer from *benzene/petroleum ether which has **m 79-80°**. The **threo-hydrochloride**, **m 204-205°**, crystallises from EtOH [Abrams & Kipping *J Chem Soc* 1480 1936, DOI: 10.1039/JR9360001480; Rebstock et al. *J Am Chem Soc* **73** 3666 1951, DOI: 10.1021/ja01152a030]. The **IRS,2RS-erythro** isomer has **m 80.5-81°** and its **hydrochloride** has **m 242°** [Hartung et al. *J Am Chem Soc* **52** 3317 1930, DOI: 10.1021/ja01371a046.] [*Beilstein* **13** II 390, **13** III 1791, **13** IV 1952.]

N-Aminophthalimide [1875-48-5] $C_8H_6N_2O_2$, **M 162.2**, **m 200-202°, 200-205°, pK_{Est} ~0**. The amino-imide is prepared by adding phthalimide (12g) to hydrazine hydrate (4.4g) in 95% EtOH (80ml), shaking for 2 minutes at ~20°, the spongy mass is rapidly heated under reflux for 3 minutes (NH₃ evolved), H₂O (50ml), is added and the clear solution is poured into H₂O (200ml) whereby the amino-imide crystallised during 1 hour. It is recrystallised from 96% EtOH (solubility is ~2% at boiling temperature) to form a yellow solution from which the amino-imide crystallises on cooling. It sublimes *in vacuo* at *ca* 150°. It solidifies after melting, and remelts at 338-341° (see **m** of phthalhydrazide in 'Heterocyclic Compounds' in this Chapter). On boiling for a few minutes with a little Ac₂O it is converted to the **N-acetyl derivative**, **m 228-230°** after recrystallisation from H₂O. **N-Benzylideneaminophthalimide**, is obtained by boiling a 96% EtOH solution (25ml) containing benzaldehyde (1g) *N*-aminophthalimide (1g) under reflux for 1 hour, allowing to crystallise overnight and recrystallising the colourless needles, **m 166-167°**, from EtOH. **N-p-Anisylideneamino-phthalimide**, is similarly obtained, and forms stout sulfur-yellow crystals **m 198-191°** from EtOH. [Drew & Hatt *J Chem Soc* 16 1937, DOI: 10.1039/JR9370000016; *Beilstein* **22/11** V 122.]

4-Aminopropiophenone (PAPP) [70-69-9] $C_9H_{11}NO$, **M 149.2**, **m 140°, b 180°/10mm**, **pK_{Est} ~2.2**. Crystallise it from water (yellow needles) or aqueous EtOH. The **hydrochloride** [6170-25-8] $C_9H_{11}NO \cdot HCl$, **M 185.7**, has **m 188-189°** (yellow needles from aqueous EtOH/HCl drops), and the **semicarbazone** has **m 139-140°** (from EtOH/H₂O). [Derick & Bornmann *J Am Chem. Soc* **35** 1269 1913, DOI: 10.1021/ja02198a014; *Beilstein* **14** H 59, **14** I 375, **14** III 146, **14** IV 139.] It is an antidote for cyanide poisoning.

4-(2-Aminopropyl)phenol (Hydroxyamphetamine) [103-86-6] $C_9H_{13}NO$, **M 151.2**, **m 125-126°, pK_{Est(1)} ~9.4 (OH)**, **pK_{Est(2)} ~9.7(NH₂)**, **pK₂₅ 9.31**. Crystallise the phenol from *benzene. The **R(-)-enantiomer** crystallises from EtOH or *C₆H₆ with **m 110.5-111.5°** and **[α]_D¹⁷ -52.0 (c 1, EtOH)**. The (\pm)-**hydrochloride** has **m 171-172°** (EtOH/Et₂O), and the **2,4-dinitrophenylhydrazone** has **m 190-192°** (EtOH). [*Beilstein* **13** I 251, **13** III 1709, **13** IV 1871.]

1-Aminopyrene [1606-67-3] $C_{16}H_{11}N$, **M 217.3**, **m 115-117°, 117-118°, pK₁²⁵ 2.91 (50% aqueous EtOH)**, **pK₂²⁵ 2.77 (50% aqueous EtOH)**. Crystallise it from hexane. [*Beilstein* **12** IV 3460.]

4-Aminosalicylic acid (PAS, p-aminosalicylic acid) [65-49-6] $C_7H_7NO_3$, **M 153.1**, **m 150-151°**(dec), **139-141°**(evolving CO₂), **150-151°**(dec), [**capillary melting points**], **220°**(dec), and **240 ±1°** [average of 5 samples on a Dennis & Shelton m.p. apparatus], **pK₁²⁵ 1.78 (CO₂H)**, **pK₂²⁵ 3.63 (NH₂)**, **pK₂₅ 13.74 (OH)**. Crystallise the acid from EtOH. Alternatively, purify it via the barium salt. **NOTE:** The melting point is not a good criterion of purity, and the most nearly correct value is **240°** [see Seaman et al. *J Am Chem Soc* **71** 2940

1949, DOI: 10.1021/ja01176a521]. Its solubility (w/v) in EtOH at $\sim 25^\circ$ is 4.8%, but is less in H₂O which is 0.2%, even less in Et₂O and almost insoluble in *C₆H₆. The *ethyl ester* forms needles from EtOH with **m 115°**. The *benzoyl derivative* (prepared from PAS and benzoyl chloride in pyridine) has **m 185-187°**. The *hydrochloride*, **m 224°(dec)**, is crystallised from EtOH/HCl. Its alkali salts are more soluble in H₂O and as such are more useful, and decompose at high temperatures. The *potassium salt* (**Paskalium**) [133-09-5] C₇H₆KNO₃, **M 191.2**, is freely soluble in H₂O and less subject to stomach irritation than the free acid or the Na salt. The *sodium salt dihydrate* (**Aminacyl**) [6018-19-5] C₇H₆NNaO₃ · 2H₂O, **M 211.2**, has a solubility in H₂O at $\sim 25^\circ$ of 50% and is sparingly or almost insoluble in most organic solvents. The *calcium salt* (**Nippas calcium**) [133-15-3] C₁₄H₁₂CaN₂O₆, **M 344.3**, has a solubility in H₂O at $\sim 25^\circ$ of 14.3%, but slowly hydrolyses and darkens in colour. [Erlenmeyer et al. *Helv Chim Acta* **31** 988 1948, DOI: 10.1002/hlca.19480310404; Sheehan *J Am Chem Soc* **70** 1665 1948, DOI: 10.1021/ja01184a516, Hassan et al. *Anal Profiles Drug Subs* **10** 1 1981, DOI: 10.1016/S0099-5428(08)60636-X; *Beilstein* **14** IV 1967.] It is an antibacterial and is used in tuberculosis.

5-Aminosalicylic acid (5-amino-2-hydroxybenzoic acid, Mesalamine, 5-AS) [89-57-6] C₇H₇NO₃, **M 153.1, m 276-280°, 283°(dec), pK₁²⁵ 2.74 (CO₂H), pK₂²⁵ 5.84 (NH₂)**. It crystallises as needles from H₂O containing a little NaHSO₃ to avoid aerial oxidation to the quinone-imine. The *Me ester* gives needles from *C₆H₆, **m 96°**, and the *hydrazide* has **m 180-182°** (from H₂O). [Fallab et al. *Helv Chim Acta* **34** 26 1951, DOI: 10.1002/hlca.19510340104; Shavel *J Am Pharm Assoc* **42** 402 1953, PMID: 13069333; *Beilstein* **14** IV 2058.] It is anti-inflammatory, and is a substrate for peroxidase used in ELISA tests.

9-Aminotrypticene [793-41-9] C₂₀H₁₅N, **M 269.3, m 221-221..5°, 223.5-224.5°, pK²⁵ ~10.68**. Prepared by catalytic reduction of 9-nitrotrypticene [797-67-1] with Pd/C [Theilacker et al. *Chem Ber* **93** 1658 1960, DOI: 10.1002/cber.19600930728; Theilacker & Beyer *Chem Ber* **94** 2968 1961, DOI: 10.1002/cber.19610941121], the aminotrypticene formed can then be recrystallised from ligroin as white prisms. The ¹H NMR (400MHz, CDCl₃, and TMS) has δ at 7.47 (1-H, 8-H, 13-H), 7.05 (2-H, 7-H, 14-H), 7.00 (3-H, 6-H, 15-H), 7.37 (4-H, 5-H, 16-H), 5.40 (10-H, bridgehead) and 2.56 (NH₃) with $J_{1,2} = 7.5\text{Hz}$, $J_{1,3} = 0.5\text{Hz}$, $J_{1,4} = 0.0\text{Hz}$, $J_{2,3} = 7.4\text{Hz}$, $J_{2,4} = 1.2\text{Hz}$ and $J_{3,4} = 7.4\text{Hz}$; the ¹³C NMR (100MHz, CDCl₃, and TMS) has δ at 119.2 (1-C, 8-C, 13-C), 124.8 (2-C, 7-C, 14-C), 125.1 (3-C, 6-C, 15-C), 123.1 (4-C, 5-C, 16-C), 144.8 (4a-C, 10a-C, 11-C), 146.5 (9a-C, 8a-C, 12-C), 64.3 (9-C) and 53.1 (10-C); and the ¹³C NMR for the solid state has been recorded but is only slightly different from the solution spectrum. [Imashiro et al. *J Am Chem Soc* **109** 729 1987, DOI: 10.1021/ja00237a016].

p-tert-Amylphenol (p-2,2-dimethylpropylphenol) [80-46-6] C₁₁H₁₆O, **M 164.3, m 88-89°, 93.5-94.2°, 94-96°, b 112-°/3mm, 138°/15mm, 262°/760mm, d₄²⁰ 0.962, pK_{Est} ~10.2**. Purify *via* its benzoate, as for phenol. After evaporating the solvent from its solution in ether, the material is recrystallised (from the melt) to a constant melting point. It is insoluble in H₂O but soluble in most organic solvents. The *benzoyl* derivative has **m 60°** (from EtOH). [Berliner et al. *J Am Chem Soc* **76** 507 1954, DOI: 10.1021/ja01631a052; Huston et al. *J Am Chem Soc* **67** 899 1945, DOI: 10.1021/ja01222a005; *Beilstein* **6** H 548, **6** I 269, **6** II 506, **6** III 1965, **6** IV 3383.]

Aniline [62-53-3] C₆H₇N, **M 93.1, m -6.0°, b 68.3/10mm, 184.4°/760mm, d₄²⁰ 1.0220, n_D²⁰ 1.585, n_D²⁰ 1.5832, pK²⁵ 4.60**. Aniline is *hygroscopic*. It can be dried with KOH or CaH₂, and distilled under reduced pressure. Treatment with stannous chloride removes sulfur-containing impurities, reducing the tendency to become coloured by aerial oxidation. It can be crystallised from Et₂O at low temperatures. More extensive purifications involve preparation of derivatives, such as the double salt of aniline hydrochloride and cuprous chloride or zinc chloride, or *N*-acetylaniline (**m 114°**) which can be recrystallised from water. Redistilled aniline is dropped slowly into a strong aqueous solution of recrystallised oxalic acid. Aniline oxalate (**m 174-175°**) is filtered off, washed several times with water and recrystallised three times from 95% EtOH. Treatment with saturated Na₂CO₃ solution regenerated aniline which was distilled from the solution, dried and redistilled under reduced pressure [Knowles *Ind Eng Chem* **12** 881 1920, DOI: 10.1021/ie50129a017]. Alternatively, after refluxing with 10% acetone for 10hrs, aniline is acidified with HCl (to Congo Red) and extracted with Et₂O until colourless. Its hydrochloride is crystallised repeatedly before aniline is liberated with cold alkali, dried over solid KOH, and distilled. This sulfur-free product remains colourless in air [Hantzsch & Freese *Chem Ber*

27 2529 1894, DOI: 10.1002/cber.189402702255; Hantzsch & Freese *Chem Ber* **27** 2966 1894, DOI: 10.1002/cber.189402703601894]. Non-basic materials, including nitro compounds, are removed from aniline in 40% H₂SO₄ by passing steam through the solution for 1 hour. Pellets of KOH are then added to liberate the aniline which is steam distilled, dried with KOH, distilled twice from zinc dust at 20mm, dried with freshly prepared BaO, and finally distilled from BaO in an all-glass apparatus [Few & Smith *J Chem Soc* 753 1949, DOI: 10.1039/JR9490000753]. Aniline is absorbed through skin and is **TOXIC**. [Beilstein **12** IV 223.] **Aniline hydrobromide** [542-11-0] C₆H₇N. HBr, **M 174.0**, has **m 286°**. Crystallise the hydrobromide from water or EtOH and dry it at 5mm over P₂O₅. Also recrystallise it four times from MeOH containing a few drops of concentrated HBr by addition of petroleum ether (b 60-70°), then dry it to constant weight over paraffin chips *in vacuo* [Gutbezahl & Grunwald *J Am Chem Soc* **75** 559 1953, DOI: 10.1021/ja01099a014]. It precipitates from EtOH solution on addition of Et₂O, and the filtered solid is recrystallised from EtOH and dried *in vacuo*. Store it in the dark. [Buchanan et al. *J Am Chem Soc* **108** 1537 1986, DOI: 10.1021/ja00267a025; Beilstein **12** III 232.] **Aniline hydrochloride** [142-04-1] C₆H₇N. HCl, **M 129.6**, has **m 196-198°, 200.5-201°**. Purification is as for aniline HBr. [Beilstein **12** IV 232.] **Aniline hydroiodide** [45497-73-2] C₆H₇N. HI, **M 220.0**, it **decomposes on heating**. Purification is as for aniline HBr; store it in the dark. [Beilstein **12** III 232.]

m-Anisaldehyde (3-methoxybenzaldehyde) [591-31-1] **M 136.2**, **b 143°/50mm**, **d₄²⁰ 1.117**, **n_D²⁰ 1.553**. Wash it with saturated aqueous NaHCO₃, then H₂O, dry it with anhydrous MgSO₄ and distil it under reduced pressure under N₂. Store it under N₂ in sealed glass ampoules. [Beilstein **8** IV 241.] **p-Anisaldehyde (4-methoxybenzaldehyde)** [123-11-5] has **m -1°, b 249°/atm, 89-90°/2mm**, **d₄²⁰ 1.119**, **n_D²⁰ 1.576**. Wash the aldehyde with saturated aqueous NaHCO₃, then H₂O, steam distil, extract the distillate with Et₂O, dry (MgSO₄) the extract, filter and distil this under a vacuum and N₂. Store it in glass ampoules under N₂ in the dark. [Beilstein **8** IV 252.] **4-Methoxybenzaldehyde dimethylacetal** [2186-92-7] C₁₀H₁₄O₃, **M182.2**, with **b 85-87°/0.1mm**, **n_D²⁰ 1.505**, has been used as a protecting reagent for diols, as in carbohydrate chemistry, which can be easily and selectively removed under mild conditions [Kloosterman et al. *Recl Trav Chim Pays-Bas* **103** 243 1984, DOI: 10.1002/recl.19841030706; Peters et al. *Tetrahedron* **43** 3803 1987, DOI: 10.1016/S0040-4020(01)86866-6; Johansson & Samuelsson *JCS Perkins I* 2371 1984, DOI: 10.1039/P19840002371]. [Beilstein **8** H 74.]

o-Anisidine (2-methoxyaniline) [90-04-0] C₇H₉NO, **M 123.2**, **m ~5°, b 109°/17mm, 119°/21mm, 225°/atm**, **d₄²⁰ 1.096**, **n_D²⁰ 1.575**, **pK²⁵ 4.52**. It is separated from the *m*- and *p*- isomers by steam distillation. It is also separated from its usual synthetic precursor *o*-nitroanisole by dissolving it in dilute HCl (pH <2.0) extracting the nitro impurity with Et₂O, adjusting the pH to ~8.0 with NaOH, extracting the amine into Et₂O or steam distilling. Extract the distillate with Et₂O, dry the extract (Na₂SO₄), filter, evaporate and fractionate the residual oil. Protect the almost colourless oil from light which turns it yellow in color. [Biggs & Robinson *J Chem Soc* 388 1961, DOI: 10.1039/JR9610000388; Nodzu et al. *Yakugaku Zasshi (J Pharm Soc Japan)* **71** 713, 715 1951, Beilstein **13** IV 806.] **m-Anisidine (3-methoxyaniline)** [536-90-3] has **m -1° to +1°, b ~5°, b 79°/1mm, 128°/17mm, 251°/atm**, **d₄²⁰ 1.101**, **n_D²⁰ 1.583**, **pK²⁵ 4.20**. *o*-Isomer impurity can be removed by steam distillation. Another possible impurity is the precursor 3-nitroanisole which can be removed as for the preceding *o*-isomer and fractionating using an efficient column. It is a yellow liquid. [Gilman & Kyle *J Am Chem Soc* **74** 3027 1952, DOI: 10.1021/ja01132a023; Bryson *J Am Chem Soc* **82** 4858 1960, DOI: 10.1021/ja01503a028; Kadaba & Massie *J Org Chem* **22** 333 1957, DOI: 10.1021/jo01354a610; Beilstein **13** IV 953.] **p-Anisidine (4-methoxyaniline)** [104-94-9] has **m 57°, pK²⁵ 5.31**. Crystallise *p*-anisidine from H₂O or aqueous EtOH. Dry it in a vacuum oven at 35° for 6 hours and store it in a dry box. It does not distil with steam (see above). [Moore et al. *J Am Chem Soc* **108** 2257 1986, DOI: 10.1021/ja00269a022.] Purify it also by vacuum sublimation [Guarr et al. *J Am Chem Soc* **107** 5104 1985, DOI: 10.1021/ja00304a015]. [Beilstein **13** IV 1015.]

Anisole [100-66-3] C₇H₈O, **M 108.1**, **m -37.5°, b 43°/11mm, 153.8°/760mm**, **d₁₅ 0.9988**, **n_D²⁰ 1.5143**, **pK⁰ -6.61 (aqueous H₂SO₄)**. Shake anisole with half its volume of 2M NaOH, and the emulsion is allowed to separate. Repeat three times, then wash twice with water, dry over CaCl₂, filter, dry over sodium wire and finally distil it from fresh sodium under N₂ using a Dean-Stark trap (samples in the trap being rejected until free from turbidity) [Caldin et al. *JCS Faraday Trans I* **72** 1856 1976, DOI: 10.1039/F19767201856]. **Alternatively**, dry it with CaSO₄ or CaCl₂, or by refluxing with sodium or BaO with crystalline FeSO₄ or by passage through an alumina column. Traces of phenols are removed by prior shaking with 2M NaOH, followed

by washing with water. It has been purified by zone refining. [*Beilstein* 6 IV 548.]

2-*p*-Anisyl-1,3-indanone (anisindione, Unidone) [117-37-3] $C_{16}H_{12}O_3$, M 252.3, m 156-157°, pK²⁰ 4.09. Crystallise anisindione from acetic acid or EtOH. [Horeau & Jacques *Bull Soc Chim Fr* 53 1948, Koelsch *J Am Chem. Soc* 58 1331 1936, DOI: 10.1021/ja01299a005; *Beilstein* 8 III 2931.] It is an anticoagulant.

Anthracene [120-12-7] $C_{14}H_{10}$, M 178.2, m 215-216°, 218°, b 342°/760mm, pK²⁵ -7.4 (aqueous H_2SO_4). Likely impurities are anthraquinone, anthrone, carbazole, fluorene, 9,10-dihydroanthracene, tetracene and bianthryl. Carbazole is removed by continuous-adsorption chromatography [see Sangster & Irvine *J Chem Phys* 24 670 1956 DOI: org/10.1063/1.1742596] using a neutral alumina column and eluting with *n*-hexane. [Sherwood in *Purification of Inorganic and Organic Materials*, Zief (ed), Marcel Dekker, New York, 1969.] The solvent is evaporated, and anthracene is sublimed under vacuum, then purified by zone refining, under N_2 in darkness or non-actinic light. It has also been purified by co-distillation with ethylene glycol (boils at 197.5°), from which it can be recovered by addition of water, followed by crystallisation from 95% EtOH, *benzene, toluene, a mixture of *benzene/xylene (4:1), or Et_2O . It has also been chromatographed on alumina with petroleum ether in a dark room (to avoid photo-oxidation of adsorbed anthracene to anthraquinone). Other purification methods include sublimation in a N_2 atmosphere (in some cases after refluxing with sodium), and recrystallisation from toluene [Gorman et al. *J Am Chem Soc* 107 4404 1985, DOI: 10.1021/ja00301a006]. Anthracene has been crystallised from EtOH, chromatographed through alumina in hot *benzene (*fume hood*) and then sublimed in a vacuum in a pyrex tube that has been cleaned and baked at 100°. (For further details see Craig & Rajikan *JCS Faraday Trans 2*, 74 292 1978, DOI: 10.1039/F29787400292; and Williams & Zboinski *JCS Faraday Trans 1* 74 618 1978, DOI: 10.1039/F29787400618.) It has been chromatographed on alumina, recrystallised from *n*-hexane and sublimed under reduced pressure. [Saltiel *J Am Chem Soc* 108 2674 1986, DOI: 10.1021/ja00270a028; Masnovi et al. *J Am Chem Soc* 108 1126 1986, DOI: 10.1021/ja00266a003.] Alternatively, recrystallise it from cyclohexane, chromatograph it on alumina with *n*-hexane as eluent, and recrystallise two more times [Saltiel et al. *J Am Chem Soc* 109 1209 1987, DOI: 10.1021/ja00238a034]. Anthracene is fluorescent and forms a *picrate complex*, m 139°, on mixing the components in $CHCl_3$ or C_6H_6 , but decomposes on attempted crystallisation. [*Beilstein* 5 IV 2281.]

Anthracene-9-carboxylic acid (anthroic acid) [723-62-6] $C_{15}H_{10}O_2$, M 222.2, m 213-217°, 214°(dec), pK²⁰ 3.65. Crystallise the acid from EtOH. It is fluorescent in EtOH with λ_{max} at 254nm (0.1% EtOH). The corresponding alcohol **9-anthracenemethanol** [1468-95-7] $C_{15}H_{12}O$, M 208.3, has m 162-164° (from MeOH); and the respective methacrylate ester **9-anthracenylmethyl methacrylate** [31645-35-9] $C_{19}H_{16}O_2$, M 276.3, has m 80-85° [from C_6H_6 /petroleum ether (b 40-60°) and λ_{max} at 249nm (EtOH).] [*Beilstein* 9 IV 2671.]

9-Anthraldehyde (anthracene-9-carboxaldehyde) [642-31-9] $C_{15}H_{10}O$, M 206.2, m 103-105°, 104-105°. Crystallise the aldehyde from acetic acid or EtOH. It forms a *charge-transfer complex* with TCNE (tetracyanoethylene) in $CHCl_3$. [Masnovi et al. *J Am Chem Soc* 108 1126 1986, DOI: 10.1021/ja00266a003; *Beilstein* 7 IV 1738.] The *syn-oxime* [34810-13-4] $C_{15}H_{11}NO$, M 221.3, has m 162-164° (from MeOH) [*Beilstein* 7 III 2528].

Anthranol (enol tautomer of anthrone, see below) [529-86-2] $C_{14}H_{10}O$, M 196.2, m 120°(rapid heating), 160-170°(dec. but sainters >100°), pK²⁵ -5.5 (aqueous H_2SO_4). Crystallise anthrol from glacial acetic acid or aqueous EtOH. It is the less stable of two tautomers and is obtained by heating anthrone in aqueous NaOH (but not in cold NaOH) whereby it dissolves to form the sodium salt of anthrol. On cautious acidification anthrol precipitates as a yellowish-brown solid. It changes into anthrone on keeping, as it does on melting (120° in a preheated bath). The melting point depends on the initial bath temperature, and recrystallisation may cause it to tautomerise to *anthrone* (see below). The *acetate* (m 134°) crystallises from petroleum ether (m 134°) or EtOH (m 135.5-137°). [Meyer *Justus Liebigs Ann Chem* 379 37 1911, DOI: 10.1002/jlac.19113790104; Barnett et al. *J Chem Soc* 885 1928, DOI: 10.1039/JR9280000885, for tautomerism see Almdal et al. *Acta Chem Scand Series B* 40 230 1986, DOI:10.3891/acta.chem.scand.09-1425.] An absolute EtOH solution consists of 89% anthrone and 11% anthranol in equilibrium.

Anthranthrone [641-13-4] $C_{22}H_{10}O_2$, M 306.3, m 300°, 415°, pK²⁵ -7.9 (aqueous H_2SO_4). It is prepared by

intramolecular *bis*-cyclisation (dehydration) of 1,1'-dinaphthyl-8,8'-dicarboxylic acid and is purified by recrystallisation from chlorobenzene, nitrobenzene or CHCl_3 (**m 340°**). [*Beilstein* 7 I 451, 7 II 783, 7 III 4406, 7 IV 2694.]

Anthraquinone [84-65-1] $\text{C}_{14}\text{H}_8\text{O}_2$, **M 208.2**, **m 284-286°**, **286°**, **m 377°/atm**, **379-381°/atm**, **pK²⁵ -8.27 (aqueous H₂SO₄)**. Crystallise anthraquinone from CHCl_3 (38ml/g), *benzene, or boiling acetic acid, wash it with a little EtOH and dry it under vacuum over P_2O_5 . Its solubility (w/w) in EtOH is 0.44% (at 25°) and 2.3% (at ~78°); in Et₂O it is 0.11% (at 25°); in CHCl_3 it is 0.63% (at ~25°) and 1.6% (at ~61°); and in *C₆H₆ it is 0.26% (at ~25°) and 1.8% (at ~80°). [*Beilstein* 7 IV 2556.]

Anthrarufin (1,5-dihydroxy-9,10-anthraquinone) [117-12-4] $\text{C}_{14}\text{H}_8\text{O}_4$, **M 240.1**, **m 279°(dec)**, **280°(dec)**, **pK₁²⁵ 9.90**, **pK₂²⁵ 11.05**. Purify anthrarufin by column chromatography on silica gel with $\text{CHCl}_3/\text{Et}_2\text{O}$ as eluent, followed by recrystallisation from acetone. Alternatively, recrystallise it from glacial acetic acid [Flom & Barbara *J Phys Chem* **89** 4489 1985, DOI: 10.1021/j100267a017]. [*Beilstein* 7 III 2359, 8 IV 3268.]

1,8,9-Anthratrionol (1,9,8-trihydroxyanthracene) [480-22-8] $\text{C}_{14}\text{H}_{10}\text{O}_3$, **M 226.2**, **m 176-181°**, **pK_{Est} ~9.5**. Crystallise it from petroleum ether and dry it *in vacuo*. [*Beilstein* 6 IV 7602.]

Anthrimide (1,1'-imino-bis-anthraquinone, 1,1'-dianthrimide) [82-22-4] $\text{C}_{28}\text{H}_{15}\text{NO}_4$, **M 429.4**, **m >250°(dec)**. Crystallise anthrimide from chlorobenzene (red needles) or nitrobenzene (red rhombs). The needles are copper-like and metallic-looking. It is poorly soluble in organic solvents. [Eckert & Steiner *Monatsh Chem* **35** 1129 1914, DOI:10.1007/BF01518035.]

Anthrone [90-44-8] $\text{C}_{14}\text{H}_{10}\text{O}$, **M 194.2**, **m 155°**, **pK²⁵ -5.5 (aqueous H₂SO₄)**. This stable keto tautomer of 9-anthranol (above) provides yellow crystals from a 3:1 mixture of *C₆H₆/petroleum ether (b 60-80°) (10-12ml/g), or successively from *C₆H₆ then EtOH. Dry it *in vacuo*. [Meyer *Org Synth Coll Vol* **1** 60 1941, DOI: 10.15227/orgsyn.008.0008; *Beilstein* 6 IV 4930.] An absolute EtOH solution consists of 89% anthrone and 11% anthranol in equilibrium.

(±)-Atrolactic acid (0.5H₂O) (2-hydroxy-2-phenylpropionic acid) [515-30-0] $\text{C}_9\text{H}_{10}\text{O}_3 \cdot 0.5\text{H}_2\text{O}$, **M 166.2**, **m 94.5-95°(anhydrous)**, **88-91° (0.5H₂O)**, **pK¹⁸ 3.53**. Crystallise the acid from water (4g/20ml boiling H₂O with charcoal, filter and cool overnight at 0-5°) and dry it at 55°/0.5mm [Eliel & Freeman *Org Synth Coll Vol* **4** 58 1963, DOI: 10.15227/orgsyn.033.0007]. The **(±)-ethyl ester** [76496-51-0] has **b 250°/~760mm**, **129-130°/13mm**. [*Beilstein* 10 H 259, 10 I 113, 10 II 157, 10 III 560, 10 IV 467.] **(±)-Atrolactamide** [2019-68-3, *S*-5908-94-1] has **m 101-102°** (leaflets from CH_2Cl_2), and is freely soluble in H₂O [prep. form the ethyl ester and ammonia see McKenzie & Smith *J Chem Soc* **121** 1348 1922, DOI: 10.1039/CT9222101348; and by hydrolysis of the nitrile with fuming HCl see Staudinger & Ruzicka *Justus Liebigs Ann Chem* **380** 291 1911, DOI: 10.1002/jlac.19113800303], and is an *anticonvulsant*.

R(-) and S(+) Atrolactic acid [*R* 3966-30-1 and *S* 13113-71-8] $\text{C}_9\text{H}_{10}\text{O}_3$, **M 166.2**, **m 115-116°**, **116-117°**, **R [α]_D^{14.6} -37.7 (c 3.3 EtOH) and [α]_D^{14.6} -51.1 (c 2.2, H₂O)**, **S [α]_D^{14.6} +37.7 (c 3.5 EtOH)**. Recrystallise them from *C₆H₆ or *C₆H₆/petroleum ether and dry them in a vacuum. [McKenzie & Clough *J Chem Soc* 1016 1910, DOI: 10.1039/CT9109701016; Meyers & Slade *Synth Commun* **6** 601 1976, DOI: 10.1080/00397917608063555; *Beilstein* 10 II 157, 10 III 560, 10 IV 657.] **S(+)-Atrolactamide** [5908-94-1], obtained from the ethyl ester and ammonia has **m 62.5-63.5°** (glassy rhombic plates from *C₆H₆), **[α]_D¹⁴ +12.6 (c 1.87, EtOH)**, **[α]_D¹⁵ +12.8 (c 2.23, Me₂CO)** and is freely soluble in H₂O [McKenzie & Smith *J Chem Soc* **121** 1348 1922, DOI: 10.1039/CT9222101348]. It is an *anticonvulsant*.

Auramine O (4,4'-bis-dimethylaminobenzophenone imine hydrochloride) [2465-27-2] $\text{C}_{17}\text{H}_{22}\text{ClN}_3$, **M 321.9**, **m >250°(dec)**, **pK²⁵ 10.71 (free base)**, **9.78 (carbinolamine)**. It crystallises from EtOH as the hydrochloride and is very slightly soluble in CHCl_3 , its UV has λ_{max} at 434nm (ε 370). The **free base** (carbinolamine) has **m 136°** (from *C₆H₆). [Goldacre & Phillips *J Chem Soc* 1724 1949, DOI: 10.1039/JR9490001724; Conrad et al. *Biochemistry* **9** 1540 1970, DOI: 10.1021/bi00809a010; *Beilstein* 14 IV 256.]

Aurin tricarboxylic acid [4431-00-9] $C_{22}H_{14}O_9$, **M 422.4, m 300°**. The acid is dissolved in aqueous NaOH, $NaHSO_3$ solution is added until the colour is discharged and then the tricarboxylic acid is precipitated with HCl. [Heisig & Lauer *Org Synth Coll Vol I* 54 1941, DOI: 10.15227/orgsyn.009.0008; *Beilstein* 10 IV 4161]. Do not extract the acid with hot water because it softens, forming a viscous mass. Make a solution in aqueous NH_3 . **Aurin tricarboxylic acid ammonium salt (Aluminon, ATA)** [569-58-4] $C_{22}H_{14}O_9 \cdot 3NH_3$, **M 473.4, m 220-225°(dec)** is the NH_4 salt which polymerises readily in aqueous solution, in which it is freely soluble, forming a stable free radical. It has a variety of physiological activities. It forms brilliantly coloured salts (lakes) with Al, Cr, Fe and Be ions, and is used for the estimation of Al in tissues, H_2O and foodstuffs due to the colour that it forms. Store dry preferably in the dark.

4-Azidoaniline hydrochloride [91159-79-4] $C_6H_6N_4 \cdot HCl$, **M 170.6, m 165°(dec)**. Purify it by dissolving it in EtOAc with dry HCl gas followed by addition of cold dry Et_2O . Dry it *in vacuo* over P_2O_5 . The **free base** has **m 66°(dec)** (MeOH) and the **picrate** has **m 64-65°(dec)** (MeOH). The IR has ν_{max} (Nujol) at $2110cm^{-1}$ (N_3). [Escher et al. *Helv Chim Acta* 62 1217 1979, DOI: 10.1002/hlca.19790620432; Maffei & Rivolta *Gazetta Chim Ital* 84 750 1954, *Beilstein* 12 H 772, 12 IV 1741.]

Azobenzene [103-33-3] $C_{12}H_{10}N_2$, **M 182.2, m 65-68°, 68°, b 293°/atm, pK^{25} 2.48**. Ordinary azobenzene is nearly all in the *trans*-form. It is partly converted into the *cis*-form on exposure to light [for isolation see Hartley *J Chem Soc* 633 1938, DOI: 10.1039/JR9380000633, and for spectra of *cis*- and *trans*-azobenzenes, see Winkel & Siebert *Chem Ber* 74B 670 1941, DOI: 10.1002/cber.19410740503]. *trans*-Azobenzene is obtained by chromatography on alumina using 1:4 C_6H_6 /heptane or petroleum ether, and it crystallises from EtOH (after refluxing for several hours) or hexane. All operations should be carried out in diffuse red light or in the dark. It is almost insoluble in H_2O but soluble in organic solvents and AcOH. [cf: Vogel *Practical Organic Chemistry* 3rd Edn Longmans, London, p 631 1959, *Beilstein* 16 IV 8.]

Azolitmin B (colouring matter from Litmus) [1395-18-2] **M ~3300, m >250°(dec)**. It is a mixture of water soluble dyes extracted from lichens. It crystallises from water as dark violet scales, or as a red powder by precipitating 0.5g from H_2O (80ml) by addition of EtOH (20ml). It is an indicator which is red at pH 4.5 and blue at pH 8.3. The chromophore of litmus components is 7-hydroxyphenoxaz-1-one. [cf. Kolthoff & Rosenblum *Acid-Base Indicators*, Macmillan, NY, pp160-162, 365-366 1937.]

***p,p'*-Azoxyanisole (4,4'-dimethoxyazoxybenzene)** [1562-94-3] **M 258.3, transition temperatures: 118.1-118.8°, 135.6-136.0°, pK^{25} -5.23 (20% aqueous EtOH + 80% aqueous H_2SO_4)**. Crystallise the dye from absolute or 95% EtOH, or acetone, and dry it by heating under vacuum or sublime it in a vacuum onto a cold finger. [*Beilstein* 16 II 326.]

Azoxybenzene (Fenazox) [495-48-7] $C_{12}H_{10}N_2O$, **M 198.2, m 36° [*trans*, 20972-43-4], m 87° [*cis*, 21650-65-7], pK^{25} -6.16 (20% aqueous EtOH + 80% aqueous H_2SO_4)**. Crystallise azoxybenzene from EtOH or MeOH, and dry it for 4 hours at $25^\circ/10^{-3}mm$. Sublime it before use. [Bigelow & Palmer *Org Synth Coll Vol II* 57 1943, DOI: 10.15227/orgsyn.011.0016; *Beilstein* 16 II 326.] *Cis*- and *trans*- azoxybenzene were respectively prepared by oxidation of the respective *cis*- and *trans*- azobenzenes with perbenzoic acid in C_6H_6 . The rates of reactions (second order rates at 25°) showed that *cis*-azobenzene was oxidised to *cis*-azoxybenzene much more rapidly than the *trans*- isomer. After the oxidation of the *trans*- isomer in the dark, the reaction mixture was washed with dilute Na_2CO_3 solution to remove acids, then H_2O , dried, the solvent was evaporated and the residue was purified by chromatography on Al_2O_3 with light petroleum which gave *trans*-azoxybenzene, **m 36°**. When this oxidation was carried out in the presence of sunlight, *cis*-azoxybenzene, **m 86°** was obtained. Similar oxidation of *cis*-azobenzene, in the dark, but without chromatographic purification gave a yellow solid which was freed from *trans*-azoxybenzene by trituration with petroleum ether (b 40-60°) *ca* 4 times to give the much less soluble *cis*-azoxybenzene, **m 87°**. When heated at 100° for 0.5hr, this *cis*-compound was converted into *trans*-azoxybenzene, **m 36°**. **Note:** that when *cis*-azoxybenzene was reduced with $LiAlH_4$ in dry Et_2O only *trans*-azobenzene, **m 68°** was obtained, and was unchanged after resolidification. [Badger et al. *J Chem Soc* 2143 1953, DOI: 10.1039/JR9530002143.] *Trans*-azoxybenzene is insoluble in H_2O but is steam volatile; and its solubility (w/w at $\sim 15^\circ$) in absolute EtOH is 17.5%, but much higher, 43.5%, in petroleum ether (b 40-60°).

***p,p*-Azoxyphenetole (4,4'-diethoxyazoxybenzene)** [4792-83-0] $C_{16}H_{18}N_2O_3$, **M 286.3**, **m 137-138°, 138.5°**, (melts to a turbid liquid crystal at *ca* **165°** which clarifies at **167.5°**). Crystallise the dye from toluene or EtOH. For the melting character see Homfray [*J Chem Soc* **97** 1669 1910, DOI: 10.1039/CT9109701669]. It has λ_{\max} at 360nm [Purvis *J Chem Soc* **107** 660 1915, DOI: 10.1039/CT9150700660]. [*Beilstein* **16** H 638, **16** I 384, **16** II 326.]

Azulene [275-51-4] $C_{10}H_8$, **M 128.2**, **m 98.5-99°, 98-100°, b 115-135°/10mm, 242°/atm, pK^{25} -1.65 (aqueous H_2SO_4)**. Crystallises from EtOH in intensely blue prisms or leaflets with the odour of naphthalene. It forms a **trinitrobenzene complex** $C_{16}H_{11}N_3O_6$, **m 166.5-167.7°**, when mixed with one equivalent of 1,3,5-trinitrobenzene which recrystallised from EtOH as fine brown needles. Azulene can be recovered from this complex by running it on a thick Al_2O_3 TLC plate and eluting with cyclohexane- $*C_6H_6$ (1:1) which separates the components. Also, sublimation of the complex at 150° in a CO atmosphere produces azulene as blue metallic-looking plates, **m 98.5-99°**. **Azulene trinitrotoluene complex** $C_{17}H_{13}N_3O_6$, **m 99.5-100°**, is prepared similarly with trinitrotoluene and recrystallised from EtOH. [Plattner & Pfau *Helv Chim Acta* **20** 224 1937, DOI: 10.1002/hlca.19370200133.] Azulene has UV with λ_{\max} ($\log \epsilon$) at 272nm (5.1), 296nm (3.9), 326nm (3.9), 340nm (4.0) and 358nm (3.5) in hexane. [Susz et al. *Helv Chim Acta* **20** 469 1937, DOI: 10.1002/hlca.19370200173, and IR: Plattner & Wyss *Helv Chim Acta* **24** 483 1941, DOI: 10.1002/hlca.19410240166; *Beilstein* **5** IV 1636.]

Benzalacetone (trans-4-phenyl-3-buten-2-one) [122-57-6] $C_{10}H_{10}O$, **M 146.2**, **m 41.5°, 42°, b 81.7°/0.1mm, 127°/10mm, 161°/40mm. 260-262°/atm, d_4^{25} 1.0079, $n_D^{45.9}$ 1.5836**. Crystallise it from petroleum ether (b 40-60°), or distil it, preferably under reduced pressure. It is steam volatile, and is soluble in most organic solvents but not in petroleum ether. [Drake & Allen *Org Synth Coll Vol I* 77 1941, DOI: 10.15227/orgsyn.003.0017; *Beilstein* **7** IV 1003.]

Benzalacetophenone (Chalcone) [94-41-7] $C_{15}H_{20}O$, **M 208.3**, **m 56-58°, 57-58°, b 208°/25mm, 345-348°/atm (some dec), d_4^{62} 1.0712, n_D^{62} 1.5836. pK^{25} -5.73 (aqueous H_2SO_4)**. Crystallise it from EtOH by warming to 50° (about 5ml/g), iso-octane, or toluene/petroleum ether, or recrystallise it from MeOH, and then twice from hexane. SKIN IRRITANT. [Kohler & Chadwell *Org Synth Coll Vol I* 78 1941, DOI: 10.15227/orgsyn.002.0001; *Beilstein* **7** IV 1658.]

Benzaldehyde [100-52-7] C_7H_6O , **M 106.1**, **m -56.5°, -26°, b 62°/10mm, 179.0°/760mm, d_4^{20} 1.044, n_D^{20} 1.5455, pK^{25} -7.1 (aqueous H_2SO_4)**. To diminish its rate of oxidation, benzaldehyde usually contains additives such as hydroquinone or catechol. It can be purified *via* its bisulfite addition compound but usually distillation (under nitrogen at reduced pressure) is sufficient. Prior to distillation it is washed with NaOH or 10% Na_2CO_3 (until no more CO_2 is evolved), then with saturated Na_2SO_3 and H_2O , followed by drying with $CaSO_4$, $MgSO_4$ or $CaCl_2$. Its solubility in H_2O is 0.3% but it is steam volatile. Benzaldehyde has the odour of almonds as it is a component of *amygdalin* which is present in bitter almonds and apricot pits. [Prepn from benzal chloride: Vogel *Practical Organic Chemistry* 3rd Edn Longmans, London, p 693 1959, *Beilstein* **7** IV 505.] **Amygdalin** (R(D)-mandelonitrile 6-O- β -D-glucosido- β -D-glucoside) [29883-15-6] $C_{20}H_{27}NO_{11}$, **M 457.4**, is the cyanohydrin glycoside of benzaldehyde which is hydrolysed by acid or enzymatically to benzaldehyde, HCN and two molecules of glucose [Conn *Int Rev Biochem (Biochemistry of Nutrition IA)* **27** 21 1979]. **anti-Benzaldoxime** [932-90-1] C_7H_7NO , **M 121.1**, has **m 33-34°**. Crystallise the oxime from diethyl ether by adding petroleum ether (b 60-80°). The *syn*-isomer [622-32-2] has **b 121-124°/12mm, m 34-36°**. [*Beilstein* **7** H 218, **7** IV 527.] It has also been crystallised from dilute aqueous NH_3 , H_2O , Me_2CO , then $*C_6H_6$ using a Soxhlet extractor. Its solubility (w/v at ~20°) in EtOH is 16.7% and in pyridine it is 30.3%. Dry it in an oven at 110° for 8 hours and store in a desiccator over 99% H_2SO_4 . It can be purified by zone melting or by sublimation. [Bates & Hobbs *J Am Chem Soc* **73** 2151 1951, DOI: 10.1021/ja01149a070; *Beilstein* **9** IV 725.]

Benzamidine [618-39-3] $C_7H_8N_2$, **M 120.2**, **m 65-70°, 64-66°, pK^{20} 11.6**. It is liberated from its hydrochloride chloride (see below) by treatment with 5M NaOH, extracted into Et_2O , dried (Na_2SO_4), filtered, evaporated and the residue was sublimed *in vacuo*. [*Beilstein* **9** H 280, **9** I 123, **9** II 199, **9** 1264, **9** IV 898.]

Benzamidine hydrochloride hydrate [*hydrate* 206752-36-5; *anhydrous* 1670-14-0] $C_7H_8N_2 \cdot HCl \cdot H_2O$, **M 156.6 (anhydrous), m 86-88° (hydrate), m 169-173° (anhydrous), pK²⁰ 11.6** (see free base above). Recrystallise it from dilute HCl (crystals contain xH_2O) or EtOH/few drops HCl; dry it in a vacuum. It is a proteolytic inhibitor [Jeffcoate & White *J Clin Endocrinol Metab* **38** 155 1974, DOI: org /10.1210/jcem-38-1-155; *Beilstein* **9** IV 898.]

Benzanilide [93-98-1] $C_{13}H_{11}NO$, **M 197.2, m 163°, 164°, b 117-119°/10mm, pK⁵⁵ 1.26**. Crystallise benzanilide from petroleum ether (b 70-90°) using a Soxhlet extractor, and dry it *in vacuo* overnight at 120°. Also crystallise it from EtOH. Its solubility (w/v) in EtOH is 1.6% (at ~20°) and 14.3% (at ~78°), and it is sparingly in Et₂O but almost insoluble in H₂O. [Webb *Org Synth Coll Vol I* 82 1941, DOI: 10.15227/orgsyn.007.0006; *Beilstein* **12** IV 417.]

Benz[a]anthracene (1,2-benzanthracene, tetraphene) [56-55-3] $C_{18}H_{12}$, **M 228.3, m 157-159°, 159-160°, 167°, b 437.6°/atm**. Crystallise 1,2-benzanthracene from AcOH, MeOH, EtOH or *C₆H₆ (charcoal), then chromatograph it on alumina from sodium-dried *benzene (twice), using vacuum distillation to remove *benzene. Final purification is by vacuum sublimation. [Fieser & Hershberg *J Am Chem Soc* **59** 2502 1937, DOI: 10.1021/ja01291a008; *Beilstein* **5** IV 2549.]

Benz[a]anthracene-7,12-dione [2498-66-0] $C_{18}H_{10}O_2$, **M 258.3, m 169.5-170.5°, 169-171°**. Crystallise the dione from MeOH (charcoal), toluene, toluene/hexane, Me₂CO, or AcOH. Alternatively, purify it by sublimation *in vacuo*, or by zone refining. [*Beilstein* **7** H 826, **7** I 440, **7** II 760, **7** III 4278, **7** IV 2644.]

Benzanthrone [82-05-3] $C_{17}H_{10}O$, **M 230.3, m 168-170°, 170°, 170-173°, pK²⁵ -3.2 (aqueous H₂SO₄)**. Crystallise benzanthrone from EtOH or xylene. The solubility (w/w at ~20°) in *PhCl is 2.05%, in *C₆H₆ is 1.61% and in AcOH it is 0.52%. On a larger scale benzanthrone (~58g) is dissolved in tetrachloroethane (175ml, b 147°/atm), charcoal (12g) is added and boiled under reflux for 15min, filtered, the carbon is washed with hot solvent (50ml), the combined hot filtrates are diluted with boiling EtOH (750ml) and set aside to crystallise. Pure benzanthrone, m 170-171°, separates as yellow needles (~50g, after drying *in vacuo*). [Macleod & Allen *Org Synth Coll Vol 2* 62 1941, DOI: 10.15227/orgsyn.014.0004; *Beilstein* **7** IV 1819.]

***Benzene** [71-43-2] C_6H_6 , **M 78.1, f 5.5°, b 80.1°/atm, d₄²⁰ 0.874, n_D²⁰ 1.50110, n_D²⁵ 1.49790**. For most purposes, *benzene can be purified sufficiently by shaking with conc H₂SO₄ until free from thiophene, then with H₂O, dilute NaOH and water, followed by drying (with P₂O₅, sodium, LiAlH₄, CaH₂, 4X Linde molecular sieve, or CaSO₄, or by passage through a column of silica gel, and for a preliminary drying, CaCl₂ is suitable), and distillation. A further purification step to remove thiophene, acetic acid and propionic acid, is crystallisation by partial freezing. The usual contaminants in dry thiophene-free *benzene are non-benzenoid hydrocarbons such as cyclohexane, methylcyclohexane, and heptanes, together with naphthenic hydrocarbons and traces of toluene. Carbonyl-containing impurities can be removed by percolation through a *Celite column impregnated with 2,4-dinitrophenylhydrazine, phosphoric acid and H₂O*. (Prepared by dissolving 0.5g DNPH in 6ml of 85% H₃PO₄ by grinding together, then adding and mixing 4ml of distilled H₂O and 10g Celite.) [Schwartz & Parker *Anal Chem* **33** 1396 1961, DOI: 10.1021/ac60178a036.] *Benzene has been freed from thiophene by refluxing with 10% (w/v) of Raney nickel for 15 minutes, after which the nickel is removed by filtration or centrifugation. Dry *benzene is obtained by doubly distilling high purity *benzene from a solution containing the blue ketyl formed by the reaction of sodium-potassium alloy with a small amount of benzophenone. Thiophene has been removed from *benzene (absence of a bluish-green coloration when 3ml of *benzene is shaken with a solution of 10mg of isatin in 10ml of concentrated H₂SO₄) by refluxing the *benzene (1.25L) for several hours with 40g HgO (freshly precipitated) dissolved in 40ml glacial acetic acid and 300ml of water. The precipitate is filtered off, the aqueous phase is removed and the *benzene is washed twice with H₂O, dried and distilled. Alternatively, *benzene dried with CaCl₂ has been shaken vigorously for 0.5 hour with anhydrous AlCl₃ (12g/L) at 25-35°, then decanted, washed with 10% NaOH, and water, dried and distilled. The process is repeated, giving thiophene-free *benzene. [Holmes & Beeman *Ind Eng Chem* **26** 172 1934, DOI: 10.1021/ie50290a011.] After shaking successively for about an hour with conc H₂SO₄, distilled water (twice), 6M NaOH, and distilled water (twice), *benzene is distilled through a 3-ft glass column to remove most of the water. Absolute EtOH is added and the *benzene-alcohol azeotrope is distilled. (This low-boiling distillation leaves any non-azeotrope-

forming impurities behind.) The middle fraction is shaken with distilled water to remove EtOH, and again redistilled. Final slow and very careful fractional distillation from sodium, then LiAlH₄ under N₂, removed traces of water and peroxides. [Peebles et al. *J Am Chem Soc* **82** 4780 1960, DOI: 10.1021/ja01503a006.]

Benzene liquid and vapour are very TOXIC, CARCINOGENIC and HIGHLY FLAMMABLE, and all operations should be carried out in an efficient fume cupboard and in the absence of naked flames in the vicinity.* [Beilstein **5 H 175, **5** I 95, **5** II 119, **5** III 469.]

Rapid purification: To dry *benzene, alumina, CaH₂ or 4A molecular sieves (3% w/v) may be used (dry for 6 hours). Then *benzene is distilled, discarding the first 5% of distillate, and stored over molecular sieves (3A, 4A) or Na wire.

[²H₆]*Benzene (*benzene-d₆) [1076-43-3] C₆D₆, M 84.2, b 80°/773.6mm, 70°/562mm, 60°/399mm, 40°/186.3mm, 20°/77.1mm, 10°/49.9mm, 0°/27.5mm, d₄²⁰ 0.9488, d₄⁴⁰ 0.9257, n_D²⁰ 1.4991, n_D⁴⁰ 1.4865. Hexadeuteriobenzene of 99.5% purity is refluxed over and distilled from CaH₂ onto Linde type 5A sieves under N₂. [Beilstein **5** III 518, **5** IV 630.]

Benzeneazodiphenylamine (4-phenylazodiphenylamine) [28110-26-1; 101-75-7] C₁₈H₁₅N₃, M 273.3, m 82°, 86°, 89-91°, pK²² 0.48. Purify the dye by chromatography on neutral alumina using dry *C₆H₆ with 1% of dry MeOH. The major component, which gave a stationary band, is cut out and eluted with EtOH or MeOH. [Högfeldt & Bigeleisen *J Am Chem Soc* **82** 15 1960, DOI: 10.1021/ja01486a005.] It crystallises from petroleum ether, EtOH or aqueous EtOH, and has λ_{max} at 420nm (ε 28,000) (aqueous EtOH), and 540nm (aqueous EtOH/H₂SO₄) [Badger et al. *J Chem Soc* 1888 1954, DOI: 10.1039/JR9540001888; Beilstein **16** H 314, **16** III 343, **16** IV 457.]

1-Benzeneazo-2-naphthol (Sudan I) [842-07-9] C₁₆H₁₂N₂O, M 248.3, m 106°, 120,5°, 132° (polymorphism) 134°, 135°, pK_{Est} ~9.5 (OH). Crystallise the dye from EtOH. It forms Cu and Ni salts. [Beilstein **16** H 162, **16** I 254, **16** II 70, **16** III 129, **16** IV 228.]

1-Benzeneazo-2-naphthylamine (1-phenylazo-2-naphthylamine, Yellow AB) CI 11380 [85-84-7] C₁₆H₁₃N₃, M 247.3, m 102-104°, 103-104°, pK_{Est} ~4.1. Crystallise the dye from glacial acetic acid, acetic acid/water or ethanol. It forms Cu, Co and Ni salts. [Beilstein **16** H 369, **16** I 328, **16** II 193, **16** III 417, **16** IV 551.]

1,2-Benzenedimethanol (1,2-bishydroxymethylbenzene, phthalyl alcohol) [612-14-6] C₈H₁₀N₂, M 138.2, m 61-64°, 63-64°, 64-65°, 65-66.5°, b 145°/3mm. Recrystallise it from *C₆H₆, H₂O, petroleum ether or pentane. It has been extracted in a Soxhlet with Et₂O, evaporated and recrystallised from hot petroleum ether. It is also purified by dissolving in Et₂O, allowing to evaporate till crystals are formed, filtering off and washing the colourless crystals with warm petroleum ether or pentane. The *diacetate* has m 35°, 35-36°. [Nystrom & Brown *J Am Chem Soc* **69** 1197 1947, DOI: 10.1021/ja01197a060; IR and UV: Entel et al. *J Am Chem Soc* **74** 441 1952, DOI: 10.1021/ja01122a048; Beilstein **6** IV 5953.]

Benzene-1,2,4,5-tetracarboxylic (Pyromellitic acid) [89-05-4] C₁₀H₆O₈, M 254.2, m 276°, 281-284°, pK₁²⁵ 1.87, pK₂²⁵ 2.72, pK₃²⁵ 4.30, pK₄²⁵ 5.52. Dissolve it in 5.7 parts of hot dimethylformamide, decolorise, filter and cool. The precipitate is collected and dried in air. The solvent is removed by heating in an oven at 150-170° for several hours. It also crystallises from water. The tetracarboxylic acid forms a *mono-S-benzylisothiuronium salt* which crystallises from aqueous EtOH with m 245-246° (244° also reported). [Beilstein **9** H 997, **9** IV 3804.] Its *tetramethyl ester* [635-10-9] C₁₄H₁₄O₈, M 310.3, crystallises in leaflets from MeOH which have m 142° (141.5°, 143-144° also reported). **1,4,5,6-Benzenetetracarboxamide** [6183-35-3] C₁₀H₁₀N₄O₄, M 250.2, decomposes >300° (from aqueous NH₃); see below. The respective *tetranitrile* [712-74-3] C₁₀H₂N₄, M 175.2, has m 265-268°. (See 1,2,4,5-tetracyanobenzene below.) [Phillippi & Thelen *Org Synth Coll Vol* **2** 551 1943, DOI: 10.15227/orgsyn.010.0090; Beilstein **9** IV 3800.]

Benzene-1,2,4,5-tetracarboxylic dianhydride (Pyromellitic dianhydride) [89-32-7] C₁₀H₂O₈, M 218.1, m 286°, b 397-400°/760mm. Crystallise the dianhydride from ethyl methyl ketone or dioxane. Dry, and sublime it *in vacuo*. It is used for the estimation of alcohols and amines in the presence of aldehydes and phenols, and with DMSO as a solvent in the pyromellitic dianhydride method for determination of the amounts of alcohols and

amines [Siggia et al. *Analyt Chem* **33** 900 1961, DOI: 10.1021/ac60175a027; Harper et al. **37** 600 1965, DOI: 10.1021/ac60223a047]. [Beilstein **19** H 196, **19/5** V 407.]

Benzene-1,2,3-tricarboxylic (Hemimellitic) acid (H₂O) [anhydrous 569-51-7; hydrate 36362-97-7] C₉H₆O₆, **M 210.1** (anhydrous), **m 190°(dec.?)**, **190-192°(dec, hydrate)**, **195°(dec, anhydrous)**, **pK₁²⁵ 2.62**, **pK₂²⁵ 3.82**, **pK₃²⁵ 5.51**. Crystallise the acid from water to obtain the hydrate. Dehydrate at high vacuum over P₂O₅. [Beilstein **9** H 976, **9** IV 3745.] **Benzene-1,2,4-tricarboxylic (Trimellitic) acid (H₂O)** [528-44-9] has **m 229-231°(dec)**, **229-234°(dec)**, **218-220°** (also reported), **pK₁²⁵ 2.64**, **pK₂²⁵ 4.15**, **pK₃²⁵ 5.67**. Crystallise the acid from aqueous EtOH or glacial AcOH and dry over P₂O₅. It is soluble (w/w) at 25° in Me₂NCHO (31%), EtOH (25.3%), Me₂CO (7.9%), H₂O (2.1%), EtOAc (1.7%), petroleum ether (0.03%), CCl₄ (0.04%) and xylene (0.006%), but insoluble in CS₂, CHCl₃ or *C₆H₆. [Beilstein **9** H 976, **9** IV 3745.] Sublimation above its melting point yields the anhydride, **4-carboxyphthalic anhydride (Trimellitic anhydride, benzene-1,2,4-benzenetricarboxylic anhydride)** [552-30-7] C₉H₄O₅, **M 192.1**, **pK_{Est}²⁵ 2.2**, which can be distilled under a vacuum with **b 240-245°/14mm**, and/or recrystallised from EtOAc/petroleum ether, Me₂CO/petroleum ether or Ac₂O and dried over P₂O₅/vacuum to give colourless crystals with **m 163-166° (161-163.5° also reported)**. It is soluble (w/w) at 25° in Me₂CO (49.6%), EtOAc (21.6%), Me₂NCHO (15.5%), xylene (0.4%) and petroleum ether (0.06%). **Trimellitic anhydride chloride (benzene-1,2,4-tricarboxylic anhydride chloride)** [1204-28-0] C₉H₃ClO₄, **M 210.6**, has **m 66-68°** (from light petroleum b 60-80°). [Beilstein **18/8** V 562.] **Benzene-1,3,5-tricarboxylic (Trimesic) acid** [554-95-0] has **m >300°**, **360°(dec)**, **pK₁²⁵ 2.64**, **pK₂²⁵ 3.71**, **pK₃²⁵ 5.01**. Crystallise the acid from water or AcOH and dry it *in vacuo*. The **trimethyl ester** has **m 144°** (from MeOH or MeOH/H₂O). [Beilstein **9** H 978, **9** IV 3747.] The **tri-acid chloride (benzene-1,3,5-tricarbonyl chloride)** [4422-95-1] C₉H₃Cl₃O₃, **M 265.5**, is purified by distillation, **b 180°/16mm**, and by crystallisation from low boiling petroleum ether, **m 34.5-36°** (32-38° also reported). [Beilstein **9** IV 3748.]

1,2,4-Benzenetriol (hydroxyhydroquinone) [533-73-3] C₆H₆O₃, **M 126.1**, **m 140.5-141°(sintering at 139°)**, **pK₁²⁰ 9.08**, **pK₂²⁰ 11.82**. Crystallise the triol from Et₂O or Et₂O/EtOH, and dry it in a vacuum. It sublimes in a vacuum. The **triacetate**, **m 96.5-97°**, forms needles from absolute EtOH, the **tri-4-nitrobenzoate** has **m 120°**, and the **trimethyl ether** has **b 247°/atm**. The **picrate** forms orange-red needles **m 96°**. [Vliet *Org Synth Coll Vol 1* 317 1941, DOI: 10.15227/orgsyn.004.0035; Beilstein **6** H 1087, **6** I 541, **6** II 1071, **6** III 6276.]

Benzethonium chloride (Hyamine 1622, [diisobutylphenoxyethoxyethyl]dimethylbenzyl-ammonium chloride, (N,N-dimethyl-N-[2-[2-[4-(1,1,3,3-tetramethylbutyl)phenoxy]-ethoxy]-ethyl]-benzenemethan-ammonium chloride) [121-54-0] C₂₇H₄₂ClNO₂, **M 448.1**, **m 164-166° (sinters at 120°, monohydrate)**. Crystallise it from boiling acetone after filtering or from CHCl₃/petroleum ether. The crystals are filtered off, washed with diethyl ether and dried for 24 hours in a vacuum desiccator. It is a cationic antiseptic surfactant which forms crystals also from a 1:9 MeOH/Et₂O mixture. It foams in water, and is a disinfectant. [Beilstein **12** IV 2187.]

Benzhydrol (diphenylmethanol) [91-01-0] C₁₃H₁₂O₃, **M 184.2**, **m 69°**, **b 297°/748mm**, **180°/20mm**. Crystallise benzhydrol from hot H₂O or petroleum ether (b 60-70°), petroleum ether containing a little *benzene, from CCl₄, or EtOH (1ml/g). (Its solubility in H₂O at 25° is 0.1% w/v.) An additional purification step includes passage of a *benzene solution through an activated alumina column. It sublimes in a vacuum. Also recrystallise it three times from MeOH/H₂O [Naguib *J Am Chem Soc* **108** 128 1986, DOI: 10.1021/ja00261a021]. [Wiselogle & Sonneborn *Org Synth Coll Vol 1* 90 1941, DOI: 10.15227/orgsyn.008.0024; Beilstein **6** IV 4648.] § A commercial polystyrene supported version is available.

Benzidine (4,4'-diaminobiphenyl) [92-87-5] C₁₂H₁₂N₂, **M 184.2**, **m 128-129°**, **pK₁²⁰ 3.85**, **pK₂²⁰ 4.95**. Its solution in *benzene is decolorised by percolating through two 2-cm columns of activated alumina, then concentrated until benzidine crystallises on cooling. Recrystallise alternately from EtOH and *benzene to constant absorption spectrum [Carlin et al. *J Am Chem Soc* **73** 1002 1951, DOI: 10.1021/ja01147a035]. It has also been crystallised from hot water (charcoal) and from diethyl ether. Its solubility (w/v) in H₂O is 0.04% at 25° and 0.9% at 100°; in EtOH it is 20% at 78° and in Et₂O it is 2% at 25°. Dry it under vacuum in an Abderhalden pistol. Store it in the dark in a stoppered container. **CARCINOGENIC**. [Beilstein **13** IV 364.] **Benzidine dihydrochloride** [531-85-1] C₁₂H₁₂N₂. 2HCl, **M 257.2**, **m >250°(dec)**. Crystallise the salt by dis-

solving in hot H₂O, and adding concentrated HCl to the slightly cooled solution. Dry it *in vacuo* over KOH. **CARCINOGENIC.** [Beilstein 13 IV 365.]

Benzil (dibenzoyl, diphenylethanedione) [134-81-6] C₁₄H₁₀O₂, M 210.2, m 94-95°, 96-96.5°, b 188°/12mm, 346-348°/atm. Crystallise benzil from *benzene after washing with alkali. (Crystallisation from EtOH did not free benzil from material reacting with alkali.) [Hine & Haworth *J Am Chem Soc* 80 2274 1958, DOI: 10.1021/ja01542a060.] It has also been crystallised from CCl₄, diethyl ether or EtOH [Inoue et al. *JCS Faraday Trans 2* 82 523 1986, DOI: 10.1039/F29868200523]. Its IR has ν_{\max} (CHCl₃) at 1686, 1608 and 1460cm⁻¹. It condenses with vicinal diamines to form aza-heterocycles. [Clarke & Dreger *Org Synth Coll Vol I* 87 1941, DOI: 10.15227/orgsyn.006.0006; Beilstein 7 IV 2502.] Benzil forms three **benzildioximes** C₁₄H₁₂N₂O₂, M 240.3, which can be separated by fractional crystallisation *viz*: an α -(*anti*)-dioxime [E,E 23873-81-6] m 237°, a β -(*syn*)-dioxime [Z,Z 572-45-2] m 206°, and a γ -(*amphi*)-dioxime [E,Z 572-43-0] m 166°, and can be distinguished by the three different reactions which they undergo. The α -isomer produces an anhydride (diphenyloxazine), the β -isomer provides oxanilide, and the γ -isomer forms *N*'-benzoyl phenylurea. α -(*anti*)-Dioxime complexes with Ni²⁺ and Fe²⁺ [Singh et al. *Talanta* 26 425 1979, DOI: 10.1016/0039-9140(79)80107-1]. The **bis-semicarbazone** C₁₆H₁₆N₆O₂, m 243-244°, crystallises in leaflets from EtOH. [Beilstein 7 III 3812, 7 IV 2504.] **Benzil monohydrazone** [5433-88-7] C₁₄H₁₂N₂O, M 224.3, has m 151°(dec). Crystallise it from EtOH. The **monoacetyl hydrazone** has m 91° (from EtOH). The **bis-phenylhydrazone** has m 225°. [Metze & Meyer *Chem Ber* 90 481 1957, DOI: 10.1002/cber.19570900405; Beilstein 7 I 394.] **Benzil monoxime** [14090-77-8], [E 574-15-2], [Z 574-16-3] C₁₄H₁₁NO₂, M 105.1, has m 140°. The **trans-isomer** (α) crystallises from *C₆H₆ (must not use animal charcoal), or 30% EtOH and has m 140° (137-138°). The **cis-isomer** (β) also crystallises from *C₆H₆ but crystals have 0.5*C₆H₆ (m 62-63°), and the solvent free compound has m 112° (113-114°). **Note** that when pure α -isomer is boiled for 15 minutes with 1/10th of its weight of animal charcoal in the minimum volume of *C₆H₆ at its boiling point, filtered from charcoal and cooled, an excellent yield of the β -isomer + 0.5 *C₆H₆ is obtained. Pure β -isomer in aqueous alcoholic Cu(OAc)₂ should not produce a colour change, but if contaminated with the α -isomer a greenish colour is produced. [Beilstein 7 III 3812, 7 IV 2504.] The α -isomer complexes with Co²⁺, Pd²⁺, Ir³⁺, Rh³⁺, and Pt⁴⁺ [Singh et al. *Talanta* 26 425 1979, DOI: 10.1016/0039-9140(79)80107-1].

Benzilic acid (diphenylglycollic acid) [76-93-7] C₁₄H₁₂O₃, M 228.3, m 149-151°, 150°, pK¹⁸ 3.06. Crystallise benzilic acid from *benzene (*ca* 6ml/g), or hot H₂O. The **methyl ester** C₁₅H₁₄O₃, distils at 187°/13mm and the distillate solidifies with m 74-75°. [Ballard & Dehn *Org Synth* 1 29 1921, DOI: 10.15227/orgsyn.001.0029; Beilstein 10 IV 1256.]

Benzo[a]biphenylene [252-47-1] C₆H₁₀, M 202.2, m 72-73° (compare with β -isomer below). It forms yellow needles from MeOH and sublimes *in vacuo* (m 72.0-72.8°). The **2,4,7-trinitrofluorenone complex** crystallises as black needles m 201.5-202.5°. [Cava & Stucker *J Am Chem Soc* 77 6022 1955, DOI: 10.1021/ja01627a066; Barton et al. *J Chem Soc(C)* 1276 1967, DOI: 10.1039/J39670001276; Beilstein 5 IV 2462.] **Benzo[b]biphenylene** [259-56-3] C₆H₁₀, M 202.2, has 242.6-243.6°. It forms yellow crystals from *C₆H₆/cyclohexane m 234-245° (sublimation). Slow evaporation of a tetrahydrofuran solution gave colourless triclinic crystals suitable for X-ray diffraction studies [Ferrara et al *Acta Cryst* C45 57 1989, DOI: 10.1107/S0108270188008765]. The **2,4,7-trinitrofluorenone complex** crystallises as red needles from *C₆H₆/MeOH m 214-216°. It has been sublimed *in vacuo*. [Jensen & Coleman *Tetrahedron Lett* No 20 7 1959, DOI: 10.1016/S0040-4039(01)99477-8; Barton et al. *JCS Perkin Trans 1* 967 1986, DOI: 10.1039/P19860000967; Beilstein 5 IV 2462.]

Benzoic acid [65-85-0] C₇H₆O₂, M 122.1, m 122.4°, 122.6-123.1°, b 132.1°/10mm, 146.7°/20mm, 162.6°/40mm, 205.8°/200mm, 249.2°/atm, pK¹⁵ 4.218, pK²⁰ 4.208, pK³⁰ 4.205, pK⁴⁰ 4.219, pK⁵⁰ 4.24. For use as a volumetric standard, analytical reagent grade benzoic acid should be carefully fused to *ca* 130° (to dry it) in a platinum crucible, and then powdered in an agate mortar. Benzoic acid has been crystallised from boiling water (charcoal), aqueous acetic acid, glacial acetic acid, *C₆H₆, aqueous EtOH, petroleum ether (b 60-80°), and from EtOH solution by adding water. The solubility (w/v) in H₂O is 0.21% at 10°, 0.34% at 25°, and 6.8% at 95°. Also its solubility (w/v) in EtOH is 43.5% at ~25° and 67% at 78°; and at ambient temperatures in Et₂O and Me₂CO it is 33%, in CHCl₃ it is 22%, in *C₆H₆ it is 10%, in turpentine oil it is 4.3%, and in CCl₄ and CS₂ it is

3.3%. It is readily purified by fractional crystallisation from its melt and by sublimation in a vacuum at 80°. The *S*-benzylisothiuronium salt has **m** 167° (from EtOH/H₂O). [Beilstein 9 IV 273.] *Benzoic anhydride* [93-97-0] C₁₄H₁₀O₃, **M** 226.2, has **m** 42°, **b** 142.8°/1mm, 180°/5mm, 218°/20mm, 252.7°/60mm, 299.1°/200mm, 360°/atm. Free it from benzoic acid by washing with NaHCO₃, then water, and drying. Crystallise it from *benzene (0.5ml/g) by adding just enough petroleum ether (b 40-60°) to cause cloudiness, then cool in ice. It can be distilled without decomposition, but preferably distilled in a vacuum. [Clarke & Rahrs *Org Synth Coll Vol* 1 91 1941, DOI: 10.15227/orgsyn.003.0021; Beilstein 19 IV 550.]

(±)-Benzoin (2-hydroxy-2-phenylacetophenone) [119-53-9, 579-44-2] C₁₄H₁₂O₂, **M** 212.3, **m** 137°, **b** 194°/12mm, 344°/768mm. Crystallise benzoin from CCl₄, hot EtOH (8ml/g, prisms), or 50% acetic acid. Also crystallise it from high purity *benzene, then twice from high purity MeOH, to remove fluorescent impurities [Elliott & Radley *Anal Chem* 33 1623 1961, DOI: 10.1021/ac60179a002]. It is almost insoluble in H₂O (0.03 w/w%), much more so in pyridine (20 w/w%); and it can be sublimed. [Adams & Marvel *Org Synth Coll Vol* 1 94 1941, DOI: 10.15227/orgsyn.001.0033; Beilstein 8 IV 1279.] (±)-α-Benzoin oxime (*E*-oxime, Cuprone) [441-38-3; E 1143-89-1; 5828-63-2, 574-13-0] C₁₄H₁₁NO₂, **M** 227.3, has **m** 151°, 151-152°, 152°. The oxime crystallises in prisms from *C₆H₆, and is soluble in EtOH, Et₂O, CHCl₃ and Me₂CO but not in H₂O. It is used for the spectroscopic determination of Cu²⁺, Pd²⁺, Pt⁴⁺, Rh³⁺ and V⁵⁺. A 0.25% solution in CHCl₃ is used for the extraction of Mo and W (in concentrated HCl) and for precipitating V (at pH 2-4), as well as Cu, Pd, Bi, and Au [Singh et al. *Talanta* 26 425 1979, DOI: 10.1016/0039-9140(79)80107-1]. The β-isomer (*Z*-oxime) [7110-50-1] can be recrystallised from Et₂O to form prisms of the etherate, which lose Et₂O on standing in air, and the ether-free solid has **m** 99°. It also crystallises from EtOH in prisms with **m** 99°. It has antiseptic properties. [Beilstein 8 IV 1282.]

R-(-)-Benzoin [5928-66-5] C₁₄H₁₂O₂, **M** 212.2, **m** 131-132.5°, 132°, 135-137°, [α]_D¹⁹ -115 (c 1.5, Me₂CO), [α]_D¹² -118 (c 1.2, Me₂CO) and the *acetate* forms needles with [α]_D¹² -217.7 (CHCl₃). The *enantiomeric S*-(+)-benzoin [5928-67-6] C₁₄H₁₂O₂, **M** 212.2, **m** 132°, 135-137°, [α]_D²⁰ +118 (c 1.5, Me₂CO), [α]_D¹² +120.5 (c 1.2, Me₂CO), and the *E*-oxime has **m** 164° (from EtOH) and [α]_D²⁴ +3 (CHCl₃). These are obtained by crystallisation of the (+)-quinidine diastereoisomeric *benzoin hydrogenphthalate* salts, followed by addition of ice-cold dilute HCl (to remove quinidine), and hydrolysis of the acid phthalates with 1N H₂SO₄/EtOH (40ml/200ml) gives the pure optically active benzoin enantiomers as needles from EtOH with the stated melting points and rotations. [Optical Resolution: Kenyon & Patel *J Chem Soc* 435 1965, DOI: 10.1039/JR9650000435.] [Beilstein 8 H 167.]

Benzonitrile [100-47-0] C₇H₅N, **M** 103.1, **m** -12.9°, **b** 28.2°/1mm, 69.2°/10mm, 94°/40mm, 191.1°/atm, **d**₄²⁰ 1.010, **n**_D²⁰ 1.528. It has been purified by steam distillation, the distillate is extracted into Et₂O, washed with dilute NaHCO₃, dried overnight with CaCl₂, and the Et₂O distilled off. The residue is then dried with CaSO₄, MgSO₄ or K₂CO₃, and distilled from P₂O₅ in an all-glass apparatus, under reduced pressure (**b** 69°/10mm), collecting the middle fraction. Distillation from CaH₂ causes some decomposition of benzonitrile. Isonitriles can be removed by preliminary treatment with concentrated HCl until the odour of isonitrile (carbylamine) has gone, followed by preliminary drying with K₂CO₃. (This treatment also removes amines.) Steam distil (to remove small quantities of carbylamine). The distillate is extracted into ether, washed with dilute Na₂CO₃, dried overnight with CaCl₂, and the ether is removed by evaporation. The residue is distilled at 40mm (**b** 96°) [Kice et al. *J Am Chem Soc* 82 834 1960, DOI: 10.1021/ja01489a020].

Conductivity grade benzonitrile (specific conductance 2 x 10⁻⁸ mho) is obtained by treatment with anhydrous AlCl₃, followed by rapid distillation at 40-50° under vacuum. After washing with alkali and drying with CaCl₂, the distillate is redistilled in a vacuum several times at 35° before fractionally crystallising several times by partial freezing. It is dried over finely divided activated alumina from which it is withdrawn when required [Van Dyke & Harrison *J Am Chem Soc* 73 402 1951, DOI: 10.1021/ja01145a132]. [Beilstein 9 IV 892.]

Benzo[ghi]perylene (1,12-benzoperylene) [191-24-2] C₂₂H₁₂, **M** 276.3, **m** 273°, 277-278.5°, 278-280°, **b** >500°/atm. It forms light green crystals on recrystallisation from *C₆H₆ or xylene and sublimes at 320-340°/0.05mm [UV: Hopff & Schweizer *Helv Chim Acta* 42 2315 1959, DOI: 10.1002/hlca.19590420704; *Clar Chem Ber* 65 846 1932, DOI: 10.1002/cber.19320650534; Fluoresc. Spectrum: Bowen & Brocklehurst *J Chem Soc* 3875 1954, DOI: 10.1039/JR9540003875]. It also recrystallises from propan-1-ol [Altman & Ginsburg *J*

Chem Soc 466 1959, DOI: 10.1039/JR9590000458]. The **1,3,5-trinitrobenzene complex** has **m 310-313°** (deep red crystals from *C₆H₆), the **picrate** has **m 267-270°** (dark red crystals from *C₆H₆), and the **styphnate** (2,4,6-trinitroresorcinol complex) has **m 234°** (wine red crystals from *C₆H₆). [*Beilstein* 5 IV 2766.]

3,4-Benzophenanthrene (benzo[c]phenanthrene) C₁₈H₁₂, [195-19-7] **M 228.3, m 65.6-66.2°, 68°**. Crystallise benzo[c]phenanthrene from EtOH, petroleum ether (colourless needles or leaflets), or EtOH/Me₂CO (fine needles). The **picrate** crystallises in red needles with **m 125.8-126.2°** (126-127°) from *C₆H₆/light petroleum. Traces of picric acid can be removed from recovered hydrocarbon by passing a *C₆H₆ solution through an Al₂O₃ column. [Newman & Joshel *J Am Chem Soc* 60 485 1938, DOI: 10.1021/ja01269a069; *Beilstein* 5 III 2378, 5 IV 2552.]

Benzophenone (diphenylketone) [119-61-9] C₁₃H₁₀O, **M 182.2, m 48.5-49°, b 108.2°/1mm, 157.6°/10mm, 197.5°/40mm, 176.8°/400mm, 305.4°/10mm, d₄²⁰ 1.0869, n_D²⁵ 1.5975, pK²⁵ -6.0(-8.4) (aqueous H₂SO₄)**. Crystallise it from MeOH, EtOH, cyclohexane, *benzene or petroleum ether, then dry in a current of warm air and store it over BaO or P₂O₅. It is also purified by zone melting and by sublimation [Itoh *J Phys Chem* 89 3949 1985, DOI: 10.1021/j100265a002; Naguib et al. *J Am Chem Soc* 108 128 1986, DOI: 10.1021/ja00261a021; Gorman & Rodgers *J Am Chem Soc* 108 5074 1986, DOI: 10.1021/ja00277a005; Okamoto & Teranishi *J Am Chem Soc* 108 6378 1986, DOI: 10.1021/ja00280a040; Naguib et al. *J Phys Chem* 91 3033 1987, DOI: 10.1021/j100295a078]. Its solubility at ambient temperature is 13% in EtOH, 17% in Et₂O and very high in CHCl₃. [Marvel & Sperry *Org Synth Coll Vol I* 95 1941, DOI: 10.15227/orgsyn.008.0026; *Beilstein* 7 H 411, 7 III 2048, 7 IV 1357.] **Benzophenone oxime** [574-66-3] C₁₃H₁₁NO, **M 197.2, has m 140-144°, 142°, 143-144°, pK²⁵ 11.18**. Crystallise the oxime from MeOH (4ml/g). [*Beilstein* 7 II 355, 7 III 2063 1370.]

Benzophenone-3,3',4,4'-tetracarboxylic dianhydride (BTDA) [2421-28-5] C₁₇H₆O₇, **M 322.2, m 218-122°, 218.5-219.5°, 225.5°**. The main impurity is the free acid formed by hydrolysis (check for OH bands in the IR). This can be converted to the dianhydride by treating with Ac₂O (molar ratio of 4 to 1 of acid), heating at 110-120° for 1.5 to 2 hours, cooling to 0–5° and collecting the dianhydride. This is then dissolved in hot dioxane or Me₂CO, filtered and cooled to 10–15°. The moisture sensitive solid is collected and dried at 120–130° *in vacuo*. It sublimes at high vacuum. [Faberov et al. *J Org Chem USSR* 4 153 (English translation) 1968.]

Benzopinacol (1,1,2,2-tetraphenyl-1,2-ethanediol) [464-72-2] C₂₆H₂₂O₂, **M 366.5, m 170-180° (depends on heating rate), 171-173°, 188-190°**. Crystallise benzopinacol from EtOH, petroleum ether (b 90-100°), or *C₆H₆ (prisms with 1 mol of *C₆H₆). Its solubility (v/v) in 95% EtOH is 2.5% (at 78°), in *C₆H₆ it is 3.8% (at 80°), and in AcOH it is 8.7% (at 118°); but in Et₂O, CHCl₃ and CS₂ it is freely soluble. Since it decomposes at its melting point it is best to place the melting point tube in a bath at 150°, then to heat slowly whereby it melts at **193-195°**. [Bachmann *Org Synth Coll Vol 2* 71 1943, DOI: 10.15227/orgsyn.014.0008; *Beilstein* 6 IV 7053.]

Benzo[a]pyrene (3,4-benzpyrene, benzo[def]chrysene) [50-32-8] C₂₀H₁₂, **M 252.3, m 177.5-178°, 179.0-179.5°, b 310-312°/10mm, 495°/atm**. A solution of 250mg of benzo[a]pyrene in 100ml of *benzene is diluted with an equal volume of hexane, then passed through a column of alumina, Ca(OH)₂ and Celite (3:1:1). The adsorbed material is developed with a 2:3 *benzene/hexane mixture. (It showed as an intensely fluorescent zone.) The main zone is eluted with 3:1 acetone/EtOH, and is transferred into 1:1 *benzene-hexane by adding H₂O. The solution is washed, dried with Na₂SO₄, evaporated and crystallised from *benzene by the addition of MeOH [Lijinsky & Zechmeister *J Am Chem Soc* 75 5495 1953, DOI: 10.1021/ja01118a009]. Alternatively, it can be chromatographed on activated alumina, eluted with a cyclohexane/*benzene mixture containing up to 8% *benzene, and the solvent is evaporated under reduced pressure [Cahnmann *Anal Chem* 27 1235 1955, DOI: 10.1021/ac60104a009], and crystallised from EtOH [Nithipatikom & McGown *Anal Chem* 58 3145 1986, DOI: 10.1021/ac00127a050]. [*Beilstein* 5 III 2517, 5 IV 2687.] It intercalates with DNA. **CARCINOGENIC**. **Benzo[e]pyrene (1,2-benzpyrene)** [192-97-2] has **m 178-179°, 178-180°**. Purify it by passage through an Al₂O₃ column (Woelm, basic, activity I), elute with *C₆H₆ and recrystallise from 2 volumes of EtOH/*C₆H₆ (4:1). It forms colourless or light yellow prisms or needles. [Campbell *J Chem Soc* 3659 1954, DOI: 10.1039/JR9540003659; Buchta & Kröger *Justus Liebigs Ann Chem* 705 190 1967, DOI: 10.1002/jlac.19677050122.] The **1,3,5-trinitrobenzene complex** has **m 253-254°** (orange needles from EtOH),

the *picrate* prepared by mixing 20mg in 1ml of $^*\text{C}_6\text{H}_6$ with 20mg of picric acid in 2ml $^*\text{C}_6\text{H}_6$, collecting the deep red crystals, and recrystallising from $^*\text{C}_6\text{H}_6$ has **m 228-229°** [NMR: Cobb & Memory *J Chem Phys* **47** 2020 1967, DOI:org/10.1063/1.1712232]. [*Beilstein* **5** III 2520, **5** IV 2689.] **CARCINOGEN.**

***p*-Benzoquinone** [106-51-4] $\text{C}_6\text{H}_4\text{O}_2$, **M 108.1, m 113°, 113-115°, 115.7°, b 293°/atm.** Purify *p*-benzoquinone in one or more of the following ways: steam distillation followed by filtration and drying (e.g. in a desiccator over CaCl_2), recrystallisation from petroleum ether (b 80-100°), $^*\text{benzene}$ (with, then without, charcoal), water or 95% EtOH, sublimation under vacuum (e.g. from room temperature to liquid N_2) and recrystallisation from tetrachloroethylene. It slowly decomposes and should be stored, refrigerated, in an evacuated or sealed glass vessel in the dark. It should be resublimed before use. It is a radical inhibitor, solvates with H_2O in the liquid and vapour forms, and has been used as a dehydrogenating agent in the Wettstein-Oppenauer oxidation [Review: Mandell *J Am Chem Soc* **78** 3199 1956, DOI: 10.1021/ja01594a061; Wolfenden et al. *J Am Chem Soc* **109** 463 1987, DOI: 10.1021/ja00236a026; *Beilstein* **7** IV 2065.]

1-Benzosuberone (6,7,8,9-tetrahydrobenzocyclohepten-5-one) [826-73-3] $\text{C}_{11}\text{H}_{12}\text{O}$, **M 160.2, b 80-85°/0.5mm, 90-93°/1mm, 138-139°/12mm, 154°/15mm, 175-175°/40mm, d_4^{20} 1.086, n_D^{20} 1.5638.** Purify it by dissolving in toluene, washing with aqueous 5% NaOH, then brine, drying (MgSO_4), and distilling. The **2,4-dinitrophenylhydrazones** has **m 210.5°, 207-208°** (from $\text{CHCl}_3/\text{MeOH}$). The ***Z*-O-picryloxime** has **m 156-157°** (from $\text{Me}_2\text{CO}/\text{MeOH}$), the ***E*-O-picryloxime** has **m 107°**. The **oxime** has **m 106.5-107.5°**. [UV: Gilmore & Horton *J Am Chem Soc* **73** 1411 1951, DOI: 10.1021/ja01148a004; Hedden & Brown *J Am Chem Soc* **75** 3744 1953, DOI: 10.1021/ja01111a040; Huisgen et al. *Chem Ber* **90** 1844 1957, DOI: 10.1002/cber.19570900923; *Beilstein* **7** IV 1029.]

Benzoylacetone (1-phenyl-1,3-butanedione) [93-91-4] $\text{C}_{10}\text{H}_{10}\text{O}_2$, **M 162.2, m 58.5-59.0°, 61°, b 128-130°/10mm.** Crystallise benzoylacetone from Et_2O or MeOH and dry it under vacuum at 40°. It is readily distilled in a vacuum and the colourless distillate solidifies. [*Beilstein* **7** IV 2151.]

2-Benzoylbenzoic acid [85-52-9] $\text{C}_{14}\text{H}_{10}\text{O}_3$, **M 226.2, m 126-129°, 129.2, 130°, pK^{25} 3.54.** Recrystallise the acid from $^*\text{C}_6\text{H}_6$ or cyclohexane, but it is best recrystallised by dissolving in a small volume of hot toluene and then adding just enough petroleum ether to cause turbidity, and cool. Dry it in a low vacuum at 80°. It can be sublimed at 230-240°/0.3mm [Bray et al. *J Chem Soc* 265 1957, DOI: 10.1039/JR9570000265]. The ***S*-benzylisothiuronium salt** has **m 177-178°** (from EtOH). [Lewenz & Serijan *J Am Chem Soc* **75** 4087 1953, DOI: 10.1021/ja01112a514; *Beilstein* **10** H 747, **10** IV 2977.] **3-Benzoylbenzoic acid** [579-18-0] has **m 164-166°, 165°, $\text{pK}_{\text{Est}} \sim 3.5$.** Crystallise the acid from EtOH and sublime it at 160°/1mm. [*Beilstein* **10** H 752, **10** III 3304, **10** IV 2982.] **4-Benzoylbenzoic acid** [611-95-0] $\text{C}_{14}\text{H}_{10}\text{O}_3$, **M 226.2, has m 196.5-198°, 197-200°, $\text{pK}_{\text{Est}} \sim 3.7$.** Dissolve the acid in hot H_2O by adding enough aqueous KOH solution till distinctly alkaline, filter and then acidify with drops of concentrated HCl. Filter off, wash the solid with cold H_2O , dry it at 100°, and recrystallise it from EtOH. [Wertheim *J Am Chem Soc* **55** 2540 1933, DOI: 10.1021/ja01333a051; *Beilstein* **10** H 753, **10** IV 3305.]

Benzoyl chloride [98-88-4] $\text{C}_7\text{H}_5\text{ClO}$, **M 140.6, m -1.0°, b 56°/4mm, 100°/35mm, 196.8°/745mm, d_4^{20} 1.2120, n_D^{10} 1.5537.** A solution of benzoyl chloride (300ml) in $^*\text{C}_6\text{H}_6$ (200ml) is washed with two 100ml portions of cold 5% NaHCO_3 solution, separated, dried with CaCl_2 and distilled [Oakwood & Weisgerber *Org Synth Coll Vol* **3** 112 1955, DOI: 10.15227/orgsyn.024.0014]. Repeated fractional distillation at 4mm Hg through a glass helices-packed column (avoiding porous porcelain or silicon-carbide boiling chips, and hydrocarbon or silicon greases on the ground joints) gave benzoyl chloride that did not darken on addition of AlCl_3 . Further purification is achieved by adding 3 mole% each of AlCl_3 and toluene, standing overnight, and distilling off the benzoyl chloride at 1-2mm [Brown & Jensen *J Am Chem Soc* **80** 2291 1958, DOI: 10.1021/ja01542a065]. Refluxing for 2 hours with an equal weight of thionyl chloride before distillation has also been used. [*Beilstein* **9** IV 721.] **Strong IRRITANT. Use in a fume cupboard.**

Benzoyl disulfide (dibenzoyl disulfide) [644-32-6] $\text{C}_{14}\text{H}_{10}\text{O}_2\text{S}_2$, **M 274.3, m 131.2-132.3°.** About 300ml of solvent is blown off from a filtered solution of dibenzoyl disulfide (25g) in acetone (350ml). The remaining acetone is decanted from the solid which is recrystallised first from 300ml of 1:1 (v/v) EtOH/EtOAc, then from

300ml of EtOH, and finally from 240ml of 1:1 (v/v) EtOH/ethyl acetate. The yield is about 40% [Pryor & Pickering *J Am Chem Soc* **84** 2705 1962, DOI: 10.1021/ja00873a012]. [Beilstein **9** H 424, **9** II 289, **9** III 1977.] Handle in a fume cupboard because of **TOXICITY** and obnoxious odour.

Benzoylformic acid (phenylglyoxylic acid) [611-73-4] $C_8H_6O_3$, **M 150.14**, **m 62-65°**, **64.5-65.5°**, **67°**, **b 84°/0.1mm**, **163-167°/15mm**, **pK²⁵ 1.39 (1.79)**. If the sample is oily, then it may contain H₂O. In this case dry it in a vacuum desiccator over P₂O₅ or KOH until crisp. For further purification dissolve 5.5g in hot CCl₄ (750ml), add charcoal (2g, this is necessary otherwise the acid may separate as an oil), filter, cool in ice-water until crystallisation is complete. Filter the acid off, and the solvent on the crystals is removed by keeping the acid (4.5g) in a vacuum desiccator for 2 days. Slightly yellow crystals are obtained. It can be recrystallised also from *C₆H₆/petroleum ether, and can be distilled in a vacuum. The acid is estimated by titration with standard NaOH. The **phenylhydrazone** is recrystallised from EtOH, **m 163-164°**, and the **semicarbazone acid** has **m 259°(dec)** (from EtOH). The **methyl ester** distils at **137°/14mm**, **110-111°/2mm**, **n_D²⁰ 1.5850**. [Baer & Kates *J Am Chem Soc* **67** 1482 1945, DOI: 10.1021/ja01225a022; Schaefer & Corey *J Org Chem* **24** 1825 1959, DOI: 10.1021/jo01093a629; Beilstein **10** H 654, **10** IV 2737.]

Benzoyl isothiocyanate [532-55-8] C_8H_5NOS , **M 163.2**, **m 25.5-26°**, **b 72.5-73°/6mm**, **88-91°/20mm**, **94-96°/21mm**, **202.5-204°/724mm**, **250-255°/atm**, **d₄²⁰ 1.213**, **n_D²⁰ 1.6354**. Distil the isothiocyanate over a small amount of P₂O₅, whereby the distillate crystallises in prisms. It is readily hydrolysed by H₂O to give benzamide and benzoylurea, but with NH₃ it gives **benzoylurea m 210°** which can be recrystallised from EtOH. [Hill & Degnan *J Am Chem Soc* **62** 1595 1940, DOI: 10.1021/ja01863a073; Terss & McEwen *J Am Chem Soc* **76** 580 1954, DOI: 10.1021/ja01631a075; Frank & Smith *Org Synth Coll Vol* **3** 735 1955, DOI:10.15227/orgsyn.028.0089, Beilstein **9** IV 777.]

Benzoyl peroxide [94-36-0] $C_{14}H_{10}O_4$, **M 242.2**, **m 95°(dec)**, **103-106°(dec)**. Dissolve dibenzoyl peroxide in CHCl₃ at room temperature and precipitate it by adding an equal volume of MeOH or petroleum ether. Similarly, it is precipitated from acetone by adding two volumes of distilled water. It has also been crystallised from 50% MeOH and from diethyl ether. Dry it under vacuum at room temperature for 24 hours. Store it in a desiccator in the dark at 0°. When purifying in the absence of water it can be **EXPLOSIVE**, and operations should be done on a very small scale with adequate protection. Large amounts should be kept moist with water and stored in a refrigerator. It is a free radical initiator. [Kim et al. *J Org Chem* **52** 3691 1987 DOI: 10.1021/jo00392a038; Beilstein **9** IV 777.] It is also available commercially blended with dibutyl phthalate, tricresyl phosphate, dicyclohexyl phthalate or moistened with H₂O. Its activity can be assessed by analysis [Braun *Org Synth Coll Vol* **I** 431 1941, DOI: 10.15227/orgsyn.008.0030].

p-Benzoylphenol (4-hydroxybenzophenone) [1137-42-4] $C_{13}H_{10}O_2$, **M 198.2**, **m 132-135°**, **133.4-134.8°**, **pK²⁵ 7.95**. Dissolve p-benzoylphenol in hot EtOH (charcoal), filter and cool. Alternatively, crystallise it once from EtOH/H₂O and twice from *benzene [Grunwald *J Am Chem Soc* **73** 4934 1951, DOI: 10.1021/ja01154a138; Dryland & Sheppard *JCS Perkin Trans 1* 125 1986, DOI: 10.1039/P19860000125]. [Beilstein **8** IV 1263.]

N-Benzoyl-N-phenylhydroxylamine [304-88-1] $C_{13}H_{11}NO_2$, **M 213.2**, **m 118-120°**, **121-122°**. Recrystallise it from hot water, *benzene Et₂O/hexane or acetic acid. It complexes with metals. [Beilstein **15** III 8, **15** IV 7.]

N-Benzoyl-o-tolylhydroxylamine [1143-74-4] $C_{14}H_{13}NO_2$, **M 227.3**, **m 104°**. Recrystallise the hydroxylamine from aqueous EtOH. [Beilstein **15** III 8, **15** IV 7.]

Benzyl acetate [140-11-4] $C_9H_{10}O_2$, **M 150.2**, **m -51°**, **b 92-93°/10mm**, **134°/102mm**, **214.9°/760mm**, **d₄²⁰ 1.0562**, **n_D²⁵ 1.4994**. Purify the acetate by fractional distillation, preferably in a good vacuum. Values of n_D²⁵ of 1.5232-1.5242 are too high and should be nearer to 1.4994. [Merker & Scott *J Org Chem* **26** 5180 1961, DOI: 10.1021/jo01070a095; Beilstein **6** IV 2262.]

Benzyl acetoacetate [5396-89-4] $C_{11}H_{12}O_3$, **M 192.2**, **b 130°/2mm**, **156-157°/10mm**, **162-167°/15mm**, **275-277°/atm**, **d₄²⁰ 1.114**, **n_D²⁰ 1.514**. Fractionate the ester and collect fractions with the expected physical prop-

erties. Otherwise add *ca* 10% by weight of benzyl alcohol and heat in an oil bath (160-170°, open vessel) for 30 minutes during which time excess of benzyl alcohol will have distilled off, then fractionate. [Baker et al. *J Org Chem* **17** 77 1952, DOI: 10.1021/jo01135a009; *Beilstein* **6** IV 2480.]

4'-Benzylacetophenone [782-92-3] $C_{15}H_{14}O$, **M 210.3**, **m** 37°, 38°, 39°, **b** 197-198°/9mm, 209-210°/15mm. Distil it in a vacuum, then recrystallise it from EtOH (*ca* 1ml/g). The *oxime* has **m** 99.5° (from 60% aqueous EtOH). [*Beilstein* **7** H 449, **7** III 2176.]

Benzyl alcohol [100-51-6] C_7H_8O , **M 108.1**, **m** -15.3°, **b** 58°/1mm, 93°/10mm, 129°/60mm, 205.5°/atm, d_4^{20} **0.981**, n_D^{20} **1.54033**, **pK²⁵ 15.4**. It is usually purified by careful fractional distillation under reduced pressure in the absence of air. Benzaldehyde, if present, can be detected by UV absorption at 283nm. It has also been purified by shaking with aqueous KOH and extracting with peroxide-free diethyl ether. After washing with water, the extract is treated with saturated NaHS solution, filtered, washed, dried with CaO and distilled under reduced pressure [Mathews *J Am Chem Soc* **48** 562 1926, DOI: 10.1021/ja01414a002]. Peroxy compounds can be removed by shaking with a solution of Fe^{2+} followed by washing the alcohol layer with distilled water and fractionally distilling it. Its solubility at ambient temperature in H_2O is 4%w/v, and in 50% aqueous EtOH it is 67%w/v. [*Beilstein* **6** IV 2222.] § A polystyrene bound (at the *p*-position) resin which is cross-linked with divinylbenzene is available commercially.

Benzylamine [100-46-9] C_7H_9N , **M 107.2**, **m** 10°, **b** 98°/12mm, 178°/742mm, 185°/768mm, d_4^{20} **0.981**, n_D^{20} **1.5392**, **pK²⁵ 9.33**. Dry it with NaOH or KOH, then distil it under N_2 , through a column packed with glass helices, taking the middle fraction. Also distil it from zinc dust under reduced pressure. The *picrate* has **m** 196° (from EtOH), and the *p-toluenesulfonamide* has **m** 116° (from MeOH). [*Beilstein* **12** IV 2155.] **Benzylamine hydrochloride** [3287-99-8] $C_7H_9N \cdot HCl$, **M 143.6**, has **m** 248° (rapid heating), 253°. Crystallise the salt from water. [*Beilstein* **12** IV 2155.]

N-Benzylaniline (N-phenylbenzylamine) [103-32-2] $C_{13}H_{13}N$, **M 183.4**, **m** 36°, 37-38°, **b** 306-307°/atm, d_4^{20} **1.061**, **pK²⁵ 4.04**. Crystallise the amine from petroleum ether (b 60-80°) (*ca* 0.5ml/g). The *picrate* has **m** 113° (from Et_2O). [*Beilstein* **12** H 1023, **12** I 449, **12** II 548, **12** III 2215, **12** IV 2172.]

Benzyl bromide [100-39-0] C_7H_7Br , **M 171.0**, **m** -4°, -3.9°, **b** 85°/12mm, 127°/80mm, 192°/760mm, 198-199°/atm, d_4^{20} **1.438**, n_D^{25} **1.575**. Wash benzyl chloride with concentrated H_2SO_4 (CARE), water, 10% Na_2CO_3 or $NaHCO_3$ solution, and again with water. Dry it with $CaCl_2$, Na_2CO_3 or $MgSO_4$ and fractionally distil it in the dark, under reduced pressure. It has also been thoroughly degassed at 10^{-6} mm and redistilled in the dark. This gives material with λ_{max} at (MeCN): 226nm (ϵ 8200) [Kosower & Mohammed *J Am Chem Soc* **93** 2709 1971, DOI: 10.1021/ja00740a021]. [*Beilstein* **5** IV 829.] *Handle in a fume cupboard, extremely LACHRYMATORY.*

Benzyl bromoacetate [5437-45-6] $C_9H_9BrO_2$, **M 229.1**, **b** 96-98°/0.1mm, 146°/12mm, 166-170°/22mm, d_4^{20} **1.444**, n_D^{25} **1.5412**. Dilute the ester with Et_2O , wash it with 10% aqueous $NaHCO_3$, H_2O , dry ($MgSO_4$) and fractionate it using a Fenske (glass helices packing) column. [Bergmann & Szinai *J Chem Soc* 1521 1956, DOI: 10.1039/JR9560001521; *Beilstein* **6** IV 2265.] **LACHRYMATORY.**

N-Benzyl-tert-butylamine (N-tert-butylbenzylamine) [3378-72-1] $C_{11}H_{17}N$, **M 163.3**, **b** 91°/12mm, 109-110°/25mm, 218-220°/atm, d_4^{20} **0.899**, n_D^{25} **1.4942**, **pK²⁵ 10.19**. Dissolve the amine in Et_2O , dry it over KOH pellets, filter and fractionate it in a N_2 atmosphere to avoid reaction with CO_2 from the air. The *hydrochloride* has **m** 245-246°(dec) (from MeOH/ Me_2CO) and the *perchlorate* has **m** 200-201°. [Freifelder et al. *J Am Chem Soc* **80** 4320 1958, DOI: 10.1021/ja01549a051; *Beilstein* **12** IV 2166.]

Benzyl carbamate [621-84-1] $C_8H_9NO_2$, **M 151.2**, **m** 86°, 86-88°, 90-91°. If it smells of NH_3 , then dry it in a vacuum desiccator and recrystallise it from 2 volumes of toluene and dry it in a vacuum desiccator again. It forms glistening plates from toluene, and can be recrystallised from H_2O [Martell & Herbst *J Org Chem* **06** 878 1941, DOI: 10.1021/jo01206a013; Carter et al. *Org Synth Coll Vol* **3** 167 1955 DOI: 10.15227/orgsyn.023.0013]. [*Beilstein* **6** IV 2278.]

Benzyl chloride [100-44-7] C_7H_7Cl , M 126.6, m -48° to 43° , b $63^\circ/8mm$, $179^\circ/atm$, d_4^{20} 1.100, n_D^{20} 1.538. Dry it with $MgSO_4$ or $CaSO_4$, or reflux it with fresh Ca turnings, then fractionally distil it under reduced pressure, collecting the middle fraction and storing it over CaH_2 or P_2O_5 . It has also been purified by passage through a column of alumina. *Alternatively*, it is dried over $MgSO_4$ and distilled in a vacuum. The middle fraction is degassed by several freeze-thaw cycles and then fractionated in an 'isolated fractionating column' (which has been evacuated and sealed off at $\sim 10^{-6}$ mm) over a steam bath. The middle fraction is retained. The final samples are distilled in a vacuum from this sample and again retaining the middle fraction. The purity is $>99.9\%$ (no other peaks are visible by GLC, and the NMR spectrum is consistent with the structure. [Kosower & Mohammed *J Am Chem Soc* **93** 2709 1971, DOI: 10.1021/ja00740a021], *Beilstein* **5** IV 809.] **IRRITANT** and *strongly* **LACHRYMATORY**.

N-Benzyl-3-chloropropionamide (Beclamide) [501-68-8, 24752-66-7] $C_{10}H_{12}ClNO$, M 197.7, m 94° , 96° . Crystallise the amide from MeOH. [*Beilstein* **12** III 2257, **12** IV 2234.] It possesses anticonvulsant activity [Ahmadi et al. *J Pharm Pharmacol* **47** 876 1995, DOI: 10.1111/j.2042-7158.1995.tb05757.x], and used as an adjunct in the treatment of schizophrenia [Raptis et al. *Acta Psych Scand* **8** 162 1990; DOI: 10.1111/j.1600-0447.1990.tb06472.x].

Benzyl cinnamate [103-41-3] $C_{16}H_{14}O_2$, M 238.3, m $34-35^\circ$, 39° , b $154-157^\circ/0.5mm$, $195-200^\circ/5mm$, $228-230^\circ/22mm$, d^{15} 1.109. Recrystallise the ester to a constant melting point from 95% EtOH. It has the odour of balsam. *Alternatively*, dissolve it in Et_2O , wash it with 10% aqueous Na_2CO_3 , H_2O , dry (Na_2SO_4), evaporate and fractionate it under reduced pressure using a short Vigreux column. It decomposes when boiled at atmospheric pressure. [Eliel & Anderson *J Am Chem Soc* **74** 547 1952, DOI: 10.1021/ja01122a081; Bender & Zerner *J Am Chem Soc* **84** 2550 1962, DOI: 10.1021/ja00872a019; *Beilstein* **9** IV 2012.]

Benzyl cyanide [140-29-4] C_8H_7N , M 117.1, m -24° , b $60^\circ/1mm$, $100^\circ/8mm$, $233.5^\circ/760mm$, d_4^{20} 1.015, n_D^{20} 1.523. Any benzyl isocyanide impurity can be removed by shaking vigorously with an equal volume of 50% H_2SO_4 at 60° , washing with saturated aqueous $NaHCO_3$, then half-saturated NaCl solution, drying and fractionally distilling under reduced pressure. Distillation from CaH_2 causes some decomposition of this compound: it is better to use P_2O_5 . Other purification procedures include passage through a column of highly activated alumina, and distillation from Raney nickel. *Precautions should be taken because of possible formation of free TOXIC cyanide, use an efficient fume cupboard.* [Adams & Thai *Org Synth Coll Vol* **1** 107 1943, DOI: 10.15227/orgsyn.002.0009; *Beilstein* **9** IV 1663.]

N-Benzyl dimethylamine (BDMA, DMBA) [103-83-3] $C_9H_{13}N$, M 135.2, b $66-67^\circ/15m$, $83-84^\circ/30mm$, $98-99^\circ/24mm$, $181^\circ/760mm$, d_4^{20} 0.898, n_D^{20} 1.516, pK^{25} 8.91. Dry the amine over KOH pellets and fractionate it over Zn dust in a CO_2 -free atmosphere. It has a pK^{25} of 8.25 in 45% aqueous EtOH. Store it under N_2 or in a vacuum. The *picrate* has m $94-95^\circ$, and the *picrolonate* has m 151° (from EtOH). [Skita & Keil *Chem Ber* **63** 34 1930, DOI: 10.1002/cber.19300630104; Coleman *J Am Chem Soc* **55** 3001 1933, DOI: 10.1021/ja01334a067; Devereux et al. *J Chem Soc* 2845 1957, DOI: 10.1039/JR9570002845.] The *tetraphenyl borate salt* has m $182-185^\circ$. [Crane *Anal Chem* **28** 1794 1956, DOI: 10.1021/ac60119a052; *Beilstein* **12** IV 2161.]

Benzyl dimethyloctadecylammonium chloride [122-19-0] $C_{27}H_{50}ClN$, M 424.2, m $150-158^\circ$ (sinters at 120°). Crystallise the salt from acetone, $EtOAc$ or $EtOAc/Et_2O$. [Sumiki et al. *J Agric Chem Soc Jpn* **26** 325 1952, *Chem Abstr* **47** 3505 1953, *Beilstein* **12** III 2212, **12** IV 2168.]

Benzyl ether (dibenzyl ether) [103-50-4] $C_{14}H_{14}O$, M 198.3, m $1.3-3.5^\circ$, b $158-160^\circ/0.1mm$, $173-174^\circ/21mm$, $298^\circ/atm$ with some dec, d_4^{20} 1.043, n_D^{20} 1.54057. Reflux the ether over sodium, then distil it under reduced pressure. It been purified by fractional freezing. [*Beilstein* **6** IV 2240.]

N-Benzyl-N-ethylaniline [92-59-1] $C_{15}H_{17}N$, M 221.3, b $170-180^\circ/14mm$, $212-222^\circ/54mm$, $285-286^\circ/710mm$, $312-313^\circ/atm$ (dec), d_4^{20} 1.029, n_D^{20} 1.595, $pK_{Est} \sim 4.6$. Dry the amine over KOH pellets and fractionate it. The *picrate* crystallises from $*C_6H_6$ as lemon yellow crystals m $126-128^\circ$ (softening at 120°). [Forrest et al. *J Chem Soc* 303 1951, DOI: 10.1039/JR9510000303; IR: Hill & Meakins *J Chem Soc* 760 1958, DOI: 10.1039/JR9580000760; *Beilstein* **12** H 1026, **12** IV 2176.]

Benzyl ethyl ether [539-30-0] $C_9H_{12}O$, M 136.2, b 65°/10mm, 186°, d_4^{20} 0.949, n_D^{20} 1.4955. Dry the ether with $CaCl_2$ or NaOH, then fractionally distil it. It is insoluble in H_2O , but steam distils with it. [Letsinger & Pollart *J Am Chem Soc* **78** 6079 1956, DOI: 10.1021/ja01604a036; *Beilstein* **6** III 1454, **6** IV 2229.]

Benzyl ethyl ketone (1-phenylbutan-2-one) [1007-32-5] $C_{10}H_{12}O$, M 148.2, b 49-49.5°/0.01mm, 66-69°/1mm, 83-85°/5mm, 101-102°/10mm, 229-233°/atm, d_4^{20} 0.989, n_D^{25} 1.5015. Purify the ketone by fractionation using an efficient column. It can be converted into the *oxime* which is distilled, b 117-118°/2mm, 145-146°/15mm, d_{25}^{25} 1.036, n_D^{25} 1.5363; decompose the oxime, and the ketone is redistilled. It can also be purified via the *semicarbazone* which has m 154-155°. [Myers et al. *J Am Chem Soc* **77** 5655 1955, DOI: 10.1021/ja01626a056; Hass et al. *J Org Chem* **15** 8 1950, DOI: 10.1021/jo01147a002; *Beilstein* **7** IV 712.]

N-Benzylhydroxylamine hydrochloride [29601-98-7] $C_7H_{10}ClNO$, M 159.6, m 108-110°(sublimes), pK_{Est} ~4.7. Crystallise it by dissolving it in EtOH containing a few drops of EtOH/HCl and adding dry Et_2O . The *free base* [622-30-0] C_7H_9NO , M 123.1, m ~105°, b 253.9°/760mm, can be obtained from the hydrochloride by treatment with ethanolic NH_3 , filter the NH_4Cl off, and evaporate the solvent. Alternatively, dissolve the salt in EtOH or MeOH, shake with a strong anion exchange resin, filter off the resin and evaporate. Strong irritant of the skin, eyes and respiratory system. All necessary precautions should be taken. [*Beilstein* **15** H 18, **15** IV 21.]

O-Benzylhydroxylamine hydrochloride [2687-43-6] $C_7H_{10}ClNO$, M 159.6, m 230-235°, 234-238°(sublimes), pK_{Est} ~5.9. Recrystallise the hydrochloride from H_2O or EtOH. [Hearn & Ward *Aust J Chem* **22** 161 1969, DOI: 10.1071/CH9690161]. It has antitubercular activity [Truitt et al. *J Am Chem Soc* **74** 3956 1952, DOI: 10.1021/ja01135a523]. *Beilstein* **6** III 1552, **6** IV 2562.]

N-Benzylideneaniline [538-51-2] $C_{13}H_{11}N$, M 181.2, m 48° (54°), 56°, b 310°/760mm. It is steam volatile and crystallises from *benzene or 85% EtOH. The *picrate* has m 159°. [Bigelow & Eatough *Org Synth Coll Vol* **1** 80 141, DOI: 10.15227/orgsyn.008.0022; *Beilstein* **12** H 195, **12** I 169, **12** II 113, **12** III 319, **12** IV 311.]

Benzylidene malononitrile [2700-22-3] $C_{10}H_6N$, M 155.2, m 83-84°, 87°. Recrystallise the nitrile from EtOH [Bernasconi et al. *J Am Chem Soc* **107** 3612 1985, DOI: 10.1021/ja00298a035]. It has λ_{max} at 307nm (EtOH). [*Beilstein* **9** H 895, **9** II 640, **9** III 4380, **9** IV 3462.]

Benzyl isocyanate [3173-56-6] C_8H_7NO , M 133.2, b 82-84°/10mm, 87°/14mm, 95°/17mm, 101-104°/33mm, d_4^{20} 1.08, n_D^{20} 1.524. Purify the isocyanate by fractionation through a two-plate column. It is a viscous liquid and is **TOXIC**. [Haworth et al. *J Chem Soc* 182 1947, DOI: 10.1039/JR9470000182; Ferstandig & Scherrer *J Am Chem Soc* **81** 4838 1959, DOI: 10.1021/ja01527a022; IR: Derkosch et al. *Monatsh Chem* **88** 35 1957, DOI: 10.1007/BF01075427; *Beilstein* **12** IV 2276.]

Benzyl isothiocyanate (Benzyl mustard oil) [622-78-6] C_8H_7NS , M 149.2, b 123-124°/1mm, 138-140°/20mm, 255-260°/atm, d_4^{20} 1.1234, n_D^{20} 1.6039. Dissolve benzyl isothiocyanate in Et_2O , filter, if there is any solid, and distil it through an efficient column at 11mm with a bath temperature at ca 150°. Characterise it by reacting (0.5ml) in EtOH (1ml) with 50% $NH_2NH_2 \cdot H_2O$ (2 ml) to give *4-benzylthiosemicarbazide* as colourless needles which are recrystallised from EtOH, m 130°. **TOXIC** [Hoggarth & Young *J Chem Soc* 1582 1950, DOI: 10.1039/JR9500001582; Schmidt et al. *Justus Liebigs Ann Chem* **612** 11 1958, DOI: 10.1002/jlac.19586120103; IR and UV: Svátek et al. *Acta Chem Scand* **13** 442 1959, DOI: 10.3891/acta.chem.scand.13-0442; *Beilstein* **12** IV 2276.]

Benzylmalonic acid (β -phenylisusuccinic acid) [616-75-1] $C_{10}H_{10}O_4$, M 194.2, m 119-120°, 121°, pK_1^{25} 2.91, pK_2^{25} 5.87. Crystallise the acid from * C_6H_6 or * C_6H_6 /petroleum ether. [Drummond & Waters *J Chem Soc* 2456 1954, DOI: 10.1039/JR9540002456]. The *bis-S-benzylisothiuronium salt* has m 161-162° (from EtOH) [Friedger & Pedersen *Acta Chem Scand* **9** 1425 1955, DOI:10.3891/acta.chem.scand.09-1425]. [*Beilstein* **9** III 4283, **9** IV 3357.] **Diethyl benzylmalonate** [607-81-8] $C_{14}H_{18}O_4$, M 250.3, b 140-140.5°/1.5mm, 150-152°/4mm, 162-163°/10mm, d_{25}^{25} 1.064, n_D^{20} 1.486, was prepared from the acid with EtOH/* C_6H_6 / H_2SO_4 /24hrs then workup [Gardner & Rydon *J Chem Soc* 42 1938, DOI: 10.1039/JR9380000042].

(-)-*N*-Benzyl-*N*-methylephedrinium bromide [benzyl(2-hydroxy-1-methyl-2-phenethyl) dimethylammonium bromide] [58648-09-2] $C_{18}H_{24}BrNO$, *M* 350.3, *m* 209-211°, 212-214°, $[\alpha]_D^{25}$ -3.8° (c 1.45, MeOH), $[\alpha]_D^{20}$ -5.3° (c 1.45, MeOH). Recrystallise the bromide from MeOH/Et₂O. [Horner & Brich *Justus Liebigs Ann Chem* 710 1978, DOI: 10.1002/jlac.197819780504.] The *chloride* is recrystallised from EtOAc/*n*-hexane, *m* 198-199° $[\alpha]_D^{25}$ -8.67° (c 1.45, MeOH). [Julia et al. *JCS Perkin Trans 1* 574 1981, DOI: 10.1039/P19810000574; *Beilstein* 13 IV 1890.]

Benzyl 4-nitrophenyl carbonate [13795-24-9] $C_{14}H_{11}NO_5$, *M* 273.2, *m* 78-80°. Dissolve the carbonate in Et₂O, wash with H₂O (3x) and saturated aqueous NaCl, dry (MgSO₄), evaporate this in a vacuum and recrystallise the residue from a small volume of MeOH, *m* 78-79°. Alternatively, dissolve it in Et₂O, wash it with N HCl (2x), 0.5N NaHCO₃ (4x) then H₂O, dry (Na₂SO₄), evaporate the Et₂O and recrystallise the residue from *C₆H₆/petroleum ether, *m* 79-80°. The 2-nitro-isomer has *m* 27-28°, *b* 151°/11mm. [Khosla et al. *Indian J Chem* 5 279 1967, Wolman et al. *J Chem Soc (C)* 689 1967, DOI: 10.1039/J39670000689; *Beilstein* 6 IV 2277.]

Benzyloxyacetyl chloride [19810-31-2] $C_9H_9ClO_2$, *M* 184.6, *b* 81°/0.2mm, 84-87°/0.4mm, 105-107°/5mm, *d*₄²⁰ 1.19, *n*_D²⁰ 1.523. Check the IR to see if there are OH bands. If so, then it may be contaminated with free acid formed by hydrolysis. Add oxalyl chloride (amount depends on contamination and needs to be judged, *ca* 3mols), heat at 50° in the absence of moisture for 1 hour and fractionate twice, *b* 81°/0.2mm (with bath temperature at 81°). Excessive heating results in decomposition to give benzyl chloride. The *anilide* is formed by adding aniline in CHCl₃ solution and has *m* 49°. It has been used for the synthesis of various β-lactams [Jayaraman et al. *J Org Chem* 59 932 1994, DOI: 10.1021/jo00083a042; van Brabandt et al. *J Org Chem* 71 7083 2006, DOI: 10.1021/jo0608319]. [Fischer & Gohlke *Helv Chim Acta* 16 1130 1933, DOI: 10.1002/hlca.193301601140; *Beilstein* 6 IV 2470.]

2-Benzyloxybenzoic acid (salicylic acid benzyl ether) [14389-86-7] $C_{14}H_{12}O_3$, *M* 228.2, *m* 73-77°, *pK*_{Est} ~4.1. It has been prepared from salicylic acid (in 95% EtOH) and benzyl chloride in the presence 5 N aqueous NaOH and refluxed for 4 hours, followed by concentration and acidification. The substance may contain salicylic acid as impurity, so the dried product should be heated at 120° under reduced pressure (2mm) to remove any salicylic acid and then recrystallised from 70% EtOH (colourless plates, *m* 70°). [Cavallito & Buck *J Am Chem Soc* 65 2140 1943, DOI: 10.1021/ja01251a034; *Beilstein* 10 III 247, 10 IV 316.] **3-Benzyloxybenzoic acid** [69026-14-8] has *m* 133-137°, 135.5-136°, *pK*_{Est} ~4.1. Recrystallise this acid from acetic acid (colourless plates, *m* 137-138°, 134° also reported). [Kipping & Wren *J Chem Soc* 3246 1957, DOI: 10.1039/JR9570003246; Jones *J Chem Soc* 430 1943, DOI: 10.1039/JR9430000430, *Beilstein* 10 II 41, 10 III 100.] **4-Benzyloxybenzoic acid** [1486-51-7] has *m* 188°, 189-192°, 193.5-194.5°, *pK*_{Est} ~4.1. Purify it by recrystallising from aqueous EtOH (charcoal, colourless plates), and by vacuum sublimation at 120°. It crystallises also from 95% EtOH. The *benzyl ester* crystallises as colourless needles, *m* 115°, from 95% EtOH [Kipping & Wren *J Chem Soc* 3246 1957, DOI: 10.1039/JR9570003246; Cavallito & Buck *J Am Chem Soc* 65 2140 1943, DOI: 10.1021/ja01251a034; *Beilstein* 10 II 93, 10 III 290, 10 IV 351.]

Benzyloxybutan-2-one [6278-91-7] $C_{11}H_{14}O_2$, *M* 178.2, *b* 90-92°/0.1mm, 88-91°/0.5mm, 121-126°/5mm, *d*₄²⁰ 1.0275, *n*_D²⁰ 1.5040. Dissolve the ketone in CHCl₃, wash with H₂O, aqueous saturated NaHCO₃, H₂O, dry (MgSO₄), evaporate the CHCl₃, and fractionate it. [Hoffman et al. *J Am Chem Soc* 79 2316 1957, DOI: 10.1021/ja01566a080; *Beilstein* 6 IV 2255.]

Benzyloxycarbonyl chloride (Cbz-Cl, BOC-Cl, benzyl chloroformate) [501-53-1] $C_8H_7ClO_2$, *M* 170.6, *b* 85-87°/7mm, 103°/20mm, 173°/724mm, *d*₄²⁰ 1.195, *n*_D²⁰ 1.5190. The commercial material is usually better than 95% pure and may contain some toluene, benzyl alcohol, benzyl chloride and HCl. After long storage, e.g. two years at 4°, Greenstein and Winitz [*The Chemistry of the Amino Acids* Vol 2 p. 890, J Wiley and Sons NY, 1961] recommended that the liquid should be flushed with a stream of dry air, filtered and stored over sodium sulfate to remove CO₂ and HCl which are formed by decomposition. It may further be distilled from an oil bath at a temperature below 85° because Thiele and Dent [*Justus Liebigs Ann Chem* 302 245 1898, DOI: 10.1002/jlac.18983020302] stated that benzyloxycarbonyl chloride decarboxylates to benzyl chloride slowly at 100° and vigorously at 155°. Redistillation at higher vacuum below 85° yields material which shows

no other peaks than those of benzyloxycarbonyl chloride by NMR spectroscopy. [Carter et al. *Org Synth Coll Vol 3* 167 1955, DOI: 10.15227/orgsyn.023.0013; *Beilstein 6* IV 2278.] **LACHRYMATORY** and **TOXIC**.

p-(Benzyloxy)phenol (monobenzene) [103-16-2] $C_{13}H_{12}O_2$, **M 200.2**, **m 122.5°**, **pK_{Est} ~10.1**. Crystallise it from EtOH or H₂O, and dry (P₂O₅) under vacuum. It causes loss of pigment. [Walba et al. *J Am Chem Soc* **108** 5210 1986, DOI: 10.1021/ja00277a027; *Beilstein 6* IV 5778.]

S-(-)-3-Benzyloxypropan-1,2-diol (1-benzyl-sn-glycerol) [17325-85-8] $C_{10}H_{14}O_3$, **M 182.2**, **m 24-26°**, **b 117-118°/10⁻⁴mm**, **115-116°/0.02mm**, **121-123°/0.2mm**, **125°/0.5mm**, **261°/atm**, **d₄²⁰ 1.1437**, **n_D²² 1.5295**, **[α]_D²⁵ -5.9 (neat)**. Purify the S-(-)-diol (obtained by starting from D-mannitol) by fractional distillation. **R-(+)-3-Benzyloxypropan-1,2-diol (3-benzyl-sn-glycerol)** [56552-80-8] was shown to have **m 25-29°** and **[α]_D²⁰ +5.5 (c 20, CHCl₃)**. The *racemate* has [4799-67-1]. [Baer et al. *J Biol Chem* **230** 447 1958, Gigg & Gigg *J Chem Soc C* 1865 1967, DOI: 10.1039/J39670001865; *Beilstein 6* IV 2247.]

2-Benzylphenol (α-phenyl-o-cresol) [28994-41-4] $C_{13}H_{12}O$, **M 184.2**, **m 20.2-20.9°**, **52°**, **54°**, **b 121-123°/1mm**, **175°/18mm**, **312°/760mm**, **n_D²⁰ 1.5994-1.5995**, **pK_{Est} ~10.0**. Distil 2-benzylphenol in a vacuum and the distillate crystallises on cooling to a solid with **m ~21°** (22-23° in a sealed capillary tube). This forms a **phenylcarbamate m 117.5-118°** (needles from ligroin), and with MeI/Me₂CO/K₂CO₃ (in a tube at 100°) provides the corresponding **2-benzylphenyl methyl ether b 159-160°/12mm**. It has a stable form **B** with **m ~52°** and an unstable form **A** with **m ~21°**. The second form is obtained by recrystallisation from EtOH, or by warming the lower melting form (an exothermic change). On cooling molten crystals of **B** provides crystals of the **A** form. The IR shows strong phenolic -OH absorption at ν_{max} at 3390cm⁻¹ (2.95μ). [Kremers et al. *Justus Liebigs Ann Chem* **442** 210 1925, DOI: 10.1002/jlac.19254420114; Kornblum & Lurie *J Am Chem Soc* **81** 2705 1959, DOI: 10.1021/ja01520a030; *Beilstein 6* H 675, **6** IV 4628.] **4-Benzylphenol (α-phenyl-p-cresol)** [101-53-1, 7563-63-5] $C_{13}H_{12}O$, **M 184.2**, has **m 79-81°**, **84°**, **84.5-85.5°**, **b 154-157°/4mm**, **198-200°/10mm**, **322°/atm**, **pK_{Est} ~10.2**. Crystallise 4-benzylphenol from water. [Ziegenbein et al. *Chem Ber* **88** 1906 1955, DOI: 10.1002/cber.19550881213; McKinney & Reynolds *Talanta* **1** 46 1958, DOI:10.1016/0039-9140(58)80007-7; *Beilstein 6* H 675, **6** I 324, **6** II 629, **6** III 3357, **6** IV 4640.]

4-N-Benzylsulfanilamide (Septazen) [1709-54-2] $C_{13}H_{14}N_2O_2S$, **M 262.3**, **m 175°**, **178°**. Crystallise Septazen from dioxane/H₂O, EtOH/H₂O or Me₂CO (**m 174.5-175.8°**). Its solubility in H₂O at 37° is 0.03-0.43mg/100ml. [*Beilstein 14* III 2026.]

Benzylthiocyanate (benzyl rhodanide) [3012-37-1] C_8H_7NS , **M 149.2**, **m 39-41°**, **43°**, **42-44°**, **b 230-235°/atm**, **256°(dec)**. Crystallise the thiocyanate from EtOH or aqueous EtOH. [*Beilstein 6* H 460, **6** I 228, **6** II 434, **6** III 1600, **6** IV 2680.]

Benzyl toluene-p-sulfonate [1024-41-5] $C_{14}H_{14}O_3S$, **M 262.3**, **m 58°**, **58.5-58.8°**. Crystallise the ester from petroleum ether (b 40-60°), CHCl₃/hexane or Et₂O/*C₆H₆. Dry it *in vacuo* but **not** in a desiccator over CaCl₂ as it causes hydrolysis of the ester. [Emmons & Ferris *J Am Chem Soc* **75** 2257 1953, DOI: 10.1021/ja01105a509; *Beilstein 11* II 48, **11** III 207, **11** IV 273.]

Benzyltributylammonium bromide [25316-59-0] $C_{19}H_{33}BrN$, **M 356.4**, **m 169-171°**, **174-175°**. Recrystallise the bromide from EtOAc/EtOH and EtOH/Et₂O. [Kantor & Hauser *J Am Chem Soc* **73** 4122 1951, DOI: 10.1021/ja01153a022; Petersen et al. *J Am Chem Soc* **81** 3264 1959, DOI: 10.1021/ja01522a027; *Beilstein 12* IV 2166.]

Benzyl 2,2,2-trichloroacetimidate [81927-55-1] $C_9H_8Cl_3NO$, **M 252.5**, **m 3-4°**, **b 106-114°/0.5mm**, **d₄²⁰ 1.349**, **n_D²⁰ 1.545**. Purify the imidate by distillation to remove up to 1% of PhCH₂OH as stabiliser. A solution in hexane can be stored for up to 2 months without decomposition. It is *hygroscopic* and has to be stored dry. Small amounts can be purified by preparative TLC on 2mm Merck plates pre-coated with silica gel 60 F-245, eluted with hexane and the spots detected by UV light (or spraying a side run with 5% H₂SO₄ in EtOH and heating). This procedure separates the *benzimidate* from any trichloroacetamide or N-benzyl-trichloroacetamide impurities; the latter could be formed on heating such as distillation *via* a rearrangement. Its IR has ν_{max} at 3380

(NH) and $1670(\text{C}=\text{N})\text{cm}^{-1}$ among other peaks. [For Synthesis and NMR see Wessel et al. *JCS Perkin Trans 1* 2247 1985, DOI: 10.1039/P19850002247, and Clizbe & Overman *Org Synth* **58** 4 1978, DOI: 10.15227/orgsyn.058.0004; Cramer et al. *Chem Ber* **91** 1049 1958, DOI: 10.1002/cber.19580910530; *Beilstein* **6** IV 2265.]

Benzyltrimethylammonium chloride [56-93-9] $\text{C}_{10}\text{H}_{16}\text{ClN}$, **M 185.7, m 238-239°(dec)**. A 60% aqueous solution of the salt is evaporated to dryness under a vacuum on a steam bath, and then left in a vacuum desiccator containing a suitable drying agent. The solid residue is dissolved in a small volume of boiling absolute EtOH and precipitated by adding an equal volume of diethyl ether with cooling. After washing, the precipitate is dried under a vacuum [Karusch *J Am Chem Soc* **73** 1246 1951, DOI: 10.1021/ja01147a110]. [*Beilstein* **12** IV 2162.] It is available commercially as a 60% solution in H_2O with d_{20}^{20} 1.072g/ml and n_D^{20} 1.470. **Benzyltrimethylammonium fluoride monohydrate and x-hydrate (BTAF)** [H_2O 329-97-5; $x\text{H}_2\text{O}$ 127582-36-9] $\text{C}_{10}\text{H}_{16}\text{FN}\cdot\text{H}_2\text{O}$, **M 169.2 (anhydr)**, has **m 181-189°**. It is a good source of *naked fluoride ions* for catalysis in methylation reaction of Me_3SiO -cycloalkanes. BTAF is most reactive when it is as free from H_2O as possible. It has been prepared by adding ~4.7% aqueous HF (ca 8.6ml) to 40% methanolic Triton B (10ml, benzyltrimethylammonium hydroxide, see below) until the pH of the solution reaches 8–7. The solvent is removed in a vacuum of ~1mm and the residue is dried at $50^\circ/0.5\text{mm}$ for 20 hours (oil pump with liquid N_2 trap). The highly hygroscopic BTAF (~3.5g) is pulverised and stored over P_2O_5 . It decomposes above 100° , but drying at $90^\circ/0.1\text{mm}/48\text{hr}$ or $30^\circ/0.1\text{mm}$ to constant weight (2 days) produces a reasonably useful catalyst which has been used together with 4A molecular sieves in dry THF, dioxane or MeCN [Kuwajima et al. *J Am Chem Soc* **104** 1025 1982, DOI: 10.1021/ja00368a018]. Aldol condensations with trimethylsilylcyclopropanes, using BTAF or dry tetra-*n*-butylammonium fluoride (TBAF) were similarly achieved [Paquette et al. *J Am Chem Soc* **106** 6442 1984, DOI: 10.1021/ja00333a068].

Benzyltrimethylammonium hydroxide (Triton B) [100-85-6] $\text{C}_{10}\text{H}_{16}\text{N}\cdot\text{OH}$, **M 167.3, d 0.91**. A 38% solution (as supplied) is decolorised (charcoal), then evaporated under reduced pressure to a syrup, with final drying at $75^\circ/1\text{mm}$ pressure. The *anhydrous* base is obtained by prolonged drying over P_2O_5 in a vacuum desiccator. [*Beilstein* **12** IV 2162.] It is commercially available as a 40wt% solution in H_2O with d_{25}^{25} 1.059g/ml, n_D^{20} 1.43; and as a 40w% solution in MeOH with d_{25}^{25} 0.92g/ml.

Bibenzyl (1,2-diphenylethane, dibenzyl) [103-29-7] $\text{C}_{14}\text{H}_{14}$, **M 182.3, m 50-53°, 52.0-52.5°, 52.5-53.5°, b 284°/atm, d_4^{20} 0.9782**. Crystallise bibenzyl from hexane, MeOH, or 95% EtOH. It has also been sublimed under vacuum, and further purified by percolation through columns of silica gel and activated alumina. It is prepared by reduction of benzoin or benzyl [Clemmensen *Chem Ber* **47** 688 1914, DOI: 10.1002/cber.191404701107], but is best obtained by catalytic reduction of stilbene. Thus a mixture of stilbene (5g, see [103-30-0]) in dioxane (200ml) and Raney Ni (22g) is refluxed with stirring for 24 hours. The catalyst is filtered off, the filtrate is evaporated *in vacuo*, and the residue is recrystallised from MeOH to give an 80% yield of bibenzyl m 52.0-52.5°. Its FT-IR (melt) has ν_{max} at 3062.6, 1601.7, 1494.9, 1030.2, 906.0, 752.1, 696.9, 580.2 and 519.3 cm^{-1} ; its ^1H NMR (300MHz, CDCl_3 , TMS) has δ at 7.27 (t, 4H, arom-H), 7.18 (t, 6H, arom-H) and 2.90 (s, 4H, benzylic-H); and its ^{13}C NMR (15MHz, CDCl_3 , CDCl_3 as internal standard with δ at ~77.0) has δ at 141.7, 128.36, 128.24, 125.83 and 37.89. [Kleiderer & Kornfeld *J Org Chem* **13** 455 1948, DOI: 10.1021/jo01161a022.] [*Beilstein* **5** IV 1868.]

(±)-1,1'-Bi-(2-naphthol) [1,1'-di-(2-naphthol), BINOL] [602-09-5, 41024-90-2] $\text{C}_{20}\text{H}_{14}\text{O}_2$, **M 286.3, m 215-217°, 218°, $\text{pK}_{\text{Est}(1)} \sim 7.1$, $\text{pK}_{\text{Est}(2)} \sim 11.2$** . Crystallise the binaphthol from toluene or *benzene (10ml/g). When crystallised from chlorobenzene it has **m 238°**. Its solubility in dioxane is 5%. [*Beilstein* **6** IV 7020.] **1,1'-Bi-(2-naphthol) [1,1'-di-(2-naphthol)]** [*R*-(+)- 18531-94-7], [*S*-(-)- 18531-99-2] have **m 207.5-208.5°, 209-211°, $[\alpha]_D^{20}$ (+) and (-) 37.4.0 (c 0.5, THF), $[\alpha]_{546}^{25}$ (+) and (-) 51 (c 0.1, THF), pK as above**. Dissolve it in cold 2.5N NaOH, extract with CH_2Cl_2 , and acidify with 5% HCl. Collect the white precipitate and recrystallise it from aqueous EtOH and dry it in a vacuum [Akimoto & Yamada *Tetrahedron* **27** 5999 1971, DOI: 10.1016/S0040-4020(01)91765-X]. It is optically stable in dioxane-water (100°/24 hours). *Racemisation*: 72% in 1.2N HCl at $100^\circ/24\text{ hours}$, and 68% in 0.67M KOH in BuOH at $118^\circ/23\text{ hours}$ [Kyba et al. *J Am Chem Soc* **95** 2692 1973, DOI: 10.1021/ja00789a051]. It was also crystallised from * C_6H_6 (solubility is 1%) using Norite, or EtOH/ H_2O after chromatography through silica gel, eluting with $\text{Me}_2\text{CO}/^*\text{C}_6\text{H}_6$. [Kyba et al. *J Org*

Chem **42** 4173 1977, DOI: 10.1021/jo00862a001; see also Brussee & Jansen *Tetrahedron Lett* **24** 3261 1983, DOI: 10.1016/S0040-4039(00)88151-4; Akimoto & Yamada *Tetrahedron* **27** 5999 1971, DOI: 10.1016/S0040-4020(01)91765-X; *Beilstein* **6** IV 7020.]

1,1'-Binaphthyl [(±)- 32507-32-7 and 604-53-5, *R*(-)- 24161-30-6, *S*(+)- 734-77-0] $C_{20}H_{14}$, **M 254.3, m 145°, 159°, b ~240°/13mm, (±, 2 forms), 153-154°, 154°, (+) and (-), $[\alpha]_D^{20}$ (-) and (+) ~220 (*C₆H₆).** Purify 1,1'-binaphthyl through a silica gel column with Me₂CO/*C₆H₆ [or Al₂O₃ with 10% *C₆H₆/petroleum ether (b 30-60°)] and recrystallise it from EtOH, pentane, or slow evaporation of *C₆H₆, Me₂CO or Et₂O solutions. Half life ~10 hours at 25° in various solvents. [Wilson & Pincock *J Am Chem Soc* **97** 1474 1975, DOI: 10.1021/ja00839a033; Akimoto & Yamada *Tetrahedron* **27** 5999 1971, DOI: 10.1016/S0040-4020(01)91765-X; *Beilstein* **5** I 358, **5** II 642, **5** III 2465, **5** IV 2634.] **2,2'-Binaphthyl (β, β'-binaphthyl)** [61-78-2] has **m 188°**. Crystallise the 2,2'-binaphthyl from *C₆H₆, or Et₂O/*C₆H₆ (**m 187-189°**). The **2,4,7-trinitrofluorenone complex** forms orange-red needles from EtOH/*C₆H₆ (**m 170.6-171°**). [*Beilstein* **5** H 727, **5** I 359, **5** II 643, **5** III 2467, **5** IV 2636.]

Biphenyl (diphenyl) [92-52-4] $C_{12}H_{10}$, **M 154.2, m 68-75°, 70-71°, b 112°/7mm, 255°/760mm, d_4^{20} 0.992.** Crystallise biphenyl from EtOH, MeOH, aqueous MeOH, petroleum ether (b 40-60°) or glacial acetic acid. Free it from polar impurities by passage through an alumina column in *benzene, followed by evaporation. The residue has been purified by distillation in a vacuum and by zone refining. Treatment with maleic anhydride removes anthracene-like impurities. It has been recrystallised from EtOH followed by repeated vacuum sublimation and passage through a zone refiner. [Taliani & Bree *J Phys Chem* **88** 2351 1984, DOI: 10.1021/j150655a033; *Beilstein* **5** H 576, **5** I 271, **5** II 479, **5** III 1726, **5** IV 1807.]

4-Biphenylcarbonyl chloride [14002-51-8] $C_{13}H_9ClO$, **M 216.7, m 110-112°, 114-115°.** Dissolve the carbonyl chloride in a large volume of petroleum ether (10 x, b 50-70°), filter it through a short column of neutral alumina, evaporate to dryness *in vacuo* and recrystallise it from petroleum ether (b 60-80°). [*Beilstein* **9** IV 2480.] **LACHRYMATORY.**

Biphenyl-2-carboxylic (2-phenylbenzoic) acid [947-84-2] $C_{13}H_{10}O_2$, **M 198.2, m 111-113°, 114°, b 199°/10mm, 343-344°/atm, pK^{25} 3.46.** Crystallise the acid from *C₆H₆/petroleum ether or aqueous EtOH. [*Beilstein* **9** IV 2472.] **Biphenyl-4-carboxylic (4-phenylbenzoic) acid** [92-92-2] has **m 220-225°, 228°, pK^{25} 5.66 (in 50% 2-butoxyethanol)** and is similarly purified. [*Beilstein* **9** IV 2479.]

2,4'-Biphenyldiamine [492-17-1] $C_{12}H_{12}N_2$, **M 184.2, m 45°, b 363°/760mm, $pK_{Est(1)}$ ~4.8, $pK_{Est(2)}$ ~3.9.** Crystallise the diamine from aqueous EtOH or petroleum ether (**m 54-54.5°**). [*Beilstein* **9** III 416, **9** IV 360.]

Biphenyl-4,4'-dicarboxylic acid [787-70-2] $C_{14}H_{10}O_4$, **M 242.2, m >300°, $pK_{Est(1)}$ ~3.5, $pK_{Est(2)}$ ~4.3.** The dicarboxylic acid is a white amorphous or microcrystalline substance which does not melt or sublime. It is best purified by precipitation of an aqueous alkaline solution with mineral acid, washing well with H₂O and drying *in vacuo* at 100°. It is characterised by conversion to **diphenyl-4,4'-dicarbonyl chloride** (with PCl₅) [Work *J Chem Soc* 1315 1940, DOI: 10.1039/JR9400001315], or by phase transfer catalysis with SOCl₂ + BuEt₃N⁺Cl⁻ in 1,2-dichloroethane [Burdett *Synthesis* 441 1991, DOI: 10.1055/s-1991-26487,] which crystallises from *C₆H₆ with **m 184°**. The di-acid chloride gives the **dimethyl ester** with MeOH which has **m 215-217°** (plates from MeOH, **m**'s of **214°** and **224°** were also reported). The **diethyl ester** is similarly prepared with EtOH and has **m 122°** (from EtOH). The **4,4'-dicarbonitrile** [1591-30-6] $C_{14}H_8N_2$, **M 204.2**, prepared from the dicarbonyl chloride and aqueous ammonia followed by dehydration, or from benzidine *via*, a *bis* Sandmeyer reaction, has **m 238-240°**. [*Beilstein* **9** II 665, **9** III 4519, **9** IV 3563.] The dicarboxylic acid was successfully used to make dendrimers with polyhydrophobic and hydrophilic arms *via* intermediate polyhydroxyphenyl esters [Zubarev et al. *Org Lett* **8** 1367 2006, DOI: 10.1021/ol060080x].

Biphenylene [259-79-0] $C_{12}H_8$, **M 152.2, m 111°, 113-114°.** Biphenylene forms yellow crystals from cyclohexane, MeOH (**m 110-111°**) or EtOH (**m 111-112°**). It sublimes *in vacuo*. The **2,4,7-trinitrofluorenone complex** has **m 154°** and the **picrate** gives red needles **m 122°** from EtOH. [*Beilstein* **5** I 298, **5** II 530, **5** III 1935, **5** IV 2137.]

(±)-**α-(4-Biphenyl)butyric acid** [959-10-4] $C_{16}H_{16}O_2$, M 240.3, m 124-125°, $pK_{Est} \sim 4.5$. Crystallise the acid from MeOH, petroleum ether or AcOH (m 123-125°). [Beilstein 9 III 3370, 9 IV 2558.] **γ-(4-Biphenyl)butyric acid** [6057-60-9] has m 118°, 120-121°, $pK_{Est} \sim 4.8$. Crystallise the acid from MeOH (m 118°) or *C_6H_6 (m 118-119°). [Beilstein 9 I 290, 9 III 3370, 9 IV 2558.]

Bis-(p-bromophenyl) ether [53563-56-7] $C_{12}H_8Br_2O$, M 328.0, m 60.1-61.7°. Crystallise the ether twice from EtOH, petroleum ether, once from * benzene and dry it *in vacuo* [Purcell & Smyth *J Am Chem Soc* 83 1063 1961, DOI: 10.1021/ja01466a013]. [Beilstein 6 III 745, 9 IV 1048.]

2R,3R-(+)-1,4-Bis-(4-chlorobenzoyloxy)-2,3-butanediol [85362-86-3] and **2S,3S-(-)-1,4-Bis-(4-chlorobenzoyloxy)-2,3-butanediol** [85362-85-2] $C_{18}H_{20}Cl_2O_4$, M 371.3, m 72-74°, 76-77°, $[\alpha]_D^{20}$ (+) and (-) 6.4 (c 3.1 $CHCl_3$). Recrystallise the diols from toluene-hexane. IR (KBr) has ν_{max} at 3250, 1598, 1493, 1085 cm^{-1} ; 1H NMR ($CDCl_3$) has 2.6-2.9 (2H, m, OH x 2), 3.4-3.7 (4H, m, CH- CH_2O x2), 3.7-4.0 (2H, m, CH x2), 4.47 (4H, s, CH_2Ar x 2), 7.05-7.35 (8H, m, aromatic protons). They are useful reagents for optical resolution of racemates. [Terashima et al. *Tetrahedron Lett* 23 4107 1982, DOI: 10.1016/S0040-4039(00)88360-4; Tamoto et al. *Tetrahedron* 40 4617 1984, DOI: 10.1016/S0040-4020(01)91522-4.]

N,N-Bis-(2-chloroethyl) 2-naphthylamine (chlornaphthazine) [494-03-1] $C_{14}H_{15}Cl_2N_4$, M 268.3, m 54-56°, b 210°/5mm, $pK_{Est} \sim 5.3$. Crystallise it from petroleum ether. At 15° it is soluble in EtOH (3.2%), Et_2O (50%), Me_2CO (84%) and *C_6H_6 (80%). It is a nitrogen mustard. [Beilstein 12 III 2996, 12 IV 3126.] **CARCINOGENIC.**

1,4-Bis-(chloromethyl)durene (1,4-bischloromethyl-2,3,5,6-tetramethylbenzene) [3022-16-0] $C_{12}H_{16}Cl_2$, M 231.2, m 197-198°. Crystallise it three times from *C_6H_6 (m 193-194°) or petroleum ether (m 195-196°), then dry it *in vacuo* in a drying pistol. [Fuson et al. *J Am Chem Soc* 75 5952 1953, DOI: 10.1021/ja01119a050; Beilstein 5 IV 1140.]

2,2-Bis-(p-chlorophenyl)-1,1-dichloroethane (p,p'-DDD) [72-54-8] $C_{14}H_{10}Cl_4$, M 320.1, m 109-110°, 109-111°, 111-112°. Crystallise DDD from EtOH and dry it *in vacuo*. The purity is checked by TLC. [Beilstein 5 III 1830.] **TOXIC INSECTICIDE.**

2,2-Bis-(p-chlorophenyl)-1,1-dichloroethylene (p,p'-DDE) [72-55-9] $C_{14}H_8Cl_4$, M 318.0, m 89-91°. Crystallise DDE from MeOH or EtOH and dry it *in vacuo*. The purity is checked by TLC. It is a metabolite of DDT. [Gätzi & Stambach *Helv Chim Acta* 29 563 1946, DOI: 10.1002/hlca.19460290308; Beilstein 5 H 639, 5 III 1891.] **POSSIBLE CARCINOGEN.**

2,2-Bis-(4-chlorophenyl)-1,1,1-trichloroethane (p,p'-DDT, 1,1,1-trichloro-2,2-bis(p-chlorophenyl)ethane) [50-29-3] $C_{14}H_9Cl_5$, M 354.5, m 108°, 108.5-109°, 107-110°. Crystallise DDT from *n*-propyl alcohol (5ml/g), then dry it in air or an air oven at 50-60°. Alternatively, crystallise it from 95% EtOH, and the purity is checked by TLC. [Beilstein 5 III 1833.] **TOXIC INSECTICIDE.**

4,4'-Bis-(diethylamino)benzophenone [90-93-7] $C_{21}H_{28}N_2O$, M 324.5, m 89-92°, 93-95°, 95-96°, b $\sim 300^\circ/10mm$, $pK_{Est(1)} \sim 1.8$, $pK_{Est(2)} \sim 3.3$. Crystallise the phenone from EtOH (25ml/g) and dry it under vacuum. Its *picrate* forms yellow needles from EtOH with m 178.5°. [Beilstein 14 II 59.]

Bis-(4-dimethylaminobenzylidene)benzidine [6001-51-0] $C_{30}H_{30}N_4$, M 446.6, m 318°, $pK_{Est} \sim 0$. Crystallise the benzidine from nitrobenzene. [Beilstein 14 H 35.]

3,4-Bis-(4-hydroxyphenyl)hexane (Hexesterol, Homoestrol, Syntrogène) [5635-50-7 (no configuration), 84-16-2 (meso-3RS,4-SR)] $C_{18}H_{22}O_2$, M 270.4, m 185-186°, 187°. Free it from diethylstilboestrol by zone refining. Crystallise meso-Hexesterol from * benzene or aqueous EtOH (m 185-188°). The meso-dibenzoyl derivative has m 236-237°. The 3RS,4RS(±)-racemate [5776-72-7] crystallises from petroleum ether, *C_6H_6 /petroleum ether, Et_2O /petroleum ether, or MeOH/ H_2O and has m 128-129°. The (±)-diacetyl derivative has m 137-139°, and the (±)-dipropionyl derivative (Retalon), m 127-128°, crystallises from light petroleum.

The (\pm)-*dibenzoyl* derivative has **m 123-124°**. *Hexesterol 4,4'-diphosphoric acid ester* (Cytostatin) [4188-82-0] **C₁₈H₂₄O₈P₂, M 430.3**, is acidic and is soluble in aqueous alkaline solutions. The *3R,4R(+)-isomer* [26614-21-1] and *3S,4S(-)-isomer* [26614-22-2] crystallise from Et₂O/petroleum ether with **m 80-80.5°** and have [α]_D¹⁷ (+) and (-) 17.7 (c 5, EtOH). Their *dibenzoyl* derivatives have **m 116.5°**. [Beilstein **6** III 5503, **6** IV 6761.] They have estrogenic activity where the optically active forms are more potent, and they have antineoplastic activity. [Aboul-Enein et al. *Anal Profiles Drug Subst* **11** 347 1982, DOI: 10.1016/S0099-5428(08)60269-5; Kharasch & Kleiman *J Am Chem Soc* **65** 491 1941, DOI: 10.1021/ja01244a001.]

4,4-Bis(4-hydroxyphenyl)valeric acid [diphenolic acid, DPA] [126-00-1] **C₁₇H₁₄O₄, M 286.3, m 168-171°, 171-172°, pK_{Est(1)} ~ 4.8 (CO₂H), pK_{Est(2)} ~ 7.55 (OH), pK_{Est(3)} ~ 9.0 (OH)**. When recrystallised from *C₆H₆, the crystals have 0.5 mol of *C₆H₆ (**m 120-122°**, lower melting modification), and when recrystallised from toluene, the crystals have 0.5 mol of toluene. Purify the acid also by recrystallisation from hot H₂O which provides the higher melting modification. It is soluble in Me₂CO, AcOH, EtOH, propan-2-ol, methyl ethyl ketone. It can also be recrystallised from AcOH, heptane/Et₂O or Me₂CO/*C₆H₆. It has λ_{\max} at 225 and 279nm in EtOH. The *methyl ester* has **m 87-89°** (aqueous MeOH to give the *trihydrate*). [Bader & Kontowicz *J Am Chem Soc* **76** 4465 1954, DOI: 10.1021/ja01646a053; Beilstein **10** IV 1890.]

1,4-Bismethylaminoanthraquinone (Disperse Blue 14) [2475-44-7] **C₁₆H₁₄N₂O₂, M 266.3, m 220-222°, C.I. 61500, λ_{\max} 640 (594)nm**. Purify the anthraquinone by thin-layer chromatography on silica gel plates, using toluene/acetone (3:1) as eluent. The main band is scraped off and extracted with MeOH. The solvent is evaporated and the dye is dried in a drying pistol [Land et al. *JCS Faraday Trans 1* **72** 2091 1976, DOI: 10.1039/F19767202091]. It crystallises from *n*-butanol with **m 221-222°** and has λ_{\max} at 539 and 644nm (EtOH). [Beilstein **14** H 198, **14** III 440, **14** IV 459.]

Bis-(1-naphthylmethyl)amine [5798-49-2] **C₂₂H₁₉N₂, M 329.4, m 62°, 63-64°, pK_{Est} ~8.4**. Crystallise the amine from petroleum ether (pale yellow crystals), Et₂O (**m 73-74°**) or *C₆H₆ (**m 62°**). The *hydrochloride* crystallises from H₂O as colourless needles **m 239°**, and the *picrate* has **m 206°(202°)**. [For the reduction of 1-cyanonaphthalene with Ni/H₂ see Rupe & Becherer *Helv Chim Acta* **6** 880 1923, DOI: 10.1002/hlca.19230060196; for the determination of nitrates see Konek *Chem Abstr* **28** 5779 1934, Beilstein **12** II 741, **12** IV 3195.]

Bis-(4-nitrophenyl) carbonate [5070-13-3] **C₁₃H₈N₂O₇, M 304.2, m 142-143°**. Dissolve the carbonate in CHCl₃, wash it with 2N NaOH (3 x) and once with concentrated HCl, dry (Na₂SO₄), evaporate and crystallise the residue from toluene (authors say prisms from 15 volumes of *benzene). [For preparation of activated 4-nitrophenyl esters of protected *H*-amino acids see Glatthard & Matter *Helv Chim Acta* **46** 795 1963, and Tolle *J Am Chem Soc* **104** 6883 1982, DOI: 10.1021/ja00388a114; and for the preparation of ureas see Izdebski & Pawlak *Synthesis* 423 1989, DOI: 10.1055/s-1989-27272; Beilstein **6** III 820.]

Bis-(4-nitrophenyl) ether [101-63-3] **C₁₂H₈N₂O₅, M 260.2, m 142-143°, 144.4-144.7°, 147-148°**. Crystallise the ether twice from *C₆H₆ or petroleum ether and dry it *in vacuo*. [Beilstein **6** II 822, **6** IV 1290.]

Bis-(4-nitrophenyl) methane [1817-74-9] **C₁₃H₁₀N₂O₄, M 258.3, m 183°, 184°, 187°**. Crystallise the methane twice from *C₆H₆, petroleum ether or AcOH (**m 188.6-189.6°**), and dry it *in vacuo*. [Beilstein **5** III 1797, **5** IV 1853.]

Bis-(trifluoroacetoxy)iodobenzene (BTI) [2712-78-9] **C₁₀H₅F₆IO₄, M 430.0, m 112-114° (dec), 120-121°, 124-126°**. Crystallise the iodo compound from warm trifluoroacetic acid and dry it over NaOH pellets. Recrystallise it also from Me₂CO/petroleum ether. Its melting point depends on the heating rate. [Spyroutis & Varvoglis *Synthesis* 445 1975, DOI: 10.1055/s-1975-23796; application: Almond et al. *Org Synth* **66** 132 1988, DOI: 10.15227/orgsyn.066.0132.] The cyclisation of styrylamines to *N*-alkyl and *N*-aryl indoles in CHCl₃ is mediated by BTI under mild conditions (30 minutes at room temperature) [Du et al. *Org Lett* **8** 5919 2006, DOI: 10.1021/ol062288o].

***N*-BOC-1,2-phenylenediamine [(2-aminophenyl)carbamic acid, *tert*-butyl ester]** [146651-75-4]

C₁₁H₁₆NO₂, M 208.3, m 109-114°. Purify the ester by recrystallisation from CHCl₃/hexane (1:1, v/v) and dry it *in vacuo*. [Seto et al. *J Am Chem Soc* **115** 1321 1993, DOI: 10.1021/ja00057a015; Sessler et al. *J Am Chem Soc* **127** 11442 2005, DOI: 10.1021/ja0522938.]

Brilliant Green (4-dimethylaminotriphenyl carbinol, Emerald Green, Malachite Green G) [633-03-4] C₂₇H₁₄N₂O₄S, M 482.7, m 209-211°(dec), pK²⁵ 4.75. Purify the dye by precipitating the *perchlorate* from aqueous solution (0.3%) after filtering, heating to 75° and adjusting to pH 1-2 with dilute H₂SO₄ to form the HSO₄⁻ salt. It has λ_{max} at 623nm, and changes colour from yellow to green at pH 0.0 to pH 2.6 respectively. Recrystallise it from EtOH/water (1:4) [Kerr & Gregory *Analyst (London)* **94** 1036 1969, DOI: 10.1039/AN9699401036]. [*Beilstein* **13** IV 2281.] It is used as an external and internal antiseptic.

4-Bromoacetanilide [103-88-8] C₈H₈FBNO, M 214.1, m 165-169°, 167°. Crystallise the anilide from aqueous MeOH or EtOH. Purify it by zone refining. [*Beilstein* **12** IV 1504.] It has analgesic and antipyretic properties.

4-Bromoacetophenone [99-90-1] C₈H₇BrO, M 199.1, m 49-51°, 54°, b 117°/7mm, 130°/15mm, 255°/atm. Crystallise it from EtOH, MeOH or from petroleum ether (b 80-100°). The *oxime* m 128.5° crystallises from aqueous EtOH. [Tanner *J Org Chem* **52** 2142 1987, DOI: 10.1021/jo00387a005; *Beilstein* **7** IV 647.]

ω-Bromoacetophenone (phenacyl bromide) [70-11-1] C₈H₇BrO, M 199.1, m 48-51°, 50°, 57-58°, 135°/18mm Crystallise the bromide from EtOH, MeOH or petroleum ether (b 80-100°). [Tanner *J Org Chem* **52** 2142 1987, DOI: 10.1021/jo00387a005; *Beilstein* **7** IV 649.] **LACHRYMATORY.** Useful for characterising acids as *phenacyl esters* by reaction with sodium salts of acids in EtOH or aqueous EtOH.

4-Bromoaniline [106-40-1] C₆H₆BrN, M 172.0, m 66°, pK²⁵ 3.86. Recrystallise the base (with much loss) from aqueous EtOH or EtOH/Et₂O. The *benzoyl* derivative has m 204° (from EtOH). [*Beilstein* **12** IV 1497.]

2-Bromoanisole (1-bromo-2-methoxybenzene) [578-57-4] C₇H₇BrO, M 187.0, m 2.5°, b 124°/40mm, 223°/atm, d₄²⁰ 1.513, n_D²⁵ 1.5717. Crystallise the anisole by repeated partial freezing, then distil it under reduced pressure. [*Beilstein* **6** IV 1037.] **4-Bromoanisole (1-bromo-4-methoxybenzene) [104-92-7]** has **M 13.4°, b 99-100°/18mm, 124°/40mm, d₄²⁰ 1.495, n_D²⁵ 1.5617.** Crystallise the anisole by repeated partial freezing, then distil it under reduced pressure. [*Beilstein* **6** III 741, **6** IV 1044.]

9-Bromoanthracene [1564-64-3] C₁₄H₉Br, M 257.1, m 98-100°. Crystallise 9-bromoanthracene from MeOH or EtOH followed by sublimation *in vacuo*. [Masnovi et al. *J Am Chem Soc* **108** 1126 1986, DOI: 10.1021/ja00266a003; *Beilstein* **5** IV 2295.]

4-Bromobenzal diacetate [55605-27-1] C₁₁H₁₁BrO₄, M 287.1, m 95°. Crystallise the diacetate from hot EtOH (3ml/g). [Liebermann & Connor *Org Synth Coll Vol* **2** 441 1948, DOI: 10.15227/orgsyn.018.0061; *Beilstein* **7** II 182, **7** IV 579.]

Bromobenzene [108-86-1] C₆H₅Br, M 157.0, m -31°, b 28°/5mm, 40°/10mm, 68.6°/40mm, 110°/200mm, 155.9°/atm, d₄²⁰ 1.495, n_D²⁰ 1.5588, n_D¹⁵ 1.56252. Wash bromobenzene with concentrated H₂SO₄ (CARE) then 10% NaOH or NaHCO₃ solutions, and H₂O. Dry it with CaCl₂ or Na₂SO₄, or pass it through activated alumina, before refluxing with, and distilling from, CaH₂, using a glass helix-packed column. [*Beilstein* **5** IV 670.]

4-Bromobenzene diazonium tetrafluoroborate [673-40-5] C₆H₄BBrF₄N₂, M 270.8, m 133°(dec), 135-140°(dec), 135°(dec). Wash the salt with Et₂O until the wash is colourless and allow it to dry by blowing N₂ over it. Store it at 0-4° in the dark. [Schiemann & Pillarsky *Chem Ber* **64** 1340 1931, DOI: 10.1002/cber.19310640622; *Beilstein* **16** III 517.]

2-Bromobenzonitrile [2042-37-7] C₇H₄BrN, M 182.0, m 52-54°, 55.8°, 56°, 53-57°, b 251-253°/754mm. The nitrile is prepared from 2-bromobenzoic acid by treatment with SOCl₂ (23 hours boiling) and distilling to give **2-bromobenzoyl chloride**, b 120-126°/15mm, n_D²⁰ 1.5925, which on treatment with 12 equivalents of NH₃

gives a 98% yield of **2-bromobenzamide** **m 159.5-161.5°** [4001-73-4]. By boiling this amide with excess of SOCl_2 for 17 hours followed by evaporation and steam distillation gives an 84% yield of the nitrile that forms needles when crystallised from H_2O . [Lutz et al. *J Org Chem* **12** 617 1947, DOI: 10.1021/jo01169a001.] It has also been prepared by the method of Miller [*Org Synth Coll Vol* **3** 646, 648 1955, DOI: 10.15227/orgsyn.029.0075] in which the amide is treated with a large excess of POCl_3 in the presence of sodium metabisulfite ($\text{Na}_2\text{S}_2\text{O}_5$) to give the nitrile (**m 55-55.5°**) [Herbst & Wilson *J Org Chem* **22** 1142 1957, DOI: 10.1021/jo01361a002]. [*Beilstein* **9** H 348, **9** II 232, **9** III 1387, **9** IV 1013.]

2-Bromobenzoic acid (o-bromobenzoic acid) [88-65-3] $\text{C}_7\text{H}_5\text{BrO}_2$, **M 201.0**, **m 148.9°, 150°, pK²⁰ 2.88**. Crystallise the acid from $^*\text{C}_6\text{H}_6$ or MeOH. The **anilide** has **m 141°** (from EtOH/ H_2O). [*Beilstein* **9** IV 1011.] **3-Bromobenzoic acid (m-bromobenzoic acid)** [585-76-2] has **m 155°, pK²⁵ 3.81**. Crystallise the acid from acetone/water, MeOH or acetic acid. The **anilide** has **m 137°** (from EtOH/ H_2O). [*Beilstein* **9** IV 1013.] **3-Bromobenzoic acid hydrazide** [39115-96-3] $\text{C}_7\text{H}_7\text{BrN}_2\text{O}$, **M 215.1**, prepared by heating the acid chloride or ester with excess of hydrazine hydrate and recrystallising from EtOH or aqueous EtOH has **m 157-159°** [*Beilstein* **9** H 351]. **4-Bromobenzoic acid (p-bromobenzoic acid)** [586-76-5] has **m 251-252°, 254-256°, 257-258°, pK²⁵ 3.96**. Crystallise the acid from MeOH, or MeOH/water mixture, 90% EtOH and Et_2O . The **methyl ester** has **m 81°** from Et_2O or dilute MeOH. The **anilide** has **m 197°** (from EtOH). [Hale & Thorp *J Am Chem Soc* **35** 262 1913, DOI: 10.1021/ja02192a008; Vandenbelt et al. *Anal Chem* **26** 726 1954, DOI: 10.1021/ac60088a031; *Beilstein* **9** IV 1017.] The acid is also obtained by oxidation of 4-bromoalkyl (Pr, *iso*-Pr, Bu, *iso*-Bu and *sec*-Bu)benzenes (5ml) in refluxing solutions of concentrated HNO_3 (100ml) and H_2O [Lamneck *J Am Chem Soc* **76** 1106 1954, DOI: 10.1021/ja01633a051]. **4-Bromobenzoic acid hydrazide** [5933-32-4] $\text{C}_7\text{H}_7\text{BrN}_2\text{O}$, **M 215.1**, prepared by heating the ester, or acid chloride (see below) with excess of hydrazine hydrate and recrystallising from EtOH or aqueous EtOH has **m 165-167°** [*Beilstein* **9** H 354].

4-Bromobenzophenone [90-90-4] $\text{C}_{13}\text{H}_9\text{BrO}$, **M 261.1**, **m 79-84°, 81°, 81-82°, b 350°/atm**. Crystallise the phenone from EtOH. The **2,4-dinitrophenylhydrazone** forms orange-red leaflets from dioxane/EtOH with **m 207-209°**. [Allen & Van Allan *J Am Chem Soc* **66** 7 1944, DOI: 10.1021/ja01229a003; *Beilstein* **7** H 422, **7** III 2079, **7** IV 1378.]

4-Bromobenzoyl acetonitrile [4592-94-3] $\text{C}_9\text{H}_6\text{BrNO}$, **M 224.1**, **m 160-164°, 162.4-163.4°**. The nitrile is purified by dissolving in slightly alkaline H_2O (charcoal), filtering and acidifying with HCl to give colourless needles (**m 162-163°**). It recrystallises from EtOH. With $\text{Me}_2\text{SO}_4/\text{KOH}$ at 130° it gives **4-bromo-β-methoxycinnamyl nitrile** **m 58.5-59.5°** (from high boiling petroleum ether) [Fuson & Wolf *J Am Chem Soc* **61** 1940 1939, DOI: 10.1021/ja01877a002; Grothaus & Dains *J Am Chem Soc* **58** 1334 1936, DOI: 10.1021/ja01299a006]. [*Beilstein* **10** III 2998.]

4-Bromobenzoyl chloride [586-75-4] $\text{C}_7\text{H}_4\text{BrClO}$, **M 219.5**, **m 36-39°, 39.8°, 41°, b 62°/0.1mm, 104.5°/6mm, 126.4-127.2°/14mm, 174°/102mm**. Check IR of a film to see if OH bands are present. If absent then recrystallise from petroleum ether and dry it *in vacuo*. If OH bands are weak, then distil it *in vacuo* and recrystallise if necessary. If OH bands are very strong, then treat with an equal volume of redistilled SOCl_2 reflux for 2 hours, then evaporate excess of SOCl_2 and distil the residual oil or low melting solid. Store it in the dark away from moisture. **LACHRYMATORY**. [Martin & Partington *J Chem Soc* 1175 1936, DOI: 10.1039/JR9360001175; *Beilstein* **9** IV 1023.]

4-Bromobenzyl bromide [589-15-1] $\text{C}_7\text{H}_6\text{Br}_2$, **M 249.9**, **m 60-61°, 61-62.5°, 62-64°, b 115-124°/12mm, 125-135°/12mm**. It has been prepared by peroxide catalysed bromination, and recrystallised from EtOH, or/and distillation in a vacuum using a Fenske-type column (e.g. 31 cm in length packed with 1/16 in glass helicies with a take-off of the cold finger type) [Goerner & Nametz *J Am Chem Soc* **73** 2940 1951, DOI: 10.1021/ja01150a509]. [*Beilstein* **5** H 308, **5** IV 836.] **LACHRYMATORY**, use an efficient fume cupboard. **p-Bromobenzyl chloride** [589-17-3] $\text{C}_7\text{H}_6\text{BrCl}$, **M 205.5**, has **m 38-39°, 40-41°, b 100-115°/12mm, 136-137°/27mm**. It has been prepared by peroxide catalysed chlorination, and recrystallised from EtOH, and/or distil it in a vacuum using a Fenske-type column (e.g. 31 cm in length packed with 1/16 inch glass helicies with a take-off of the cold finger type) [Goerner & Nametz *J Am Chem Soc* **73** 2940 1951, DOI: 10.1021/ja01150a509]. [*Beilstein* **5** IV 832.] **LACHRYMATORY**, use an efficient fume cupboard.

2-Bromobiphenyl [2052-07-5] $C_{12}H_9Br$, M 233.1, m 1.5-2.0°, b 140°/11mm, 148-150°/10mm, 297-298°/atm, d_4^{25} 1.352, n_D^{20} 1.6248. 2-Bromobiphenyl is prepared from 2-aminobiphenyl (12g) in hot constant-boiling aqueous HBr (21g, 3-equivalents) by diazotisation at 5° with a solution of NaNO₂ (6g) in H₂O (10ml), then excess of HNO₂ is removed at the end of the reaction by addition of urea with stirring for 20 minutes. This solution is added dropwise into a solution of CuBr (from 15g of CuSO₄) in constant-boiling aqueous HBr (20ml) with stirring, set aside for 20 minutes, after which the brown complex is decomposed by heating on a steam bath for 2 hours. The dark oil is extracted into Et₂O, the extract is filtered through glass wool, washed with H₂O, dilute NaOH, H₂O again, dried (CaCl₂), filtered, evaporated and the residual oil is distilled to give a pale yellow oil (7g, b 146-152°/12mm). Further purification can be achieved by dissolving it in *C₆H₆, passing through an alumina column and eluting with *C₆H₆, evaporating to dryness and distilling the residual oil (b 148-150°/10mm) to yield pure colourless 2-bromobiphenyl. This has been recrystallised from pentane at -40° to give a liquid with a freezing point of 1.5-2.0°. [de la Mare & Hassan *J Chem Soc* 3004 1957, DOI: 10.1039/JR9570003004; Augood et al. *J Chem Soc* 3412 1953, DOI: 10.1039/JR9530003412; NMR: Brownstein *J Am Chem Soc* 80 2300 1958, DOI: 10.1021/ja01150a509; *Beilstein* 5 H 580, 5 II 485, 5 III 1742, 5 IV 1818.] It is used for the preparation of John-Phos and related catalytic ligands (see Chapter 5, Catalysis—Part 2). **3-Bromobiphenyl** [2113-57-7] has b 103-104°/0.2mm, 110°/1mm, 158-167°/11mm, 169-173°/17mm, d_4^{25} 1.3976, n_D^{20} 1.6380. 3-Bromobiphenyl can be prepared by bromination of 2-acetamidobiphenyl (m 118-119° from aqueous AcOH) with one molecular equivalent of Br₂ in AcOH; the monobromo derivative (m 127-127.5° from EtOH) is hydrolysed with 95% EtOH/conc HCl (1.4:1, 4 hours reflux) poured into excess of H₂O, basified and filtered off to give 2-amino-5-bromobiphenyl (m 53-56°). The base is subsequently diazotised and the diazonium salt is deaminated by warming with Cu bronze or with hypophosphorous acid [cf. Kornblum in *Organic Reactions* J. Wiley & Sons NY, II 294 1944] to yield crude 3-bromobiphenyl which is isolated by steam distillation, extraction of the distillate with *C₆H₆, washing the extract successively with, dilute NaOH, H₂O, concentrated H₂SO₄, H₂O, drying (K₂CO₃), evaporating and distilling. This can be purified further by passing a solution of it in *C₆H₆ through an Al₂O₃ column, eluting with *C₆H₆, and the combined eluates are distilled. The distillate is dissolved in petroleum ether (b 40-60°), washed several times with concentrated H₂SO₄, H₂O again, aqueous NaHCO₃, H₂O again, dried (CaCl₂), filtered, the solvent is evaporated off and the residue is fractionally distilled to give pure 3-bromobiphenyl. [de la Mare & Hassan *J Chem Soc* 3004 1957, DOI: 10.1039/JR9570003004; Lichtin & Leftin *J Am Chem Soc* 74 4207 1952, DOI: 10.1021/ja01136a510; Huber et al. *J Am Chem Soc* 68 1109 1946, DOI: 10.1021/ja01210a060]. It has also been prepared in low yield (13%) from diazotized *m*-bromoaniline and *C₆H₆ according to Gomberg and purified as above [Marvel et al. *J Am Chem Soc* 61 77 1939, DOI: 10.1021/ja01870a022, also see 4-bromobiphenyl below]. [*Beilstein* 5 II 485, 5 III 1742, 5 IV 1818.] **4-Bromobiphenyl** [92-66-0] has m 82-86°, 88.8-89.2°, 89.5°, 90°, b 170-175°/8mm, 310°/atm. This biphenyl is prepared by the method of Gomberg & Bachmann [*Org Synth Coll Vol* 1 113 1944, DOI: 10.15227/orgsyn.008.0042] from diazotised 4-bromoaniline and benzene, then extract into Et₂O, wash with base, acid, brine, dry (Na₂SO₄), filter, evaporate and recrystallise to constant melting point from absolute EtOH (89.7°, after drying *in vacuo*). [Elks et al. *J Chem Soc* 1284 1940, DOI: 10.1039/JR9400001284; Augood et al. *J Chem Soc* 3412 1953, DOI: 10.1039/JR9530003412; *Beilstein* 5 H 580, 5 I 275, 5 II 485, 5 III 1742, 5 IV 1819.]

1-Bromo-2-chlorobenzene [694-80-4] C_6H_4BrCl , M 191.5, m -12°, b 79-82°/14mm, 200-202°/atm, 204°/atm, d_4^{25} 1.638, n_D^{20} 1.582. *o*-Chlorobromobenzene has been prepared in at least three different ways. It is obtained in 54% yield by carrying out a Sandmeyer reaction from *o*-chloroaniline *via* diazotisation (using NaNO₂/HBr) and decomposing it with CuBr in 48% aqueous HBr. Steam distil off the product, basify the distillate with 20% aqueous NaOH (extract it into Et₂O if quantities are small), dry the organic layer (Na₂SO₄), filter and distil [Fry & Grote *J Am Chem Soc* 48 710, 1926, DOI: 10.1021/ja01414a025]. The second procedure involves adding Br (32g) in CCl₄ (50ml) during 20 minutes to refluxing silver 2-chlorobenzoate in CCl₄ (250ml), boil for a further 30 minutes to complete the decarboxylation, filter off the AgBr, wash the filtrate with aqueous Na₂SO₃ (until red colour of Br is discharged), aqueous 0.5M Na₂CO₃ (2 x 25ml), evaporate the solvent and distil bromochlorobenzene (38-48% yield) [Dauben & Tilles *J Am Chem Soc* 72 3185 1950, DOI: 10.1021/ja01163a105; Barnes & Prochaska *J Am Chem Soc* 72 3188 1950, DOI: 10.1021/ja01163a106]. The third procedure involves phase transfer catalytic radical bromination. *o*-Chlorobenzene free radical is generated from the respective *o*-benzenediazonium⁺ BF₄⁻ with KOAc/THF and a

catalytic amount of 18-crown-6 using CBrCl_3 as brominating agent to produce *o*-bromochlorobenzene in 70% yield, a great improvement on the Sandmeyer method [Korzeniowski & Gokel *Tetrahedron Lett* 3519 1977, DOI: 10.1016/S0040-4039(01)83281-0]. [Beilstein 5 H 209, 5 I 115, 5 II 161, 5 III 562, 5 IV 680.]

4-Bromo-4'-chlorobenzophenone [27428-57-5] $\text{C}_{13}\text{H}_8\text{BrClO}$, M 295.6, m 150°. Crystallise the phenone from EtOH or $^*\text{C}_6\text{H}_6$ and further purify it by zone refining (100 passes) [Groves & Turner *J Chem Soc* 509 1929, DOI: 10.1039/JR9290000509; Lin & Hanson *J Phys Chem* 91 2279 1987, DOI: 10.1021/j100293a014]. [Beilstein 7 II 360, 7 III 2081.]

2'-Bromo-2,6-dimethoxybiphenyl [755017-61-9] $\text{C}_{14}\text{H}_{13}\text{BrO}_2$, M 293.2, m 141-142°, 143-146°. This intermediate, which is used for the preparation of the ligand S-Phos, is synthesised by adding *n*-BuLi (9.60ml, 1.6M solution in hexanes, 15.4mmol, 1.2 equivalents) *via* a syringe over 5 minutes to a cold (0°) solution of 1,3-dimethoxybenzene (2.00ml, 15.3mmol, 1.2 equivalents, [151-10-0]) in dry THF (30ml), allowing the temperature to rise to ~25°, then it is stirred for 5 hours. The mixture is re-cooled (0°) and 2-bromochlorobenzene (1.50ml, 12.8mmol, 1.0 equivalents, [694-80-4]) is added dropwise *via* a syringe over 15 minutes while stirring vigorously; and the burgundy coloured solution is stirred for a further 15 minutes at 0°. MeOH (0.25ml) is added *via* syringe to decompose excess BuLi, the whole is evaporated to dryness *in vacuo*, the residue is then stirred with Et_2O (50ml) and H_2O (50ml), the layers are separated, the aqueous phase is extracted with Et_2O (2 x 25ml), the combined Et_2O solutions are dried (MgSO_4), filtered and evaporated *in vacuo*. The yellow residue is recrystallised from MeOH to provide the analytically pure biphenyl (3.03g, 81%) as pale yellow crystals. The IR (film) has ν_{max} at 2964, 1584, 1472, 1432, 1248, 1108, 1025, 783 cm^{-1} ; the ^1H NMR [100MHz, CDCl_3] has δ at 7.69 (dd, $J = 6.9, 1.1\text{Hz}$, 1H), 7.34-7.40 (m, 2H), 7.20-7.28 (m, 2H), 6.68 (d, $J = 8.5\text{Hz}$, 2H), 3.76 (s, 6H) from TMS; the ^{13}C NMR [75MHz, ^1H decoupled, CDCl_3] has δ at 157.8, 136.25, 132.52, 132.47, 129.6, 128.8, 127.1, 125.4, 119.0, 104.2, 56.2 from TMS. [Barder et al. *J Am Chem Soc* 127 4685 2005, DOI: 10.1021/ja042491j.]

4-Bromo-*N,N*-dimethylaniline [586-77-6] $\text{C}_8\text{H}_{10}\text{BrN}$, M 200.1, m 55°, b 264°/atm, pK 19.2 (acidic), pK²⁵ 4.23. Reflux it for 3 hours with two equivalents of Ac_2O , then fractionally distil it. [Beilstein 12 IV 1499.]

1-Bromo-2,4-dinitrobenzene [584-48-5] $\text{C}_6\text{H}_3\text{BrN}_2\text{O}_4$, M 247.0, m 72.5-73°, 75°. Crystallise it from ethyl ether, isopropyl ether, 80% EtOH or absolute EtOH. [Beilstein 5 III 640, 5 IV 749.]

***N*-(2-Bromoethyl)phthalimide** [574-98-1] $\text{C}_{10}\text{H}_8\text{BrNO}_2$, M 254.1, m 81-83°, 82.5-83.5°. The following is to be carried out in an efficient FUME HOOD. Dissolve the compound (180g) in CS_2 (500 ml) by refluxing for 15 minutes (to cause the separation of the most likely impurity, 1,2-diphthalimidoethane), filter and evaporate under reduced pressure. The product forms light tan crystals (m 78-80°). Recrystallise it from EtOH (charcoal) [the compound (50g) is dissolved in hot 75% EtOH (200ml), boiled for *ca* 10 minutes, carbon is added (5g, Norite), filter and cool to 0°], to give white crystals (40g) which can be recrystallised (m 80-81°); and further recrystallisation gives m 82-83°. [Salzberg & Supniewski *Org Synth Coll Vol I* 119 1932, DOI: 10.15227/orgsyn.007.0008; Landini & Rolla *Synthesis* 389 1976, DOI: 10.1055/s-1976-24051; Beilstein 21/10 V 275.]

3-Bromo-5-hydroxybenzoic acid [140472-69-1] $\text{C}_7\text{H}_5\text{BrO}_3$, M 217.0, m 233.5°, 237-241°, pK_{Est(1)} ~2.3, pK_{Est(2)} ~13.0. The acid crystallises from H_2O (m 238-239°), and with Me_2SO_4 it yields the 5-methoxy derivative with m 190-191° (from EtOH). The 5-methoxy-methyl ester, prepared by reaction of the acid chloride with MeOH, has b 156-157°/4mm. [Baddar et al. *J Chem Soc* 469 1955, DOI: 10.1039/JR9550000465; Beilstein 10 IV 333.]

2-Bromomethylanthraquinone (MAQ-Br) [7598-10-9] $\text{C}_{15}\text{H}_9\text{BrO}_2$, M 301.1, m 200-202°, 203°(dec). Recrystallise the quinone from AcOH, wash the crystals with a little Et_2O , dry it in air and then in a vacuum at 100°. It is prepared by bromination of 2-methylanthraquinone with $\text{Br}_2/\text{PhNO}_2$ at 145-150°, or *N*-bromosuccinimide in CCl_4 containing a trace of $(\text{PhCOO})_2$. [Beilstein 7 IV 2576.]

2-(Bromomethyl)benzonitrile [22115-41-9] $\text{C}_8\text{H}_6\text{BrN}$, M 196.0, m 72-74°, 72-73°, 79°, b 152-155°/15mm. Purify the nitrile by steam distillation. Extract the distillate with Et_2O , dry the extract (Na_2SO_4), evaporate and

distil the residue. The solidified distillate can be recrystallised from petroleum ether or cyclohexane. ^1H NMR (CDCl_3) with δ at 7.8-7.2 (m 4H), 4.62 (s, 2H); the IR has ν_{max} at 2238 cm^{-1} . [Drory *Chem Ber* **24** 2563 1891, DOI: 10.1002/cber.18910240254; Borsche et al. *Chem Ber* **67** 675 1934, DOI: 10.1002/cber.19340670428; Buckley et al. *Aust J Chem* **22** 577 1969, DOI: 10.1071/CH9690577; Beilstein **9** III 2312.] **LACHRYMATORY**, use an efficient fume cupboard.

4-Bromo- α -methylbenzyl alcohol [(\pm) 5391-88-8, 25675-29-0, *R*-(+) 76155-78-7, *S*-(-) 100760-04-1] **C₈H₉BrO**, **M 201.1**. The (\pm)-racemate is purified by distillation in a vacuum (**b 90°/1mm, 119-121°/7mm, d 1.46**) and it solidifies on cooling (**m 36-37°**) [Overberger et al. *Org Synth Coll Vol* **3** 200 1955, DOI: 10.15227/orgsyn.028.0028]. The (\pm)-*tosyl* derivative [114200-15-6] has **m 56-57°**. The *R*-(+)-*enantiomer* is also purified by distillation in a vacuum (**b 110°/3mm, d₄²⁵ 1.322, n_D²⁰ 1.569**) and has $[\alpha]_{\text{D}}^{20} +39$ (c 1, CHCl_3), +32.9 (c 1.39, MeOH). The *S*-(-)-*enantiomer* is similarly purified, and has respective negative optical rotations. [Stein et al. *Can J Chem* **63** 3442 1985, DOI: 10.1139/v85-565; Cervinka et al. *Col Czech Chem Commun* **51** 401 1986, DOI: 10.1135/ccccc19860401; Beilstein **6** II 447.]

2-(Bromomethyl)-naphthalene [939-26-4] **C₁₁H₉Br**, **M 221.1**, **m 52-54°, 56°, 56-57°, b 133-136°/0.8mm, 214°/100mm**. Dissolve the bromo compound in toluene, wash it with saturated aqueous NaHCO_3 , dry (Mg SO_4), evaporate, fractionally distil the residue and recrystallise the solidified distillate from EtOH. [Chapman & Williams *J Chem Soc* 5035, 1952, DOI: 10.1039/JR9520005035; Bergmann & Szmuszkowicz *Bull Soc Chim Fr* **20** 566 1953, Beilstein **5** IV 1698.] **LACHRYMATORY**.

1-Bromonaphthalene [90-11-9] **C₁₀H₇Br**, **M 207.1**, **m -2 to -1° and m 6.2°** (to modifications), **b 84.2°/1mm, 118°/6mm, 133.6°/10mm, 172.2°/40mm, 281.1°/760mm, d₄²⁰ 1.4834, n_D^{16.5} 1.66011**. Purify 1-bromonaphthalene by passage through activated alumina, and three vacuum distillations. It darkens in air on standing when distilled at atmospheric pressure but not so when distilled in a vacuum below 15mm. It is used in *refractometry*, e.g. determination of H_2O in EtOH by the cloud method, determination of refractive index of crystals when immersed in it, for determination of fat content; and in microscopy as a general immersion oil when mixed with polymerised castor oil. The **1,3,5-trinitrobenzene complex**, **m 137°**, crystallises in yellow needles from EtOH. [Clarke & Brethen *Org Synth Coll Vol* **I** 121 1941, DOI: 10.15227/orgsyn.001.0035; Beilstein **5** H 547, **5** IV 1665.] **2-Bromonaphthalene** [580-13-2] has **m 52-55°, 54-56°, 59°, 123-127°/5mm, 281.1°/760mm, d₄²⁵ 1.605g/ml, n_D²⁰ 1.663**. Purify 2-bromonaphthalene by fractional elution from a chromatographic column of activated alumina. Crystallise it from EtOH. [Beilstein **5** IV 1667.]

1-Bromo-2-naphthol [573-97-7] **C₁₀H₇BrO**, **M 223.1**, **m 76-78°, 83°, 84°, pK_{Est} ~8.0**. Distil the naphthol at 10mm then recrystallise it from $^*\text{C}_6\text{H}_6$ /petroleum ether (b 30-60°) **m 80-81°**. The *benzoyl* derivative has **m 98.5-99.5°** (from MeOH). [Hazlet *J Am Chem Soc* **62** 2156 1940, DOI: 10.1021/ja01865a065; Beilstein **6** H 650, **6** II 604, **6** III 2994.] **6-Bromo-2-naphthol** [15231-91-1] has **m 122-124°, 122-126°, pK_{Est} ~9.1**. Crystallise the naphthol from EtOH or $^*\text{C}_6\text{H}_6$ /petroleum ether (**m 128°**). The *benzoyl* derivative has **m 122°**, (from EtOH). [Ruggli & Michels *Helv Chim Acta* **14** 779 1931, DOI: 10.1002/hlca.19310140411; Beilstein **6** H 651, **6** II 605, **6** III 2996.]

ω -Bromo-4-nitroacetophenone [99-81-0] **C₈H₆BrO₃**, **M 244.1**, **m 94-99°, 98°**. Crystallise it from $^*\text{C}_6\text{H}_6$ /petroleum ether. [Beilstein **7** IV 661.] **IRRITANT**.

2-Bromonitrobenzene (2-bromo-1-nitrobenzene) [577-19-5] **C₆H₄BrNO₂**, **M 202.1**, **m 40 to 42°, 43°, b 261°/atm**. Crystallise it twice from petroleum ether, using charcoal before the first crystallisation. [Beilstein **5** III 618, **5** IV 728.] **3-Bromonitrobenzene (3-bromo-1-nitrobenzene)** [585-79-5] has **m 51° to 54°, 53.5-55.5°, 55-56°, b 256°/atm**. Crystallise it twice from petroleum ether, using charcoal before the first crystallisation. [Beilstein **5** III 618, **5** IV 729.]

4-Bromonitrobenzene (4-bromo-1-nitrobenzene) [586-78-7] **C₆H₄BrNO₂**, **M 202.1**, **m 124 to 126°, 127°, b 255-256°/atm**. Crystallise it twice from petroleum ether, using charcoal before the first crystallisation. [Beilstein **5** III 619, **5** IV 729.]

4-Bromophenacyl bromide [99-73-0] $C_8H_6Br_2O$, M 277.9, m 108 to 110°, 109-110°, 110-111°. Crystallise the bromide from EtOH (*ca* 8ml/g). Useful for making phenacyl esters for characterising carboxylic acids by reacting with the Na salt of the acid. [Langley *Org Synth Coll Vol* 1 127 1941, DOI: 10.15227/orgsyn.009.0020; *Beilstein* 7 IV 652.]

2-Bromophenol [95-56-7] C_6H_5BrO , M 173.0, m 3 to 7°, 6°, b 195°/atm, d_4^{20} 1.492, n_D^{20} 1.589, pK^{20} 8.508, pK^{30} 8.400, pK^{35} 8.352, pK^{40} 8.308, pK^{50} 8.231. Purify the phenol by at least two passes through a chromatographic column and distil it. [*Beilstein* 6 IV 1037.] **3-Bromophenol** [591-20-8] has m 28 to 32°, 31°, b 88-89°/3mm, 236°/atm, pK^{20} 9.099, pK^{30} 8.967, pK^{35} 8.907, pK^{40} 8.853, pK^{50} 8.753, pK^{60} 8.666. Purify the phenol by at least two passes through a chromatographic column and distil it. [Carpenter et al, *J Org Chem* 16 586 1951, DOI: 10.1021/jo01144a011; *Beilstein* 6 IV 1037.] **4-Bromophenol** [106-41-2] has m 64°, 235-236°/atm, pK^{25} 9.36, $pK^{31.9}$ 9.244, $pK^{39.7}$ 9.152, $pK^{45.3}$ 9.073, $pK^{50.8}$ 9.019, $pK^{56.0}$ 8.971, $pK^{58.6}$ 8.947. Crystallise the phenol from $CHCl_3$, CCl_4 , petroleum ether (b 40-60°), Et_2O or water (14.3wt% at ~25°), and dry it at 70° under vacuum for 2 hours. It is used as a disinfectant. [*Beilstein* 6 IV 1043.]

4-Bromophenoxy)acetic acid [1878-91-7] $C_8H_7BrO_3$, M 231.1, m 157°, 158°, pK^{25} 3.13. Crystallise the acid from EtOH or H_2O (m 161.4-161.8°). [Hayes & Branch *J Am Chem Soc* 65 1555 1943, DOI: 10.1021/ja01248a031; *Beilstein* 6 III 747, 6 IV 1052.]

3-(4-Bromophenoxy)propionic acid [93670-18-9] $C_9H_9BrO_3$, M 245.1, m 146°, pK_{Est} ~4.2. Crystallise the acid from EtOH, MeOH or $*C_6H_6$ /hexane (m 144-145°). [*Beilstein* 6 III 748, 6 IV 1052.]

4-Bromophenylacetic acid [1878-68-8] $C_8H_7BrO_2$, M 215.0, m 112-113°, 113-115°, 114°, pK^{25} 4.19. The acid crystallises from H_2O as needles. The *acid chloride* has b 238°/760mm, m 50°, the *4-nitrobenzyl ester* forms elongated prisms from EtOH with m 128-129°, and the *anilide* has m 174-175°. [Dippy & Williams *J Chem Soc* 161 1934, DOI: 10.1039/JR9340000161; Campbell & McKail *J Chem Soc* 1251 1948, DOI: 10.1039/JR9480001251; Schwenk & Papa *J Org Chem* 11 798 1946, DOI: 10.1021/jo01176a023; *Beilstein* 9 III 2275.]

4-Bromophenylhydrazine [589-21-9] $C_8H_7BrO_2$, M 187.1, m 108-109°, pK^{20} -5.6 (aqueous H_2SO_4), pK^{25} 5.05. Crystallise the hydrazine from H_2O . The *hydrochloride* crystallises from EtOH/ H_2O with m 213-214°, and the *tosylate* has m 212° (from EtOH). [*Beilstein* 15 H 434, 15 I 117, 15 II 160, 15 III 289, 15 IV 282.]

4-Bromophenyl isocyanate [2493-02-9] C_7H_4BrNO , M 189.0, m 41-42°, 42-44°, b 158°/14mm. Crystallise the isocyanate from petroleum ether (b 30-40°). It has a pungent odour. [*Beilstein* 12 H 647, 12 I 321.]

4-Bromophenyl isothiocyanate [1985-12-2] C_7H_4BrNO , M 214.1, m 56-58°. Recrystallise the isothiocyanate from boiling *n*-hexane. Any insoluble material is most probably the corresponding urea. It is also purified by steam distillation, cool the receiver, add NaCl and extract in Et_2O , wash the extract with N H_2SO_4 , dry ($MgSO_4$), evaporate and recrystallise the residual solid. [Cymerman-Craig et al. *Org Synth Coll Vol* 4 700 1963, DOI: 10.15227/orgsyn.036.0056; cf. Dains et al. *Org Synth Coll Vol* 1 447 1941, DOI: 10.15227/orgsyn.006.0072; *Beilstein* 6 IV 1051, 12 II 354, 12 III 1463p, 12 IV 1519.]

N-(3-Bromopropyl)phthalimide [5460-29-7] $C_{11}H_{10}BrNO_2$, M 268.1, m 72-74°, 74°. Place it in a Soxhlet and extract it with Et_2O , whereby the bis-phthalimido impurity is not extracted. Evaporate the Et_2O and recrystallise the residue from EtOH, aqueous EtOH or petroleum ether. [Gabriel & Weiner *Chem Ber* 21 2669 1888, DOI: 10.1002/cber.18880210288; Gaudry *Can J Chem* 31 1060 1953, DOI: 10.1139/v53-139; *Beilstein* 21/10 V 1277.]

5-Bromosalicyl hydroxamic acid [5798-94-7] C_7H_6BrNO , M 210.1, m 232°(dec), $pK_{Est(1)}$ ~ 1.5, $pK_{Est(2)}$ ~ 7.0, $pK_{Est(3)}$ ~ 8.7. Crystallise the hydroxamic acid from H_2O (m 249°) or from EtOH (m 232° dec). It sublimes *in vacuo* and sublimate has m 235°. It complexes with metals and is tuberculostatic. [*Beilstein* 10 IV 221.]

4-Bromostyrene [2039-82-9] C_8H_7Br , M 183.1, b 49.5-50°/2.5mm, 87-88°/12mm, 89°/16mm, 102-104°/12mm.

20mm, d_4^{20} 1.3984, n_D^{20} 1.5925. It polymerises above 75° in the presence of benzoyl peroxide. To purify, if it has not gone to a solid resin, dissolve it in Et₂O, dry (MgSO₄) and add *ca* 0.1g of 4-*tert*-butylcatechol (polymerisation inhibitor) per 100g of bromostyrene. Filter, evaporate this under reduced pressure (use as high a vacuum as possible) and distil the residue. Store it in dark bottles in the presence of the inhibitor (at above concentration). [Overberger & Saunders *Org Synth Coll Vol* **3** 204 1955, DOI: 10.15227/orgsyn.028.0031; *Beilstein* **5** IV 1349.]

2-Bromotoluene [95-46-5] C₇H₇Br, M 171.0, m -27°, b 58-60°/10mm, 74°/19mm, 181.7°/760mm, d_4^{20} 1.422, n_D^{20} 1.556. Fractionally distil it through an efficient column. It can be separated from its isomers by gas chromatography on a column of ‘Sil-o-cel’ firebrick (30-40mesh, 80 parts) coated with 5% (20 parts) of ICI E301 con rubber with N₂ carrier gas at 170°/atm and 100ml/minute and using a conductivity cell detector. [Cowley et al. *J Chem Soc* 1799 1959, DOI: 10.1039/JR9590001799; *Beilstein* **5** H 304, **5** I 153, **5** II 234 **5** III 704, **5** IV 825.] **4-Bromotoluene** [106-38-7] has m 28°, b 184°/atm, d_4^{20} 1.390. Crystallise it from EtOH [Traylor & Stewart *J Am Chem Soc* **108** 6977 1986, DOI: 10.1021/ja00282a023]. [*Beilstein* **5** IV 827.]

α-Bromo-4-toluic acid [4-(bromomethyl)benzoic acid] [6232-88-8] C₈H₇BrO₂, M 215.0, m 224 to 229°, 229-230°, pK_{Est} ~3.2. Crystallise the acid from Me₂CO. [*Beilstein* **9** IV 1745.]

α-Bromo-*p*-xylene (4-methylbenzyl bromide) [104-81-4] C₈H₉Br, M 185.1, m 35°, 38°, b 120°/15mm, 218-220°/740mm. Crystallise the bromide from EtOH or pentane. [*Beilstein* **5** H 385, **5** IV 969.] **STRONG LACHRYMATOR.**

2-*tert*-Butoxycarbonyloxyimino-2-phenylacetoneitrile (BOC-ON) [58632-95-4] C₁₃H₁₄N₂O₃, M 246.3, m 87-89°. Triturate the solid with 90% aqueous MeOH, filter, wash with 90% aqueous MeOH and dry it in a vacuum. Recrystallise it from MeOH (needles or plates), but use warm MeOH and cool to crystallise; *do not boil as it decomposes slowly*. Its IR has ν_{\max} at 1785 (C=O) cm⁻¹ and NMR (CDCl₃) usually shows two *tert*-butyl singlets for *syn* and *anti* isomers. Store it in a brown bottle (fridge). It evolves CO₂ at room temperature (stoppered bottle can explode!), but can be stored over silica gel which may extend its useful life to more than a year. [Itoh et al. *Org Synth* **59** 95 1980, DOI: 10.15227/orgsyn.059.0095.]

4-Butoxyphenylacetic acid [4547-57-3] C₁₂H₁₆O₃, M 208.3, m 86-87°, 88.5°, pK_{Est} ~4.4. Recrystallise it from petroleum ether (b 40-60°). [McElvain & Carney *J Am Chem Soc* **68** 2592 1946, DOI: 10.1021/ja01216a051; *Beilstein* **10** IV 545.]

***n*-Butyl 4-aminobenzoate (Butamben)** [94-25-7] C₁₁H₁₅NO₂, M 193.2, m 57-59°, b 174°/8mm, pK_{Est} ~2.5. Crystallise Butamben from EtOH, or/and distil it in a high vacuum. It is poorly soluble in H₂O (0.014w/v%). It is used as a local anaesthetic. [*Beilstein* **14** IV 1130.]

***tert*-Butylammonium bromide** [60469-70-7] C₄H₁₂BrN, M 154.1, m >250°(dec). Recrystallise the salt several times from absolute EtOH or by dissolving in absolute EtOH and adding Et₂O slowly to crystallise the salt. Dry it thoroughly at 105°. [IR: Chenon & Sandorfy *Can J Chem* **36** 1181 1958, DOI: 10.1139/v58-173; *Beilstein* **4** IV 659.]

2-*tert*-Butylanthracene [13719-97-6] C₁₈H₁₈, M 234.3, m 148-149°. Recrystallise the anthracene from EtOH and finally purify it by TLC. [*Beilstein* **5** IV 2364.]

***n*-Butylbenzene** [104-51-8] C₁₀H₁₄, M 134.2, m -88°, b 22.7°/1.0mm, 48.8°/5mm, 76.3°/20mm, 102.6°/60mm, 136.9°/200mm, 183.3°/atm, d_4^{20} 0.860, n_D^{20} 1.4897, n_D^{25} 1.487. Distil butylbenzene from sodium. Wash it with small portions of conc H₂SO₄ until the acid is no longer coloured, H₂O then aqueous Na₂CO₃(care). Dry it (MgSO₄), and distil it twice from Na, collecting the middle fraction [Vogel *J Chem Soc* 607 1948, DOI: 10.1039/JR9480000607]. [*Beilstein* **5** IV 1033.] ***tert*-Butylbenzene** [98-06-6] has m -51.8°, b 13.0°/1.0mm, 39.0°/5mm, 65.5°/20mm, 90.6°/60mm, 123.7°/200mm, 169.1°/atm, d_4^{20} 0.8669, n_D^{20} 1.493, n_D^{25} 1.490. Wash it with cold concentrated H₂SO₄ until a fresh portion of acid is no longer coloured, then with 10% aqueous NaOH (care-effervescence), followed by distilled water until neutral. Dry it (CaSO₄), and distil it in a

glass helices-packed column, taking the middle fraction. [*Beilstein* 5 IV 1045.]

4-tert-Butyl benzoyl chloride [1710-98-1] $C_{11}H_{13}ClO$, M 196.7, b 135°/10mm, 149.9-150.5°/14mm, 266-268°(dec), d_4^{20} 1.082, n_D^{20} 1.536. Distil it in a vacuum. If IR shows OH group, then treat it with thionyl chloride or oxalyl chloride at ca 50° for 30 minutes, evaporate and fractionate it in a vacuum using a short column. Strongly **LACHRYMATORY**; use a good fume hood. [Fuson & Tull *J Am Chem Soc* 71 2544 1949, DOI: 10.1021/ja01175a086; Tsuno et al. *Bull Chem Soc Jpn* 32 960 1959, DOI: org/10.1246/bcsj.32.960; Swain et al. *J Am Chem Soc* 72 5426 1950, DOI: 10.1021/ja01168a017; *Beilstein* 9 III 2526.]

4-tert-Butylcatechol [98-29-3] $C_{10}H_{14}O_2$, M 166.22, m 47-48°, 55-56°, 58°, b 285°/atm, $pK_{Est(1)}$ ~9.5, $pK_{Est(2)}$ ~13.0. Distil it in a vacuum, then recrystallise it from pentane or petroleum ether (or $*C_6H_6$). [*Beilstein* 6 IV 6014.]

6-tert-Butyl-1-chloro-2-naphthol [525-27-9] $C_{14}H_{15}ClO$, M 234.7, m 76°, b 185°/15mm, pK_{Est} ~8.0. Recrystallise the naphthol from petroleum ether. Its *methyl ether* has m 115° (from EtOH/petroleum ether). [Buu-Hoi et al. *J Org Chem* 15 1060 1950, DOI: 10.1021/jo01151a024; *Beilstein* 6 IV 4367.]

2-tert-Butyl-4,5-dimethylphenol [1445-23-4] $C_{12}H_{18}O$, M 178.3, b 144-150°/20mm, pK_{Est} ~11.8. It is obtained by placing 3,4-dimethylphenol (170g, 1.39mol, m ~65°, see [95-65-8]), concentrated H_2SO_4 (1.5ml) and a magnetic stirrer bar in a medium pressure vessel, then purging the closed vessel with isobutylene while heating and stirring vigorously at 70° (which keeps the phenol in a molten state) under an isobutylene pressure of 20 psi. The reaction is complete when the liquid stops expanding. The mixture is cooled, Et_2O is added and the mixture is washed with saturated aqueous $NaHCO_3$ (3 x 150ml), the combined organic layers are dried ($MgSO_4$), filtered, the solvent is evaporated and the residual oil is distilled to give 2-tert-butyl-4,5-dimethylphenol, and its 1H NMR (400MHz, $CDCl_3$, TMS) has δ at 7.02 (s, 1H, aromatic CH), 6.49 (s, 1H, aromatic CH), 2.19 (s, 3H, CH_3), 2.17 (s, 3H, CH_3), 1.40 (s, 9H, $C(CH_3)_3$). *Note tert-butyl 3,4-dimethylphenyl ether is formed instead when lower temperatures and very small amounts of acid are used* [Stevens *J Org Chem* 20 1232 1955, DOI: 10.1021/jo01126a010; Alexander et al. *J Am Chem Soc* 120 4041 1998, DOI: 10.1021/ja974353i; Albert *J Am Chem Soc* 76 4983 1954, DOI: 10.1021/ja01648a068; for acetyl derivative see Fischer & Teo *Can J Chem* 56 258 1978, DOI: 10.1139/v78-041.]

2-tert-Butyl hydroquinone [1948-33-0] $C_{10}H_{18}O$, M 166.2, m 125-127°, 127-128°, 129°, $pK_{Est(1)}$ ~10.5, $pK_{Est(2)}$ ~11.6. Recrystallise the hydroquinone from H_2O or MeOH and dry it in a vacuum at 70°. Store it in a dark container. [Stroh et al. *Angew Chem* 69 699 1957, DOI: 10.1002/ange.19570692202; *Beilstein* 6 IV 6013.]

2-tert-Butyl-4-methoxyphenol (2-tert-butyl-4-hydroxyanisole, BHA) [121-00-6] $C_{11}H_{16}O$, M 180.3, m 64.1°, pK_{Est} ~10.8. Fractionally distil the phenol *in vacuo*, then pass it as a solution in $CHCl_3$ through alumina, and evaporate the eluate. Recrystallise the residue from petroleum ether. [*Beilstein* 6 IV 6013.]

4-tert-Butylnitrobenzene [3282-56-2] $C_{10}H_{13}NO_2$, M 179.2, m 28.4°, b 135°/10mm, 140-142°/15mm, n_D^{20} 1.5230. Recrystallise it three times by partially freezing a mixture of the mono-nitro isomers, then recrystallise it twice from MeOH and dry it *in vacuo* [Brown *J Am Chem Soc* 81 3232 1959, DOI: 10.1021/ja01522a018]. [*Beilstein* 5 H 418, 5 I 203, 5 II 321, 5 III 943, 5 IV 1052.]

tert-Butyl perphthalic acid (monoperoxyphthalic acid 1-tert-butyl ester) [15042-77-0] $C_{12}H_{14}O_5$, M 238.2, m 104-104.5° (dec), pK_{Est} ~6.2. Crystallise the per acid-ester from Et_2O or Et_2O /petroleum ether and dry it over H_2SO_4 . The ester was prepared from tert-butylhydroperoxide and phthalic anhydride [Davies et al. *J Chem Soc* 1541 1953, DOI: 10.1039/JR9530001541]. Possibly **EXPLOSIVE**. [*Beilstein* 9 IV 3260.]

4-tert-Butylphenol [98-54-4] $C_{10}H_{14}O$, M 150.2, m 96 to 101°, 98°, 99°, b 236-238°/atm, pK^{25} 10.39. Crystallise the phenol to constant melting point from petroleum ether (b 60-80°). It sublimes *in vacuo*. Also purify it *via* the benzoate, as for phenol. The *salicylate ester* [87-18-30] has m 63-64° (from aqueous EtOH, or EtOH). [*Beilstein* 6 IV 3296.]

4-*tert*-Butylphenoxyacetic acid [1798-04-5] $C_{12}H_{16}O_3$, M 208.3, m 86.5°, 88-89°, 94°, 96.5°, $pK_{Est} \sim 2.9$. Crystallise the acid from petroleum ether or petroleum ether/* C_6H_6 mixture. [Beilstein 6 H 524, 6 III 1869.]

***tert*-Butyl phenyl carbonate** [6627-89-0] $C_{11}H_{14}O_3$, M 194.2, b 74-78°/0.5mm, 83°/0.6mm, d_4^{20} 1.05, n_D^{20} 1.480. If IR is free from OH, then purify it by redistillation; otherwise dissolve it in Et_2O , wash it with 5% HCl, then H_2O , dry it ($MgSO_4$), evaporate and distil it through a Claisen head under vacuum. Care should be taken as distillation of large quantities can lead to decomposition with liberation of CO_2 and isobutylene; **take the necessary precautions**. Used for mono-Boc protection of some α,ω -diamines [Pittelkow et al. *Org Synth* 84 209 2007, DOI: 10.15227/orgsyn.084.0209]. [Carpino *J Am Chem Soc* 79 98 1957, DOI: 10.1021/ja01558a026; Beilstein 6 IV 629.]

***n*-Butyl phenyl ether** [1126-79-0] $C_{10}H_{14}O$, M 150.2, b 95°/17mm, 210.20°/760mm, d_4^{20} 0.935, n_D^{20} 1.4969. Dissolve it in diethyl ether, wash first with 10% aqueous NaOH to remove traces of phenol, then repeatedly with distilled water, followed by evaporation of solvent and distillation under reduced pressure [Arnett & Wu *J Am Chem Soc* 82 5660 1960, DOI: 10.1021/ja01506a027]. [Beilstein 6 H 143, 6 I 82, 6 II 145, 6 III 550, 6 IV 558.]

***N*-*tert*-Butyl α -phenyl nitron (PBN)** [3376-24-7] $C_{11}H_{15}NO$, M 177.2, m 73-74°. Crystallise PBN from hexane. It is a free radical trap. Its solubility in H_2O is 2.9% and slightly less so in aqueous saline (2.3%), but more soluble in organic solvents. Its UV has λ_{max} in EtOH at 224 (ϵ 7,240) and 293.5 (ϵ 17,700) nm; and in MeCN at 298 (ϵ 17,2716) nm. [cf. Janzen *Methods Enzymology* 105 188 1984, DOI: 10.1016/S0076-6879(84)05025-4; Beilstein 7 IV 519.]

4-*tert*-Butyltoluene [98-51-1] $C_{11}H_{16}$, M 148.2, m -53.2°, b 91°/28mm, 189-192°/atm, d_4^{20} 0.854, n_D^{20} 1.4920. A sample containing 5% of the *meta*-isomer is purified by selective mercuration. Fractional distillation of the solid arylmercuric acetate, after removal from the residual hydrocarbon, gives pure *p*-*tert*-butyltoluene [Stock & Brown *J Am Chem Soc* 81 5615 1959, DOI: 10.1021/ja01530a024]. [Beilstein 5 H 439, 5 III 1003, 5 IV 1079.]

***tert*-Butyl 2,4,5-trichlorophenyl carbonate** [16965-08-5] $C_{11}H_{11}Cl_3O_3$, M 297.6, m 64-66°, 67-68.5°. Crystallise the carbonate from a mixture of MeOH (90ml) and water (6ml) using charcoal [Broadbent et al. *J Chem Soc (C)* 2632 1967, DOI: 10.1039/J39670002632; Fieser 2 55].

Caffeic acid (3,4-dihydroxycinnamic acid) [331-39-5] $C_9H_8O_4$, M 180.2, m 195°, 211 to 213°(dec) 223-225° (softens at ca 195°), pK_1^{25} 4.62, pK_2^{25} 9.07. Recrystallise this antioxidant from strong aqueous solutions as yellow crystals, but forms a *monohydrate* from dilute solutions. Yellow alkaline solutions become orange probably because of oxidation to quinones. It is much more soluble in hot than in cold H_2O , and its solubility in EtOH is 5w/v%. The **methyl ester** $C_{10}H_{10}O_4$, m 152-153°, forms colourless needles from H_2O . It was found in green coffee and is an inhibitor of leukotrienes. [Herrmann *Pharmazie* 11 433 1956, PMID: 13370323; Beilstein 10 IV 1776.]

Capsaicin (*E*-*N*-[(4-hydroxy-3-methoxyphenyl)-methyl]-8-methyl-6-nonenamide) [404-86-4] $C_{18}H_{27}NO_3$, M 305.4, m 64-66°, 65°, 66.1°, b 210-220°/0.01mm, ϵ 7000 (281nm, EtOH). Recrystallise capsaicin from petroleum ether (b 40-60°), or petroleum ether/ Et_2O (9:1). Also purify it by chromatography on neutral Al_2O_3 (grade V) and elute successively with * C_6H_6 , * C_6H_6 /EtOAc (17:3) then * C_6H_6 /EtOAc (7:3), and distil it at 120°/10⁻⁵mm, then repeatedly recrystallise the needles from isopropanol (charcoal). [Crombie et al. *J Chem Soc* 1007 1955, DOI: 10.1039/JR9550001007; Bennett & Kirby *J Chem Soc(C)* 442 1968, DOI: 10.1039/J39680000442.] It is the active principle in cayenne pepper; and a burning taste is experienced at concentrations as low as 1 in 100,000. It **causes pain**, used in Police work as a deterrent but is **neurotoxic**. **STRONG IRRITANT**. [Bevan & Szolcsanyi *Trends in Pharmacol Sci* 11 331 1990, DOI:10.1016/0165-6147(90)90237-3; Beilstein 13 IV 2588].

4-(Carbamoylmethoxy)acetanilide [14260-41-4] $C_{10}H_{12}N_2O_3$, M 208.2, m 208°. Crystallise the anilide from water.

N-Carboethoxyphthalimide (N-ethoxycarbonylphthalimide) [22509-74-6] $C_{10}H_9NO_4$, M 219.2, m 87-89°, 90-92°. Crystallise the imide from toluene/petroleum ether (or *benzene/petroleum ether). It is partly soluble in Et_2O , *benzene and $CHCl_3$. [Heller & Jacobsohn *Chem Ber* **54** 1107 1921, DOI: 10.1002/cber.19210540529; *Beilstein* **21/10** V 428.]

3-Carboxybenzaldehyde (3-formylbenzoic acid, isophthalaldehydic acid) [619-21-6] $C_8H_6O_3$, M 150.1, m 173°, 173-175°, pK^{25} 3.84. The acid was prepared in several ways including the following two. *m*-Bromomethylbenzoic acid (3g) is allowed to react with hexamine (3.9g) in $CHCl_3$ (30ml) by refluxing for 3 hours, cooled, the hexamine quaternary salt (4.7g) is collected, dried *in vacuo*, and decomposed by boiling in 50% aqueous acetic acid for 2 hours. The solution is cooled, diluted with an equal volume of H_2O and chilled to give a crop of the desired acid as needles, which are washed with aqueous $NaHCO_3$ and dried. A further crop can be obtained by extracting the acidic filtrate with Et_2O to give total yield of 48%. [Dyer et al. *J Chem Soc* 4778, 4781 1952, DOI: 10.1039/JR9520004778]. Alternatively, *m*-carboxycinnamic acid (m 268-270°, 8.6g) in 1N Na_2CO_3 (50ml) and H_2O (400ml) is treated slowly at 4-5° with 5% of aqueous $KMnO_4$ (80ml) with vigorous stirring. MnO_2 is removed by filtration, the filtrate is concentrated to 150ml, acidified with 5N HCl and the mixture of acids that separated, together with the solution, are extracted thoroughly with Et_2O , which on evaporation give 3-formylbenzoic acid (3.6g) which is purified by recrystallisation from H_2O . [Berner *Acta Chem Scand* **10** 1208 1956, DOI: 10.1016/0165-6147(90)90237-3]. The *acid chloride* [75650-38-3] has **b** 130°/20mm, the *methyl ester* [52178-50-4] has **m** 53°, the *ethyl ester* [33745-47-0] has **d**¹⁸ 1.093 and **b** 278°/atm, the *amide* has **m** 190°(dec), the *oxime* has **m** 188°, the *semicarbazone* has **m** 275°(dec) (from aqueous $EtOH$), the *bis(2-hydroxyethyl)amine salt* has **m** 167° (yellow crystals from aqueous $EtOH$), and the *phenylhydrazone* has **m** 164°. [Irreverre et al. *J Biol Chem* **236** 1093 1961, Davies et al. *J Chem Soc* 2202 1922, DOI: 10.1039/CT9222102202; Simonis *Chem Ber* **45** 1584 1912, DOI: 10.1002/cber.19120450224; *Beilstein* **10** H 671, **10** I 317, **10** II 465, **10** III 2990, **10** IV 2750.]

4-Carboxybenzaldehyde (4-formylbenzoic acid, terephthalaldehydic acid) [619-66-9] $C_8H_6O_3$, M 150.1, m 247°, 248°(dec), ~250°, 248-250°, 256°, pK^{25} 3.77. Of the many syntheses of this acid, the more convenient one is by refluxing a solution of the commercially available 4-bromomethylbenzoic acid (21.0g, 6232-88-8) with 10% $Cu(NO_2)_2$ (300ml) for 6 hours, cooling, filtering off the solid and recrystallising from glacial $AcOH$ (100ml) to give fine plates, which on further recrystallisation from H_2O , provide pure 4-formylbenzoic acid as fine needles (2.2g 56%). The *phenylhydrazone* has **m** 221-222° [Stewart & Walker *Can J Chem* **35** 1561 1957, DOI: 10.1139/v57-204], the *2,4-dinitrophenylhydrazone* has **m** 319.5-320.5°, after crystallisation from pyridine then $PhNO_2$ [Bowen & Wilkinson *J Chem Soc* 747 p 750 1950, DOI: 10.1039/JR9500000747], the *oxime* has **m** 216-217° [Wheeler et al. *J Org Chem* **22** 547 1957, DOI: 10.1021/jo01356a022]. The acid is also obtained by hydrolysis of the *methyl ester*, **m** 61-61° (from petroleum ether), **b** 135°/12mm, by aqueous H_2SO_4 ; the ester having been prepared from methyl 4-cyanobenzoate by reaction with $ZnCl_2/HCl$ [Slotta & Kethur *Chem Ber* **71** 335 1938, DOI: 10.1002/cber.19380710224]. The UV in hexane of the *formyl acid* has λ_{max} (ϵ) at 249 (17,500), 257 (15,500) [B-band], and 279 (1,650), 288 (1,950), 298 (1,600) [C-band] nm [Dearden & Forbes *Can J Chem* **36** 1362 1958, DOI: 10.1139/v58-201]. [for pK see Humffray et al. *JCS Chem Comm* 610 1965, DOI: 10.1039/C19650000610; *Beilstein* **10** H 671, **10** I 317, **10** II 465, **10** III 2989, **10** IV 2752.]

4-Carboxyphenylacetonitrile [6627-91-4] $C_9H_7NO_2$, M 161.2, m 114-115°. Crystallise the nitrile (with considerable loss) from *benzene, glacial acetic acid or H_2O . The *methyl ester* has **m** 47-48° (from C_6H_6). [Price & Rogers *Org Synth Coll Vol* **3** 174 1955, DOI: 10.15227/orgsyn.022.0030; *Beilstein* **9** H 859, **9** II 618, **9** III 4267.]

Catechol (1,2-dihydroxybenzene, pyrocatechol) [120-80-9] $C_6H_6O_2$, M 110.1, m 100-103°, 105°, b 104°/5mm, b 134°/20mm, b 161°/60mm, 197.7°/100m, 245.5°/atm, **d**²⁵ 1.344, pK_1^{25} 9.45, pK_2^{25} 12.8. Crystallise catechol from *benzene or toluene (prisms) and sublime it *in vacuo*. It darkens in air, but more rapidly in aqueous solution. It is steam volatile with solubility in H_2O of ~43.5w/v%. It is an antiseptic. [Rojo et al. *Anal Chem* **58** 2988 1986, DOI: 10.1021/ac00127a020; *Beilstein* **6** IV 5557.]

Cation exchange resin. The resin should be conditioned before use by successive washing with water, $EtOH$ and water, and taken through two $H^+-Na^+-H^+$ cycles by successive use of M NaOH, water and M HCl then

washed with water until neutral.

Chamazulene (1,4-dimethyl-7-ethylazulene, see also Guaiazulene below) [529-05-5] $C_{14}H_{16}$, **M 184.3**, **b 145°/11mm, 161°/12mm, d_4^{20} 0.9883**. The blue oil occurs in flower oil of chamomile, *Artemisia. Matricaria* and tansy. It can be purified by chromatography on Al_2O_3 (Basic I) and eluting the blue band with petroleum ether. The hydrocarbon is then redistilled at high vacuum. Its UV has λ_{max} at 370nm (log ϵ 3.7, heptane). The **trinitrobenzene complex**, crystallises as violet-black needles from absolute EtOH with **m 131.5-132.5°**. The **picrate** has **m 115-116°** (from EtOH), and the **styphnate** has **m 92-93°** (from EtOH). [Ruzicka & Rudolph *Helv Chim Acta* **9** 118 1926, DOI: 10.1002/hlca.19260090114; Meisels & Weizmann *J Am Chem Soc* **75** 3865 1953, DOI: 10.1021/ja01111a531; Mukherjee, Dunn and Houk *J Am Chem Soc* **101** 251 1979, DOI: 10.1021/ja00495a058; *Beilstein* **5** III 1677, **5** IV 1751.]

p-Chloranil (2,3,5,6-tetrachloro-1,4-benzoquinone) [118-75-2] $C_6Cl_6O_2$, **M 245.9**, **m 289°(dec), 290°, 294.2-294.6°(sealed tube)**. Crystallise *p*-chloranil from acetic acid, acetone, *benzene, EtOH or toluene, dry it in a vacuum over P_2O_5 , or from acetic acid and drying over NaOH in a vacuum desiccator. It can be sublimed under vacuum at 290°. A sample may contain significant amounts of the *o*-chloranil isomer as impurity. Purify it by triple sublimation under vacuum and recrystallise before use. It is soluble in $CHCl_3$, CCl_4 and CS_2 . **It is a skin and mucous membrane irritant, and is TOXIC**. It has been used as a fungicide. [UV: Pummerer et al. *Chem Ber* **85** 535 1952, DOI: 10.1002/cber.19520850608; Brook *J Chem Soc* 5035 p 5040 1952, DOI: 10.1039/JR9520005035; *Beilstein* **7** IV 2083.]

Chloranilic acid (2,5-dichloro-3,6-dihydroxy-1,4-benzoquinone) [87-88-7] $C_6H_2Cl_2O_4$, **M 209.0**, **m 283-284°, ~300°, pK_1^{25} 1.22, pK_2^{25} 3.01**. A solution of 8g of quinone in 1L of boiling water is filtered while hot, then extracted twice at about 50° with 200ml portions of *benzene. The aqueous phase is cooled in ice-water. The red crystals are filtered off, washed with three 10ml portions of water, and dried at 115°. It can be sublimed *in vacuo*. [Weissbart & Rysselberghe *J Phys Chem* **61** 765 1957, DOI: 10.1021/j150552a015.] The **diacetate** has **m 182-185°** [Conant & Fieser *J Am Chem Soc* **46** 1858 1924, DOI: 10.1021/ja01673a014; Thamer & Voigt *J Phys Chem* **56** 225 1952, DOI: 10.1021/j150494a018]. [*Beilstein* **8** IV 2707.] It is a moderately strong acid and forms salts. The **barium salt** [32458-20-1] $C_6BaCl_2O_4 \cdot 3H_2O$, **M 398.3**, is used for sulfate determination.

4-Chloroacetanilide [539-03-7] C_8H_8ClNO , **M 169.6**, **m 176 to 178°, 179°**. Crystallise the anilide from EtOH or aqueous EtOH. [*Beilstein* **12** IV 1178.]

ω -Chloroacetophenone (phenacyl chloride) [532-27-4] C_8H_7ClO , **M 154.6**, **m 54 to 56°, 58-59°, b 244-245°/atm, d^{15} 1.324**. Crystallise it from MeOH, aqueous EtOH, CCl_4 , or petroleum ether. It is a skin, eyes and respiratory irritant — take all necessary precautions when preparing or using it. [Tanner et al. *J Org Chem* **52** 2142 1987, DOI: 10.1021/jo00387a005; *Beilstein* **7** IV 641.]

***o*-Chloroaniline** [95-51-2] C_6H_7ClN , **M 127.6**, **m -1.9°, b 84.5°/10mm, 108.4°/30mm, 208.8°/760mm, d_4^{20} 1.213, n_D^{20} 1.588, pK^{25} 2.66**. Free it from small amounts of the *p*-isomer by dissolving in one equivalent of H_2SO_4 and steam distilling. The *p*-isomer remains behind as the sulfate. [Sidgwick & Rubie *J Chem Soc* 1013 1921, DOI: 10.1039/CT9211901013.] An alternative method is to dissolve it in warm 10% HCl (11ml/g of amine) and on cooling, ***o*-chloroaniline hydrochloride** separates out. The latter can be recrystallised until the acetyl derivative has a constant melting point (**m 90°**). (In this way, yields are better than *via* the recrystallisation of the **picrate** from EtOH or of the acetyl derivative from petroleum ether.) [King & Orton *J Chem Soc* 1377 1911, DOI: 10.1039/CT9119901377]. [*Beilstein* **12** III 1281, **12** IV 1115.] ***p*-Chloroaniline** [106-47-8] has **m 70-71°, b 106.8-107.3-5°/12mm, 116°/17mm, d_4^{20} 1.175, pK^{25} 3.98**. Crystallise the aniline from MeOH, petroleum ether (b 30-60°), or 50% aqueous EtOH, then *benzene/petroleum ether (b 60-70°), and then dry it in a vacuum desiccator. It can be distilled under vacuum (**b 75-77°/3mm**). It sublimes in a very high vacuum. The **acetate** crystallises from aqueous MeOH (**m 178°, 180°**) or EtOH or AcOH (**m 173-174°**) and has **b 331.3°/760mm**. [*Beilstein* **12** III 1325, **12** IV 1116.]

***p*-Chloroanisole** [623-12-1] C_7H_7ClO , **M 142.6**, **b 79°/11.5mm, 196.6°/760mm, d_4^{20} 1.164, $n_D^{25.5}$ 1.5326**. Wash the anisole with 10% (by volume) of aqueous H_2SO_4 (three times), 10% aqueous KOH (three times), and

then with water until neutral. Dry it (MgSO_4), and fractionally distil it from CaH_2 through a glass helices-packed column under reduced pressure. [Beilstein 16 IV 822.]

1-Chloroanthracene [4985-70-0] $\text{C}_{14}\text{H}_9\text{Cl}$, M 212.9, has m 79°, 80.5-81.5°, 81°, 81-82°, 83.5°. It has been prepared by Zn/NH_3 reduction of 1-chloroanthraquinone and purified by recrystallisation from AcOH (leaflets). It is soluble in EtOH and Et_2O . Store it in the dark as it polymerises in the presence of light. Its UV (CHCl_3) has bands at λ_{max} nm(ϵ) 255.5 (153,000), 328(1,900), 342(3,300), 360(5,750) and 380(4,900). The *picrate* has m 101-102°. [Gore *J Chem Soc* 1616 1959, DOI: 10.1039/JR9590001613; Richardson et al. *J Org Chem* 21 890 1956, DOI: 10.1021/jo01114a018; Camenzind & Rickborn *J Org Chem* 51 1914 1986, DOI: 10.1021/jo00360a054; Moursounidis & Wege *Aust J Chem* 41 235 1988, DOI: 10.1071/CH9880235; UV: Ohno & Iguchi *Bull Chem Soc Japan* 41 2264 1968, DOI: org/10.1246/bcsj.41.2246; Beilstein 5 I 324, 5 II 574, 5 III 2133, 5 IV 2292.] **2-Chloroanthracene** [17135-78-3] has m 215°, 215-217°, 217-218°, 223°. It has been prepared by Zn/NH_3 reduction of 2-chloroanthraquinone and purified by chromatography on Al_2O_3 (with $^*\text{C}_6\text{H}_6$) and followed by recrystallisation from AcOH (needles or leaflets). A solution in $^*\text{C}_6\text{H}_6$ exhibits blue fluorescence. It has UV (CHCl_3) bands at λ_{max} nm(ϵ) 257 (105,000), 348(4,500), 365(6,400) and 384(5,750). [Gore *J Chem Soc* 1616 1959, DOI: 10.1039/JR9590001613, Richardson et al. *J Org Chem* 21 890 1956, DOI: 10.1021/jo01114a018; UV: Ohno & Iguchi *Bull Chem Soc Japan* 41 2264 1968, DOI: org/10.1246/bcsj.41.2246; Beilstein 5 I 324, 5 II 575, 5 III 2132, 5 IV 2292.] **9-Chloroanthracene** [716-53-0] has m 104-106°, 105-107°. 9-Chloroanthracene crystallises from EtOH or petroleum ether (b 60-80°) as yellow needles. [Nonhebel *Org Synth Coll Vol* 5 206 1973, Masnovi *J Am Chem Soc* 108 1126 1986, DOI: 10.1021/ja00266a003; UV: Ohno & Iguchi *Bull Chem Soc Japan* 41 2264 1968, DOI: org/10.1246/bcsj.41.2246; Beilstein 5 H 663, 5 III 2133, 5 IV 2292.]

10-Chloro-9-anthraldehyde [10527-16-9] $\text{C}_{15}\text{H}_9\text{ClO}$, M 240.7, m 218°, 217-219°. The aldehyde crystallises as yellow needles from EtOH, AcOH or toluene. [Beilstein 7 III 2529.]

2-Chlorobenzaldehyde [89-98-5] $\text{C}_7\text{H}_5\text{ClO}$, M 140.6, m 11°, b 209-215°/atm, 213-214°/atm, d_4^{20} 1.248, n_D^{20} 1.566. Wash it with 10% Na_2CO_3 solution, then fractionally distil it in the presence of a small amount of catechol as stabiliser. [Beilstein 7 H 233, 7 IV 561.] **3-Chlorobenzaldehyde** [587-04-2] has m 18°, b 213-214°/atm, d_4^{20} 1.241, n_D^{20} 1.564. Purify it by low temperature crystallisation from petroleum ether (b 40-60°) and distillation. [Beilstein 7 H 234, 7 IV 566.] **4-Chlorobenzaldehyde** [104-88-1] has m 45°, 47°, 50°, b 213-214°/atm. Crystallise it from EtOH/water (3:1), then sublime it twice at $\sim 50^\circ/2\text{mm}$. [Beilstein 7 H 235, 7 IV 568.]

Chlorobenzene [108-90-7] $\text{C}_6\text{H}_5\text{Cl}$, M 112.6, m -45°, b 131.7°/atm, 131-132°/atm, d_4^{20} 1.107, n_D^{20} 1.52480. The main impurities are likely to be chlorinated impurities originally present in the $^*\text{benzene}$ used in the synthesis of chlorobenzene, and also unchlorinated hydrocarbons. A common purification procedure is to wash it several times with concentrated H_2SO_4 then with aqueous NaHCO_3 or Na_2CO_3 , and water, followed by drying with CaCl_2 , K_2CO_3 or CaSO_4 , then with P_2O_5 , and distilling. It can also be dried with Linde 4A molecular sieve. Passage through, and storage over, activated alumina has been used to obtain low conductance material. TOXIC vapour, causes CNS depression. [Flaherty & Stern *J Am Chem Soc* 80 1034 1958, DOI: 10.1021/ja01538a004; Beilstein 5 H 199, 5 IV 640.]

2-Chlorobenzoic acid [118-91-2] $\text{C}_7\text{H}_5\text{ClO}_2$, M 156.6, m 139° to 140°, 142°, d_4^{25} 1.544, pK^{25} 2.91. Crystallise the acid successively from glacial acetic acid, aqueous EtOH, and petroleum ether (b 60-80°). Other solvents include hot water (0.9w/v%, cold) or toluene (ca 25w/v%). The crude material can be initially purified by dissolving 30g in 100ml of hot water containing 10g of Na_2CO_3 , boiling with 5g of charcoal for 15 minutes, then filtering and adding 31ml of 1:1 aqueous HCl. The precipitate is washed with a little water and dried at 100°. [Clarke & Taylor *Org Synth Coll Vol* 2 135 1943, DOI: 10.15227/orgsyn.010.0020; Beilstein 9 IV 956.] **3-Chlorobenzoic acid** [535-80-8] has m 154-156°, 158°, d_4^{25} 1.496, pK^{25} 3.82 (5.25 in 50% dimethylacetamide). Crystallise the acid successively from glacial acetic acid, aqueous EtOH and petroleum ether (b 60-80°). Its solubility in cold H_2O is 0.035w/v% but is more soluble at 100°. It also recrystallises from $^*\text{C}_6\text{H}_6$ or $\text{Et}_2\text{O}/\text{hexane}$, and sublimates at 55° *in vacuo*. [Vandenbelt et al. *Anal Chem* 26 726 1954, DOI: 10.1021/ac60088a031.] The *methyl ester* has m 21°, b 231°/760mm. The *S-benzylisothiuronium salt* has

m 164-165° (from EtOH) [Friediger & Pedersen *Acta Chem Scand* **9** 1425 1955, DOI: 10.3891/acta.chem.scand.09-1425; Samuel *J Chem Soc* 1318 1960, DOI: 10.1039/JR9600001318]. [*Beilstein* **9** IV 969.] **4-Chlorobenzoic acid** [74-11-3] has **m 238° to 241°, 243°, pK²⁵ 3.99**. Purify it as for *m*-chlorobenzoic acid. It has also been crystallised from hot water (solubility in cold H₂O is 0.019w/v%), and from EtOH. The **methyl ester** has **m 44°** (from aqueous MeOH). [*Beilstein* **9** IV 973.] **4-Chlorobenzhydrazide** [536-40-3] **C₇H₇ClN₂O**, **M 170.6**, has **m 162°, 164°, 165°**. It was prepared from the acid chloride and hydrazine hydrate (with cooling). Recrystallise it from H₂O. [*Beilstein* **9** III 1368.]

2-Chlorobenzonitrile [873-32-5] **C₇H₄ClN**, **M 137.6**, **m 43°, 45-46°, b 232°/atm**. Crystallise the nitrile to a constant melting point from *benzene/petroleum ether (b 40-60°), and/or distil it. [*Beilstein* **9** IV 965.]

4-Chlorobenzophenone [134-85-0] **C₁₃H₉ClO**, **M 216.7**, **m 74°, 75-76°, b 195-196°/17mm**. Recrystallise it from EtOH, and/or distil it under vacuum. [Wagner et al. *J Am Chem Soc* **108** 7727 1986, DOI: 10.1021/ja00284a041; *Beilstein* **7** H 419, **7** I 227, **7** II 359, **7** III 2072, **7** IV 1375.]

4-Chlorobenzotrifluoride (*o*-chlorotrifluoromethylbenene) [88-16-4] **C₇H₄ClF₃**, **M 180.6**, **m -6.37°, b 19.6°/3mm, 152.3°/760mm, d₄²⁵ 1.364, n_D²⁵ 1.4533**. Dry the trifluoride over CaSO₄, and distil it at high reflux ratio through a silvered vacuum-jacketed glass column packed with one-eighth inch glass helices [Potter & Saylor *J Am Chem Soc* **73** 90 1951, DOI: 10.1021/ja01145a032]. [*Beilstein* **5** H 302, **5** III 692, **5** IV 814.] **3-Chlorobenzotrifluoride** [98-15-7] has **m -56.49°, b 50°/31mm, 137.6°/760mm, d₄²⁰ 1.3345, n_D²⁰ 1.4432**. Purify it as for *o*-chlorobenzotrifluoride above. [*Beilstein* **5** III 692, **5** IV 814.] **4-Chlorobenzotrifluoride** [98-56-6] has **m -33.18°, b 19.3°/5mm, 138.6°/760mm, d₄³⁰ 1.3278, n_D³⁰ 1.4430**. Purify it as for *o*-chlorobenzotrifluoride above. It is useful as a dielectric fluid. [*Beilstein* **5** IV 815.]

4-Chlorobenzyl chloride [104-83-6] **C₇H₆Cl₂**, **M 161.0**, **m 28-29°, b 96°/15mm, 216-222°/atm**. Dry it over CaSO₄, then fractionally distil it under reduced pressure. Crystallise it from heptane or dry diethyl ether at low temperature. [*Beilstein* **5** IV 816.] **LACHRYMATORY**.

trans-4-Chlorocinnamic acid [1615-02-7] **C₉H₇ClO₂**, **M 182.6**, **m 243°, 248-250°, 249-251°, pK²⁵ 4.41**. Recrystallise the acid from EtOH or aqueous EtOH (charcoal). Its UV has λ_{max} at 275nm (EtOH). [Walling & Wolfstirn *J Am Chem Soc* **69** 852 1947, DOI: 10.1021/ja01196a033; *Beilstein* **9** H 596, **9** II 395, **9** III 2727, **9** IV 2033.]

4-Chloro-3,5-dimethylphenol [88-04-0] **C₈H₉ClO**, **M 156.6**, **m 114°, 115.5°, 116°, b 246°/atm, pK²⁵ 9.70**. Crystallise the phenol from *benzene or toluene. It is steam volatile, has low solubility in cold H₂O (0.03w/v%) but is much more soluble in EtOH (50%). It is antibacterial and a strong antiseptic and germicide. [*Beilstein* **6** IV 3152.]

1-Chloro-2,4-dinitrobenzene [97-00-7] **C₆H₃ClN₂O₄**, **M 202.6**, **m 48-50°, 51°, 52-54°, 54°, b 315°/atm, d₄²⁰ 1.697**. Usually it is recrystallised from EtOH or MeOH. It has also been crystallised from Et₂O, *C₆H₆, *C₆H₆/petroleum ether or isopropyl alcohol. A preliminary purification step is to pass its solution in *benzene through an alumina column. It has also been purified by zone refining. It exists in three forms: one stable and two unstable. The stable form crystallises as yellow needles from Et₂O, **m 51°, b 315°/760mm** with some decomposition, and is soluble in EtOH. A labile form also crystallises from Et₂O, **m 43°**, and is more soluble in organic solvents. The second labile form has **m 27°**. [Hoffman & Dame, *J Am Chem Soc* **41** 1013 1919, DOI: 10.1021/ja02227a012; Welsh *J Am Chem Soc* **63** 3276 1941, DOI: 10.1021/ja01857a014; Littlejohn & Smith *J Chem Soc* 2476 1957, DOI: 10.1039/JR9570002476; *Beilstein* **5** IV 744.]

4-Chloro-3,5-dinitrobenzoic acid [118-97-8] **C₇H₃ClN₂O₆**, **M 246.6**, **m 159-161°, 163°, pK_{Est} ~2.5**. Crystallise the acid from EtOH/H₂O, EtOH or *C₆H₆. The 1:1 **naphthalene complex** (by fusing various ratios of ingredients and recrystallising from EtOH) has **m 122°**. [*Beilstein* **9** H 416, **9** III 1953, **9** IV 1360.]

Chlorogenic [1-(3,4-dihydroxycinnamoyloxy)-D-quinic] acid [327-97-9] **C₁₆H₁₈O₉**, **M 354.3**, **m 208°, [α]_D²⁵ -36 (c 1, H₂O), pK₁²⁵ 3.59, pK₅²⁵ 8.59**. Recrystallise the acid from water (solubility is 4% at room temp-

erature), and dry it at 110°. It gives a yellow solution in aqueous base and complexes with Fe. [Beilstein 10 H 537, 10 I 271, 10 II 378, 10 III 2408, 10 IV 2259.]

Chlorohydroquinone (2-chloro-1,4-dihydroxybenzene) [615-67-8] $C_6H_5ClO_2$, M 144.6, m 100°, 104°, 106°, b 263°/atm, pK_1^{25} 8.81, pK_2^{25} 10.78. Crystallise the hydroquinone from $CHCl_3$ or toluene. [Beilstein 6 IV 5767.]

1-Chloro-4-iodobenzene [637-87-6] C_6H_4ClI , M 238.5, m 53-54°, 56.2°, b 104.2°/16mm, 226-227°/atm, d_4^{25} 1.886. Distil it in a vacuum then recrystallise it from EtOH. [Sugden *J Chem Soc* 1167 1924, DOI: 10.1039/CT9242501167; Beilstein 5 H 221, 5 III 579, 5 IV 695.]

5-Chloro-2-methoxyaniline (2-amino-4-chloroanisole, 5-chloro-o-anisidine) [95-03-4] C_7H_8ClNO , M 157.6, m 81-83°, 82-84°, 84°, pK^{25} 3.56. Purify the aniline by steam distillation and recrystallisation from H_2O or 40% aqueous EtOH. The *N*-acetate forms needles from hot H_2O with m 104°, the *N*-benzoyl derivative forms needles from aqueous EtOH with m 77-78°, and the *picrate* has m 194°(dec). [Raiford & Colbert *J Am Chem Soc* 48 2652 1926, DOI: 10.1021/ja01421a022; Beilstein 13 IV 879.]

3-Chloro-4-methoxyphenethylamine [7569-87-1] $C_9H_{12}ClNO$, M 185.7, b 140°/0.6mm, d_4^{25} ~1.081, n_D^{20} ~1.553, pK^{25} ~9.8. This strong base is prepared by reduction of *3-chloro-4-methoxyphenylacetone nitrile* (176mmol, b 140-145°/0.1-0.1mm, m 55-56°, [7569-58-6]) in THF (100ml) with $LiAlH_4$ (8.0g, 210mmol) suspended in THF (250ml) under reflux for 5 hours. The greenish coloured solution is carefully decomposed with ice cold H_2O , the solids are filtered off, washed with Et_2O , the combined organic liquids are dried (Na_2SO_4), filtered, evaporated and the crude brown residual amine is distilled in as high a vacuum as possible to give a clear oil (11.4g, 35%). It absorbs CO_2 from air and is best stored in an inert atmosphere. Its 1H NMR [60MHz, $CDCl_3$] has δ at 7.20 (d, 1H, ArH2), 7.04 (q, 1H, ArH6), 6.83 (d, 1H, ArH5), 3.88 (s, 3H, OCH_3), 3.12-2.20 (m, 4H, $PhCH_2CH_2N$), 1.28 (s, 2H, NH_2) ppm from TMS [Charifson et al. *J Med Chem* 31 1941 1988, DOI: 10.1021/jm00118a012]. Alternatively, the amine is prepared from the respective phenethyl bromide and dry NH_3 in EtOH followed by dilution with Et_2O , washing with aqueous NaOH, and evaporation. The residue is dissolved in 5% aqueous HCl and evaporated to dryness to give *3-chloro-4-methoxyphenethylamine hydrochloride* [7569-60-0] M 186.1, m 192-195°, as an apparently amorphous white powder when crystallisation from EtOH/ Et_2O is attempted [Fosdick et al. *J Am Chem Soc* 68 840 1946, DOI: 10.1021/ja01209a038]. The *N*-benzoyl derivative [115514-67-5] m 137-140° is a colourless solid which is insoluble in Et_2O . [Beilstein 13 III 1650.]

9-Chloromethyl anthracene [24463-19-2] $C_{15}H_{11}Cl$, M 226.7, m 138°, 140°, 141-142°(dec), 141-142.5°. If it is free from OH in the IR then recrystallisation from hexane/ C_6H_6 or $*C_6H_6$ (as needles). If OH is present, then some solvolysis has occurred. In this case treat 8.5g of it with $SOCl_2$ (4.8g) in dioxane (60ml) and reflux for 5 hours, then evaporate to dryness and wash the residue with cold $*C_6H_6$ and recrystallise it. It has been used as a protecting agent for phenols, thiophenols, mercaptans and carboxylic acids. It has a low fluorescence emission maximum at 412nm and is soluble in $CHCl_3$ and hexane. With KI/Me_2CO it forms the *iodomethyl* derivative. [Fierens et al. *Helv Chim Acta* 38 2009 1955, DOI: 10.1002/hlca.19550380740; Hunter et al. *J Org Chem* 21 1512 1956, DOI: 10.1021/jo01118a633; Beilstein 5 III 3152, 5 IV 2313.]

4-Chloro-2-methylphenol [1570-64-5] C_7H_7ClO , M 142.6, m 49°, b 112-114°/18mm, 225°/760mm, pK^{25} 9.71. Purify the phenol by crystallisation from petroleum ether (m 51°) and by zone melting. [Beilstein 6 H 359, 6 I 174, 6 II 332, 6 III 1264, 6 IV 1987.] **4-Chloro-3-methylphenol** [59-50-7] has m 63-65°, 66°, 67°, b 238°/760mm, pK^{25} 9.55. Crystallise the phenol from petroleum ether (m 66°) or $*C_6H_6$. It is dimorphic, with the second form having m 55.5°. Its solubility in H_2O is 0.38w/v% at 20°, but is much more at 100°, and in most organic solvents. A clear dilute aqueous solution turns yellow on standing in air, probably due to oxidation. Store in dark containers. It is a strong antiseptic, and disinfectant. [Kornholm et al. *Environ Sci Technol* 35 3247 2001, DOI: 10.1021/es000275e; Beilstein 6 H 381, 6 I 187, 6 II 355, 6 III 1315, 6 IV 2064.]

4-Chloro-2-methylphenoxyacetic acid (MCPA, Agroxone) [94-74-6] $C_9H_9ClO_3$, M 200.6, m 113-117°, 120°, 122-123°, pK^{20} 3.62(3.05). It is insoluble in H_2O (solubility is 0.55g/L at 20°) and recrystallises from

*C₆H₆ or chlorobenzene as plates [Jönsson et al. *Acta Chem Scand* **6** 993 1952, DOI: 10.3891/acta.chem.scand.06-0993]. The *S-benzylisothiuronium salt* has **m 164-165°**, and the Cu²⁺ salt has **m 247-249°(dec)** [Armarego et al. *Nature* **183** 1176 1959, DOI:10.1038/1831176a0; UV: Grabe *Acta Chem Scand* **4** 806 1950, DOI: 10.3891/acta.chem.scand.04-0806, IR: Sjöberg *Acta Chem Scand* **4** 798 1950, DOI: 10.3891/acta.chem.scand.04-0798]. [Beilstein **6** IV 1991.] It forms a considerably more water soluble **Sodium salt (Chiptox) [3653-48-3] C₉H₈ClNaO₃, M222.6** It is a plant growth substance and a herbicide.

2-Chloromethyl-2-phenylpropane (neophyl chloride, 1-chloro-2-methyl-2-phenylpropane, β-chloro-tert-butylbenzene) [515-40-2] C₁₀H₁₃Cl, M 168.7, b 53°/1.0mm, 95.1-95.2°/10mm, 97°/13.0mm, 104°/18.0mm, 120°/30mm, 111°/90.0mm, 222°/741mm (dec), d₄²⁵ 1.5228, n_D²⁰ 1.5250. It is prepared by adding β-methylal chloride (603g, 6.66 moles, CH₂=C(Me)-CH₂Cl, 3-chloro-2-methyl-1-propene, prepared by the chlorination of butylene, b 71.5-72.5°/760mm, d₄²⁰ 0.9165, n_D²⁰ 1.4274 [563-47-3], Beilstein **1** IV 803) into a vigorously stirred mixture of *benzene (1404g, 18.5 moles, washed twice with concentrated H₂SO₄ and used as such) and concentrated H₂SO₄ (104g, 1 mole) at 20° which required 12 hours, and stirring is continued for a further 11 hours at room temperature. The organic layer is collected, excess of *C₆H₆ is distilled off and the residue is fractionated through an 8-plate column to give pure (99.1 ±0.3% by acetolysis) *neophyl chloride* (765.5g, 68%). It is less reactive than neopentyl chloride towards Na metal and less reactive still towards EtNa, and both are inert towards most basic reagents. [Whitmore et al. *J Am Chem Soc* **65** 1469 1943, DOI: 10.1021/ja01248a010; Smith & Sellas *Org Synth Coll Vol* **4** 702 1963, DOI: 10.15227/orgsyn.032.0090; Beilstein **5** IV 1048.] It readily forms the Grignard reagent *neophyl magnesium chloride* [35293-35-7] **M 193.0**, with Mg in Et₂O; and a 0.5M solution of this reagent in Et₂O is available commercially. It reacts with solid CO₂ to give an 82% yield of *β-phenylisovaleric acid* [1010-48-6] (**m 58-59.5°**, from petroleum ether b 60-90°); and oxidation provides a 72% yield of *2-methyl-2-phenylpropan-1-ol* [2173-69-5] (**b 131°/30mm, n_D²⁰ 1.5261**) whose *phenylurethane* has **m 59.5-60.5°**, *α-naphthylurethane* has **m 91.5-92.5°**, and its *p-toluenesulfonate* has **m 74-75°**. [Whitmore et al. *J Am Chem Soc* **65** 1469 1943, DOI: 10.1021/ja01248a010; Fainberg & Winstein *J Am Chem Soc* **78** 2763 1956, DOI: 10.1021/ja01125a001; Winstein et al. *J Am Chem Soc* **74** 1113 1952, DOI: 10.1021/ja01125a001.]

N-(Chloromethyl)phthalimide [17564-64-6] C₉H₆ClNO₂, M 195.6, m 131-135°, 134-135°, 136.5°. Purify the imide by recrystallisation from EtOAc or CCl₄ or via the 1:1 complex with pyridine [Sakellarios *J Am Chem Soc* **70** 2822 1948, DOI: 10.1021/ja01188a516; Böhme et al. *Chem Ber* **92** 1258 1959, DOI: 10.1002/cber.19590920604]. [Beilstein **21/10** V 372.]

1-Chloronaphthalene [90-13-1] C₁₀H₇Cl, M 162.6, f -2.3°, b 81°/1.0mm, 111-113°/5mm, 136-136.5°/20mm, 259.3°/760mm, d₄²⁰ 1.194, n_D²⁰ 1.6326. Wash the chloronaphthalene with dilute NaHCO₃, then dry it with Na₂SO₄ and fractionally distil it *in vacuo*. It is steam volatile. Alternatively, before distillation, it is passed through a column of activated alumina, or dried with CaCl₂, then distilled from sodium. It can be further purified by fractional crystallisation by partial freezing or by crystallisation of its *picrate* to constant melting point (**m 132-133°**) from EtOH, and recovering it from the picrate. The *styphnate complex* C₁₀H₇Cl·C₆H₃N₃O₈, (**m 112°**) crystallises in yellow needles. [Beilstein **5** III 1570, **5** IV 1658.] It is a useful liquid in refractometry (cf. 1-bromonaphthalene above). **2-Chloronaphthalene [91-58-7]** has **m 59.5-60°, 61°, b 121-122°/12mm. 264-266°/760mm.** Distil 2-chloronaphthalene in a vacuum, then crystallise it from 25% EtOH/water, then dry it *in vacuo* (see the 1-isomer above). It is steam volatile. [Beilstein **5** III 1573, **5** IV 1660.]

1-Chloro-2-naphthol [633-99-8] C₁₀H₇ClO, M 178.6, m 70°, 71°, pK_{Est} ~8.3. Crystallise the naphthol from petroleum ether. The *acetate* has **m 42-43°**. [Beilstein **6** I 315, **6** II 603, **6** III 2990, **6** IV 4289.] **2-Chloro-1-naphthol [606-40-6]** has **m 64-65°, 65°, pK₂₀ 9.9 (aqueous EtOH)**. Crystallise the naphthol from petroleum ether. [Beilstein **6** I 308, **6** II 581, **6** III 2933, **6** IV 4230.] **4-Chloro-1-naphthol [604-44-4]** has **m 116-117°, 120-121°, pK₂₅ 8.86, 10.7 (aqueous EtOH)**. Crystallise the naphthol from EtOH or CHCl₃. It is a useful chromogenic peroxidase substrate for enzyme-linked detection methods. [Elias *Am J Clin Pathol* **73** 797 1980 DOI: 10.1093/ajcp/73.6.797; Beilstein **6** H 611, **6** II 582, **6** III 2933, **6** IV 4233.]

4-Chloro-2-nitroaniline [89-63-4] 242°/atm, M 172.6, m 114-115°, 116-116.5°, 117°, 119°, pK₂₅ -0.99. Crystallise the aniline from hot H₂O (**m 115.8-116°**), EtOH, EtOH/H₂O or *C₆H₆, and dry it for 10 hours at 60°

in vacuo. It has **m 115.5-116°** after sublimation. [Beilstein **12** I 355, **12** II 396, **12** III 1649, **12** IV 1669.]

2-Chloro-4-nitrobenzamide (Aklomix) [3011-89-0] $C_7H_5ClN_2O_3$, **M 200.6**, **m 170-171°, 172°**. Crystallise the amide from EtOH (grey scales). It is used against protozoal and bacterial (tuberculostatic) infections. [Jensen & Ploug *Acta Chem Scand* **3** 13 1949, DOI: 10.3891/acta.chem.scand.03-0013; Grohmann *Chem Ber* **24** 3808 (3813) 1891, DOI: 10.1002/cber.189102402258; Beilstein **9** H 404, **9** III 1768.]

2-Chloro-1-nitrobenzene (o-chloronitrobenzene) [88-73-3] $C_6H_4ClNO_2$, **M 157.6**, **m 31°, 32.8-33.2°, b 245-246°/atm, 246°/atm d²⁵ 1.348**. Crystallise the yellow solid from EtOH, MeOH or pentane (charcoal). It can be distilled at atmospheric pressure. [Beilstein **5** IV 721.] **3-Chloro-1-nitrobenzene (m-chloronitrobenzene)** [121-73-3] has **m 43°, 45.3-45.8°, 46°, 47°, b 117°/12mm, 236°/atm, d²⁵ 1.534**. Crystallise the nitrobenzene from MeOH or 95% EtOH (charcoal), then pentane. [Hartman & Brethen *Org Synth Coll Vol* **1** 126 1964, Beilstein **5** IV 722.] **4-Chloro-1-nitrobenzene (p-chloronitrobenzene)** [100-00-5] has **m 80-83°, 83.5-84°, b 113°/8mm, 242°/atm, d^{100.5} 1.2914**. Crystallise the nitrobenzene from 95% EtOH (charcoal) and sublime it *in vacuo*. [Emmons *J Am Chem Soc* **76** 3470 1954, DOI: 10.1021/ja01642a030; Newman & Fones *J Am Chem Soc* **69** 1221 1947, DOI: 10.1021/ja01197a514; Beilstein **5** IV 723.]

3-Chloroperbenzoic acid (MCPBA) [937-14-4] $C_7H_5ClO_3$, **M 172.6**, **m 92-94°(dec), pK²⁵ 7.57**. Recrystallise MCPBA from CH_2Cl_2 [Traylor & Miksztal *J Am Chem Soc* **109** 2770 1987, DOI: 10.1021/ja00243a033]. Peracid of 99+% purity can be obtained by washing commercial 85% material with phosphate buffer pH 7.5 and drying the residue under reduced pressure. Alternatively, the peracid can be freed from *m*-chlorobenzoic acid by dissolving 50g/L of *benzene and washing with an aqueous solution buffered at pH 7.4 ($NaH_2PO_4/NaOH$) (5 x 100ml). The organic layer is dried over $MgSO_4$ and carefully evaporated under vacuum. Necessary care should be taken in case of **EXPLOSION**. The solid is recrystallised twice from CH_2Cl_2/Et_2O and stored at 0° in a plastic container as glass catalyses the decomposition of the peracid. The acid is assayed iodometrically [Schwartz & Blumbergs *J Org Chem* **29** 1976 1964, DOI: 10.1021/jo01030a078]. [Bortolini et al. *J Org Chem* **52** 5093 1987, DOI: 10.1021/jo00232a006; McDonald et al. *Org Synth Coll Vol* **6** 276 1988, DOI: 10.15227/orgsyn.050.0015; Beilstein **9** IV 972.] A most useful oxidant in organic chemistry. Commercial peracid is ~80-85% pure and the contaminants are 3-chlorobenzoic acid (a stronger acid, pK²⁵ 3.83) and H_2O . It is exceedingly stable, with less than 1% loss of activity in a year at ~25°. The solubility of pure peracid at 25° (w/w%) is in H_2O (0.154), hexane (1.4), CCl_4 (2.1), $*C_6H_6$ (8.0), $CHCl_3$ (9.8), CH_2Cl_2 (11.2), EtOAc (51.0), *t*-BuOH (69.0), Et_2O (89.4) and EtOH (113.0). [Fieser **1** 135, **2** 68.]

2-Chlorophenol [95-57-8] C_6H_5ClO , **M 128.6**, **m 8.8°, b 61-62°/10mm, 176°/760mm, d²⁵ 1.241, n_D²⁰ 1.558, pK²⁵ 8.34**. Pass 2-chlorophenol at least twice through a gas chromatography column. It has also been purified by fractional distillation. [Beilstein **6** IV 782.] **3-Chlorophenol** [108-43-0] has **m 32°, 33°, 34°, b 44.2°/1mm, 214°/760mm, d²⁵ 1.218, n_D²⁰ 1.563, pK²⁵ 9.13**. It could not be obtained solid by crystallisation from petroleum ether. It is best purified by distillation under reduced pressure. [Beilstein **6** IV 810.] **4-Chlorophenol** [106-48-9] has **m 40°, 43°, 45°, 100-101°/10mm, 220°/atm, d²⁵ 1.306, pK²⁵ 9.38**. Distil the phenol, then crystallise it from petroleum ether (b 40-60°) or hexane, and dry it under vacuum over P_2O_5 at room temperature. [Bernasconi & Paschalis *J Am Chem Soc* **108** 2969 1986, DOI: 10.1021/ja00271a027; Beilstein **6** IV 820.]

4-Chlorophenoxyacetic acid (4-PCA) [122-88-3] $C_8H_7ClO_3$, **M 186.6**, **m 157°, 159°, pK²⁰ 3.00, 4.15 (50% aqueous EtOH)**. Crystallise the acid from EtOH or aqueous AcOH. It is a plant growth substance and a herbicide. [Beilstein **6** IV 845.] **4-Chlorophenoxyacetyl chloride** [4122-68-3] $C_8H_6Cl_2O_2$, **M 205.0**, has **m 18.8° and b 142°/17mm, d²⁵ 1.314, n_D²⁰ 1.5468**, and typical of acid chlorides, is an irritant and is corrosive.

(±)-α-4-Chlorophenoxypropionic acid [3307-39-9] $C_9H_9ClO_3$, **M 200.6**, **m 116°, pK_{Est} ~3.2**. Crystallise the acid from EtOH or $HCOOH$ (m 114.5-115.5°). It is a plant growth substance. The *R*(+)- and *S*(-)-*enantiomers* have **m 103-104°** (from petroleum ether) and $[\alpha]_D^{25}$ (+) and (-) 41 (c 1, EtOH). [Beilstein **6** III 695, **6** IV 850.] **β-4-Chlorophenoxypropionic acid** [3284-79-5] has **m 136°, 138°, 140°, pK_{Est} ~4.2**. Crystallise the acid from EtOH. It is a plant growth substance. [Beilstein **6** III 696, **6** IV 851.]

3-Chlorophenylacetic acid [1878-65-5] $\text{C}_8\text{H}_7\text{ClO}_2$, **M 170.6**, **m** 74°, 76°, 79°, **pK²⁵ 4.11**. Crystallise the acid from EtOH/water, or as needles from C_6H_6 or H_2O (charcoal). The *acid chloride* (prepared by boiling with SOCl_2) has **b 127-129°/15mm**. [Dippy & Williams *J Chem Soc* 161 1934, DOI: 10.1039/JR9340000161; Misra & Shukla *J Indian Chem Soc* 28 480 1951, *Beilstein* 9 III 2263, 9 IV 1674.] **4-Chlorophenylacetic acid** [1878-66-6] has **m 102°, 105°, 106°, pK²⁵ 4.12**. Purify it as for 3-chlorophenylacetic acid. The *acid chloride* [25025-34-0] $\text{C}_8\text{H}_6\text{Cl}_2\text{O}$, **M 189.0**, (prepared by boiling with SOCl_2) has **b 85°/1mm** and **d²⁵ 1.292**, **n_D²⁰ 1.5510**. [*Beilstein* 9 III 2263, 9 IV 1675.]

4-Chloro-1-phenylbutan-1-one (γ -chlorobutyrophenone) [939-52-6] $\text{C}_{10}\text{H}_{11}\text{ClO}$, **M 182.7**, **m** 19° to 20°, **b 120-121°/3mm**, **134-137°/5mm**, **d₄²⁰ 1.149**, **n_D²⁰ 1.55413**. Fractionate the ketone several times using a short column. It recrystallises from petroleum ether at -20° in glistening white rosettes and is filtered at 0°, and dried in a vacuum desiccator over H_2SO_4 . The *semicarbazone* has **m 136-137°**. [Conant et al. *J Am Chem Soc* 46 1882 1924, DOI: 10.1021/ja01673a015; Cloke *J Am Chem Soc* 51 1174 1929, DOI: 10.1021/ja01379a028; Hart & Curtis *J Am Chem Soc* 79 931 1957, DOI: 10.1021/ja01561a042; *Beilstein* 7 IV 711.]

1-(2-Chlorophenyl)-1-(4-chlorophenyl)-2,2-dichloroethane (Mitotane, *o,p'*-DDD) [53-19-0] $\text{C}_{14}\text{H}_{10}\text{Cl}_4$, **M 320.1**, **m 75.8-76.8°, 76-78°**. Purify Mitotane by recrystallisation from pentane, MeOH or EtOH. It is soluble in isooctane and CCl_4 . [Haller et al. *J Am Chem Soc* 67 1591 1945, DOI: 10.1021/ja01225a058; *Beilstein* 5 IV 1883.]

3-(4-Chlorophenyl)-1,1-dimethylurea (Monuron, CMU) [150-68-5] $\text{C}_9\text{H}_{11}\text{ClN}_2\text{O}$, **M 198.7**, **m 171°, 170.5-171.5°** (176-177° also reported). Crystallise monuron from MeOH. The pH of a saturated solution in H_2O is *ca* 6.3, and it forms a more soluble *trichloroacetate salt* (Urox), $\text{C}_9\text{H}_{11}\text{ClN}_2\text{O} \cdot \text{C}_2\text{HCl}_3\text{O}_2$, **m 78-81°**, which is an effective herbicide against monocotyledonous plants. [*Beilstein* 12 IV 1191.]

4-Chloro-1,2-phenylenediamine [95-83-0] $\text{C}_6\text{H}_7\text{ClN}_2$, **M 142.6**, **m 69-70°, 73°**, **pK₁²⁵ -0.27** (aqueous H_2SO_4), **pK₂²⁵ 3.35** (3.67). Recrystallise the diamine from petroleum ether. [*Beilstein* 13 IV 68.]

4-Chlorophenyl isocyanate [104-12-1] $\text{C}_7\text{H}_4\text{ClNO}$, **M 153.6**, **m 26°, 28-31°, 31-32°, 32°, 32.5°**, **b 80.6-80.9°/9.5mm**, **115-117°/45mm**, **d²⁵ 1.20**, **n_D²⁰ 1.5618**. Purify the isocyanate by recrystallisation from petroleum ether (b 30-40°) or better by fractional distillation. **TOXIC irritant**. [*Beilstein* 12 III 1376, 12 IV 1213.]

4-Chlorophenyl isothiocyanate [2131-55-7] $\text{C}_7\text{H}_4\text{ClNS}$, **M 169.6**, **m 42°, 44°, 43-45°, 45°, 46°, 47°**, **b 110-115°/4mm**, **135-136°/24mm**. Check the IR first to see if free from OH frequencies. Triturate it with petroleum ether (b 30-60°) and decant the solvent. Repeat this 5 times. The combined extracts are evaporated under reduced pressure to give almost pure compound as a readily crystallisable oil with a pleasant anise odour. It can be recrystallised from the minimum volume of EtOH at 50° (do not boil too long as it could react). It can be purified by vacuum distillation. [van der Kerk et al. *Org Synth Coll Vol* 5 223 1973, DOI: 10.15227/orgsyn.045.0019; *Beilstein* 12 IV 1214.] **It is an IRRITANT and causes dermatitis; use gloves.**

4-Chlorophenyl 2-nitrobenzyl ether [109669-56-9] $\text{C}_{13}\text{H}_{10}\text{ClNO}_3$, **M 263.7**, **m 44.5°**, **b 154-156°/3mm**, **208°/11mm**. Distil it under reduced pressure, and it crystallises from EtOH (**m 44-45°**) or MeOH (**m 46°**) as yellow needles. [*Beilstein* 6 II 210, 6 III 801, 6 IV 1253.] **4-Chlorophenyl 4-nitrobenzyl ether** [5442-44-4] has **m 77°**, **b 215°/12mm**. Distil it in a vacuum and crystallise it from EtOH, MeOH (**m 75.5-76°**) or petroleum ether (**m 76°, 77°**). Its UV has λ_{max} at 222 and 302nm (EtOH). [*Beilstein* 6 II 222, 6 III 821, 6 IV 1288.]

4-Chlororesorcinol [95-88-5] $\text{C}_6\text{H}_5\text{ClO}_2$, **M 144.6**, **m 105°, 106°, 108°**, **b 147°/18mm**, **pK_{Est(1)} ~9.2**, **pK_{Est(2)} ~10.1**. Crystallise it from boiling CCl_4 (10g/L, charcoal) and dry it in air. [*Beilstein* 6 II 818.] **IRRITANT**.

5-Chlorosalicylaldehyde [635-93-8] $\text{C}_7\text{H}_5\text{ClO}_2$, **M 156.6**, **m 98.5-99°, 99.5°, 101°, 102°**, **pK²⁵ 7.4**. Steam distil it, then crystallise it from aqueous EtOH or C_6H_6 (**m 100°**). It forms complexes with Cu^{2+} and Fe^{2+} . [*Beilstein* 8 H 53, 8 II 45, 8 III 181, 8 IV 224.]

4-Chlorothiophenol [106-54-7] $\text{C}_6\text{H}_5\text{ClS}$, **M 144.6**, **m 49°, 51-52°, 53.5-54°**, **b 205-207°/atm**, **pK²⁵ 6.14**.

Recrystallise the thiophenol from aqueous EtOH. The *SMe ether* has **m** 129° and the *SEt ether* has **m** 64°. [D'Souza et al. *J Org Chem* **52** 1720 1987, DOI: 10.1021/jo00385a014; *Beilstein* **6** H 326, **6** I 149, **6** III 1034.]

2-Chlorotoluene [95-49-8] C_7H_7Cl , **M** 126.6, **m** -36°, **b** 159°/atm, d_4^{20} 1.083, n_D^{20} 1.5255. Dry 2-chlorotoluene for several days with $CaCl_2$, then distil it from Na using a glass helices-packed column. It is steam volatile. [cf: Marvel & McElvain *Org Synth Coll Vol* **1** 170 1941, DOI: 10.15227/orgsyn.003.0033; *Beilstein* **5** IV 805.] **3-Chlorotoluene** [108-41-8] has **m** -48°, **b** 161-163°/atm, d_4^{20} 1.072, n_D^{20} 1.522. Purify it as for 2-chlorotoluene above. [cf: Marvel & McElvain *Org Synth Coll Vol* **1** 170 1941, DOI: 10.15227/orgsyn.003.0033; *Beilstein* **5** IV 806.] **4-Chlorotoluene** [106-43-4] has **m** 7.2°, **b** 162.4°/atm, d_4^{20} 1.07, n_D^{20} 1.521. Dry it with BaO, fractionally distil it, then fractionally crystallise it by partial freezing. [cf: Marvel & McElvain *Org Synth Coll Vol* **1** 170 1941, DOI: 10.15227/orgsyn.003.0033; *Beilstein* **5** IV 806.]

Chrysene (1,2-benzanthracene) [218-01-9] $C_{18}H_{12}$, **M** 228.3, **m** 254°, 255-256°, **b** 254°/atm, d_4^{20} 1.274. Purify chrysene by chromatography on alumina from petroleum ether in a darkened room. Its solution in C_6H_6 is passed through a column of decolorising charcoal, then crystallised by concentrating the eluate. It has also been purified by crystallising from C_6H_6 or C_6H_6 /petroleum ether, and by zone refining. [Gorman et al. *J Am Chem Soc* **107** 4404 1985, DOI: 10.1021/ja00301a006]. It is freed from 5*H*-benzo[*b*]carbazole by dissolving it in *N,N*-dimethylformamide and successively adding small portions of alkali and iodomethane until the fluorescent colour of the carbazole anion no longer appears when alkali is added. The chrysene (and alkylated 5*H*-benzo[*b*]carbazole) separate on addition of water. Final purification is by crystallisation from ethylcyclohexane and/or from 2-methoxyethanol [Bender et al. *Anal Chem* **36** 1011 1964, DOI: 10.1021/ac60212a018]. It can be sublimed in a vacuum. [*Beilstein* **5** IV 2554.]

Chrysoidine G (4-phenylazo-1,3-benzenediamine monohydrochloride, CI 11270, basic orange 2) [532-82-1] $C_{12}H_{13}ClN_4$, **M** 248.7, **m** 235° (dec) (118-118.5° also reported? free base) λ_{max} 449nm, pK_1 3.32, pK_2 5.21. It is a red-brown powder which is recrystallised from H_2O . It gives a yellow solution in concentrated H_2SO_4 which turns orange on dilution. Its solubility at 15° is 5.5% (H_2O), 4.75% (EtOH), 6.0% (cellosolve), 9.5% (ethylene glycol), 0.005% (xylene) and is insoluble in C_6H_6 . The *hydroiodide* has **m** 184° (from EtOH) and the *picrate* forms red needles with **m** 196°. [Muramatsu *Bull Chem Soc Jpn* **31** 864 1958, *Beilstein* **6** IV 561.] The *free base* [9495-54-5] $C_{12}H_{12}N_4$ is CI Solvent Orange 3. The *citrate salt* [5909-04-6] is an antiseptic. The dye and its salts yield orange solutions in EtOH, and are chiefly used for colouring cotton and silk. They are also used in microscopy for staining microorganisms, and they complex with Cu^{2+} ions.

trans-Cinnamaldehyde [14271-10-9] C_9H_8O , **M** 132.2, **m** -4°, -7.5°, -9°, **b** 80°/0.4mm, 85.8°/1.1mm, 125-128°/11mm, 152.2°/40mm, 163.7°/60mm, 199.3°/200mm, 246°/760mm (dec), d_4^{20} 1.0510, n_D^{20} 1.623. Purify the aldehyde by steam distillation (solubility is 1 in 700 parts H_2O) followed by distillation *in vacuo*. The *cis*-isomer has **b** 67-69°/40mm and d_4^{20} 1.0436 and n_D^{20} 1.5937. The *trans-semicarbazone* has **m** 210°(dec) from $CHCl_3$ /MeOH, (*cis-semicarbazone* has **m** 196°), the *trans-phenylsemicarbazone* has **m** 177° from $CHCl_3$ /MeOH (the *cis-phenylsemicarbazone* has **m** 146°), the *trans*- and *cis*-2,4-dinitrophenylhydrazones have **m** 250°(dec) from MeOH [Gamboni et al. *Helv Chim Acta* **38** 255 1955, DOI: 10.1002/hlca.19550380130; Holum *J Org Chem* **26** 4814 1961, DOI: 10.1021/jo01070a009]. [*Beilstein* **9** IV 984.]

cis-Cinnamic acid (Z-3-phenyl-2-propenoic acid) [102-94-3] $C_9H_8O_2$, **M** 148.2, **m** 68° (for *allo*-form), pK^{25} 3.93. The *cis*-acid is prepared by catalytic reduction of phenylpropionic acid and after distillation in a high vacuum at ~95° it gives the most stable *allo*-isomer **m** 68°. Recrystallisation from petroleum ether yields **Liebermann's iso-cinnamic acid m** 58°. When the *allo*-acid (**m** 68°) is heated at 20° above its melting point in a sealed capillary for 0.5 hours and allowed to cool slowly, **Erlenmyer's iso-cinnamic acid m** 42° is formed. This form can also be obtained in larger amounts by heating the *allo*-acid at 80° for 3 hours, and on cooling it remains liquid for several weeks but gives the **m** 42° acid on inoculation with the crystals from the capillary tube. This form is unchanged in 6 weeks when kept in a dark cupboard. All three forms have the same pK values and the same rate of bromination. There is also a very labile form with **m** 32°. [Liebermann, *Chem Ber* **26** 1571 1893, DOI: 10.1002/cber.18930260275; Claisen & Crismer *Justus Liebigs Ann Chem* **218** 129 1883, DOI: 10.1002/jlac.18832180203; Robinson & James *J Chem Soc* 1453 1933, DOI: 10.1039/JR9330001453; Berthoud & Urech *Helv Chim Acta* **13** 437 1930, DOI: 10.1002/hlca.19300130403; McCoy & McCoy *J Org*

Chem **33** 2354 1968, DOI: 10.1021/jo01270a037; *Beilstein* **9** IV 2001.]

trans-Cinnamic (E-3-phenyl-2-propenoic) acid [140-10-3, 621-82-9 for E-Z mixture] $C_9H_8O_2$, **M 148.2**, **m 132°**, **134.5-135°**, **pK²⁵ 4.42 (4.50)**. Crystallise the acid from *benzene, CCl_4 , hot water, water/EtOH (3:1), or 20% aqueous EtOH. Dry it at 60° *in vacuo*. It is steam volatile. [*Beilstein* **9** IV 2002.] **trans-Cinnamoyl chloride** [102-92-1] C_9H_7ClO , **M 166.6**, has **m 35-37°**, **b 101°/2mm**, **154°/25mm**, **256-258°/atm**, **d₄^{37.6} 1.6202**, **n_D^{37.6} 1.1632**. Refractionate it in a vacuum until the distillate solidifies on cooling, and recrystallise the yellow distillate from petroleum ether. The **trans-amide** has **m 145-150°** (from H_2O) [*Beilstein* **9** III 2711]. [Adams & Ulich *J Am Chem Soc* **42** 605 1920, DOI: 10.1021/ja01448a024; Bergmann et al. *J Chem Soc* 2522 1952, DOI: 10.1039/JR9520002522; *Beilstein* **9** H 587, **9** I 233, **9** II 390, **9** III 2710, **9** IV 2020.] **trans-Cinnamic anhydride** [538-56-7] $C_{18}H_{14}O_3$, **M 278.4**, has **m 136°**. Crystallise the anhydride from * C_6H_6 or toluene/petroleum ether (b 60-80°) or EtOH (**m 135-136°**). It is insoluble in H_2O . [*Beilstein* **9** III 2703, **9** IV 2018.]

N-Cinnamoyl-N-phenylhydroxylamine [7369-44-0] $C_{15}H_{13}NO_2$, **M 239.3**, **m 158-163°**. Recrystallise the hydroxylamine from EtOH.

Cinnamyl alcohol [104-54-1] $C_9H_{10}O$, **M 134.2**, **m 30°**, **33°**, **b 143.5°/14mm**, **λ_{max} 251nm** (ϵ 18,180 $M^{-1}cm^{-1}$). Crystallise the alcohol from diethyl ether/pentane. [*Beilstein* **6** I 281.]

Coniferyl alcohol [4-hydroxy-3-methoxy-cinnamyl alcohol, 3-(4-hydroxy-3-methoxy-phenyl)-2-propen-1-ol] [458-35-5] $C_{10}H_{12}O_3$, **M 180.2**, **m 73°**, **74°**, **75°**, **80°**, **b 163-165°/3mm**, **pK²⁵ 9.54**. It is soluble in EtOH and insoluble in H_2O . It can, however, be recrystallised from EtOH and distilled in a vacuum. It polymerises in dilute acid. The **benzoyl** derivative has **m 95-96°** (from petroleum ether), and the **tosylate** has **m 66°**. [Derivatives: Freudenberg & Achtzehn *Chem Ber* **88** 10 1955, DOI: 10.1002/cber.19550880103; UV: Herzog & Hillmer *Chem Ber* **64** 1288 1931, DOI: 10.1002/cber.19310640614; *Beilstein* **6** II 1093.]

Coronene [191-07-1] $C_{24}H_{12}$, **M 300.4**, **m 438-440°**, **442°**, **b 525°/atm**, **λ_{max} 345nm** ($\log \epsilon$ 4.07). Crystallise coronene from *benzene or toluene, then sublime it in a vacuum. [*Beilstein* **5** III 2651.] It is an *n*-channel organic semiconductor [Newman et al *Chem Mater* **16** 4436 2004, DOI: 10.1021/cm049391x], and used to prepare MBE-grown layered superconductors [Schuerlein, Schmidt et al. *Japanese Journal of Applied Physics, Part 1* **34** 3837 1995, DOI: 10.1143/JJAP.34.3837].

o-Cresol [95-48-7] C_7H_8O , **M 108.1**, **m 30.9°**, **b 191°/760mm**, **n_D⁴¹ 1.536**, **n_D⁴⁰ 1.534**, **pK²⁵ 10.22**. It can be freed from *m*- and *p*-isomers by repeated fractional distillation. It crystallises from *benzene by addition of petroleum ether. It has been fractionally crystallised by partial freezing of its melt. The **3,5-dinitrobenzoate** (prepared from 3,5-dinitrobenzoyl chloride in dry pyridine, and recrystallised from EtOH or aqueous Me_2CO) has **m 138°**. [*Beilstein* **6** IV 1940.] **m-Cresol** [108-39-4] has **f 12.0°**, **b 202.7°/atm**, **d₄²⁰ 1.034**, **n_D²⁰ 1.544**, **pK²⁵ 0.09**. Separation of the *m*- and *p*-cresols requires chemical methods, such as conversion to their sulfonates [see Englund et al. *J Am Chem Soc* **45** 189 1953, DOI: 10.1021/ie50517a055]. An equal volume of H_2SO_4 is added to *m*-cresol, stirred with a glass rod until solution is complete. Heat for 3 hours at 103-105°. Dilute carefully with 1-1.5 volumes of water, heat to boiling point and steam distil until all unsulfonated cresol has been removed. Cool and extract the residue with ether. Evaporate the solution until the boiling point reaches 134° and steam distil off the *m*-cresol. Another purification method involves distillation, fractional crystallisation from the melt, then redistillation. Free from *p*-cresol by solution in glacial acetic acid and bromination by about half of an equivalent amount of bromine in glacial acetic acid. The acetic acid is distilled off, then fractional distillation of the residue under vacuum gives bromocresols from which 4-bromo-*m*-cresol is obtained by crystallisation from hexane. Addition of the bromocresol in glacial acetic acid slowly to a reaction mixture of HI and red phosphorus or (more smoothly) of HI and hypophosphorus acid, in glacial acetic acid, at reflux, removes the bromine. After an hour, the solution is distilled at atmospheric pressure until layers are formed. Then it is cooled and diluted with water. The cresol is extracted with ether, washed with water, $NaHCO_3$ solution and again with water, dried with a little $CaCl_2$ and distilled [Baltzly et al. *J Am Chem Soc* **77** 2522 1955, DOI: 10.1021/ja01614a049]. *m*-Cresol is the exclusive phenolic formed from *o*- and *p*-toluic acids when heated with $CuCO_3$, MgO , H_2O and air at ~230° [Kaeding et al. *J Am Chem Soc* **53** 805 1961,

DOI: 10.1021/ie50622a023]. The **3,5-dinitrobenzoate** (prepared from 3,5-dinitrobenzoyl chloride in dry pyridine, and recrystallised from EtOH or aqueous Me₂CO) has **m 165°**. [Beilstein 6 IV 2035.] Separation of the *m*- and *p*-cresols have been achieved by extractive crystallisation [Chivate & Shah *Chem Eng Sci* 5 232 1956, DOI:10.1016/0009-2509(56)80035-3]. *p*-Cresol [106-44-5] C₇H₈O, **M 108.1, m 34.8°, b 201.9°, n⁴¹ 1.531, n⁴⁶ 1.529, pK²⁵ 10.27**. It can be separated from *m*-cresol by fractional crystallisation of its melt. Purify it by distillation, by precipitation from *benzene solution with petroleum ether, and *via* its benzoate, as for phenol. Dry it under vacuum over P₂O₅. It has also been crystallised from petroleum ether (b 40-60°) and by conversion to sodium *p*-cresoxyacetate which, after crystallisation from water is decomposed by heating with HCl in an autoclave [Savard *Ann Chim (Paris)* 11 287 1929]. *p*-Cresol, free from its isomers, can be prepared by fusing sodium *p*-toluenesulfonate with KOH (but not with NaOH) at 300-320° followed by treatment with H₂SO₄, steam distillation, drying and distillation (**b 94-95°/15mm, m 31.4°**) in 63-72% yield [Hartman *Org Synth Coll Vol* 1 175 1941, DOI: 10.15227/orgsyn.003.0037]. *p*-Cresol can be freed from contaminating *o*- and *m*- isomers by reaction with a tenth of its weight of 2,6-dichloroquinone chloroimide (which reacts with the latter two isomers to form a blue colour) [Gibbs *J Am Chem Soc* 49 839 1927, DOI: 10.1021/ja01402a027]. The **3,5-dinitrobenzoate** (prepared from 3,5-dinitrobenzoyl chloride in dry pyridine, and recrystallised from EtOH or aqueous Me₂CO) has **m 189°**. [For UV see: Sreeramamurthy *Trans Faraday Soc* 47 1256 1951, DOI: 10.1039/TF9514701256; Beilstein 6 II 2093.]

***o*-Cresolphthalein [3,3-bis(4-hydroxy-3-methylphenyl)-1(3*H*)isobenzofura-1-one] [596-27-0] C₂₂H₁₈O₄, M 346.4, m 219° to 221°, 223°, 221.3-223.6°, λ_{max} 381nm and 566nm, pK_a 9.4**. It is prepared by heating *o*-cresol (2 mols), phthalic anhydride (1 mol) and anhydrous zinc chloride (0.5 mol) at 120°/3hrs [Hubacher et al. *J Am Pharm Assoc* 42 23 1953, DOI: 10.1002/jps.3030420108], and finally purified by recrystallisation from EtOH. It is soluble in alcohol, but only slightly soluble in H₂O. Hence dissolve it in the smallest volume of EtOH, add H₂O to turbidity, then allow the EtOH to evaporate off and collect the white solid, dry it and recrystallise it from 95% EtOH. It is used in analytical chemistry as an indicator where in aqueous solution at pH ~8 it is colourless but turns red in alkaline solution, e.g. at pH 9.8 (cf: pK_a at 9.4). Unlike some other phthaleins it is inactive as a laxative. [Beilstein 18 V/4 193.]

***o*-Cresolphthalein Complexone {Metalphthalein, Phthalein purple, 3,3-bis[5-(*N,N*-di(carboxymethyl)-aminomethyl)-4-hydroxy-3-methylphenyl)-1(3*H*)isobenzofuran-1-one]} [2411-89-4] C₃₂H₃₂N₂O₁₂, M 636.6, m 181° to 185°(dec), 186°(dec), λ_{max} 575nm, pK₁ 2.2, pK₂ 2.9, pK₃ 7.0, pK₄ 7.8, pK₅ 11.4, pK₆ 12.0**. *o*-Cresolphthalein (see previous entry, a complexone precursor without the two bis-carboxymethylaminomethyl groups) is a contaminant and is one of the starting materials. It can be removed by dissolving the reagent in H₂O and adding a 3-fold excess of sodium acetate and fractionally precipitating it by dropwise addition of HCl to the clear filtrate. Wash the white precipitate with cold H₂O and dry the *monohydrate* at 30° in a vacuum (0.01mm). The pure material gives a single spot on paper chromatography (eluting solvent EtOH/water/phenol, 6:3:1, and developing with NaOH). [Anderegg et al. *Helv Chim Acta* 37 113 1954, DOI: 10.1002/hlca.19540370114.] It complexes with Mg²⁺, Ba²⁺, Sr²⁺, Ca²⁺, Cd²⁺ and Zn²⁺, and is an excellent indicator for the complexometric titration of alkaline earth metals which show colour changes (e.g. red/pink, red/rose, red/colourless) in NH₃-buffer at pH 10–11 [cf. Bishop *Indicators* Pergamon Press 1972, Library of Congress Cat Card No 78-171464]. It has also been used, in the presence of Ba²⁺, for the titration of sulfate ions. [Beilstein 18 III/IV 8141.]

***o*-Cresotic acid (3-methylsalicylic acid) [83-40-9] C₈H₈O₃, M 152.2, m 163-164°, 165°, 165-166°, pK₁²⁵ 3.32**. Crystallise the acid from water. It is steam volatile. [Beilstein 10 H 220, 10 II 131, 10 III 505, 10 IV 601.] ***m*-Cresotic acid (4-methylsalicylic acid) [50-85-1] has m 173°, 176°, 177°, (182-183°), pK₁²⁵ 3.15, pK₂²⁵ 13.35**. Crystallise the acid from water. It is steam volatile. It sublimes at 130°/11mm. [Beilstein 10 H 233, 10 II 137, 10 III 521, 10 IV 617.] ***p*-Cresotic acid (5-methylsalicylic acid) [89-56-5] has m 150°, 151°, 152°, 151-154°, pK₁²⁵ 3.40, pK₂²⁵ 13.45**. Crystallise the acid from H₂O. It is steam volatile, and decomposes somewhat on sublimation. [Beilstein 10 H 227, 10 II 134, 10 III 516, 10 IV 610.]

Crystal Violet Chloride {Gentian violet, *N*-4[bis[4-(dimethylaminophenyl)methylene]-2,5-cyclohexadien-1-ylidene]-*N*-methylmethaninium chloride} [548-62-9] C₂₅H₃₀ClN₃, M 408.0, m 205°(dec), pK 9.36. Crystallise the greenish dye (with a metallic lustre) from water (20ml/g), the crystals being separated from the chilled solution by centrifugation, then wash them with chilled EtOH (solubility is 1g in 10 ml of hot EtOH) and

diethyl ether and dry under vacuum. It is soluble in CHCl_3 but insoluble in Et_2O . The carbinol is precipitated from an aqueous solution of the dye-hydrochloride, using excess NaOH , then dissolve in HCl and recrystallise it from water as the chloride [UV and kinetics: Turgeon & La Mer *J Am Chem Soc* **74** 5988 1952, DOI: 10.1021/ja01143a050]. The **carbinol base** has **m 195°** (needles from EtOH). The **diphthalate** (blue and turns red in H_2O) crystallises from H_2O , **m 153-154° (dec at 185-187°)** [Chamberlain & Dull *J Am Chem Soc* **50** 3088 1928, DOI: 10.1021/ja01398a035]. [Beilstein **13** H 233, **13** IV 2284.] Its solubility in glycerin is 6.7w/v%. It is used as a biological stain for Gram bacteria, with iodine for chromosome and nucleoli stain, and is an amyloid stain. It is anthelmintic and is a topical antibacterial.

Cumene (isopropyl benzene) [98-82-8] C_9H_{12} , **M 120.2**, **m -96°**, **b 69-70°/41mm**, **152.4°/760mm**, **d₄²⁰ 0.864**, **n_D²⁰ 1.49146**, **n_D²⁵ 1.48892**. Usual purification is by washing it with several small portions of concentrated H_2SO_4 (until the acid layer is no longer coloured), then with water, 10% aqueous Na_2CO_3 , again with water, and drying with MgSO_4 , MgCO_3 or Na_2SO_4 , followed by fractional distillation. It can then be dried with, and distilled from, Na , NaH or CaH_2 . Passage through columns of alumina or silica gel removes oxidation products. It has also been steam distilled from 3% NaOH , and azeotropically distilled with 2-ethoxyethanol (which is subsequently removed by washing out with water). [Beilstein **5** IV 985.]

Cumene hydroperoxide [80-15-9] $\text{C}_9\text{H}_{11}\text{O}_2$, **M 152.2**, **b 60°/0.2mm**, **100-101°/8mm**, **d₄²⁰ 1.028**, **n_D²⁴ 1.5232**. Purify the hydroperoxide by adding 100ml of 70% material slowly and with agitation to 300ml of 25% NaOH in water, keeping the temperature below 30°. The resulting crystals of the sodium salt are filtered off, washed twice with 25 ml portions of *benzene, then stirred with 100ml of *benzene for 20 minutes. After filtering off the crystals and repeating the washing, they are suspended in 100ml of distilled water and the pH is adjusted to 7.5 by addition of 4M HCl . The free hydroperoxide is extracted into two 20ml portions of *n*-hexane, and the solvent is evaporated under vacuum at room temperature, the last traces being removed at 40-50°/1mm [Fordham & Williams *Can J Res* **27B** 943 1949, DOI: 10.1139/cjr49b-096]. Petroleum ether, **but not diethyl ether**, can be used instead of *benzene, and powdered solid CO_2 can replace the 4M HCl . [Beilstein **6** IV 3221.] *The material is potentially EXPLOSIVE.*

Cuminaldehyde (4-isopropylbenzaldehyde) [122-03-2] $\text{C}_{10}\text{H}_{12}\text{O}$, **M 148.2**, **b 82-84°/3.5mm**, **120°/23mm**, **131-135°/35mm**, **235-236°/760mm**, **d₄²⁰ 0.978**, **n_D²⁰ 1.5301**. A likely impurity is the benzoic acid. Check the IR for the presence of OH from CO_2H , and the CO frequencies. If the acid is present, then dissolve the aldehyde in Et_2O , wash it with 10% NaHCO_3 until effervescence ceases, then with brine, dry over CaCl_2 , evaporate and distil the residual oil, preferably under vacuum. It is almost insoluble in H_2O , but soluble in EtOH and Et_2O . The **thiosemicarbazone** has **m 147°** after recrystallisation from aqueous EtOH , MeOH or $^*\text{C}_6\text{H}_6$. [Crounse *J Am Chem Soc* **71** 1263 1949, DOI: 10.1021/ja01172a035; Bernstein et al. *J Am Chem Soc* **73** 906 1951, DOI: 10.1021/ja01147a007; Gensler & Berman *J Am Chem Soc* **80** 4949 1958, DOI: 10.1021/ja01551a048; Beilstein **7** H 318, **7** II 347, **7** III 1095, **7** IV 723.]

9-Cyanoanthracene (anthracene-9-carbonitrile) [1210-12-4] $\text{C}_{15}\text{H}_9\text{N}$, **M 203.2**, **m 173-174°**, **173-177°**. Recrystallise the nitrile from EtOH (until a single sharp maximum at 256.5nm is obtained) or toluene, and sublime it in a vacuum in the dark under N_2 [Ebied et al. *JCS Faraday Trans 1* **76** 2170 1980, DOI: 10.1039/F19807602170; Kikuchi et al. *J Phys Chem* **91** 574 1987, DOI: 10.1021/j100287a017]. [Beilstein **9** I304.] The dimer below can be thermally monomerised at *ca* 120°.

9-Cyanoanthracene photodimer [33998-38-8] $\text{C}_{30}\text{H}_{18}\text{N}_2$, **M 406.4**, **dec to monomer above ~147°**. Purify the dimer (obtained by UV irradiating the above monomer at λ 350-370nm) by dissolving it in the minimum amount of CHCl_3 followed by addition of EtOH at 5° [Ebied et al. *JCS Faraday Trans 1* **75** 1111 1979, DOI: 10.1039/F19797501111; Ebied et al. *JCS Faraday Trans 1* **76** 2170 1980, DOI: 10.1039/F19807602170].

4-Cyanobenzoic acid [619-65-8] $\text{C}_8\text{H}_5\text{NO}_2$, **M 147.1**, **m 219°**, **219-221° (dec)**, **pK²⁵ 3.55**. Crystallise the acid from water and dry it in a vacuum desiccator over Sicapent. [Beilstein **9** IV 3324.] **4-Cyanobenzoyl chloride** [6068-72-0] $\text{C}_8\text{H}_4\text{ClNO}$, **M 165.6**, has **m 68-70°**, **69-70°**, **73-74°**, **b 132°/8mm**, **150-151°/25mm**. If the IR shows the presence of OH, then treat it with SOCl_2 boil for 1 hour, evaporate and distil it in a vacuum. The distillate solidifies and can be recrystallised from petroleum ether. It is moisture sensitive and an **IRRITANT**.

[Ashley et al. *J Chem Soc* 103 1942, DOI: 10.1039/JR9420000103; Fuson et al. *J Org Chem* **16** 648 1951, DOI: 10.1021/jo01144a018.] [*Beilstein* **9** III 4255, **14** IV 3327.]

***p*-Cyanophenol** (*p*-hydroxybenzonitrile) [767-00-0] C_7H_5NO , **M 119.1**, **m 110°, 113°, pK²⁵ 7.97**. Crystallise the phenol from petroleum ether, *benzene or water and keep it under vacuum over P_2O_5 . [Bernasconi & Paschelis *J Am Chem Soc* **108** 2969 1986, DOI: 10.1021/ja00271a027.] [*Beilstein* **10** H 167, **10** IV 441.]

Cyclohexylbenzene (phenylcyclohexane) [827-52-1] $C_{12}H_{16}$, **M 160.3**, **m 6.8°, b 237-239°/atm, 239-240°/atm, d₄²⁰ 0.950, n_D²⁰ 1.5258**. Purify it by fractional distillation, and by fractional freezing. [*Beilstein* **5** IV 1424.]

Cyclopropyldiphenylcarbinol (cyclopropyldiphenylmethanol) [5785-66-0] $C_{16}H_{16}O$, **M 224.3**, **m 86-87°**. Crystallise the carbinol from *n*-heptane or * C_6H_6 /pentane (**m 82-83°**). It sublimes at 60°/0.001mm. The **2,4-dinitrobenzoate** has **m 140°**. [*Beilstein* **6** III 3517, **6** IV 4888.]

***p*-Cymene** (**4-isopropyltoluene**) [99-87-6] $C_{10}H_{14}$, **M 134.2**, **m -67.9°, b 177.1°/760mm, 176-178°/atm, d₄²⁰ 0.8569, n_D²⁰ 1.4909, n_D²⁵ 1.4885**. It occurs in essential oils of plants. Wash *p*-cymene with cold, concentrated H_2SO_4 until there is no further colour change, then repeatedly with H_2O , 10% aqueous Na_2CO_3 and H_2O again. Dry it over Na_2SO_4 , $CaCl_2$ or $MgSO_4$, and distil it. Further purification steps include steam distillation from 3% $NaOH$, percolation through silica gel or activated alumina, and a preliminary reflux for several days over powdered sulfur. Store it over CaH_2 . [*Beilstein* **5** IV 1060.] ***o*-Cymene** (**2-isopropyltoluene**) [527-84-4] $C_{10}H_{14}$, **M 134.2**, has **m -71.5° (-81.5° and -75.2°, unstable forms) b 178.15°/760mm, 176-178°/atm, d₄²⁰ 0.8766, n_D²⁰ 1.5006**; and ***m*-Cymene** (**3-isopropyltoluene**) [535-77-3] $C_{10}H_{14}$, **M 134.2**, has **m -64°, b 175.14°/760mm, d₄²⁰ 0.8610, n_D²⁰ 1.4930** and can be purified as described above for the *p*-isomer.

Deoxybenzoin (**2-phenylacetophenone**) [451-40-1] $C_{14}H_{12}O$, **M 196.3**, **m 54-55°, 60°, b 177°/12mm, 320°/760mm**. Crystallise deoxybenzoin from EtOH and/or distil it in a vacuum. [*Beilstein* **7** II 368, **7** III 2098, **7** IV 1393.]

(±)-Desyl bromide (**α-bromo-desoxybenzoin, ω-bromo-ω-phenyl acetophenone**) [484-50-0] $C_{14}H_{11}BrO$, **M 275.2**, **m 56°, 57.1-57.5°, 56°**. Crystallise it from 95% EtOH. It can be purified like the following (±)-desyl chloride. Store it in dark containers as it is more sensitive to light than the following chloride. [*Beilstein* **7** H 436, **7** II 370, **7** III 2122.] **(±)-Desyl chloride** (**α-chloro-desoxybenzoin, ω-chloro-ω-phenyl acetophenone**) [447-31-4] $C_{14}H_{11}ClO$, **M 230.7**, has **m 62-64°, 66-67°, 67.5°, 68°**. For the purification of small quantities recrystallise it from petroleum ether (b 40-60°), but use MeOH or EtOH for larger quantities. For the latter solvent, dissolve 12.5g of chloride in 45ml of boiling EtOH (95%), filter and the filtrate yields colourless crystals (7.5g) on cooling. A further crop (0.9g) can be obtained by cooling in an ice-salt bath. It turns brown on exposure to sunlight but it is stable in sealed dark containers. The ***R*(+)-enantiomer** has **m 75-76°** (from petroleum ether) and $[\alpha]_{546} +168.4$ (c 0.6, Me_2CO) [Roger & Wood *J Chem Soc* 811 1954, DOI: 10.1039/JR9540000811]. [Henley & Turner *J Chem Soc* 1182 1931, DOI: 10.1039/JR9310001182; Ward *Org Synth Coll Vol 2* 159 1943, DOI: 10.15227/orgsyn.012.0020; *Beilstein* **7** H 436, **7** I 234, **7** II 369, **7** III 2106, **7** IV 1396.]

Diacetoxyiodobenzene (iodobenzenediacetate) [3240-34-4] $C_{10}H_{11}IO_4$, **M 322.1**, **m 158°, 163-165°**. The purity of diacetoxyiodobenzene can be checked by treatment with H_2SO_4 then KI, and the liberated I_2 is estimated with standard thiosulfate. It has been recrystallised from 5M acetic acid and dried overnight in a vacuum desiccator over $CaCl_2$. The surface of the crystals may become slightly yellow but this does not affect its usefulness. [Sharefkin & Saltzman *Org Synth Coll Vol 5* 660 1973, DOI: 10.15227/orgsyn.043.0062; *Beilstein* **5** IV 693.] It is a very useful reagent for the synthesis of a variety of heterocyclic compounds [Pradash & Singh *Aldrichimica Acta* **27** 15 1994, Guthikonda et al. *Tetrahedron* **62** 11331 2006, DOI:10.1016/j.tet.2006.07.099].

1,2-Diacetyl benzene [704-00-7] $C_{10}H_{10}O_2$, **M 162.2**, **m 39° to 41°, 41-42°, b 110°/0.1mm, 148°/20mm**.

Purify it by distilling and by recrystallising from petroleum ether. The **bis-2,4-dinitrophenylhydrazone** has **m 221° (dec)**. [Halford & Weissmann *J Org Chem* **17** 1646 1952, DOI: 10.1021/jo50012a012; Riemschneider & Kassahn *Chem Ber* **92** 1705 1959, DOI: 10.1002/cber.19590920735; *Beilstein* **7** III 3501, **7** IV 2155.] **1,4-Diacetyl benzene** [1009-61-6] has **m 111° to 113°, 113-5-114.2°, b 128-130°/3mm**. Crystallise it from EtOH (**m 114°**) or *benzene and dry it in a vacuum over CaCl₂. Also purify it by dissolving it in acetone, treating with Norit, evaporating and recrystallising from MeOH. The **dioxime** has **m 248-259°**. [Wagner et al. *J Am Chem Soc* **108** 7727 1986, DOI: 10.1021/ja00284a041]. [*Beilstein* **7** H 686, **7** II 624, **7** III 3504, **7** IV 2156.]

1,4-Diaminoanthraquinone [128-95-0] **C₁₄H₁₀N₂O₂**, **M 238.3**, **m 265°, 268°, 269°**. Purify the anthraquinone by thin-layer chromatography on silica gel using toluene/acetone (9:1) as eluent. The main band is scraped off and extracted with MeOH. The solvent is evaporated, and the quinone is dried in a drying pistol [Land et al. *JCS Faraday Trans 1* **72** 2091 1976, DOI: 10.1039/F19767202091]. It crystallises from EtOH (**m 269°**) in dark violet crystals. Store it in sealed ampoules in the dark. [*Beilstein* **14** H 197, **14** II 113, **14** III 437, **14** IV 458.] **1,5-Diaminoanthraquinone** [129-44-2] has **m 319°**. Recrystallise it from aniline (**m 313-314°**), EtOH or acetic acid [Flom & Barbara *J Phys Chem* **89** 4489 1985, DOI: 10.1021/j100267a017]. [*Beilstein* **14** H 303, **14** I 467, **14** II 116, **14** III 466, **14** IV 479.] **2,6-Diaminoanthraquinone** [131-14-6] has **m 310-320°**. Crystallise it from pyridine or nitrobenzene (red needles). Column-chromatography on Al₂O₃/toluene is used to remove a fluorescent impurity, then it is recrystallised from EtOH. [*Beilstein* **14** I 471, **14** II 120, **14** III 480, **14** IV 486.]

3,3'-Diaminobenzidine [91-95-2] **C₁₂H₁₄N₄**, **M 214.3**, **m 175°, 177°, pK_{Est(1)} ~3.3, pK_{Est(2)} ~4.7 (free base)**. Dissolve the tetrahydrochloride (below) in H₂O (e.g. 1g in 20ml), cool the solution to 20°, then add it with stirring to a mixture of crushed ice (20g) and aqueous 10N NaOH (2ml), check that the pH is above 10, otherwise add more NaOH (keep cool). Filter off the free base and wash it thoroughly with cold H₂O. Recrystallise it from H₂O or EtOH and dry it over KOH *in vacuo*. Alternatively, liberate the free base with concentrated aqueous NH₃, in which case a dry product can be obtained more readily. [*Beilstein* **13** IV 530.] The free base or its hydrochloride (below) are substrates for peroxidase [Herzog & Fahimi *Anal Biochem* **55** 554 1973, DOI: 10.1016/0003-2697(73)90144-9], stain nucleic acids and are reagents for the spectrophotometric determination of Se [Cheng *Anal Chem* **28** 1738 1956, DOI: 10.1021/ac60119a027]. **3,3'-Diaminobenzidine tetrahydrochloride (2H₂O)** [2H₂O: 7411-49-6; xH₂O: 868272-85-9] **C₁₂H₁₄N₄ · 4HCl · xH₂O**, **M 396.1, 360.1 (anhydr)**, has **m >300°(dec)**, **pK_{Est(1)} ~3.3, pK_{Est(2)} ~4.7 (free base)**. Dissolve the salt in hot water, cool slightly and precipitate it with conc HCl, then dry it over solid NaOH. [*Beilstein* **13** IV 530.]

3,4-Diaminobenzoic acid [619-05-6] **C₉H₈N₂O₂**, **M 152.2**, **m 208°(dec), 210°(dec), 213°(dec), 228-229°, pK₁²⁵ 2.57 (4-NH₂), pK₂²⁵ 3.39 (3-NH₂), pK_{Est(3)} ~5.1 (CO₂H)**. Crystallise it from H₂O or toluene. [*Beilstein* **15** IV 1503.] **3,5-Diaminobenzoic acid** [535-87-5] has **m 228°(dec), 235°(dec), 238°(dec), 240°(dec), pK₂₅ 5.13 (CO₂H), pK₂₅ 7.12 (in 80% aqueous 2-MeOCH₂CH₂OH)**. Crystallise the acid from water to give needles of the *monohydrate*, which loses H₂O above ~100°. The *dihydrochloride* [535-87-5] **C₉H₈N₂O₂ · 2HCl**, **M 225.1**, has **m 226-228°(dec)** [>300°(dec) also reported]. It is used for the detection of *nitrites*. The *ethyl ester*, **m 84°**, crystallises from aqueous EtOH or Et₂O. [*Beilstein* **14** H 453, **14** III 1179, **14** IV 1304.]

3,4-Diaminobenzophenone (4-benzoyl-o-phenylenediamine) [39070-63-8] **C₁₃H₁₂N₂O**, **M 212.3**, **m 116-117°, pK_{Est(1)} ~<0, pK_{Est(2)} ~2.5**. Crystallise it from *C₆H₆/petroleum ether and sublime it *in vacuo* [Ayyanger et al. *Org Prep Proced Int* **23** 627 1991, DOI: 10.1080/00304949109457915; *Beilstein* **14** II 225]. **4,4'-Diaminobenzophenone** [611-98-3] has **m 242-244°, 243-245°, 246.5-247.5° (after sublimation at 0.0006mm), pK₁²⁵ 1.37, pK₂²⁵ 2.92**. Purify the phenone by recrystallisation from EtOH and by sublimation in high vacuum. The *dihydrochloride* has **m 260°(dec)** (from EtOH), and the *thiosemicarbazone* has **m 207-207.5°(dec)** (from aqueous EtOH). [Kuhn et al. *Chem Ber* **75** 711 1942, DOI: 10.1002/cber.19420750614; *Beilstein* **14** IV 255.]

1,2-Diamino-4,5-dichlorobenzene [5348-42-5] **C₆H₆Cl₂N₂**, **M 177.0**, **m 158°, 162°, 162-163°, 163°, 164°, pK_{Est(1)} ~1.0, pK_{Est(2)} ~2.9**. Reflux the amine with activated charcoal in CH₂Cl₂, followed by recrystallisation from Et₂O/petroleum ether or petroleum ether [Koola & Kochi *J Org Chem* **52** 4545 1987, DOI: 10.1021/jo00229a022]. Alternatively, recrystallise the diamine from hexane, *C₆H₆, petroleum ether or H₂O (Na₂SO₄), and sublime it at 150°/15mm. [*Beilstein* **13** IV 72.]

4,4'-Diamino-3,3'-dinitrobiphenyl (3,3'-dinitrobenzidine) [6271-79-0] $C_{12}H_{10}N_4O_4$, M 274.2, m 275°, $pK_{Est} \sim -0.2$. Recrystallise the biphenyl from aqueous EtOH, pyridine/EtOH or nitrobenzene to give scarlet needles. Diazotisation of this benzidine gives the tetrazo-biphenyl which on decomposition in boiling EtOH provided **3,3'-dinitrobiphenyl** [958-96-3] $C_{12}H_8N_2O_4$, M 244.2, m 200°, as orange-yellow needles, after crystallisation from 50% aqueous AcOH [Hodgson & Holt *J Chem Soc* 1431 1934, DOI: 10.1039/JR9340001431; *Beilstein* 13 H 236, 13 I 68, 13 II 108, 13 III 415, 13 IV 387.]

4,4'-Diaminodiphenylamine [537-65-5] $C_{12}H_{13}N_3$, M 199.3, m 158°, $pK_{Est} \sim 5.0$. Crystallise the amine from water (its solubility at 25° is 0.42%). [*Beilstein* 13 H 110, 13 I 36, 13 II 55, 13 III 256, 13 IV 203.]

4,4'-Diaminodiphenyl ether (4,4'-oxydianiline, DADPE) [101-80-4] $C_{12}H_{12}N_2O$, M 200.2, m 185°, 186-187°, 188°, 190°, 192°, 240°/10mm, pK^{25} 5.12 (50% EtOH). This base has been prepared by several syntheses including, condensation of *p*-nitrophenol with *p*-chloroaniline, nitration of diphenyl ether to 4,4'-dinitrodiphenyl ether and reduction with hydrazine and Raney Ni [Shamis & Dashevski *J Org Chem, USSR Engl Transl* 3 1005 1967], a Friedel-Crafts reaction of diphenyl ether (AcCl, $AlCl_3$) to 4,4'-diacetyldiphenyl ether (m 99-100°), conversion to the dioxime (m 180-183°), and a bis-Beckman rearrangement (AcOH, Ac_2O , HCl gas) gives 4,4'-bis-acetamidodiphenyl ether (90%). This amide (46g) in H_2O (600ml) and concentrated HCl (60ml) is refluxed for 2 hours until it all dissolved, filtered (charcoal), the filtrate is basified with Na_2CO_3 solution, the precipitate is filtered off, washed with H_2O , and dried at 100-110° to give the **free base** (32g). It is purified by zone melting and/or by sublimation at high vacuum. In the UV it has λ_{max} at 248 and 300nm in EtOH. [Franovskii et al. *J Org Chem, USSR Engl Transl* 6 2315 1970.] The **benzenesulfonate salt** crystallises from MeOH with m 286-288°(dec). [*Beilstein* 13 H 441, 13 I 148, 13 III 106, 13 IV 1038.]

4,4'-Diaminodiphenylmethane (MDA) [101-77-9] $C_{13}H_{14}N_2$, M 198.3, m 91.6-92°, 92-93°, b 232°/9mm, 398°/atm, $pK_{Est} \sim 4.9$. Crystallise the amine from water, 95% EtOH or *benzene. [*Beilstein* 13 IV 390.] Care, as it can cause liver damage. It inhibits corrosion.

2,7-Diaminofluorene [524-64-4] $C_{13}H_{12}N_2$, M 196.3, m 160°, 162°, 165°, $pK_{Est} \sim 4.6$. Recrystallise it from H_2O . [*Beilstein* 13 IV 449.] It is used as an analytical reagent for bromide, chloride, nitrate, Cd^{2+} , Cu^{2+} , Co^{2+} and Zn^{2+} .

1,4-Diaminonaphthalene [2243-61-0] $C_{10}H_{10}N_2$, M 158.2, m 114°, 116°, pK_1^{25} 5.87, pK_1^{25} 2.78. Purify as for the 1,5-somer below. It forms dark green to black crystals which are insoluble in H_2O . On prolonged standing in air it oxidises (to the quinon-imine then to 1,4-naphthoquinone?). It is prepared commercially from 1,4-dibromonaphthalene (40g), 25% ammonia (350ml), and a catalyst [e.g. anhydrous CuCl (6g)] in an autoclave at 150°/6hrs in ~60% yield. [Zechlin et al. US Pat 6,538,158 B2 2003, to Bayer Aktiengesellschaft.] **1,5-Diaminonaphthalene** [2243-62-1] has m 185°, 187°, 190°, pK^{25} 4.12, $pK_{Est(2)}$ ~2.0. Recrystallise the amino-naphthalene from boiling H_2O , but this is wasteful due to poor solubility. Boil it in chlorobenzene (charcoal), filter hot and cool the filtrate (preferably under N_2). This gives colourless crystals. Dry it in a vacuum till free from chlorobenzene (odour), and store it in sealed ampoules under N_2 away from light. [*Beilstein* 13 IV 340.] **1,8-Diaminonaphthalene** [479-27-6] has m 60°, 65°, 66.5°, b 205°/12mm, pK^{25} 4.44 (in 1-0% aqueous EtOH). Crystallise 1,8-diaminonaphthalene from water or aqueous EtOH, and sublime it in a vacuum. It is an antioxidant used in lubricating oils, and for the detection of nitrite and Se. The **dihydrochloride** m 280°, forms leaflets from aqueous HCl. The N,N' -dimethyl derivative [20734-56-9] has m 103-104° and pK^{25} 5.61, the N,N,N',N' -trimethyl- derivative [20734-57-0] has m 29-30° and pK^{25} 6.43. [Hodgson et al. *J Chem Soc* 202 1945, DOI: 10.1039/JR9450000202; *Beilstein* 13 IV 344.] **2,3-Diaminonaphthalene** [771-97-1] has m 199°, pK^{21} 3.54 (in 50% aqueous EtOH). Crystallise the diamine from water, or dissolve it in 0.1M HCl, by heating to 50°. After cooling, the solution is extracted with decalin to remove fluorescent impurities, and centrifuged. [*Beilstein* 13 IV 346.]

2,5-Di-tert-amylhydroquinone [79-74-3] $C_{16}H_{26}O_2$, M 250.4, m 179-180°, 185.8-186.5°. Crystallise the hydroquinone under N_2 from boiling AcOH (7ml/g) plus boiling water (2.5ml/g). [Stolow & Bonaventura *J Am Chem Soc* 85 3636 1963, DOI: 10.1021/ja00905a023]. Store it in sealed ampoules under N_2 away from light. It is an antioxidant. [*Beilstein* 6 H 952, 6 III 4748.]

Di-n-amyl phthalate (dipentyl phthalate) [131-18-0] $C_{18}H_{26}O_4$, **M 306.4**, **b 204-206°/11mm**, d_4^{25} **1.023**, n_D^{20} **1.489**. Wash the ester with aqueous Na_2CO_3 , then distilled water. Dry it with $CaCl_2$ and distil it in a vacuum. Store it in a vacuum desiccator over P_2O_5 . [Beilstein **9** IV 3178.]

Diazoaminobenzene (1,3-diphenyltriazene) [136-36-6] $C_{12}H_{11}N_3$, **M 197.2**, **m 99°, 100°**. Crystallise the triazene from petroleum ether (b 60-80°) (**m 94-96°**), 60% MeOH/water or 50% aqueous EtOH (charcoal) containing a small amount of KOH. Its solubility in petroleum ether (b 60-80°) is ~6%. Also purify it by chromatography on alumina/toluene and elute with toluene/petroleum ether. Store the pale yellow needles in the dark. [Hartman & Dickey *Org Synth Coll Vol 2* 163 1943, DOI: 10.15227/orgsyn.014.0024; Beilstein **16** H 687, **16** I 404, **16** II 351, **16** III 643, **16** IV 904.]

Dibenzalacetone [DBA, bda, trans-trans-1,5-diphenyl-1,4-dien-3-one, 1,5-(bisphenyl)-penta-1E,4E-diene-3-one] [538-58-9] $C_{17}H_{14}O$, **M 234.3**, **m 107°, 111°, 112°, 113°, 120-122°**. Purify the ketone by flash chromatography (150 mesh Al_2O_3 deactivated with 6% v/w H_2O) using petroleum ether 40-60°/EtOAc (4/1, v/v), and the yellow solid is recrystallised by layering a concentrated CH_2Cl_2 solution with hexane (i.e. CH_2Cl_2 /hexane, 1:4, v/v). Also recrystallise the ketone from hot ethyl acetate (2.5ml/g) or EtOH. It has been also crystallised from petroleum ether b 30-60°. The IR (CH_2Cl_2) has ν_{max} at 1657m, 1651m (C=O), 1627vs (C=C), 1591w (C=C aromatic), 1574w (C=C aromatic), 983m (C=C trans) cm^{-1} ; and UV has λ_{max} (THF) at 233 ($\pi-\pi^*$) and 321 (n- π^*) nm; the 1H NMR (400MHz, $CDCl_3$) has δ at 7.65-7.15 (m, 10 H, ArH), 4.4 (dd, J = 5.8, 10.0 Hz, 1 H, C3a-H), 3.45-3.30 (m, 1 H, C6-H), 3.15-3.00 (m, 1 H, C6-H), 1.90-1.55 (m, 3 H, C4-H, C5-H₂), 0.95-0.75 (m, 1 H, C4-H), 0.40 (s, 3 H, B-CH₃). The **trans-trans-2,4-dinitrophenylhydrazone** has **m 181°**. [Conrad & Dolliver *Org Synth Coll Vol 2* 167 1943, DOI: 10.1002/9780470132487.ch37; Beilstein **7** IV 1747.] **Cis-cis-1,5-diphenyl-1,4-dien-3-one (Z,Z-isomer)**, prepared by catalytic reduction of 1,5-diphenylpentadiyne-3-one using Lindlar's catalyst in 60% yield, is a yellow oil **b 130°/0.02mm**, whose **2,4-dinitrophenylhydrazone**, prepared under mild conditions (in the absence of light), has **m 92°**; whereas under the usual conditions it gave the **trans-trans-2,4-dinitrophenylhydrazone**. Acid hydrolysis of the **cis-cis-hydrazone** m 92° gave the **trans-trans**-ketone. Catalytic reduction of 1,5-diphenylpent-1-ene-4-yn-3-one with Lindlar's catalyst gave **cis-trans-1,5-diphenyl-1,4-dien-3-one (Z,E-isomer)**, in 50% yield which crystallised from EtOH in yellow needles **m 60°**, and its **2,4-dinitrophenylhydrazone** prepared in the usual manner has **m 159°**. Exposing the **cis-cis**-ketone to normal light for 24 hours caused complete conversion to the **trans-trans**-ketone, whereas no isomerisation occurred during one month in the absence of light. **Cis-trans-1,5-diphenylpentadiene-3-one** remained unchanged after several days exposure to normal light. On heating the **cis-cis**-ketone in MeOH with concentrated hydrochloric acid at 100° for 5 minutes produced an ~1:1 mixture of **cis-trans**- and **trans-trans**- isomers. There is no evidence of light catalysed isomerisations. The three geometrical isomers have different NMR, IR and UV spectra, e.g. in the IR (KBr?) the **Z,Z**-isomer has ν_{max} at 971 (10.3 μ) cm^{-1} for **trans**-C=C [compare with above 983 cm^{-1} in CH_2Cl_2 solution], a band which is absent in the **E,E**-isomer and replaced by a band at ν_{max} at 761 (13.15 μ) cm^{-1} for **cis**-C=C frequency which is also present in the **E,Z**-isomer. Furthermore in the UV the **Z,Z**-isomer has λ_{max} at 330nm (ϵ 34,000), the **E,Z**-isomer has λ_{max} at 295nm (ϵ 20,000) and the **E,E**-isomer has λ_{max} at 287nm (ϵ 11,000). [Dinwiddie et al. *J Org Chem* **27** 327 1962, DOI: 10.1021/jo01048a529.] These dienones readily form carbocyclic and heterocyclic compounds, and are useful ligands for making stable zero-valent neutral complexes with Pd(0), e.g. $Pd(dba)_2$ and $Pd(dba)_3$ [Rettig & Maitlis *Inorg Synth* **17** 134 1977, Ukai et al. *J Organomet. Chem* **65** 253 1974, DOI:10.1016/S0022-328X(00)91277-4,] which are important catalysts [see Chapter 5 in this book, and J. Tsuji *Organic Synthesis with Palladium Compounds* Springer Verlag, Berlin, 1980 E-I. Negishi (Ed), *Handbook of Organopalladium for Organic Synthesis, Vol 1 and 2*, J. Wiley & Sons, NY, 2002. ISBN 471315060.]

Dibenz[*a,h*]anthracene (1,2:5,6-dibenzanthracene) [53-70-3] $C_{22}H_{14}$, **M 278.4**, **m 262°, 265°, 266-267°, b 524°/atm**. The yellow-green colour (due to other pentacyclic impurities) is removed from it by crystallising from *benzene or by selective oxidation with lead tetraacetate in acetic acid [Moricon et al. *J Am Chem Soc* **82** 3441 1960, DOI: 10.1021/ja01498a051]. It can be recrystallised from AcOH and sublimed. [Beilstein **5** IV 2722.]

trans-1,2-Dibenzoyl ethylene (trans-1,4-diphenyl-2-butene-1,4-dione) [959-28-4] $C_{16}H_{12}O_2$, **M 236.3**, **m 109°, 111°, 112°**. It crystallises from MeOH or EtOH as yellow needles [Keller et al. *Helv Chim Acta* **29** 512

1946, DOI: 10.1002/hlca.19460290304]. The *dioxime* has **m 210-211°(dec)** from AcOH. [IR: Kuhn et al. *J Am Chem Soc* **72** 5058 1950, DOI: 10.1021/ja01167a066; Yates *J Am Chem Soc* **74** 5376 1952, DOI: 10.1021/ja01141a047; Erickson et al. *J Am Chem Soc* **73** 5301 1951, DOI: 10.1021/ja01155a086; [Lutz *Org Synth Coll Vol 3* 248 1955, DOI: 10.15227/orgsyn.020.0029; *Beilstein 7 IV* 2578.]

Dibenzoylmethane (1,3-diphenyl-1,3-propanedione) [120-46-7] $C_{15}H_{12}O_2$, **M 224.3**, **m 77°, 79°, 80°, 219-221°/18mm**. Crystallise dibenzoylmethane from petroleum ether or MeOH. The *oxime*, **m 165°**, crystallises from Et₂O. [Allen et al. *Org Synth Coll Vol 1* 205 1941, DOI: 10.15227/orgsyn.008.0060; *Beilstein 7 IV* 2512.]

2,3,6,7-Dibenzphenanthrene (pentaphene) [222-93-5] $C_{22}H_{14}$, **M 276.3**, **m 255-256°, 257°**. Purify pentaphene through Al₂O₃ with *C₆H₆ or xylene as eluent. It crystallises from xylene as yellow plates and sublimes in high vacuum. The *dipicrate* forms orange needles **m 184°** from *C₆H₆. (Clar & John *Ber* **64** 986 1931, Clar & Stewart *J Chem Soc* 3215 1951, DOI: 10.1039/JR9450000202; Marsili & Isola *Tetrahedron Lett* 3023 1965, DOI: 10.1016/S0040-4039(01)89252-2; Franck & Zander *Chem Ber* **99** 396 1966, DOI: 10.1002/cber.19660990205.]

Dibenzyl amine [103-49-1] $C_{14}H_{15}N$, **M 197.3**, **m -26°, b 113-114°/0.1mm, 174-175°/6mm, 270°/250mm, 300° (partial dec)**, **d₄²⁰ 1.027**, **n_D²⁰ 1.576**, **pK²⁵ 8.52**. Purify the amine by distillation in a vacuum. **It causes burns to the skin.** The *dihydrochloride* has **m 265-266°** (from MeOH/HCl), and the *tetraphenyl boronate* has **m 129-133°**. [Bradley & Maisey *J Chem Soc* 247 1954, DOI: 10.1039/JR9540000247; Hall *J Phys Chem* **60** 63 1956, DOI: 10.1021/j150535a017; Donetti & Bellora *J Org Chem* **37** 3352 1972, DOI: 10.1021/jo00986a036; *Beilstein 12 IV* 2179.] It is used for the detection of cobalt, iron and cyanate ion.

N,N'-Dibenzylethylenediamine (benzathine, DBED) [140-28-3] $C_{16}H_{30}N_2$, **M 240.4**, **m 26°, b 195°/4mm, d₄²⁰ 1.02**, **n_D²⁰ 1.563**, **pK_{Est(1)} ~ 5.9**, **pK_{Est(2)} ~ 8.9**. Dissolve DBED in acid, extract with toluene, basify, extract it with Et₂O, dry over solid KOH, evaporate and fractionate it *in vacuo*. The *diacetate* crystallises from H₂O by addition of EtOH and has **m 111°** (solubility in H₂O is ~25%). The *dihydrochloride* has **m 306-308°** (from H₂O) and the *dipicrate* has **m 212°(dec)** (from H₂O). [Frost et al. *J Am Chem Soc* **71** 3842 1949, DOI: 10.1021/ja01179a510; *Beilstein 12 H* 1067, **12 III** 2304.]

Dibenzyl ketone (1,3-diphenyl-2-propanone) [102-04-5] $C_{15}H_{14}O$, **M 210.3**, **m 32°, 34°, 330°/atm**. Fractionally crystallise it from its melt, then crystallise it from petroleum ether. It has been purified by recrystallisation from anhydrous Et₂O at -70°. Store it in the dark. [*Beilstein 7 IV* 1420.]

Dibenzyl malonate [15014-25-2] $C_{17}H_{16}O_4$, **M 284.3**, **b 188-190°/0.2mm, 193-196°/1mm, d₄²⁰ 1.158**, **n_D²⁰ 1.5452**. Dissolve the ester in toluene, wash it with aqueous NaHCO₃, H₂O, dry over MgSO₄, filter, evaporate and distil it at high vacuum. [Ginsburg & Pappo *J Am Chem Soc* **75** 1094 1953, DOI: 10.1021/ja01101a025; Baker et al. *J Org Chem* **17** 77 1952, DOI: 10.1021/jo01135a009; *Beilstein 6 IV* 2270.]

2,4-Dibromoaniline [615-57-6] $C_6H_5Br_2N$, **M 250.9**, **m 79°, 80°, pK²⁵ 1.87**. Crystallise the aniline from aqueous EtOH. The *picrate* has **m 124°**. [*Beilstein 12 H* 655, **12 I** 326, **12 II** 356, **12 III** 1471, **12 IV** 1532.]

9,10-Dibromoanthracene [523-27-3] $C_{14}H_8Br_2$, **M 336.0**, **m 222-224°, 226°**. Recrystallise it from toluene, xylene or CCl₄ (yellow needles), and sublime it in a vacuum [Johnston et al. *J Am Chem Soc* **109** 1291 1987, DOI: 10.1021/ja00239a001]. [Heilbron & Heaton *Org Synth Coll Vol 1* 207 1941, DOI: 10.15227/orgsyn.003.0041; *Beilstein 5 H* 665.]

1,4-Dibromobenzene [106-37-6] $C_6H_4Br_2$, **M 235.9**, **m 87.8°**. Steam distil the dibromobenzene, then crystallise it from EtOH or MeOH and dry it in the dark under vacuum. Purify it by zone melting. [*Beilstein 5 IV* 683.]

2,5-Dibromobenzoic acid [610-71-9] $C_7H_4Br_2O_2$, **M 279.9**, **m 156°, 157°, 159°, pK_{Est} ~1.5**. Crystallise the acid from water or EtOH. [*Beilstein 9 H* 358, **9 I** 147, **9 II** 237, **9 III** 1428, **9 IV** 1027.]

(±)-2,2'-Dibromo-1,1'-binaphthyl [74866-28-7, (±) 76284-65-6] $C_{20}H_{12}Br_2$, **M 412.1, m 178-183°, 180.5-181°**. Purify the binaphthyl by chromatography through a silica gel column (70-230mesh) using hexane as eluent. It gives pale yellow crystals from EtOH with **m 187.3-187.9°**. The ***R*-(+)-enantiomer** [86688-08-6] crystallises from hexane with **m 157-157.5°** and $[\alpha]_D^{25} +32.9$ (c 1, pyridine) [Brown et al. *J Org Chem* **50** 4345 1985, DOI: 10.1021/jo00222a029]. [Okamoto et al. *J Am Chem Soc* **103** 6971 1981, DOI: 10.1021/ja00413a038; *Beilstein* **5** III 2465.]

4,4'-Dibromobiphenyl [92-86-4] $C_{12}H_8Br_2$, **M 312.0, m 163°, 164°, 165°, b 355-360°/760mm**. Crystallise it from MeOH. [*Beilstein* **5** IV 1820.]

α,α -Dibromodeoxybenzoin (α,α -dibromobenzyl phenyl ketone) [15023-99-1] $C_{14}H_{10}Br_2O$, **M 354.0, m 111.8-112.7°**. Crystallise α,α -dibromo deoxybenzoin from acetic acid, 50% aqueous EtOH (prisms, **m 109-112°**) or Et₂O (**m 112°**). [Curtius & Lang *J Prakt Chem* **44** 547 1891, DOI: 10.1002/prac.18910440147; *Beilstein* **7** H 436, **7** III 2114.]

2,5-Dibromonitrobenzene [3460-18-2] $C_6H_3Br_2NO_2$, **M 280.9, m 84°, 85-86°**. It crystallises from Me₂CO or EtOH. [*Beilstein* **5** H 250, **5** II 190, **5** III 621, **5** IV 732.]

2,6-Dibromo-4-nitrophenol [99-28-5] $C_6H_3Br_2NO_3$, **M 296.9, m 143-144°, pK²⁵ 3.39**. Crystallise the phenol from aqueous EtOH, 50% aqueous AcOH (**m 144-145°**, dec varies with rate of heating). Dry it in an oven at 40-60° or *in vacuo* over NaOH. [Hartman & Dickey *Org Synth Coll Vol* **2** 173 1943, DOI: 10.15227/orgsyn.015.0006; *Beilstein* **6** H 246, **6** I 123, **6** II 234, **6** III 849, **6** IV 1366.]

2,4-Dibromophenol [615-58-7] $C_6H_4Br_2O$, **M 251.9, m 35°, 37°, 38°, 41-42°, b 154°/10mm, 239°/760mm, pK²⁵ 7.79**. Crystallise the phenol from CHCl₃ at -40°, or distil it in a vacuum. [*Beilstein* **6** H 202, **6** I 106, **6** II 188, **6** III 753, **6** IV 1061.] **2,6-Dibromophenol** [608-33-3] has **m 53°, 56-57°, b 138°/10mm, 255-256°/740mm, pK²⁵ 6.67**. Distil the phenol under vacuum (at 10mm), then crystallise it from cold CHCl₃ or from EtOH/water. [*Beilstein* **6** H 202, **6** I 106, **6** II 188, **6** III 755, **6** IV 1064.]

α,α' -Dibromo-*o*-xylene [91-13-4] $C_8H_8Br_2$, **M 264.0, m 91°, 94°, 95°, 95-96°, 98-99°, b 129-130°/4.5mm**. Crystallise it from CHCl₃ or petroleum ether, and/or distil it under vacuum. [Wenner *J Org Chem* **17** 523 1952, DOI: 10.1021/jo01138a003; *Beilstein* **5** H 366, **5** I 180, **5** II 285, **5** III 819, **5** IV 929.] **α,α' -Dibromo-*m*-xylene** [626-15-3] has **m 75°, 77°, 78-79°, b 156-160°/12mm, 135-140°/20mm** (also reported). Crystallise it from acetone or *benzene, and fractionally distil it under vacuum. [Wenner *J Org Chem* **17** 523 1952, DOI: 10.1021/jo01138a003; *Beilstein* **5** H 374, **5** I 184, **5** II 294, **5** III 839, **5** IV 946.] **α,α' -Dibromo-*p*-xylene** [623-24-5] has **m 143.4°, 142-144°, 145-147°, b 155-158°/12-15mm, 245°/760mm**. Distil it under a vacuum and recrystallise it from EtOH, *benzene or chloroform. [Wenner *J Org Chem* **17** 523 1952, DOI: 10.1021/jo01138a003; *Beilstein* **5** H 385, **5** I 187, **5** II 301, **5** III 859, **5** IV 970.]

α -Dibutylamino- α -(4-methoxyphenyl)acetamide (Ambucetamide) [519-88-0] $C_{17}H_{28}N_2O_2$, **M 292.4, m 134°**. Crystallise ambucetamide from EtOH containing 10% diethyl ether. [Janssen *J Am Chem Soc* **76** 6192 1954, DOI: 10.1021/ja01652a098; *Beilstein* **14** IV 2101.] Antispasmodic.

2,5-Di-*tert*-butyl aniline [21860-03-7] $C_{14}H_{23}N$, **M 205.4, m 103°, 104°, 103-106°, pK²⁵ 3.34 (50% aqueous MeOH), 3.58 (90% aqueous MeOH)**. The aniline recrystallises from EtOH in fine needles after steam distillation. It has a pK_a²⁵ of 3.58 (50% aqueous EtOH). The ***tosylate*** has **m 164°** (from AcOH). [Bell & Wilson *J Chem Soc* 2340 1956, DOI: 10.1039/JR9560002340; Carpenter et al. *J Org Chem* **16** 586 1951, DOI: 10.1021/jo01144a011; Bartlett et al. *J Am Chem Soc* **76** 2349 1954, DOI: 10.1021/ja01638a019; *Beilstein* **12** IV 2891.]

***p*-Di-*tert*-butylbenzene** [1012-72-2] $C_{14}H_{22}$, **M 190.3, m 77°, 78°, 80°, 236°/760mm**. Crystallise it from Et₂O or EtOH and dry it under vacuum over P₂O₅ at 55°. [Tanner et al. *J Org Chem* **52** 2142 1987, DOI: 10.1021/jo00387a005; *Beilstein* **5** II 344.]

2,6-Di-*tert*-butyl-1,4-benzoquinone [719-22-2] $C_{14}H_{20}O_2$, M 220.3, m 65°, 66°, 67°. It can be recrystallised from MeOH and sublimes in a vacuum. [Beilstein 7 IV 2116.] **3,5-Di-*tert*-butyl-1,2-benzoquinone** [3383-21-9] has m 112-114°, 113-114°. It can be recrystallised from MeOH or petroleum ether, and forms fine red plates or rhombs. [Flaig et al. *Justus Liebigs Ann Chem* 597 196 1955, DOI: 10.1002/jlac.19555970304; IR: Ley & Müller *Chem Ber* 89 1402 1956, DOI: 10.1002/cber.19560890607; Beilstein 7 IV 2113.]

3,5-Di-*tert*-butyl catechol [1020-31-1] $C_{14}H_{22}O_2$, M 222.3, m 96°, 97°, 99°, 99-100°, $pK_{Est(1)} \sim 11.0$, $pK_{Est(2)} \sim 13.1$. Recrystallise the catechol from petroleum ether. [Ley & Müller *Chem Ber* 89 1402 1956, DOI: 10.1002/cber.19560890607; UV Flaig et al. *Z Naturforschung* 10b 668 1955.] Also purify it by crystallising three times from pentane [Funabiki et al. *J Am Chem Soc* 108 2921 1986, DOI: 10.1021/ja00271a022].

2,6-Di-*tert*-butyl-*p*-cresol (**2,6-di-*tert*-butyl-4-methylphenol**, **butylatedhydroxytoluene**, **BHT** or **DBPC**) [128-37-0] $C_{15}H_{24}O$, M 220.4, m 71.5°, pK^{25} 12.23. Dissolve BHT in *n*-hexane at room temperature, then cool with rapid stirring, to -60°. The precipitate is separated, redissolved in hexane, and the process is repeated until the mother liquor is no longer coloured. The final product is stored under N_2 at 0° [Blanchard *J Am Chem Soc* 82 2014 1960, DOI: 10.1021/ja01493a041]. It has also been recrystallised from EtOH, MeOH, *benzene, *n*-hexane, methylcyclohexane or petroleum ether (b 60-80°), and is dried in a vacuum. [Beilstein 6 IV 3511.] It is a very effective antioxidant.

2,6-Di-*tert*-butyl-4-dimethylaminomethylphenol [88-27-7] $C_{17}H_{29}NO$, M 263.4, m 93-94°, b 172°/30mm, $pK_{Est} \sim 12.0$. Crystallise it from *n*-hexane. [Beilstein 13 IV 2014.]

Di-*tert*-butyldiperphthalate [2155-71-7] $C_{16}H_{22}O_6$, M 310.3, m (48°) 57-57.5°, decomposes at 108°. Crystallise the perphthalate from Et_2O or petroleum ether and dry it over H_2SO_4 . The IR has ν_{max} at 1772cm^{-1} in CCl_4 . [Milas & Surgenor *J Am Chem Soc* 68 642 1946, DOI: 10.1021/ja01208a033; Klein & Milas *J Org Chem* 36 2900 1971, DOI: 10.1021/jo00818a044; Beilstein 9 III 4190, 9 IV 3260.] **CARE**, potentially **EXPLOSIVE**.

2,6-Di-*tert*-butyl-4-ethylphenol [4130-42-1] $C_{16}H_{26}O$, M 234.4, m 42-44°, $pK_{Est} \sim 12.3$. Crystallise the phenol from aqueous EtOH or *n*-hexane. [Beilstein 6 IV 3529.]

2,5-Di-*tert*-butylhydroquinone [88-58-4] $C_{14}H_{22}O_2$, M 222.3, m 222-223°. Crystallise the hydroquinone from * C_6H_6 or AcOH. [Beilstein 6 III 4741.]

2,6-Di-*tert*-butyl-4-isopropylphenol [5427-03-2] $C_{17}H_{28}O$, M 248.4, m 39-41°, 38-42°, b 105-106°/0.3mm, $pK_{Est} \sim 12.3$. Crystallise from *n*-hexane or aqueous EtOH. It is used for making the respective *phenoxyl radical*. [Cook & Norcross *J Am Chem Soc* 78 3797 1956, DOI: 10.1021/ja01596a064; Beilstein 6 III 3534.]

2,6-Di-*tert*-butylphenol [128-39-2] $C_{14}H_{22}O$, M 206.3, m 37-38°, pK^{25} 11.70. Crystallise the phenol from aqueous EtOH or *n*-hexane. [Beilstein 6 III 2061.]

Dibutyl phthalate (**DBP**, **butyl phthalate**) [84-74-2] $C_{16}H_{22}O_4$, M 278.4, m -35°, b 44°/2.5x10⁻⁴mm, 182°/5mm, 206°/20mm, 340°/760mm, d_4^{20} 1.4929, d_5^{25} 1.0426, n_D^{25} 1.490. Wash DBP with H_2O (to free it from alcohol), then dilute NaOH (to remove any butyl hydrogen phthalate or acid), aqueous $NaHCO_3$ (charcoal), then distilled water. Dry it ($CaCl_2$), distil it at 10torr or less, and store it in a desiccator over P_2O_5 . [Beilstein 9 II 586, 9 III 4102, 9 IV 3175.] It is an insect repellent.

2,4-Dichloroaniline [554-00-7] $C_6H_5Cl_2N$, M 162.0, m 59°, 60°, 63°, b 245°/atm, pK^{25} 2.02. Crystallise the aniline from EtOH/water. It also crystallises from EtOH and is dried *in vacuo* for 6 hours at 40° [Moore et al. *J Am Chem Soc* 108 2257 1986, DOI: 10.1021/ja00269a022; Edidin et al. *J Am Chem Soc* 109 3945 1987, DOI: 10.1021/ja00247a019]. [Beilstein 12 IV 1241.] **3,4-Dichloroaniline** [95-76-1] has m 69°, 70°, 71.5°, b 272°/atm, pK^{25} 2.97. Crystallise the aniline from MeOH, and/or distil it. [Beilstein 12 IV 1257.]

9,10-Dichloroanthracene [605-48-1] $C_{14}H_8Cl_2$, M 247.1, m 214-215°. Purify it by recrystallising it from

MeOH, EtOH, *C₆H₆ or Me₂CO (m 210-211°) followed by subliming *in vacuo*. [Masnori & Kochi *J Am Chem Soc* **107** 7880 1985, DOI: 10.1021/ja00312a014; *Beilstein* **5** H 664, **5** I 324, **5** II 575, **5** III 2134, **5** IV 2293.]

2,4-Dichlorobenzaldehyde [874-42-0] C₇H₄Cl₂O, M **175.0**, m **64°**, **67°**, **69°**, b **233°/atm**. Crystallise the aldehyde from EtOH or ligroin. It has been distilled at atmospheric pressure. [*Beilstein* **7** IV 575.] **2,6-Dichlorobenzaldehyde** [83-38-5] has m **69°**, **70.5-71.5°**. Crystallise the aldehyde from EtOH/H₂O or petroleum ether (b 30-60°). [*Beilstein* **7** IV 576.]

1,2-Dichlorobenzene (ODCB) [95-50-1] C₆H₄Cl₂, M **147.0**, m **-18° to -17°**, b **81-82°/31-32mm**, **180.5°/760mm**, d₄²⁰ **1.306**, n_D²⁰ **1.551**, n_D²⁵ **1.549**. Contaminants may include the *p*-isomer and trichlorobenzene [Suslick et al. *J Am Chem Soc* **106** 4522 1984, DOI: 10.1021/ja00328a036]. It should be shaken with concentrated or fuming H₂SO₄(CARE), washed with water, dried with CaCl₂, and distilled from CaH₂ or sodium in a glass-packed column. Low conductivity material (*ca* 10⁻¹⁰ mhos) has been obtained by refluxing with P₂O₅, fractionally distilling and passing it through a column packed with silica gel or activated alumina: it is stored in a dry-box under N₂ or with activated alumina. [*Beilstein* **5** IV 654.] **1,3-Dichlorobenzene** [541-73-1] has m **-25° to -22°**, b **173.0°/atm**, d₄²⁰ **1.289**, n_D²⁰ **1.54586**, n_D²⁵ **1.54337**. Wash it with aqueous 10% NaOH, then with water until neutral, dry and distil it. Conductivity material (*ca* 10⁻¹⁰ mhos) has been prepared by refluxing over P₂O₅ for 8 hours, then fractionally distilling, and storing with activated alumina. *m*-Dichlorobenzene dissolves rubber stoppers. [*Beilstein* **5** IV 657.] **1,4-Dichlorobenzene** [106-46-7] has m **52°, 53°, 54°**, b **174.1°/atm**, d₄²⁰ **1.241**, n_D⁶⁰ **1.52849**. *o*-Dichlorobenzene is a common impurity. The *p*-isomer has been purified by steam distillation, crystallisation from EtOH or boiling MeOH, air-dried and dried in the dark under vacuum. It has also been purified by zone refining. [*Beilstein* **5** IV 658.]

2,2'-Dichlorobenzidine [84-68-4] C₁₂H₁₀Cl₂N₂, M **253.1**, m **165°**, pK_{Est(1)} **~3.0**, pK_{Est(2)} **~4.0**. Crystallise the benzidine from EtOH or H₂O. [*Beilstein* **13** H 234, **13** I 66, **13** II 106, **13** III 477, **13** IV 384.] **3,3'-Dichlorobenzidine** [91-94-1] has m **132-133°**, pK_{Est(1)} **~4.8**, pK_{Est(2)} **~5.7**. Crystallise the benzidine from EtOH, petroleum ether (m 133°) or *benzene. [*Beilstein* **13** H 234, **13** I 67, **13** II 106, **13** III 477, **13** IV 384.] **CARCINOGEN**.

2,3-Dichlorobenzoic acid [50-45-3] C₇H₄Cl₂O₂, M **191.0**, m **168°, 168.3°, 169-170°**, pK²⁵ **2.67**. Aromatic acid impurities (to <0.05%) can be removed *via* the (±)-α-methylbenzylamine salt as described for 2,4-dichlorobenzoic acid [Ley & Yates *Organic Process Research & Development* **12** 120 2008, DOI: 10.1021/op7001547.] Crystallise the acid from H₂O, aqueous EtOH, or 30% aqueous AcOH and several times from *C₆H₆, then dry it *in vacuo* at 40° overnight. The *methyl ester* has m **35-39°**. [Hope & Riley *J Chem Soc* **123** 2470 1923, DOI: 10.1021/op7001547; Lock *Monatsh Chem* **90** 683 1959, DOI: 10.1007/BF00902394; Mather & Shorter *J Chem Soc* 4744 1961, DOI: 10.1039/JR9610004744; *Beilstein* **9** II 228, **9** IV 998.] **2,4-Dichlorobenzoic acid** [50-84-0] has m **160°, 163-164°**, pK²⁵ **2.68**. Crystallise the acid from aqueous EtOH (charcoal), then *benzene (charcoal). It can also be recrystallised from water. [*Beilstein* **9** IV 998.] It can be freed from isomeric acids (to <0.05%) *via* the (±)-α-methylbenzylamine salt as follows: dissolve the dichloro-acid (10g, 50.2mmol) in isopropanol (200ml), heat to 60° and add the (±)-benzylamine (5.49g, 45.3mmol), then stir it at 60° for 1 hour. Cool the mixture to room temperature, filter the slurry, wash it with isopropanol (25ml) and dry it *in vacuo* at 40° overnight to give 79% of the *salt* with m **185.2°**. Dissolve the salt (5g) in H₂O (50ml) and MeOH (20ml), then heat to 60° and add concentrated HCl to pH <2.0. Cool the solution to room temperature add H₂O (12ml), filter it, wash it with H₂O (30ml) and dry it *in vacuo* at 40° overnight to give 94% of the *acid* with m **162.0°**. [Ley & Yates *Organic Process Research & Development* **12** 120 2008, DOI: 10.1021/op7001547.] **2,5-Dichlorobenzoic acid** [50-79-3] has m **152°, 154°**, b **301°/760mm**, pK²⁵ **2.47**. Crystallise the acid from water. [*Beilstein* **9** IV 1005.] Aromatic acid impurities (to <0.05%) can be removed *via* the (±)-α-methylbenzylamine salt as described for 2,4-dichlorobenzoic acid [Ley & Yates *Organic Process Research & Development* **12** 120 2008, DOI: 10.1021/op7001547.] **2,6-Dichlorobenzoic acid** [50-30-6] has m **139°, 140°, 141-142°**, pK²⁵ **1.59**. Crystallise the acid from EtOH and sublime it *in vacuo*. [*Beilstein* **9** IV 1005.] Aromatic acid impurities (to <0.05%) can be removed *via* the (±)-α-methylbenzylamine salt as described for 2,4-dichlorobenzoic acid [Ley & Yates *Organic Process Research & Development* **12** 120 2008, DOI: 10.1021/op7001547.] **3,4-Dichlorobenzoic acid** [51-44-5] has m **204°, 205°, 206-207°**, pK²⁵ **3.64**. Recrystallise the acid from aqueous EtOH (charcoal) or acetic acid. [*Beilstein* **9** IV 1006.] Aromatic acid

impurities (to <0.05%) can be removed *via* the (±)-α-methylbenzylamine salt as described for 2,4-dichlorobenzoic acid [Ley & Yates *Organic Process Research & Development* **12** 120 2008, DOI: 10.1021/op7001547.] **3,5-Dichlorobenzoic acid** [51-36-5] has **m 185°, 187°, 188°, pK²⁵ 3.54**. Crystallise the acid from EtOH and sublime it in a vacuum. [Beilstein **9** IV 1008.] Aromatic acid impurities (to <0.05%) can be removed *via* the (±)-α-methylbenzylamine salt as described for 2,4-dichlorobenzoic acid [Ley & Yates *Organic Process Research & Development* **12** 120 2008, DOI: 10.1021/op7001547.]

2,6-Dichlorobenzonitrile (Diclobenil) [1194-65-6] **C₇H₃Cl₂N**, **M 172.0**, **m 143°, 145°, 146°**. Crystallise the nitrile from acetone or petroleum ether. It is very poorly soluble in H₂O, being 18mg/L at 20°. [Beilstein **9** IV 1006.] It is used as a herbicide.

4,4'-Dichlorobenzophenone (4,4'-DBP) [90-98-2] **C₁₃H₈Cl₂O**, **M 251.1**, **m 144°, 145-146°, b 353°/atm**. Recrystallise it from EtOH [Wagner et al. *J Am Chem Soc* **108** 7727 1986, DOI: 10.1021/ja00284a041]. The *oxime* has **m 135.2-136.9°** (from MeOH, Sieger & Klein *J Org Chem* **22** 951 1957, DOI: 10.1021/jo01359a026). The *semicarbazone* has **m 192-193°** (from H₂O). [Beilstein **7** H 420, **7** I 228, **7** II 359, **7** III 2076, **7** IV 1376.]

2,5-Dichloro-1,4-benzoquinone [615-93-0] **C₆H₂Cl₂O₂**, **M 177.0**, **m 160°, 161-162°, 163°**. Recrystallise it twice from 95% EtOH to give yellow needles [Beck et al. *J Am Chem Soc* **108** 4018 1986, DOI: 10.1021/ja00274a027]. The *dioxime* has **m 278°(dec)**. [Beilstein **7** H 632, **7** I 346, **7** II 580, **7** III 3376, **7** IV 2081.] **2,6-Dichloro-1,4-benzoquinone** [697-91-6] has **m 122-124°**. Recrystallise the quinone from petroleum ether (b 60-70°). It sublimes at 41°/17.6μ. [Carlson & Miller *J Am Chem Soc* **107** 479 1985, DOI: 10.1021/ja00288a035; Beilstein **7** II 580, **7** III 3376.]

2,6-Dichlorobenzoyl chloride [4659-45-4] **C₇H₃Cl₃O**, **M 209.5**, **m 15-17°, b 122-124°/15mm, 142-143°/21mm, d₄²⁰ 1.464. n_D²⁰ 1.560**. Reflux the acid chloride for 2 hours with excess of acetyl chloride (3 volumes), distil off AcCl followed by the benzoyl chloride. Store it away from moisture. It is an **IRRITANT**. [Beilstein **9** III 1377.]

2,4-Dichlorobenzyl alcohol (Dybenal) [1777-82-9] **C₇H₆Cl₂O**, **M 177.0**, **m 55°, 57°, 58°, 59.5°**. Crystallise the alcohol from EtOH or water. [Beilstein **6** IV 2597.] It is an antiseptic. **3,4-Dichlorobenzyl alcohol** [1805-32-8] has **m 34°, 38°, 39°, 148-151°/760mm**. Crystallise the alcohol from EtOH (**m 32-34°**) or water (**m 38°**, needles). Its solubility at 20° is 1g in 1250ml of H₂O. [Beilstein **6** H 445, **6** III 1558, **6** IV 2598.]

2,3-Dichloro-5,6-dicyano-*p*-benzoquinone (DDQ) [84-58-2] **C₈Cl₂N₂O₂**, **M 227.0**, **m 203°(dec), 213-216°, (210-215°)(dec)**. Crystallise DDQ from CHCl₃, CHCl₃/*benzene (4:1), or *benzene and store it at 0°. It is soluble in dioxane but decomposes in H₂O. [Pataki & Harvey *J Org Chem* **52** 2226 1987, DOI: 10.1021/jo00387a023; Beilstein **10** H 902, **10** II 635, **10** IV 3521.] It is a useful dehydrogenating and oxidising agent [Fieser **1** 128, **1** 215]. It was also used for the mild and efficient photochemical and thermal deprotection of thioacetals and thioketals in >90% yields [Mathew & Sankararaman *J Org Chem* **58** 7576 1993, DOI: 10.1021/jo00078a044], deprotection of acetals and ketals [McDonald et al. *Tetrahedron Lett* **35** 57 1994, DOI: 10.1016/0040-4039(94)88161-8]. DDQ mediates carbon-carbon bonds in annulation reactions, as in the formation of quinolines [Bartolotti et al. *Tetrahedron* **49** 10157 1993, DOI: 10.1016/S0040-4020(01)80210-6], and the preparation of 1,2-benzisoxazoles from 2-hydroxyphenyl aldioximes and ketoximes in high yields [Iranpoor et al. *Tetrahedron Lett* **47** 8247 2006, DOI: 10.1016/j.tetlet.2006.09.120].

2,4-Dichloro-6-methylphenol (2,4-dichloro-*o*-cresol) [1570-65-6] **C₇H₆Cl₂O**, **M 177.0**, **m 55°, b 111-113°/16mm, 129-132°/40mm, 231.6-231.9°/760mm, pK²⁰ 8.14**. Crystallise the cresol from Et₂O (**m 55°**) or water; it has **m 53-54°** after sublimation. [Beilstein **6** H 359, **6** II 332, **6** III 1267, **6** IV 2001.]

2,4-Dichloro-1-naphthol [2050-76-2] **C₁₀H₆Cl₂O**, **M 213.1**, **m 104°, 106-107°, pK_{Est} ~7.7**. Crystallise the naphthol from MeOH. The *1-methyl ether* has **m 58°** (from EtOH). [Beilstein **6** H 612, **6** I 308, **6** II 582, **6** III 2934, **6** IV 4233.]

2,3-Dichloro-1,4-naphthoquinone (Dichlone, Phygon) [117-80-6] $\text{C}_{10}\text{H}_4\text{Cl}_2\text{O}_2$, M 227.1, m 193°. Crystallise the quinone from EtOH (golden yellow needles), and sublime it *in vacuo*. [Beilstein 7 IV 2426.] It is a herbicide and fungicide.

2,5-Dichloro-4-nitroaniline [6627-34-5] $\text{C}_6\text{H}_4\text{Cl}_2\text{N}_2\text{O}_2$, M 207.0, m 153-154°, 157-158°, $\text{pK}^{25}_{\text{a}} -0.74$ (aqueous H_2SO_4). Recrystallise it from EtOH, then sublime it *in vacuo*. [Beilstein 12 H 735, 12 II 400.] **2,6-Dichloro-4-nitroaniline (Dichloran)** [99-30-9] has m 190°, 192°, 193°. Crystallise Dichloran from aqueous EtOH or *benzene/EtOH. [Beilstein 12 IV 1681.] It is a pesticide.

2,5-Dichloro-1-nitrobenzene (1,4-dichloro-4-nitrobenzene) [89-61-2] $\text{C}_6\text{H}_3\text{Cl}_2\text{NO}_2$, M 192.0, m 52°, 54°, 56°, 266-269°/atm. Crystallise the nitrobenzene from absolute EtOH. [Beilstein 5 IV 726.] **3,4-Dichloro-1-nitrobenzene (1,2-dichloro-4-nitrobenzene)** [99-54-7] has m 43°. Crystallise it from absolute EtOH. [Beilstein 5 IV 726.]

2,4-Dichloro-6-nitrophenol [609-89-2] $\text{C}_6\text{H}_4\text{Cl}_2\text{NO}_3$, M 208.0, m 118°, 120°, 122-123°, $\text{pK}_{\text{Est}} \sim 5.0$. Crystallise the chloronitrophenol from AcOH. Keep the solid moist with water. [Beilstein 6 IV 1358.] **2,6-Dichloro-4-nitrophenol** [618-80-4] has m 125°(dec), $\text{pK}^{25}_{\text{a}} 3.55$. Crystallise the chloronitrophenol from EtOH and dry it *in vacuo* over anhydrous MgSO_4 . [Beilstein 6 IV 1361.]

Dichlorophen [2,2'-methylenebis(4-chlorophenol)] [97-23-4] $\text{C}_{13}\text{H}_{10}\text{Cl}_2\text{O}_2$, M 269.1, b 168°, 172°, 177-178° $\text{pK}_{\text{Est}} \sim 9.7$. Crystallise dichlorophen from toluene. It is quite soluble in 95% EtOH (1:1), but less so in Et_2O . It dissolves in aqueous alkaline solutions where some decomposition may occur. [Beilstein 6 III 5406.] It is a useful fungicide, germicide, antiprotozoan, anthelmintic, and because of its antibacterial properties it is used in shampoos and soaps.

2,3-Dichlorophenol [576-24-9] $\text{C}_6\text{H}_4\text{Cl}_2\text{O}$, M 163.0, m 55 to 57°, 57°, 206°/atm, $\text{pK}^{25}_{\text{a}} 7.70$. Crystallise it from ether, and/or distil it. [Beilstein 6 IV 883.] **2,4-Dichlorophenol** [120-83-2] has m 42-43°, 45°, 209-211°/atm, $\text{pK}^{25}_{\text{a}} 7.89$. Crystallise it from petroleum ether (b 30-40°). Purify it also by repeated zone melting, using a P_2O_5 guard tube to exclude moisture. It is steam volatile. It is very *hygroscopic* when dry. [Beilstein 6 IV 885.] **2,5-Dichlorophenol** [583-78-8] has m 54° to 57°, 58°, b 211°/744mm, $\text{pK}^{25}_{\text{a}} 7.51$. Crystallise it from ligroin and sublime it in a vacuum or distil it. [Beilstein 6 IV 942.] **2,6-Dichlorophenol** [87-65-0] has m 64°, 64.5-65.5°, 66°, b 218-220°/atm, $\text{pK}^{25}_{\text{a}} 6.786$. The white solid crystallises from petroleum ether. It is a sex pheromone of the lone star tick [Sonenshine et al *J Chem Ecology* 2 201 1976, DOI: 10.1007/BF00987743]. [Tarbell et al. *Org Synth Coll Vol* 3 267 955, DOI: 10.15227/orgsyn.029.0035; Beilstein 6 IV 949.] **3,4-Dichlorophenol** [95-77-2] has m 64° to 67°, 68°, b 145-146°/767mm, $\text{pK}^{25}_{\text{a}} 8.58$. Crystallise 3,4-dichlorophenol from petroleum ether/*benzene mixture and/or distil it. [Beilstein 6 IV 952.] **3,5-Dichlorophenol** [591-35-5] has m 65° to 67°, 68°, b 122-124°/8mm, 233-234°/760mm, $\text{pK}^{25}_{\text{a}} 8.81$. Crystallise 3,5-dichlorophenol from petroleum ether/*benzene mixture and/or distil it. [Beilstein 6 IV 957.]

2,4-Dichlorophenoxyacetic acid (2,4-D) [94-75-7] $\text{C}_8\text{H}_6\text{Cl}_2\text{O}_3$, M 221.0, m 136°-138°, 140°, 146°, 160°/0.4mm, $\text{pK}^{25}_{\text{a}} 2.90$. Crystallise 2,4-D from MeOH or * C_6H_6 . It is a plant growth substance, a herbicide and is **TOXIC**. [Beilstein 6 IV 908.] Many salts and esters are available.

(±)-α-(2,4-Dichlorophenoxy)propionic acid (2,4-DP, Dichloroprop) [120-36-5] $\text{C}_9\text{H}_8\text{Cl}_2\text{O}_3$, M 235.1, m 117°, 117-118°, $\text{pK}^{20}_{\text{a}} 2.86$. Crystallise 2,4-DP from MeOH or * C_6H_6 . The acid has low solubility in H_2O (350mg/L at 20°) but is soluble alkaline solution. The *R*(+)- and *S*(-)- enantiomers have m 124° (from * C_6H_6) and $[\alpha]_{\text{D}}^{25} +35.2$ and -35.2 (c 1, Me_2CO), and $[\alpha]_{\text{D}}^{21} +26.2$ (c 1.23, EtOH). [Beilstein 6 H 189, 6 III 708, 6 IV 922-923.] It is a plant growth substance, a herbicide and is **TOXIC**. Note that only the *dextro*-enantiomer is physiologically active.

2,4-Dichlorophenylacetic acid [19719-28-9] $\text{C}_8\text{H}_6\text{Cl}_2\text{O}_2$, M 205.0, m 129°, 131°, 132-133°, $\text{pK}_{\text{Est}} \sim 4.0$. Crystallise the acid from aqueous EtOH. [Beilstein 9 III 2271, 9 IV 1678.] **2,6-Dichlorophenylacetic acid** [6575-24-2] has m 157-158°, 161°, $\text{pK}_{\text{Est}} \sim 3.8$. Crystallise the acid from aqueous EtOH. [Beilstein 9 III 2272, 9 IV 2679.]

3-(3,4-Dichlorophenyl)-1,1-dimethyl urea (Diuron) [330-54-1] $C_9H_{10}Cl_2N_2O$, M 233.1, m 153-154°, 155°, 158-159°. Recrystallise it from 95% EtOH [Beck et al. *J Am Chem Soc* **108** 4018 1986, DOI: 10.1021/ja00274a027]. It is poorly soluble in H_2O (42mg/L at 25°). It is a herbicide. [Beilstein **12** IV 1263.]

4,5-Dichlorophthalic acid [56962-08-4] $C_8H_4Cl_2O_4$, M 235.0, m 198°, 200° (dec to anhydride), $pK_{Est(1)}$ 2.2, $pK_{Est(2)}$ ~4.7. Crystallise the acid from water. It has been purified by converting to the anhydride, reacting it with boiling EtOH to form the *monoethyl ester* (m 133-134°), and hydrolysing it back to the diacid (see next entry). [Beilstein **9** III 4205.] **4,5-Dichlorophthalic anhydride** [942-06-3] $C_8H_2Cl_2O_3$, M 217.0, has m 185° to 187° and is separated from the acid by extraction with $CHCl_3$. [Beilstein **17** H 483.] **4,5-Dichlorophthalimide** [15997-89-4] $C_8H_3Cl_2NO_2$, M 216.0, has m 217° to 219°.

3,6-Dichlorophthalic anhydride [4466-59-5] $C_8H_2Cl_2O_3$, M 188°, 190°, 191-191.5°, b 339°/atm. Boil the anhydride in xylene (allowing any vapours which would contain H_2O to be removed, e.g. Dean and Stark trap), which causes any acid present to dehydrate to the anhydride, and cool. Recrystallise it from xylene [Villiger *Chem Ber* **42** 3529 1909, DOI: 10.1002/cber.19090420393; Fedorow *Izv Akad Nauk SSSR Otd Khim Nauk* 397 1948, *Chem Abstr* **48** 1585 1948]. [Beilstein **17/11** V 260.]

2,6-Dichlorostyrene [28469-92-3] $C_8H_6Cl_2$, M 173.0, m 8°, b 72-73°/2mm, 88-90°/8mm, d_4^{20} 1.4045, n_D^{20} 1.5798. Purify the styrene by fractional crystallisation from the melt and by distillation *in vacuo*. [Beilstein **5** III 1174, *Beilstein* **5** IV 815.]

2,4-Dichlorotoluene [95-73-8] $C_7H_6Cl_2$, M 161.1, m -13.5°, b 61-62°/3mm, 200°/atm, d_4^{20} 1.250, n_D^{20} 1.5513. Recrystallise 2,4-dichlorotoluene from EtOH at low temperature or fractionally distil it. [Beilstein **5** IV 815.]

2,6-Dichlorotoluene [118-69-4] $C_7H_6Cl_2$, M 161.1, b 196-203°/atm, 199-200°/atm, d_4^{20} 1.254, n_D^{20} 1.548. Fractionally distil it and collect the middle fraction. [Beilstein **5** IV 815.] **3,4-Dichlorotoluene** [95-75-0] has m -16°, b 205°/atm, d_4^{20} 1.2541, n_D^{20} 1.549. Recrystallise it from EtOH at very low temperature or fractionally distil it. [Beilstein **5** IV 815.]

α,α' -Dichloro-*p*-xylene [1,4-bis(chloromethyl)benzene] [623-25-6] $C_8H_8Cl_2$, M 175.1, m 98° to 101°, 100°, 254°/atm. Crystallise the xylene from *benzene and dry it under vacuum. [Beilstein **5** IV 967.]

Dicinnamalacetone (1,9-diphenyl-1,3,6,8-nonatetraen-5-one) [622-21-9] $C_{21}H_{18}O$, M 314.4, m 144°, 146°. Crystallise the ketone from *benzene/isooctane (1:1). The **1,3,5-trinitrobenzene complex**(1:1) has m 113° (from EtOH), and the **1,3,5-trinitrobenzene complex**(1:2) has m 110° (from EtOH). [Beilstein **7** H 524, **7** I 293, **7** II 484, **7** III 2756, **7** IV 1834.] The **2,4-dinitrophenylhydrazone** has m 199-200°.

Dicumyl peroxide [80-43-3] $C_{18}H_{22}O_2$, M 270.4, m 39° to 40°. Crystallise the peroxide from 95% EtOH (charcoal). Store it at 0°. **Potentially EXPLOSIVE**. [Beilstein **6** IV 3220.]

9,10-Dicyanoanthracene [1217-45-4] $C_{16}H_8N_2$, M 228.3, m 340°. Recrystallise the dinitrile twice from pyridine [Mattes & Farid *J Am Chem Soc* **108** 7356 1986, DOI: 10.1021/ja00283a034]. [Beilstein **9** IV 3667.]

1,4-Dicyanobenzene (terephthalonitrile) [623-26-7] $C_8H_4N_2$, M 128.1, m 221°, 222°, 225°. Crystallise the dinitrile from EtOH or AcOH, and has m 221.5-222.5° after sublimation. [Beilstein **9** H 846, **9** I 376, **9** II 613, **9** III 4255, **9** IV 3328.]

1,4-Dicyanonaphthalene [3029-30-9] $C_{12}H_6N_2$, M 178.2, m 206°. Purify it by recrystallisation from EtOH (charcoal), m 208°, and sublime it *in vacuo*. [Bradbrook & Linstead *J Chem Soc* 1739 1936, DOI: 10.1039/JR9360001739; *Beilstein* **9** H 917, **9** II 651, **9** III 464.]

***trans*-4-(Diethylamino)azobenzene** [3588-91-8] $C_{16}H_{19}N_3$, M 253.3, m 171°, pK^{26} 3.08 (50% aqueous EtOH) Purify the azobenzene by column chromatography on alumina, elute with, and then recrystallise it from

toluene [Albini et al. *JCS Perkin Trans 2* 1393 1982, DOI: 10.1039/P29820001393; Flamigni & Monti *J Phys Chem* **89** 3702 1985, DOI: 10.1021/j100263a025]. The *deoxycholate (1:4) complex* has **m 194-194°** (from EtOH). [*Beilstein* **16** H 341, **16** I 311, **16** III 342, **16** IV 455.]

N,N-Diethylaniline (DEA) [91-66-7] $C_{10}H_{15}N$, **M 149.2**, **m -38°**, **b 62-66°/3mm**, **216.5°/atm**, d_4^{20} **0.938**, n_D^{20} **1.5409** pK^{25} **6.57**. Reflux the base for 4 hours with half its weight of acetic anhydride, then fractionally distil it under reduced pressure (**b 92°/10mm**). Its solubility in H_2O is ~1.4w/v% at 10°. [*Beilstein* **12** IV 252.]

Di-(2-ethylhexyl)phthalate (DEHP) ('di-iso-octyl' phthalate) [117-81-7] $C_{24}H_{38}O_4$, **M 390.6**, **m -47°**, **b 231°/5mm**, **256-257°/1mm**, **384°/atm**, d_4^{20} **0.9803**, n_D^{20} **1.4863**. Wash the ester with Na_2CO_3 solution, then shake it with water. After the resulting emulsion has been broken by adding ether, the ethereal solution is washed twice with water, dried ($CaCl_2$), and evaporated. The residual liquid is distilled several times under reduced pressure, then stored in a vacuum desiccator over P_2O_5 [French & Singer *J Chem Soc* 1424 1956, DOI: 10.1039/JR9560001424]. [*Beilstein* **9** IV 3184.] It is a useful plasticiser.

Diethyl phenyl orthoformate (diethoxy phenoxy ethane) [14444-77-0] $C_{11}H_{16}O_3$, **M 196.3**, **b 103-104°/10mm**, **111°/11mm**, **122°/13mm**, d_4^{20} **1.0099**, n_D^{20} **1.4799**. Fractionate the ortho-ester through an efficient column under vacuum [Smith *Acta Chem Scand* **10** 1006 1956, DOI: 10.3891/acta.chem.scand.10-1006.] A reagent for the diethoxymethylation of Grignard compounds [Stetter & Reske *Chem Ber* **103** 643 1970, DOI: 10.1002/cber.19701030236]. [*Beilstein* **6** IV 610.]

Diethyl phthalate (DEP) [84-66-2] $C_{12}H_{14}O_4$, **M 222.2**, **m -3°**, **b 172°/12mm**, **295°/760mm**, **298-299°/atm**, d_4^{25} **1.1160**, n_D^{20} **1.5022**. Wash the ester with aqueous Na_2CO_3 , then distilled water, dry ($CaCl_2$), and distil it under reduced pressure. Store it in a vacuum desiccator over P_2O_5 . [*Beilstein* **9** IV 3172.] It is used as an industrial solvent for varnishes and in the manufacture of varnishes.

Diethyl 2-phthalimidomalonate [5680-61-5] $C_{15}H_{15}NO_6$, **M 305.3**, **m 72-74°**, **73-74°**, **76°**, pK^{25} **9.17**. Dissolve it in xylene and when the temperature is 30° add petroleum ether (b 40-60°) and cool to 20° whereby the malonate separates as a pale brown powder [Booth et al. *J Chem Soc* 666 1944, DOI: 10.1039/JR9440000666]. Alternatively, dissolve it in C_6H_6 , dry it over $CaCl_2$, filter, evaporate and the residual oil solidifies. Grind this with Et_2O , filter and wash it with Et_2O until white in colour, and dry it in a vacuum. It forms a yellow *sodium salt m 280°(dec)*. The anion has λ_{max} at 254nm (ϵ 18.5K) [Osterberg *Org Synth Coll Vol* **1** 271 1941, DOI: 10.15227/orgsyn.007.0078; UV of Na salt: Nnadi & Wang *J Am Chem Soc* **92** 4421 1970, DOI: 10.1021/ja00717a045]. [*Beilstein* **21** H 487, **21** I 379, **21** III/IV 5264.]

Diethylstilboesterol [DES, stilbesterol, stilboesterol, (*E*)-3,4-bis(4-hydroxyphenyl)-3-hexene] [56-23-1] $C_{18}H_{20}O_2$, **M 268.4**, **m 169°**, **170°**, **172°**. Crystallise stilbesterol from *benzene. [*Beilstein* **6** IV 6856.]

Diethyl terephthalate (diethyl 1,4-benzenedicarboxylic acid) [636-09-0] $C_{12}H_{14}O_4$, **M 222.2**, **m 44°**, **142°/2mm**, **302°/760mm**. Crystallise the ester from toluene and distil it under reduced pressure. [*Beilstein* **9** H 8, **9** I 374, **9** III 4250, **9** IV 3304.]

4,4'-Di-*n*-heptyloxyazoxybenzene [2587-42-0, 2635-26-9] $C_{24}H_{34}N_2O_3$, **M 426.6**, **m 75°**, **95°** (smectic Ø nematic) and **127°** (nematic Ø liquid), pK_{Est} ~ **-5**. Purify azoxybenzene by chromatography on Al_2O_3 (*benzene), recrystallise it from hexane or 95% EtOH and dry it by heating under vacuum. The liquid crystals can be sublimed *in vacuo*. [Mellifiori et al. *Spectrochim Acta Part A* **37(A)** 605 1981, DOI:10.1016/0584-8539(81)80057-8; Dewar & Schroeder *J Am Chem Soc* **86** 5235 1964, DOI: 10.1021/ja01077a039; Weygand & Glaber *J Prakt Chem* **155** 332 1940, *Beilstein* **16** III 600, **16** IV 5264.] It exhibits a negative low frequency dielectric anisotropy [McLemore & Carr *J Chem Phys* **57** 3245 1972, DOI:org/10.1063/1.1678746].

9,10-Dihydroanthracene [613-31-0] $C_{14}H_{12}$, **M 180.3**, **m 103° to 107°**, **110-110.5°**, **b ~312°/atm**, d_4^{20} **0.880**. Crystallise it from EtOH [Rabideau et al. *J Am Chem Soc* **108** 8130 1986, DOI: 10.1021/ja00286a002]. [*Beilstein* **5** H 641, **5** IV 2182.]

Dihydrochloranil (tetrachloro-1,4-hydroquinone) [87-88-6] $C_6H_2Cl_4O_2$, **M 247.9, m 240.5°**. Crystallise the hydroquinone from EtOH, AcOH/EtOH, AcOH (platelets), Me_2CO , $*C_6H_6$ or toluene (prisms). Sublime it at $77^\circ/0.6 \times 10^{-3}mm$. The *dibenzoyl* derivative has **m 233°**. [Conant & Fieser *J Am Chem Soc* **45** 2194 1923, DOI: 10.1021/ja01662a027; Rabideau et al. *J Am Chem Soc* **108** 8130 1986, DOI: 10.1021/ja00286a002; *Beilstein* **6** H 851, **6** I 417, **6** II 846, **6** III 4436, **6** IV 5775.]

1,4-Dihydro-1,4-epoxynaphthalene [573-57-9] $C_{10}H_8O$, **M 144.2, m 53-54.5°, 54° to 56°, 55-56°**. Dissolve it in Et_2O , wash it with H_2O , dry it over K_2CO_3 , filter, evaporate and dry the residue at 15mm, then recrystallise it from petroleum ether (b 40-60°), dry it at $25^\circ/0.005mm$ and sublime it (sublimes slowly at room temperature)[Wittig & Pohmer *Chem Ber* **89** 1334 1956, DOI: 10.1002/cber.19560890539; Gilman & Gorsich *J Am Chem Soc* **79** 2625 1957, DOI: 10.1021/ja01567a072]. [*Beilstein* **17** III/IV 548.] It is an electron-rich dienophile for Diels-Alder and cycloaddition reactions [Hanson & Wren *JCS Perkin Trans 1* 2089 1990, DOI: 10.1039/P19900002089; Hisano et al. *Chem Pharm Bull Japan* **38** 605 1990, DOI: org/10.1248/cpb.38.605; Wegener & Mullen *Chem Ber* **124** 2101 1991, DOI: 10.1002/cber.19911240935].

1,8-Dihydroxyanthraquinone (Danthrone, Chrysazine) [117-10-2] $C_{14}H_8O_4$, **M 240.1, m 191° to 193°, 194-197°, pK_1^{25} 8.30, pK_2^{25} 12.46**. Crystallise Danthrone from EtOH and sublime it in a vacuum. It complexes with Ca, Ba and Pb to form ‘lakes’, and is used as a purgative for animals. POSSIBLE CARCINOGEN. [*Beilstein* **8** IV 3217.]

2,4-Dihydroxyazobenzene (Sudan orange G) [2051-85-6] $C_{12}H_{10}N_2O_2$, **M 214.2, m 143-146°, 228°, $pK_{Est(1)} <0$, $pK_{Est(2)} \sim 7.3$, $pK_{Est(3)} \sim 9.3$** . Crystallise the dye from hot EtOH (charcoal). Has UV max at 350nm. [*Beilstein* **16** IV 264.]

2,3-Dihydroxybenzaldehyde [24677-78-9] $C_7H_6O_3$, **M 138.1, m 105-108°, 135-136°, b $120^\circ/16mm$, pK_1^{20} 7.73, pK_2^{20} 10.91**. Crystallise the aldehyde from water. It is air-sensitive, if possible store under N_2 in the dark. [*Beilstein* **8** III 1979.]

2,4-Dihydroxybenzoic acid (β -resorcylic acid) [89-86-1] $C_7H_6O_4$, **M 154.1, m 208 to 211°(dec), 226-227°(dec), pK_1^{25} 3.30, pK_2^{25} 9.12, pK_3^{25} 15.6**. Crystallise the acid from water which forms hydrated crystals. They dehydrate at 100° and melt with decomposition at 213° when heated rapidly. It is used as a spot test reagent for Fe. [Nierenstein & Clibbens *Org Synth Coll Vol 2* 557 1943, DOI: 10.15227/orgsyn.010.0094; *Beilstein* **10** IV 1420.] **2,5-Dihydroxybenzoic acid (Gentisic acid, 5-hydroxysalicylic acid)** [490-79-9] has **m 204.5-205°, 204° to 208°, pK^{25} 2.95**. Crystallise gentisic acid from hot water or $*benzene/acetone$. Dry it in a vacuum desiccator over silica gel. It is dimorphic and its phase alters at $\sim 200^\circ$ before melting. It is soluble in hot H_2O and has low solubility in cold H_2O ($\sim 0.5\%$ at 6°). The sodium salt, *sodium gentisate* [4955-90-2] $C_7H_5NaO_4$, **M 176.1 (anhydr)**, crystallises with 5.5 H_2O which effloresces in air losing $\sim 3 H_2O$ but holds on to the last 0.5 H_2O even at 100° . It is anti-inflammatory and an analgesic. [*Beilstein* **10** H 384, **10** IV 1441.] **2,6-Dihydroxybenzoic acid (γ -resorcylic acid)** [303-07-1] $C_7H_6O_4$, **M 154.1, m 167°(dec), pK^{25} 1.05**. Dissolve the acid in aqueous $NaHCO_3$ and the solution is washed with ether to remove non-acidic material. The acid is precipitated by adding H_2SO_4 , and recrystallised from water. Dry it *in vacuo*, and store it in the dark [Lowe & Smith *JCS Faraday Trans 1* **69** 1934 1973, DOI: 10.1039/F19736901934]. [*Beilstein* **10** IV 1456.]

2,4-Dihydroxybenzophenone (DHB, benzo-resorcinol) [131-56-6] $C_{13}H_{10}O_3$, **M 214.2, m 145.5°, 144-145°, 147°, $pK_{Est(1)} \sim 7.0$, $pK_{Est(2)} \sim 12.0$** . Recrystallise it from MeOH or H_2O . [*Beilstein* **8** IV 2442.]

4,4'-Dihydroxybenzophenone [611-99-4] $C_{13}H_{10}O_3$, **M 214.2, m 210°, 213-214°, 213-215°, 216.6-217.1°, $pK_{Est} \sim 7.0$** . The benzophenone was prepared by a Friedel-Crafts reaction between 4-methoxybenzoyl chloride and anisole with anhydrous $AlCl_3$ in anhydrous CS_2 , followed by demethylation with $AlCl_3$ in boiling toluene. It could also be made directly from 4-hydroxybenzoic acid and an equivalent of phenol by heating with three parts of freshly fused anhydrous $ZnCl_2$ at $125-140^\circ$ for 45 minutes; the cooled mass is treated with dilute HCl, the solid is filtered off, washed thoroughly with aqueous $NaHCO_3$ solution, H_2O , dried *in vacuo* and recrystallised from H_2O , aqueous EtOH, or dilute HCl. [Russell & Butler *J Am Chem Soc* **71** 3663 1949, DOI: 10.1021/ja01179a024.] The *oxime* has **m 266-267° (dec)** [Zigeuner & Ziegler *Monatsh Chem* **80** 359 1949,

DOI: 10.1007/BF00897769], and the **2,4-dinitrophenylhydrazones** has **m 190-192°** (from EtOH). [Beilstein 8 H 316, 8 I 641, 8 II 355, 8 III 2648, 8 IV 2452.] It is used in paints and plastics as absorber of UV light.

2,5-Dihydroxybenzyl alcohol (Gentisyl alcohol) [495-08-9] $C_7H_8O_3$, **M 140.1**, **m 47-48°**, **pK_{Est(1)} ~9.3**, **pK_{Est(2)} ~11.3**. Crystallise the alcohol from ligroin, $CHCl_3$, AcOH or H_2O . Sublime it at ~70° under high vacuum. It dissolves in aqueous NaOH solution and oxidises in air, as observed by the colour changing from yellow to red then brown. [Beilstein 6 II 1084, 6 III 6326.]

2,2'-Dihydroxybiphenyl [1806-29-7] $C_{12}H_{10}O_2$, **M 186.2**, **m 108.5-109.5°**, **110°**, **b 315°/atm**, **pK₁²⁵ 7.56**, **pK₂²⁵ 11.80**. Crystallise the biphenyl repeatedly from toluene, then sublime it at 60°/10⁻⁴mm. [Beilstein 6 IV 6645.] **4,4'-Dihydroxybiphenyl (4,4'-biphenol)** [92-88-6] has **m 280.5°**, **280-285°**, **pK_{Est(1)} ~3.6**, **pK_{Est(2)} ~11.8**. Recrystallise the biphenol from aqueous EtOH preferably under N_2 to avoid oxidation to the extended quinone. It is characterised as the *dimethyl derivative (4,4'-dimethoxybiphenyl)* from which it is prepared by demethylation. The dimethoxy derivative has **m 176.5-177°** (from AcOH, EtOH or hexane), and sublimes *in vacuo*. [Williamson & Rodebush *J Am Chem Soc* 63 3018 1941, DOI: 10.1021/ja01856a044; Beilstein 6 I 485, 6 II 962, 6 III 6389, 6 IV 6651.]

1,8-Dihydroxy-3-methylantraquinone (chrysophanic acid) [481-74-3] $C_{15}H_{10}O_4$, **M 245.3**, **m 196°**, **pK_{Est(1)} ~8.2**, **pK_{Est(2)} ~12.4**. Crystallise chrysophanic acid from EtOH or *benzene and has **m 195.6-196.2°**, after sublimation it in a vacuum. The yellow *mono-acetate* has **m 188-190°** (from MeOH or Me_2CO). It forms Ni^{2+} , Co^{2+} and Cu^{2+} complexes. [Beilstein 8 H 470, 8 I 725, 8 II 510, 8 III 3808, 8 IV 3277.]

1,2-Dihydroxynaphthalene [574-00-5] $C_{10}H_8O_2$, **M 160.2**, **m 101-103°**, **108°(anhydrous)**. It crystallises from CS_2 (needles), H_2O (leaflets, as hydrate) or aqueous EtOH (as hydrate). It is hygroscopic, air sensitive and oxidises readily in solution to the quinone. The *diacetate* [6336-79-4] has **m 109° (158-161°)**, $C_{14}H_{12}O_4$, **M 244.2**. Methylation produces a mixture of 1-hydroxy-2-methoxy-, 1-methoxy-2-hydroxy and 1,2-dimethoxy naphthalenes which have to be separated [Bezdik & Friedlaender *Monatsh Chem* 30 271 1909, DOI: 10.1007/BF01519684]. **1-methoxy-2-hydroxynaphthalene** (colourless plates **m 92°** from light petroleum, Bell & Duewell *Aust J Chem* 16 101 1963, DOI: 10.1071/CH9630101) was best prepared, in 44% yield, from 1-bromo-2-naphthol by conversion to the Grignard reagent followed by oxidation using the general procedure [Kharasch & Reynolds *J Am Chem Soc* 65 501 1943, DOI: 10.1021/ja01244a005]. The *dimethylether* [57189-64-7] has **b 278-280°/atm**, **m 31°**, and its *picrate* forms red needles with **m 97°**. [Beilstein 6 H 975.] 1,2-Dihydroxynaphthalene is an **IRRITANT**, causing sneezing. For its use as a biomarker for naphthalene exposure in humans see Klotz et al. [*Int J Hyg Environ Health* 214 110 2011, DOI: 10.1016/j.ijheh.2010.11.003, PMID: 21147027]. A useful GC-MS method for measuring 1,2-dihydroxynaphthalene and 1,4-dihydroxynaphthalene after enzymatic digestion of conjugates in urine was developed by Rapaport and coworkers [Wu et al. *J Chromatogr B*, 826 206 2005, DOI: 10.1016/j.jchromb.2005.08.022].

1,3-Dihydroxynaphthalene (naphthoresorcinol) [132-86-5] $C_{10}H_8O_2$, **M 160.2**, **m 118-120°**, **123-125°**, **124-125°**, **pK₁²⁵ 7.22**, **pK_{Est(2)} ~10.0**. It crystallises from H_2O (leaflets), a $H_2O/*C_6H_6$ (80/20), Et_2O /petroleum ether, or $CHCl_3$. It was prepared by cyclisation of ethyl γ -phenylacetoacetate in concentrated H_2SO_4 and kept overnight before pouring onto ice and isolating the product by Et_2O extraction [Soliman & West *J Chem Soc* 53 1944, DOI: 10.1039/JR9440000053]. It gave a *diacetate* **m 56°** (prisms from aqueous AcOH). On a larger scale diethyl phenylacetylmalonate gave naphthoresorcinol in three steps in good yield. It was purified by dissolving in hot H_2O , decolorising with aqueous sodium hydrosulfite and norit, adding NaCl, and allowing to stand at 5° for 24hrs. The cold mixture should be shaken, after seeding, to avoid separation of the product as an oil and to form transparent plates. It is best purified by sublimation at 120-130°/5x10⁻⁴mm. [Meyer & Bloch *Org Synth* 25 73 1945, *Org Synth Coll Vol* 3 637 1955, DOI: 10.1522/orgsyn.025.0073.]

The **3-methoxy-1-naphthol**, single product obtained by methylation with 30 parts of MeOH/HCl at ~25°, forms colourless stout needles **m 99-100°** from hot petroleum ether, and has ¹³C NMR ($CDCl_3$) with δ at 157.7 (CO), 152.9 (CO), 135.5, 132.2, 127.1, 123.0, 121.7, 101.4, 99.1, 55.3 (OCH_3). **3-Methoxy-1-naphthyl acetate** also crystallises from light petroleum in colourless needles with **m 74°**. [Bell & McCaffrey *Aust J Chem* 46 731 1993, DOI: 10.1071/CH9930731.] **1-methoxy-3-naphthol** crystallises from light petroleum (b 40-60°, charcoal) in long fine needles **m 55-56°** and the derivative **1-methoxy-4-phenylazo-3-naphthol** has **m 145-146°** (orange-

red needles from MeOH) and is insoluble in alkali [Iskander et al. *J Chem Soc (C)* 1701 1970, DOI: 10.1039/J39700001701]. **1,3-Dimethoxynaphthalene** [10076-61-3] $C_{12}H_{12}O_2$, **M 188.2**, has **b 128-132°/1mm** and **b 145-150°/3mm**. [For IR spectra see van Gemert *Aust J Chem* **21** 2203 1968, DOI: 10.1071/CH9682203.] For the determination of glycolaldehyde in the presence of hydroxypyruvate and tartronate semialdehyde, whereby λ_{max} shifts from 660nm to 440nm, see Windt & van der Drift [*Analyt Biochem* **106** 342 1980, DOI: 10.1016/0003-2697(80)90530-8, PMID: 7447001], and for use as a colour test reagent for aldoses, ketoses, pentoses and glucuronic in urine see Forsyth [*Nature* **161** 239 1948, DOI:10.1038/161239a0].

1,4-Dihydroxynaphthalene [571-60-8] $C_{10}H_8O_2$, **M 160.2**, **m 191-192°(dec)**, **pK₁²⁵ 9.37**, **pK₂²⁵ 10.93**. The diol obtained by reduction of 1,4-naphthaquinone with $SnCl_2$ is slightly pink and further recrystallisation from EtOH did not remove the colour. However, when dissolved in Et_2O , shaken with alumina and filtered, then addition of ligroin gives white crystals of 1,4-dihydroxynaphthalene. The UV (H_2O) has λ_{max} nm(ϵ_{max} $M^{-1}.cm^{-1}$) at 322(5350) for neutral species, 346(5700) for mono-anion and 363(6050) for di-anion. [Baxendale & Hardy *Trans Faraday Soc* **49** 1140 1953, DOI: 10.1039/TF9534901140]. It crystallises from hot EtOH in colourless needles where its solubility in EtOH is 50mg/ml at ~25°.

The **monoacetate** [70662-30-5] $C_{12}H_{10}O_3$, **M 202.2**, has **m 127°** (from cyclohexane), the **diacetate** [5697-00-7] $C_{14}H_{12}O_4$, **M 244.2**, has **m 128-130°** (plates from EtOH). **4-Methoxy-1-naphthol** [84-85-5] $C_{11}H_{10}O_2$, **M 174.2**, was isolated from *Galium mollugo* roots [by extraction with light petroleum (b 60-80°), $*C_6H_6$ then MeOH], purified by TLC (silica gel G/ $*C_6H_6$ -EtOAc 4:1; R_F 0.54), and recrystallised (needles) from light petroleum (b 60-80°). It had **m 124-126°**; the UV (EtOH) had λ_{max} nm(log ϵ) at 248(4.39), 317(3.78) and 331(3.77) and MS with m/z 174 [Burnett & Thomson *J Chem Soc (C)* 857 1968, DOI: 10.1039/J39680000854]. [For 1H NMR see Smith & Chiranjeevi *J Phys Chem* **70** 3505 1966, DOI: 10.1021/j100883a023; for synthesis see Russig *J Prakt Chem* **62** 30 1900, DOI: 10.1002/prac.19000620102; *Beilstein* **6** IV 6545.] [For IR spectra see van Gemert *Aust J Chem* **21** 2203 1968, DOI:10.1071/CH9682203.]

A useful GC-MS method for measuring 1,2-dihydroxynaphthalene and 1,4-dihydroxynaphthalene after enzymatic digestion of conjugates in urine was developed by Rapaport and coworkers [Wu et al. *J Chromatogr B*, **826** 206 2005, DOI: 10.1016/j.jchromb.2005.08.022].

1,5-Dihydroxynaphthalene (1,5-naphthalenediol) [83-56-7] $C_{10}H_8O_2$, **M 160.2**, **m 259°, 260°, 261°, 265°**, **pK_{Est(1)} ~9.0**, **pK_{Est(2)} ~11.0**. The diol (~30g) is purified by making into a thick paste with H_2O and suspending this in 3L of H_2O containing 200ml of EtOH, boiling under reflux for 3 hours, cooling to 30°, saturating with SO_2 , digesting below the boiling point for 1 hour and filtering fast through a large hot filter paper. The hot filtrate is poured onto crushed ice whereby the diol (15-20g) separates as colourless needles (**m 258°**) [Wheeler & Ergle *J Am Chem Soc* **52** 4872 1930, DOI: 10.1021/ja01375a032]. Recrystallise it from H_2O or nitromethane under N_2 to avoid oxidation. The **dibenzoyl** derivative has **m 245°** (from EtOH). The **1,5-naphthyl-bistriflate**, obtained from Tf_2O /pyridine in 76% yield (purified by column chromatography SiO_2 /hexane) has **m 114-115°** and 1H NMR ($CDCl_3$ /TMS): 8.18 (2H, d $J = 7.8Hz$, C4 & 8-H's), 7.68 (4H, m, C-2, C3, C6 & C7-H's) [Takeuchi et al. *J Org Chem* **58** 7388 1993, DOI: 10.1021/jo00078a016]. The **5-methoxy-1-naphthol** derivative [prepared from the diol in MeOH/HCl (1:30 weight to volume ratio) and set aside at 25° for 9-10 days] crystallised from petroleum ether (**m 135-136°**) or from CH_2Cl_2 /hexane (needles **m 140°**) [Bell & McCaffrey *Aust J Chem* **46** 731 1993, DOI:10.1071/CH9930731]. **1,5-Dimethoxynaphthalene**, prepared by methylation (excess Me_2SO_4 /aqueous KOH) has **b 179°/13mm** [Buu-Hoi & Lavit *J Org Chem* **20** 1191 1955, DOI: 10.1021/jo01126a007]. [IR spectra see van Gemert *Aust J Chem* **21** 2203 1968, DOI:10.1071/CH9682203; ^{13}C NMR see Ernst *Chem Ber* **108** 2030 1975, DOI: 10.1002/cber.19751080622; *Beilstein* **6** I 477, **6** II 950, **6** III 5265, **6** IV 6554.]

1,6-Dihydroxynaphthalene [575-44-0] $C_{10}H_{18}O_2$, **M 160.2**, **m 130°, 133°, 137-138°, 138-139°** (with previous softening), **pK_{Est(1)} ~9.4**, **pK_{Est(2)} ~11**. Crystallise it from *benzene or *benzene/EtOH after treatment with charcoal. [For fluorinated derivatives see Liu et al. *Bioorg Med Chem Lett* **11** 2903 2001, DOI: 10.1016/S0960-894X(01)00595-9]. [For IR spectra see van Gemert *Aust J Chem* **21** 2203 1968, DOI:10.1071/CH9682203; *Beilstein* **6** H 981, **6** I 480, **6** II 952, **6** III 5279, **6** IV 6557.] For catalytic reduction to 6-hydroxy-1-tetralone [3470-50-6] see below.

1,7-Dihydroxynaphthalene [575-38-2] $C_{10}H_8O_2$, M 160.2, m 178-182°, 180-184°, $pK_{Est(1)} \sim 9.4$, $pK_{Est(2)} \sim 11$. The diol is soluble in H_2O and crystallises as needles from $*C_6H_6$. The *diacetate* [51850-49-8] $C_{14}H_{12}O_4$, M 244.2, m 108°, forms plates from $*C_6H_6$. *1-Methoxy-7-naphthol* [91344-50-2] $C_{11}H_{10}O_2$, M 174.1, m 71°, crystallises in plates from hexane, the *7-benzoate* has m 92° (from aqueous EtOH) and the *7-acetate* has m 152° (from light petroleum, b 100-120°). *7-Methoxy-1-naphthol* [67247-13-6] $C_{11}H_{10}O_2$, M 174.1, prepared from *7-methoxy-1-tetralone* by reduction with elemental sulphur (heating with S at 240-250° until H_2S evolution ceased), has b 145-148°(bath temp)/0.01mm, and the solidified distillate has m 104.5-105° (103-105° also reported) after crystallising in plates from hexane. *1,7-Dimethoxynaphthalene* [5309-18-2] $C_{12}H_{12}O_2$, M 188.2, prepared from the 1,7-diol and Me_2SO_4 , has b 134-137°(bath temp)/0.01mm, b 123-130°/0.4mm, 170°/20mm, and the solidified distillate has m 68-68° (103-105° also reported) after crystallising in plates from hexane. [Shand & Thomson *Tetrahedron* **19** 1919 1963, DOI: 10.1016/0040-4020(63)85007-3.] [For IR spectra see van Gemert *Aust J Chem* **21** 2203 1968, DOI: 10.1071/CH9682203; *Beilstein* **6** IV 6559.]

1,8-Dihydroxynaphthalene [569-42-6] $C_{10}H_8O_2$, M 160.2, m 137-143°, 142-143°, 145-146°, pK_1^{20} 6.71, $pK_2^{20} > 13$. It crystallises from H_2O in leaflets. It was obtained by de-sulfonating 1,8-dihydroxynaphthalene-4-sulfonic acid (concentrated H_2SO_4 in an autoclave at 150°/1hr). It gives a golden-yellow solution in concentrated H_2SO_4 [Heller & Kretschmann *Chem Ber* **54** 1098 1921, DOI: 10.1002/cber.19210540528]. Alternatively, it was prepared from 1-naphthol-8-sulfonic acid (50g), KOH (150g), NaOH (50g) and H_2O (75ml) which were fused slowly, with vigorous stirring, to 230° in a stainless steel beaker until the melt was encrusted with black tar. After cooling, the solid was crushed in portions and stirred vigorously with H_2O (2.5L) containing concentrated HCl (700ml), boiled for 10 minutes, filtered and cooled. This was extracted with Et_2O , the extract was dried, evaporated, and the residue was recrystallised from $*C_6H_6$ /ligroin to give pure *1,8-diol* (4.3g) with m 142-143°. It crystallises in leaflets from H_2O . [Lurie et al. *J Am Chem Soc* **83** 5015 1961, DOI: 10.1021/ja01485a031]. The *diacetate* [6566-25-2] $C_{14}H_{12}O_4$, M 244.2, has m 147-148°. [For pKa see Musso & Matthies *Chem Ber* **94** 356 1961, DOI: 10.1002/cber.19610940211; and for IR spectra see van Gemert *Aust J Chem* **21** 2203 1968, DOI: 10.1071/CH9682203]. *1,8-Methoxynaphthalene* [10075-66-8], prepared by methylation (excess Me_2SO_4 /KOH), has b 167-168°/12mm, and the solidified distillate was crystallised from Et_2O , EtOH or aqueous EtOH giving the less soluble leaflets m 157° (whose picrate has m 172°, brown-red needles from EtOH), and an equal amount of the more soluble *8-methoxy-1-naphthol* [3588-75-6] with m 55-56°(57°) (which is less soluble in the methylating solvent, and recrystallised from petroleum ether). On methylation in a large volume of EtOH the ratio of dimethoxy to monomethoxy changes from 8:7 to 15.5:3. [Buu-Hoï & Lavit *J Chem Soc* 1956 2412, DOI: 10.1039/JR9560002412]. Methylation with diazomethane apparently yields only the monomethyl ether. [Staudinger et al. *Helv Chim Acta* **4** 334 1921, DOI: 10.1002/hlca.19210040134.]

2,3-Dihydroxynaphthalene [92-44-4] $C_{10}H_8O_2$, M 160.2, m 161-165°, 163°, 162-164°, 163-165°, b 353°/atm, d^{25}_4 1.81g/ml, pK_1^{25} 8.68, pK_2^{25} 12.5. It is purified by recrystallisation from $*C_6H_6$, $CHCl_3$ or hot H_2O (leaflets). Also purified by sublimation. Useful for the photometric estimation of Fe^{3+} , V^{5+} , Ti and Mo [Patrovský *Coll Czech Chem Commun* **35** 1599 1970, DOI: 10.1135/cccc19701599], Th, U^{6+} , also for the extraction and estimation of boron with the 2,3-diol and crystal violet at 610nm [Sato & Uchikawa *Analyt Chim Acta* **143** 283 1982, DOI: 10.1016/S0003-2670(01)95513-8]; and Ti^{4+} interaction with serum albumin [Tinoco et al. *Inorg Chem* **47** 8380 2008, DOI: 10.1021/ic800529v].

2,3-Dibenzoyloxynaphthalene [91201-70-6] $C_{24}H_{16}O_4$, M 368.3, has m 154-155° (needles from CH_2Cl_2 , or CH_2Cl_2 /petroleum ether).

2,3-Dimethoxynaphthalene [10103-06-7] $C_{12}H_{12}O_2$, is obtained by methylation (excess Me_2SO_4 /aq KOH) of the diol, has b 182°/17mm, m 115-116°, together with a small amount of *2-hydroxy-3-methoxynaphthalene* [18515-11-2] m 108° (colourless needles from $*C_6H_6$) [Buu-hoi & Lavit *J Org Chem* **21** 21 1956, DOI: 10.1021/jo01107a003]. [For the synthesis of 2,3-dimethoxynaphthalene from 2,5-diethoxy-tetrahydrofuran and veratrole in a Diels-Alder reaction (ice-cold 70% H_2SO_4 /2.5hrs then steam distil) to form 1,4-dihydroxy-6,7-dimethoxy-1,2,3,4-tetrahydronaphthalene which dehydrates to 2,3-dimethoxy-naphthalene as driven by aromatization see Arcoletto et al. *Chem Ind (London)* 128 1977]. [For derivatives see Hewgill & Middleton *JCS (C)* 2316 1967, DOI: 10.1039/J39670002316; and Howell & Taylor *J Chem Soc* 4252 1956, DOI: 10.1039/JR9560004252; for the synthesis of butterflyshaped compounds from 2,3-dihydroxynaphthalene see Nakanishi et al. *J Org Chem* **79** 2625 2014, DOI: 10.1021/jo500085a; and for IR spectra see van Gemert

Aust J Chem **21** 2203 1968, DOI: 10.1071/CH9682203; *Beilstein* **6** III 5291, **6** IV 6564.]

2,6-Dihydroxynaphthalene [581-43-1] $C_{10}H_8O_2$, **M 160.2**, **m 211-217°(2H₂O)**, **218°(ahydr)**, **223-225°** (also reported), **pK_{Est(1)} ~9.9**, **pK_{Est(2)} ~11**. Purify it by recrystallisation from EtOH (plates) or aqueous EtOH or hot H₂O (with 2H₂O), and sublime preferably *in vacuo*. Willstätter & Parnas [*Chem Ber* **40** 1406 1907; DOI: 10.1002/cber.19070400224] originally prepared the 2,6-diol by fusing disodium β-naphtholsulfonate (80g) with molten KOH (240g at 320°) at 320-350° for 15minutes, cooled *in vacuo*, poured into 1.5L of H₂O, acidified with concentrated HCl (charcoal) and filtered off the crude acid, which was then recrystallised from H₂O (10g of rhombic crystals). For a convenient synthesis from the readily available 6-bromo-2-naphthol in 52% overall yield and 95.7% purity *via* 6-methoxymethoxy-2-naphthol formate involving a Baeyer-Villiger oxidation-rearrangement see Cui & Li [*J Chem Res* **36** 675 2012, DOI: 10.3184/174751912X13497085947943].

The **diacetate** [22426-47-7] $C_{14}H_{12}O_4$, **M 244.2**, has **m 175°**, forms aqueous AcOH. The **mono-methylether** [5111-66-0] $C_{11}H_{10}O_2$, **M 174.2**, has **b 148-150°/0.15mm** and **m 149°**; and the **di-methylether** [5486-55-5] $C_{12}H_{12}O_2$, **M 188.2**, has **m 150°** (plates from EtOH and leaflets from *C₆H₆, with a fragrant odour). [For IR spectra see van Gemert *Aust J Chem* **21** 2203 1968, DOI: 10.1071/CH9682203; for ¹³C NMR see Ernst *Chem Ber* **108** 2030 1975, DOI: 10.1002/cber.19751080622; and for ESR see Dixon et al. *Tetrahedron Lett* **17** 623 1976, DOI: 10.1016/S0040-4039(00)77929-9; *Beilstein* **6** IV 6566.] Reduction of the **di-methylether** with Na/isoamyl alcohol gave a demethylated-dihydrotetralone [Robinson & Weygand *J Chem Soc* 386 1941, DOI: 10.1039/JR9410000386] that was shown to be **6-methoxy-2-tetralone** [2472-22-2] $C_{11}H_{12}O_2$, **M 176.2**, **n_D²⁰ 1.5645**, and had **m 36°** after recrystallisation from light petroleum (b 40-60°) [Cornforth et al. *J Chem Soc* 689 1942, DOI: 10.1039/JR9420000689]. **6-Methoxy-2-tetralone 2,4-dinitrophenylhydrazone** crystallised from EtOH in orange prisms and had **m 132°** [Crowley & Robinson *J Chem Soc* 2001 1938, DOI: 10.1039/JR9380002001].

2,7-Dihydroxynaphthalene [582-17-2] $C_{10}H_8O_2$, **M 160.2**, **m 188°, 185-190°, 190-194°, b 375.4°/760mm, d₂₅ 1.33g/cm³**, **pK_{Est(1)} ~9.8**, **pK_{Est(2)} ~11**. It is prepared commercially by fusing disodium 2,7-naphthalenedisulfonate with sodium hydroxide at high temperature. In a continuous process an aqueous mixture of the disodium disulfonate and NaOH (2.5:10) is rolled as a film over a moving hot sheet at 290-300° during 5 hours where rapid evaporation occurs. Acidification provides 2,7-dihydroxynaphthaene in 92% yield. [Matsura et al. JP 02 11,530 (90 11,530) to Kawasaki steel, *Chem Abstr* **112** P197839t 1984.] Purify it by recrystallisation from EtOH (plates) or aqueous EtOH (needles) or hot H₂O (with 2H₂O, needles), and sublime preferably at high vacuum. **2,7-Diacetoxynaphthalene** [22472-26-0] has **m 136°** (from AcOH). **2-Hydroxy-7-methoxy-naphthalene** [5060-82-2] $C_{11}H_{10}O_2$, **M 174.2**, has **b 145-148°/0.01mm, m 122-124°** (needles from EtOH or light petroleum), and **2-acetoxy-7-methoxynaphthalene** $C_{13}H_{12}O_3$, **M 216.2**, crystallises from MeOH with **m 124-124.5°**. **2,7-Dimethoxynaphthalene** [3469-26-9] $C_{12}H_{12}O_2$, **M 188.2**, prepared from reaction with Me₂SO₄/aqueous NaOH, has **b 319°/731mm, m 138°** (cream coloured needles from AcOH) [Wilson *Tetrahedron* **3** 236 1958, DOI: 10.1016/0040-4020(58)80019-8; Ullmann *Justus Liebigs Ann Chem* **327** 117 1903, DOI: 10.1002/jlac.19033270105; *Beilstein* **6** H 986.] [For sulfonation see Ansink et al. *JCS Perkin Trans* **721** 1993, DOI: 10.1039/P29930000721; for ¹³C NMR see Ernst *Chem Ber* **108** 2030 1975, DOI: 10.1002/cber.19751080622; and for IR spectra see van Gemert *Aust J Chem* **21** 2203 1968, DOI: 10.1071/CH9682203; *Beilstein* **6** IV 6570.]

2,5-Dihydroxyphenylacetic acid (homogentisic acid) [451-13-8] $C_8H_8O_4$, **M 168.2**, **m 152°, 154-152°, pK²⁰ 4.14 (COOH)**. Crystallise homogentisic acid from EtOH/CHCl₃ (anhydrous crystals) or H₂O (solubility is 85% at 25°) as monohydrate crystals. The **dimethyl ether**, $C_{10}H_{12}O_4$, has **m 124.5°**, and the **dimethyl ether methyl ester**, $C_{11}H_{14}O_4$, has **m 124.5°**. [*Beilstein* **10** IV 1506.]

3,4-Dihydroxytoluene (4-methylcatechol) [452-86-8] $C_7H_8O_2$, **M 124.1**, **m 65-66°, 68°, 69°, b 112°/3mm, 241°/760mm, 251°/760mm, pK₁²⁵ 9.44 (9.7), pK₂²⁵ 10.90 (11.9)**. Crystallise the catechol from *C₆H₆. Alternatively, crystallise it from high-boiling petroleum ether and distil it in a vacuum. The purity can be checked by TLC. [*Beilstein* **6** IV 5878.]

1,4-Diiodobenzene [624-38-4] $C_6H_4I_2$, **M 329.9**, **m 131°, 132-133°, b 185°/atm**. Crystallise it from EtOH or boiling MeOH, then dry it in air. Store it in dark bottles away from light. [*Beilstein* **5** IV 700.]

trans-4,4'-Dimethoxyazobenzene [501-58-6] $C_{14}H_{14}N_2O_2$, M 242.3, m 162.7-164.7°, 165-166°, $pK_{Est} \sim 0$. Chromatograph it on basic alumina and elute with *benzene. Then crystallise the residue from 2:2:1 (v/v) methanol/ethanol/*benzene or Me_2CO . [Beilstein 16 H 112, 16 I 237, 16 II 43, 16 III 93, 16 IV 172.]

3,5-Dimethoxybenzaldehyde [7311-34-4] $C_9H_{10}O_3$, M 166.2, m 45.5°, 46-47°, 45-48°, b 115-118°/1mm, 151°/16mm. The aldehyde was prepared by reduction of the corresponding acid chloride with 5%Pd/CaCO₃ in xylene, and was purified by distillation in a vacuum and by recrystallisation from hexane (m 48°), ligroin, petroleum ether (m 45-46°) or 70% aqueous EtOH. The *oxime* (m 115°) crystallises from *C₆H₆, the *thiosemicarbazone* (m 211-212°) crystallises from aqueous EtOH, and the *2,4-dinitrophenylhydrazone* (m 260-261°) forms orange needles from aqueous EtOH. [Mongolusk et al. *J Chem Soc* 2231 1957, DOI: 10.1039/JR9570002231; Lambooy *J Am Chem Soc* 76 133 1954, DOI: 10.1021/ja01630a037; Beilstein 8 II 291, 8 III 2073, 8 IV 1786.]

1,2-Dimethoxybenzene (veratrole) [91-16-7] $C_8H_{10}O_2$, M 137.2, m, 15°, 23°, b 208.5-208.7°atm, d_4^{20} 1.085, n_D^{25} 1.53232. Steam distil veratrole, then fractionally distil it from BaO, CaH₂ or Na. Crystallise it from *benzene or low-boiling petroleum ether at 0°. Fractionally crystallise it from its melt. Store it over anhydrous Na₂SO₄. [Beilstein 6 IV 5564.] **1,3-Dimethoxybenzene** [151-10-0] has m -52°, b 85-87°/7mm, 212-213°/atm, 217-218°/atm, d_4^{20} 1.056, n_D^{20} 1.5215. Extract it with aqueous NaOH, and water, then dry it. Fractionally distil it from BaO or Na. [Beilstein 6 IV 5663.] **1,4-Dimethoxybenzene** [150-78-7] has m 54°, 56°, 57.2-57.8°, 59°, 60°, b 213°/atm, d^{25} 1.053. Steam distil 1,4-dimethoxybenzene, then crystallise it from hexane or *benzene, and from MeOH or EtOH, but these are wasteful due to high solubilities. Dry it under vacuum. It also sublimes under vacuum. [Beilstein 6 IV 5718.]

2,4-Dimethoxybenzoic acid [91-52-1] $C_9H_{10}O_4$, M 182.2, m 107.3°, 109°, pK^{25} 4.36. Crystallise the acid from water and dry it in a vacuum desiccator over H₂SO₄. The *S-benzylisothiuronium salt* has m 158-159° (from CHCl₃). [Beilstein 10 H 379, 10 I 177, 10 II 252, 10 III 1371, 10 IV 1422.] Aromatic acid impurities (to <0.05%) can be removed via the (±)-α-methylbenzylamine salt as described for 2,4-dichlorobenzoic acid [Ley & Yates *Organic Process Research & Development* 12 120 2008, DOI: 10.1021/op7001547.] **2,6-Dimethoxybenzoic acid** [1466-76-8] has m 185°, 186-187°, 188-191°, pK^{25} 3.44. Crystallise the acid from water or 1,1-dichloroethane (m 187.5-188.5°). [Beilstein 10 H 388, 10 I 185, 10 II 259, 10 III 1401, 10 IV 1456.] **3,4-Dimethoxybenzoic acid (veratric acid)** [93-07-2] has m 179°, 181-187°, 185-186°, pK^{25} 4.43. Crystallise the acid from Et₂O, H₂O (solubility is 0.05w/v% at ~25°, and 0.6w/v% at ~100°) or aqueous acetic acid. It has m 180-181° after sublimation at 80°/1mm. [Beilstein 10 H 393, 10 I 188, 10 II 261, 10 III 1404, 10 IV 1406.] **3,5-Dimethoxybenzoic acid** [1132-21-4] has m 178°, 180°, 185-186°, pK^{25} 3.97. Crystallise the acid from water, EtOH or aqueous acetic acid and dry it in a vacuum. [Beilstein 10 H 405, 10 I 195, 10 III 1446, 10 IV 1501.]

4,4'-Dimethoxybenzophenone [90-96-0] $C_{15}H_{14}O_3$, M 242.3, m 90-93° (also reported), 141°, 143°, 144.5°, b 200°/17mm. Crystallise the ketone from absolute EtOH, aqueous EtOH or EtOH/AcOH. The *2,4-dinitrophenylhydrazone* has m 199-200°. [Beilstein 8 H 317, 8 I 641, 8 II 355, 8 III 2649, 8 IV 2453.]

2,6-Dimethoxy-1,4-benzoquinone (DMBQ) [530-55-2] $C_8H_8O_4$, M 168.1, m 253°, 255-256°, 257°(dec), 253°, 262-263°. Crystallise the quinone from H₂O or acetic acid (golden-yellow prisms). It is soluble in aqueous alkaline solutions, is steam volatile and sublimes readily at 175-180°/1mm. It has UV with λ_{max} at 287 and 377nm (CHCl₃). [Beilstein 8 H 385, 8 I 683, 8 II 433, 8 III 3354, 8 IV 2710.]

5,6-Dimethoxy-1-indanone [2107-69-9] $C_{11}H_{12}O_3$, M 192.2, m 118° to 120°. Crystallise the indanone from MeOH, then sublime it in a vacuum. [Beilstein 8 IV 1985.]

1,4-Dimethoxynaphthalene [10075-62-4] $C_{12}H_{12}O_2$, M 188.2, m 87-88°, 90°. Crystallise the naphthalene from EtOH, MeOH (m 85-86°) or petroleum ether. [Beilstein 6 H 984, 6 III 5261, 6 IV 6546.] **1,5-Dimethoxynaphthalene** [10075-63-5] has m 180°, 183-184°, b 179°/13mm. Crystallise it from EtOH, AcOH (m 178-180°) or *C₆H₆. Also distil it in a vacuum. [Beilstein 6 III 5266, 6 IV 6554.]

3,4-Dimethoxy-6-nitrobenzaldehyde (6-nitroverataldehyde) [20357-25-9] $C_9H_9NO_5$, M 211.2, m 131°, 132°, 133.5-134.5°. The aldehyde is purified by dissolving 9g in 200ml of boiling 95% EtOH, and set aside overnight to crystallise. It is then dried *in vacuo* at 50° and recrystallised from 110ml of 95% EtOH to give 6-7g of aldehyde with m 132-133°. Crystallisation from aqueous EtOH provides light yellow needles. It is light sensitive and should be stored in the dark [Fetscher *Org Synth Coll Vol* 4 735 1963, DOI: 10.15227/orgsyn.033.0065]. [Beilstein 8 H 262, 8 I 610, 8 II 290, 6 III 2065, 6 IV 1785.]

2,6-Dimethoxyphenol (pyrogallol-1,3-diethylether) [91-10-1] $C_8H_{10}O_3$, M 154.2, m 50°, 54-56°, 57°, b 261°/atm, $pK_{Est} \sim 9.6$. Purify the phenol by zone melting or sublimation in a vacuum. Its solubility in H_2O is 2w/v% at ~13°. [Beilstein 6 IV 7329.] It is a food-grade flavor ingredient. **3,5-dimethoxyphenol (phloroglucinol dimethylether)** [500-99-2] has m 42-43°, 45-47°, b 115°/0.04mm, 172-175°/17mm, $pK^{25}_{9.35}$. Purify the phenol by distillation followed by sublimation in a vacuum. [Beilstein 6 IV 7362.]

3,4-Dimethoxyphenyl acetic acid (homoveratric acid) [93-40-3] $C_{10}H_{12}O_4$, M 196.2, m 96°, 97-99°, $pK^{25}_{4.33}$. Recrystallise homoveratric acid from H_2O or $*C_6H_6$ /ligroin. **Homoveratryl chloride** [10131-60-7] $C_{10}H_{11}ClO_3$, M 214.7, has m 40-43°, b 119-119°/0.2mm, d_4^{24} 1.245, n_D^{20} 1.5489, the *amide* has m 142° (from H_2O), and **homoveratronitrile** [93-17-4] $C_{10}H_{11}NO_2$, M 177.2, has m 62-63.5°, b 171-178°/10mm, which fluoresces at λ_{ex} 430nm, λ_{em} 511 (H_2O pH 7.1). [Beilstein 10 H 409, 10 I 197, 10 II 268, 10 III 1459, 10 IV 1509.]

3,5-Dimethoxyphenylacetoneitrile [13388-75-5] $C_{10}H_{11}NO_2$, M 177.2, m 54°, 53°, 57°. Crystallise the nitrile from MeOH or petroleum ether (b 90-110°). [Adams et al. *J Am Chem Soc* 70 664 1948, DOI: 10.1021/ja01182a068; Sankaraman et al. *J Am Chem Soc* 109 5235 1987, DOI: 10.1021/ja00251a032; Beilstein 10 I 198, 10 II 269, 10 III 1470.]

2,6-Dimethoxytoluene [5673-07-4] $C_9H_{12}O_2$, M 152.2, m 39-41°, b 97-99°/15mm, 219-220°/731mm. Sublime 2,6-dimethoxytoluene *in vacuo*. Distil it (preferably under reduced pressure) and/or recrystallise it from pentane, EtOH or aqueous MeOH. [Sankaraman et al. *J Am Chem Soc* 109 5235 1987, DOI: 10.1021/ja00251a032]. [Beilstein 6 H 872, 6 III 4513, 6 IV 5877.]

4,4'-Dimethoxytrityl chloride (DMT) [40615-36-9] $C_{21}H_{19}ClO_2$, M 338.8, m 114°, 119-123°, 123-125°. DMT crystallises from cyclohexane/acetyl chloride as the hydrochloride. Dry it over KOH pellets in a desiccator. When dissolved in $*C_6H_6$ and air is blown through, HCl is removed. It crystallises from Et_2O . [Baeyer & Villiger *Chem Ber* 36 2774 1903, DOI: 10.1002/cber.19030360327; Smith et al. *J Am Chem Soc* 84 430 1962, DOI: 10.1021/ja00862a023; Schaller et al. *J Am Chem Soc* 85 3821 1963, DOI: 10.1021/ja00906a021.] If it has hydrolysed considerably (see OH in IR), then repeat the crystallisation from cyclohexane/acetyl chloride — excess of AcCl is removed in a vacuum over KOH, then recrystallise it from Et_2O . [Beilstein 6 IV 1042.] It is a useful protecting group [Amarnath & Broom *Chem Rev* 77 183 1977, DOI: 10.1021/cr60306a002; Kohli et al. *Tetrahedron Lett* 21 2683 1980, DOI: 10.1016/S0040-4039(00)78579-0; Reddy et al. *Tetrahedron Lett* 28 23 1987, DOI: 10.1016/S0040-4039(00)95639-9] and is removed under mild detritylation conditions, using $ZnBr_2$ in $MeNO_2$ at 18° for ~1min. [Matteucci & Caruthers *Tetrahedron Lett* 21 3243 1980, DOI: 10.1016/S0040-4039(00)78657-6].

4-Dimethylaminoazobenzene (DAB, Dimethyl Yellow, Butter yellow) [60-11-7] $C_{14}H_{15}O_3$, M 225.3, m 111-116°, 118-119°(dec), pK^{15}_2 -5.34 (aqueous H_2SO_4), pK^{15}_2 2.96. Crystallise the dye from acetic acid or isooctane, or from 95% EtOH by adding hot water and cooling. Dry it over KOH under vacuum at 50°. [Beilstein 6 IV 448.] It has indicator properties, viz: it is red between pH 2.9 and 4 and turns yellow above pH 4.0 (cf. pK values). **CARCINOGEN, handle with care.**

4-Dimethylaminobenzaldehyde (Ehrlich's Reagent, DMAB) [100-10-7] $C_9H_{11}NO$, M 149.2, m 73-75°, 73-75°, 74-75°, $pK_{Est} \sim 2.6$. Crystallise DMAB from water, hexane, or from EtOH (2ml/g), after charcoal treatment, by adding excess of water. Alternatively, dissolve it in aqueous acetic acid, filter it, and precipitate it with ammonia. Finally recrystallise it from EtOH. It has been used for the detection of primary amino groups

[Gargiulo et al. *J Am Chem Soc* **116** 3760 1944, DOI: 10.1021/ja00088], pyrroles [Iyer et al. *J Org Chem* **59** 6038 1994, DOI: 10.1021/jo00099a039], and is also used for diagnostic purposes, e.g. differentiating between true scarlet fever and serum eruptions. [Campaigne & Archer *Org Synth Coll Vol* **4** 331 1963, DOI: 10.15227/orgsyn.033.0027 *Beilstein* **14** IV 51.] The *hydrochloride* has **m** 109° (from EtOH/HCl/Et₂O).

4-Dimethylaminobenzoic acid [619-84-1] C₉H₁₁NO₂, **M** 165.2, **m** 241-243°(dec), 242.5-243.5°(dec), **pK₁** 2.51, **pK₂** 6.03. Crystallise the acid from H₂O or EtOH/water. [*Beilstein* **14** IV 1164.]. **4-Dimethylaminobenzoyl chloride** [4755-50-4] C₉H₁₀ClNO, **M** 183.6, has **m** 145-149°, **b** 180°/15mm, is corrosive and moisture sensitive. The *2-ethylhexyl ester* (Padimate O) [21245-02-3] C₁₇H₂₇NO₂, **M** 277.4, has **b** 325°/atm, **d**₄²⁵ 0.995, **n**_D²⁰ 1.552, and is used as a sun screen.

4-Dimethylaminobenzophenone [530-44-9] C₁₅H₁₅NO, **M** 225.3, **m** 89°, 90°, 92.5°, 92-93°, **pK_{Est}** ~2.7. Crystallise the pale green *p*-dimethylaminobenzophenone from EtOH. Dissolve 100g in 600ml of boiling EtOH, add 5g of charcoal, cool, isolate the solid by centrifugation and similarly wash the pale crystals with ice-cold EtOH. When filtered by suction, EtOH solution remains on the crystals and turns deep green in air. Dry it in a vacuum and store it in the dark. The *hydrazone* has **m** 128-130° and forms a ketyl with potassium. [Hurd & Webb *Org Synth Coll Vol* **1** 217 1941, DOI: 10.15227/orgsyn.007.0024; *Beilstein* **14** H 82, **14** I 288, **14** III 218, **14** IV 248.]

N,N-Dimethylamino-*p*-chlorobenzene (*p*-chloro-*N,N*-dimethylaniline) [698-69-1] C₈H₁₀ClN, **M** 155.6, **m** 32-33.5°, 35.5°, **b** 114-115°/12m, 231°/atm, 264.2-265.6°/760mm, **pK₂₅** 4.395. Purify it by vacuum sublimation, or distillation preferably under vacuum. [Guarr et al. *J Am Chem Soc* **107** 5104 1985, DOI: 10.1021/ja00304a015]. The *picrate* has **m** 126-128° (from methanol).

4-Dimethylamino cinnamaldehyde [6203-18-5] C₁₁H₁₃NO, **M** 175.2, **m** 138-140°, 141°. The aldehyde crystallises from EtOH or ligroin and is dried *in vacuo*. Its solubility in CHCl₃/EtOH (1:1), and in dioxane is 5w/v%. Store below 0°. The *oxime* has **m** 157° (from ligroin). The *phenylhydrazone* has **m** 169° (from MeOH). It is used as a reagent for amines (with NH₃, UV has λ_{max} at 630nm). [König et al. *Chem Ber* **61** 2074 1928, DOI: 10.1002/cber.19280610903; Qureshi & Kahn *Anal Chim Acta* **86** 309 1976, DOI:10.1016/S0003-2670(01)83053-1; *Beilstein* **14** III 184, **14** IV 197.]

4-Dimethylamino trans-cinnamic acid [1552-96-1] C₁₁H₁₃NO₂, **M** 191.2, **m** 218-220°(dec), 222°(dec), 225° (vigorous effervesces), 227-228°(dec), **pK_{Est(1)}** ~2.2, **pK_{Est(2)}** ~4.6. Recrystallise the yellow acid from EtOH and sublime it at high vacuum. The *methyl ester* has **m** 134-135° (golden yellow plates from Me₂CO), and forms a colourless crystalline perchlorate salt [C₁₂H₁₅NO₂.HClO₄] with **m** 169-170° (from 70% aqueous HClO₄) [Pfeiffer & Haefelin *Chem Ber* **55** 1769 1922, DOI: 10.1002/cber.19220550632]. The *ethyl ester* has **m** 77-78° (from aqueous EtOH or *C₆H₆). [Shoppee *J Chem Soc* 982 1930, DOI: 10.1039/JR9300000968; Galat *J Am Chem Soc* **68** 376 1946, DOI: 10.1021/ja01207a009; for its photophysics in various solvents see Bangal & Chakravorti *J Photophys Photobiol A: Chemistry* **116** 191 1998, DOI:10.1016/S1010-6030(98)00303-7]. The high dipole moment of 4.6D (in *C₆H₆) for the ethyl ester is consistent with a zwitterionic-type *trans* structure [Weizmann *Trans Faraday Soc* **35** 329 1940, DOI: 10.1039/TF9403500329]. [*Beilstein* **14** H 522, **14** II 318, **14** III 1306, **14** IV 1766.]

2S,3R-(+)-4-Dimethylamino-1,2-diphenyl-3-methyl-2-butanol [(+) 38345-66-3; (±) 4125-61-5] C₁₉H₂₅NO, **M** 283.4, **m** 54-55°, 55-57°, for free base, [α]₅₄₆²⁰ +9.3 (c 9.6, EtOH), [α]_D²⁰ +7.7 (c 9.6, EtOH), **pK_{Est}** ~10.0. Purify the *hydrochloride* by dissolving 1.5g in 13.5 ml of 5N HCl, heat to boiling and evaporate in a vacuum. Recrystallise the *hydrochloride* three times from MeOH/EtOAc giving **m** 189-190°, [α]_D -33.7 (c 1, H₂O) {the enantiomer [72541-03-8] has +34.2°}. The *hydrochloride* in the minimum volume of water is basified with aqueous 5N NaOH and extracted with Et₂O. The extract is dried (K₂CO₃) and evaporated, leaving a residue which is stored in a desiccator over solid KOH, as a low melting solid. It can be recovered using these procedures from asymmetric reductions with LAH, and re-used. [Pohland & Sullivan *J Am Chem Soc* **77** 3400 1955, DOI: 10.1021/ja01617a081; Pohland et al. *J Org Chem* **28** 2483 1963, DOI: 10.1021/jo01044a530; *Beilstein* **13** IV 2221.] ;

***N,N*-Dimethylaniline** [121-69-7] $C_8H_{11}N$, M 121.2, m 2°, b 84°/15mm, 192°/760mm, d_4^{20} 0.956, n_D^{25} 1.5556, pK^{25} 5.07. Primary and secondary amines (including aniline and monomethylaniline) can be removed by refluxing for 4-5 hours with excess acetic anhydride, and then fractionally distilling. Crocker and Jones (*J Chem Soc* 1808 1959, DOI: 10.1039/JR9590001808) used four volumes of acetic anhydride, then distilled off the greater part of it, and dissolved the residue in ice-cold dilute HCl. Non-basic materials were removed by ether extraction, then the dimethylaniline was liberated with ammonia, extracted with ether, dried, and distilled under reduced pressure. Metzler and Tobolsky (*J Am Chem Soc* 76 5178 1954, DOI: 10.1021/ja01649a072) refluxed it with only 10% (w/w) of acetic anhydride, then cooled and poured it into excess 20% HCl, which, after cooling, was extracted with diethyl ether. (The amine hydrochloride remains in the aqueous phase.) The HCl solution was cautiously made alkaline to phenolphthalein, and the amine layer was drawn off, dried over KOH and fractionally distilled under reduced pressure, under nitrogen. Suitable drying agents for dimethylaniline include NaOH, BaO, CaSO₄, and CaH₂.

Other purification procedures include the formation of the *picrate* (m 163° from Me₂CO or EtOH/H₂O), prepared in *benzene solution and recrystallised to constant melting point, then decomposed with warm 10% NaOH and extracted into ether: the extract was washed with water and distilled under reduced pressure. The *oxalate* salt has also been used for purification. The base has been fractionally crystallised by partial freezing and also from aqueous 80% EtOH then from absolute EtOH. It has been distilled from zinc dust, under nitrogen. Its solubility in H₂O is 2v/v% at 20°. [Beilstein 12 H 141, 12 I 151, 12 II 2, 12 III 245, 12 IV 243.] **TOXIC.**

2,3-Dimethylaniline [2,3-xylidine (*vic*, *o*)] [87-59-2] $C_8H_{11}N$, M 121.2, m 2°, 2.5°, b 106°/15mm, 223°/760mm, d_4^{20} 1.570, n_D^{20} 0.991, pK^{25} 4.70. Purify *vic*-xylidine by conversion into a derivative (see below), recrystallise the derivative, decompose the derivative with aqueous NaOH and fractionally distil the liquid base. The *acetyl* derivative has m 135° (from EtOH), and the *formyl* derivative has m 102° (from EtOH). [Beilstein 12 H 1101, 12 III 2438, 12 IV 2497.] **TOXIC.** **2,4-Dimethylaniline** [2,4-xylidine (*uns*, *m*)] [95-68-1] has m 11°, 16°, b 212°/736mm, 213-214°/760mm, 218°/760mm, d_4^{20} 0.974, n_D^{20} 1.5604, pK^{25} 4.89. Convert *uns*-xylidine to a derivative (see below) which, after recrystallisation, is decomposed with alkali to give the free base. Dry it over KOH and fractionally distil. The *acetyl* derivative has m 130°, the *benzoyl* derivative has m 192°, and the *picrate* has m 209°. [Beilstein 12 H 1111, 12 IV 2545.] **TOXIC.** **2,5-Dimethylaniline** [2,5-xylidine (*p*)] [95-78-3] has m 6°, 11°, 15.5°, b 104°/15mm, 215°/739mm, 218°/760mm, d_4^{20} 0.974, n_D^{20} 1.5604, pK^{25} 4.53. Convert *p*-xylidine to a derivative (see below) which, after recrystallisation, is decomposed with alkali to give the free base. Dry over KOH and fractionally distil. The *acetyl* derivative has m 142° (from H₂O or toluene), and the *benzoyl* derivative has m 140° (from EtOH). [Beilstein 12 H 1135, 12 IV 2567.] **TOXIC.** **2,6-Dimethylaniline** [2,6-xylidine (*vic*, *m*)] [87-62-7] has m 10-12°, 11°, b 98°/14mm, 210-211°/736mm, 214°/739mm, 215°/760mm, d_4^{20} 0.974, n_D^{20} 1.5604, pK^{25} 3.95. Convert *vic*-xylidine to a derivative (see below) which, after recrystallisation, is decomposed with alkali to give the free base. Dry it over KOH and fractionally distil. The *formyl derivative* [*N*-(2,6-dimethylphenyl)formamide] M 149.2, [607-92-1] has m 168-170°, the *acetyl* derivative has m 177°, the *benzoyl* derivative has m 168°, and the *picrate* has m 180°. [Beilstein 12 H 1107, 12 I 482, 12 II 604, 12 III 2462, 12 IV 2521.] **TOXIC.** **3,4-Dimethylaniline** [3,4-xylidine] [95-64-7] has m 49-51°, 51°, b 116-118°/25mm, b 226°/760mm, pK^{25} 5.17. Crystallise it from ligroin and distil it under vacuum. [Beilstein 12 H 1103, 12 IV 2502.] **TOXIC.** **3,5-Dimethylaniline** [3,5-xylidine (*sym*, *m*)] [108-69-0] has m 7-9°, 10°, 9-11°, 105-106°/15mm, 220-221°/atm, 221-222°/atm, d_4^{20} 0.974, n_D^{20} 1.557, pK^{25} 4.91. Convert *sym*-xylidine to a derivative (see below) which, after recrystallisation, is decomposed with alkali to give the free base. Dry it over KOH and fractionally distil it. The *acetyl* derivative has m 144°, the *benzoyl* derivative has m 136° and the *picrate* has m 209° (from H₂O, EtOH or 10% AcOH). [Beilstein 12 H 1131, 12 IV 2561.]

9,10-Dimethylantracene [781-43-1] $C_{16}H_{14}$, M 206.3, m 180-181°, 182-184°. Purify 9,10-dimethylantracene by crystallising from EtOH, and by recrystallising from the melt. [Beilstein 5 IV 2329.]

7,12-Dimethylbenz[*a*]anthracene (DMBA) [57-97-6] $C_{20}H_{16}$, M 256.4, m 122-123°, b 480°/atm. Purify DMBA by chromatography on alumina/toluene or *benzene in which it is freely soluble. Crystallise it from acetone/EtOH. Its solubility in H₂O is greatly increased in the presence of purines and purine nucleosides. [Beilstein 5 IV 2587.] **TUMOUR INITIATOR.**

2,3-Dimethylbenzoic acid [603-79-2] $C_9H_{10}O_2$, M 150.2, m 144°, 146°, 147°, pK^{25} 3.72. Crystallise the acid from EtOH. It is volatile in steam. The *amide* has m 156° (from H_2O). [Beilstein 9 H 531, 9 III 2434, 9 IV 1797.] **2,4-Dimethylbenzoic acid** [611-01-8] has m 125°, 125-126°, 126-127°, b 267°/727mm, pK^{25} 4.22. Recrystallise the acid from EtOH or H_2O , and sublime it in a vacuum. [Beilstein 9 H 531, 9 III 2436, 9 IV 1801.] **2,5-Dimethylbenzoic acid** [610-72-0] has m 134°, b 268°/760mm, pK^{25} 4.00. Steam distil the acid, then crystallise it from EtOH or H_2O (m 134-134.5°). [Beilstein 9 H 534, 9 IV 1802.] **2,6-Dimethylbenzoic acid** [632-46-2] has m 117°, b 118-119°/2.5mm, pK^{25} 3.35. Steam distil the acid, and crystallise it from EtOH or H_2O (m 116.3-116.7°). The *N*-dimethylamide has m 62-63° (from Et_2O). [Beilstein 9 H 531, 9 IV 1798.] **3,4-Dimethylbenzoic acid** [619-04-5] has m 163-165°, 166°, 166-167°, pK^{25} 4.50. Crystallise it from EtOH or H_2O (m 168-168.5°), and sublime it *in vacuo*. The *phenyl ester* has m 68° (from EtOH or petroleum ether) and b 155-157°/2mm. [Beilstein 9 II 353, 9 III 2441, 9 IV 1803.] **3,5-Dimethylbenzoic acid** [499-06-9] has m 170°, 171.1°, 169-172°, pK^{25} 4.30. Distil the acid in steam, crystallise it from H_2O (m 171.2-171.7°) or EtOH, and sublime it *in vacuo*. [Beilstein 9 H 536, 9 III 244, 9 IV 1806.]

4,4'-Dimethylbenzophenone [611-97-2] $C_{15}H_{14}O$, M 210.3, m 95°, 93-97°, b 150-152°/2mm, 200°/17mm, 333-334°/725mm. Purify the benzophenone by zone refining or distillation, preferably in a vacuum. [Beilstein 7 III 2181, 7 IV 1434.]

2,5-Dimethyl-1,4-benzoquinone [137-18-8] $C_8H_8O_2$, M 136.2, m 123°, 124-125°, 126-127°. Crystallise the quinone from EtOH. [Beilstein 7 IV 2090.] **2,6-Dimethyl-1,4-benzoquinone (Phloron)** [527-61-7] has m 71° to 72°, 72° (sealed tube). Crystallise the quinone from water/EtOH (8:1). [Beilstein 7 IV 657, 7 III 3402, 7 IV 2096.]

3,3'-Dimethylcarbanilide [*N,N'*-bis-(*m*-tolyl)urea] [620-50-8] $C_{15}H_{16}N_2O$, M 240.3, m 220°, 225°. Crystallise urea from ethyl acetate. The UV has λ_{max} at 258nm (EtOH). The *hydrochloride* has m 162°. [Beilstein 12 III 1970, 12 IV 1829.]

1,1-Dimethyl-1H-indene [18636-55-0] $C_{11}H_{14}$, M 144.2, b 57°/4.8mm, 115°/20mm. Purify the oily indene by gas chromatography or by fractional distillation. [Bosch & Brown *Can J Chem* 42 1718 1964, DOI: 10.1139/v64-256, cf. Beilstein 5 I 251, 5 III 1377.]

1,5-Dimethylnaphthalene [571-61-9] $C_{12}H_{12}$, M 156.2, m 78-82°, 82°, b 265-266°/atm. Crystallise it from 85% aqueous EtOH. [Beilstein 5 IV 1709.] **2,3-Dimethylnaphthalene** [581-40-8] has m 103°, 104-104.5°, 106°, 265-268°/atm, 269°/atm. Steam distil the naphthalene and crystallise it from EtOH. [Beilstein 5 IV 1713.] **2,6-Dimethylnaphthalene** [581-42-0] has m 110-111°, b 122.5-123.5°/10mm, 261-262°/760mm. Distil it in steam and crystallise it from EtOH. [Beilstein 5 IV 1714.]

3,3'-Dimethylnaphthidine (4,4'-diamino-3,3'-dimethyl-1,1'-binaphthyl) [13138-48-2] $C_{22}H_{20}N_2$, M 312.4, m 213°. Recrystallise the naphthidine from EtOH or petroleum ether (b 60-80°). It is used as an indicator for metal titration, e.g. with Zn it undergoes colour change from red-violet to greyish-green at pH 5 in acetate buffer [Brown & Hayes *Anal Chim Acta* 9 6 1953, DOI: 10.1016/S0003-2670(01)80726-1; Flaschka & Frausnitz *Z Anal Chem* 114 421 1955]. [Beilstein 13 IV 493.]

***N,N*-Dimethyl-3-nitroaniline** [619-31-8] $C_8H_{10}N_2O_2$, M 166.1, m 57°, 60°, 61°, 280°/atm, pK^{25} 2.63. Crystallise the aniline from EtOH. The *picrate* has m 119° (from EtOH). [Beilstein 12 H 701, 12 III 1544, 12 IV 1591.] ***N,N*-Dimethyl-4-nitroaniline** [100-23-2] has m 163-166°, 164°, 165°, 164.5-165.2°, pK^{25} 0.61 (0.92). Crystallise the nitroaniline from aqueous EtOH, EtOH or MeOH (m 163.5-164°). Dry it *in vacuo*. The *N*-methiodide has m 161°(dec) (from H_2O). [Beilstein 12 H 714, 12 III 1584, 12 IV 1616.]

***N,N*-Dimethyl-*p*-nitrosoaniline (4-nitroso-*N,N*-dimethylaniline)** [138-89-6] $C_8H_{10}N_2O$, M 150.2, m 85-87°, 86-87°, 92.5-93.5°(also reported), b 191-192°/100mm, pK^{25} 4.54. Recrystallise the nitroso-aniline from petroleum ether or $CHCl_3/CCl_4$ and dry it in air. It is steam volatile. Alternatively, suspend it in H_2O , heat to boiling and add HCl until it dissolves. Filter, cool and collect the *hydrochloride* [42344-05-8] with m 177° after recrystallisation from H_2O containing a small amount of HCl. The *hydrochloride* (e.g. 30g) is made into a

paste with H₂O (100ml) in a separating funnel. Add cold aqueous 2.5 NaOH or Na₂CO₃ to a pH of ~ 8.0 (green color due to the free base) and extract with toluene, CHCl₃ or Et₂O. Dry the extract (K₂CO₃), filter, distil off the solvent, cool the residue and collect the crystalline *free base*. Recrystallise it as above and dry it in air. [Beilstein 12 IV 1558.]

R-(+)-N,N'-Dimethyl-1-phenethylamine [19342-01-9] and **S-(-)-N,N'-Dimethyl-1-phenethylamine** [17279-31-1] C₁₀H₁₅N, M 149.2, b 81°/16mm, [α]_D²⁰ (+) and (-) 50.2 (c 1, MeOH), d₄²⁰ 0.908, n_D 1.502-1.504, pK_{Est} ~9.0 (for RS). The amine is mixed with aqueous 10N NaOH and extracted with toluene. The extract is washed with saturated aqueous NaCl, dried over K₂CO₃, and transferred to fresh K₂CO₃ until the solution is clear, and is filtered. The filtrate is distilled. If a short column packed with glass helices is used, the yield is increased but a purer product is obtained. [For the optical resolution of phenethylamine see Ingersoll *Org Synth Coll Vol* 2 506 1943, DOI: 10.15227/orgsyn.017.0080; Snyder & Brewster *J Am Chem Soc* 71 291 4165 1949, DOI:10.1021/ja01169a077; for methylation of amines with HCHO/HCO₂H see Clarke et al *J Am Chem Soc* 55 71 4571 1933, DOI: 10.1021/ja01338a041; Cope et al. *J Am Chem Soc* 71 3929 1949, DOI: 10.1021/ja01180a014.] The (-)-*picrate* has m 140-141° (from EtOH). The *racemate* [1126-71-2] has b 88-89°/16mm, 92-94°/30mm, 194-195°/760mm, d₄²⁰ 0.908. [Beilstein 13 III 2392.]

2,3-Dimethylphenol [526-75-0] C₈H₁₀O, M 122.2, m 70-73°, 75°, b 120°/20mm, 216.9°/760mm, 218°/atm, pK₂₅ 10.54. Crystallise 2,3-xyleneol from aqueous EtOH. [Beilstein 6 IV 3096.] **2,5-Dimethylphenol** [95-87-4] has m 73°, 74.8°, 76°, 74-78°, b 211.5°/762mm, 211-213°/atm, pK₂₅ 10.41. Crystallise 2,5-xyleneol from EtOH/ether. [Beilstein 6 IV 3164.] **2,6-Dimethylphenol** [576-26-1] has m 43-45°, 49°, b 203°/760mm, pK₂₅ 10.61. Fractionally distil 2,6-xyleneol under nitrogen, crystallise it from *benzene or hexane, and sublime it at 38°/10mm. It is used for the determination of nitrate in water samples for assessing water quality [ISO International Organisation for Standardization, 1986]. [Beilstein 6 IV 3122.] **3,4-Dimethylphenol** [95-65-8] has m 62.5°, 63°, 65°, 65-68°, 67°, b 142°/50mm, 225°/757mm, 227°/atm, pK₂₅ 10.36. Heat 3,4-xyleneol with an equal weight of concentrated H₂SO₄ at 103-105° for 2-3 hours, then dilute it with four volumes of water, reflux it for 1 hour, and either steam distil or extract it repeatedly with diethyl ether after cooling to room temperature. The steam distillate is also extracted and evaporated to dryness. (The purification process depends on the much slower sulfonation of 3,4-dimethylphenol than most of its likely contaminants.) It can also be crystallised from water, hexane or petroleum ether, and sublimed in a vacuum. [Kester *Ind Eng Chem* 24 770 1932, DOI: 10.1021/ie50271a010; Bernasconi & Paschalis *J Am Chem Soc* 108 2969 1986, DOI: 10.1021/ja00271a027; Beilstein 6 IV 3099.] **3,5-Dimethylphenol** [108-68-9, 50356-23-5] has m 61-65°, 62.5°, 63.6°, 65.5°, 68°, b 219°/atm, 222°/atm, pK₂₅ 10.19. Purify it as for 3,4-dimethylphenol. [Beilstein 6 IV 3141.]

2,6-Dimethylphenylisocyanide [2769-71-3] C₉H₉N, M 131.2, m 72-76°, 72-75°. Like most isocyanides (isonitriles, carbylamines), it reacts with transition metals to form aldimines [Malatesta & Bonati *Isonitrile Complexes of Metals* Wiley Interscience, NY 1969, SBN 470 565551, Treichel in *Advances in Organometallic Chemistry Vol 11*, (Stone & West Eds) Academic Press NY 1973], and because of the crowded area around the nitrogen atom it coordinates with the metal in metal catalysts to enhance their stereoselectivity. The isocyanide can function as a nucleophile, electrophile, carbene, radical acceptor or pseudohalogen [Ugi (ed.) *Isonitrile Chemistry* Academic Press NY 1971, Library of Congress No 73-84156]. Among the general methods for preparing isocyanides the one that uses the **Vilsmeier reagent** {(chloromethylene)dimethyliminium chloride, [ClCH=NMe₂]⁺ Cl⁻, M 128.0, m 132°(dec), [3724-43-4]} has been commonly used, and in this case it is prepared *in situ*. Thus a solution of redistilled SOCl₂ (0.5mol) in DMF (150ml) is added slowly to a stirred solution of ***N-(2,6-dimethylphenyl)formamide** (0.5mol, see below) {M 149.2, m 165°, see [607-92-1]} under N₂ and the temperature is kept below -50°. On completion, the cooling bath is momentarily removed to allow the temperature to rise to -45°, then the cooling bath is replaced while anhydrous Na₂CO₃ (1mol) is added very carefully with rapid stirring till evolution of CO₂ ceases. Stirring is continued while the mixture is heated to 35°, then kept at ~25° for 1 hour, diluted with ice-water and extracted into pentane. The extract is separated, dried (Na₂SO₄), and evaporated to give **2,6-dimethylphenylisocyanide** as a crystalline colourless solid in 74% yield. It can be recrystallised from pentane, and should be stored under N₂ between 2° to 8°. It is **toxic**. [Review: Periasamy and Walborsky *Org Prep Proc Int* 11 295 1979, DOI:10.1080/00304947909355415.]

***N-(2,6-Dimethylphenyl)formamide**, [607-92-1] M 149.2, m 164° (164-5°, 168-170° also reported, and 176-177° on rapid heating) is prepared by heating 2,6-dimethylaniline (see [87-62-7]) with formic acid, pouring into

H₂O, filtering, drying and recrystallising from EtOH. [Hodgkinson & Limpach *J Chem Soc* **77** 65 (67) 1900, DOI: 10.1039/CT9007700065; *Beilstein* 12 H 1109, 12 I 604.] It forms a 2:1 complex with gold.

Dimethyl phthalate [131-11-3] C₁₀H₁₀O₄, M 194.2, m 2°, b 100°/1mm, 148°/10mm, 194°/60mm, 210°/100mm, 283°/760mm, d₄²⁰ 1.190, d₄²⁵ 1.1865, n_D²⁰ 1.5149. Wash the ester with aqueous Na₂CO₃, then distilled water, dry (CaCl₂) and distil it under reduced pressure (b 151-152°/0.1mm). It is used as a plasticiser, and an insect repellent. [*Beilstein* 7 IV 3170.] TOXIC.

2,4-Dimethylresorcinol [634-65-1] C₈H₁₀O₂, M 138.1, m 111°, 112°, b 286.4°/760mm, d 1.2, n_D 1.582, pK_{Est(1)} ~9.8, pK_{Est(2)} ~11.7. Crystallise the resorcinol from petroleum ether (b 60-80°), *C₆H₆/petroleum ether or toluene/BuOAc. It sublimes at 100-110°/1mm. [*Beilstein* 6 H 918, 6 II 891, 6 III 4588, 6 IV 5955.]

Dimethyl terephthalate (DMT) [120-61-6] C₁₀H₁₀O₄, M 194.2, m 139-141°, 141.5-141.8°, 142°, b 288°/atm, pK_a -7.1, pK_b -6.6. Purify it by recrystallisation from aqueous EtOH, MeOH or CCl₄; or by zone melting. [*Beilstein* 6 H 843, 6 III 4250, 6 IV 3303.] Plasticiser used for the preparation of polyesters, e.g. for plastic bottles.

N,N-Dimethyl-o-toluidine [609-72-3] C₉H₁₃N, M 135.2, b 68°/10mm, 76°/18mm, 211-211.5°/760mm, d₄²⁰ 0.937, n_D²⁰ 1.53664, pK²⁵ 6.11. Isomers and other bases are removed by heating in a water bath for 100 hours with two equivalents of 20% HCl and two and a half volumes of 40% aqueous formaldehyde, then making the solution alkaline and separating the free base. After washing well with water, it is distilled at 10mm pressure and redistilled at ambient pressure [Braun & Aust *Chem Ber* **47** 260 1914, DOI: 10.1002/cber.19140470138]. Other procedures include drying with NaOH, distilling from zinc in an atmosphere of nitrogen under reduced pressure, and refluxing with excess of acetic anhydride in the presence of concentrated H₂SO₄ as catalyst, followed by fractional distillation in a vacuum. The *picrate* has m 124-125°. [*Beilstein* 12 H 857, 12 III 1843, 12 IV 1747.] **N,N-Dimethyl-m-toluidine (m-methyl-N,N-dimethylaniline)** [121-72-2] has b 72-74°/5mm, 128°/57mm, 211.5-212.5°/760mm, 215°/atm, d₄²⁰ 0.93, n_D²⁰ 1.5500, pK²⁵ 5.34. Reflux it for 3 hours with 2 molar equivalents of Ac₂O, then fractionally distil it under reduced pressure. Alternatively, dry over BaO, distil and store it over KOH. The *hydrochloride* has m 176° (from EtOH) and the *picrate* has m 131°. Methods described for N,N-dimethylaniline are applicable. [*Beilstein* 12 H 857, 12 III 1953, 12 IV 1815.] **N,N-Dimethyl-p-toluidine (p-methyl-N,N-dimethylaniline)** [99-97-8] has b 76.5-77.5°/4mm, 93-94°/11mm, b 211°/760mm, d₄²⁰ 0.937, n_D²⁰ 1.5469, pK^{25.5} 5.56, pK²⁵ 5.63. Reflux for 3 hours with 2 molar equivalents of Ac₂O, then fractionally distil it under reduced pressure. Alternatively, dry it over BaO, distil and store it over KOH. The *picrate* has m 128° (from EtOH). Methods described for N,N-dimethylaniline are applicable here. [*Beilstein* 12 H 902, 12 III 2026, 12 IV 1874.]

2,4-Dinitroaniline [97-02-9] C₆H₅N₃O₄, M 183.1, m 176-178°, 187.5-188°, 180°, ε₃₄₈ 12,300 in diluteaqueous HClO₄, pK²⁵ -4.27 (aqueous H₂SO₄). Crystallise the nitroaniline from boiling EtOH by adding one-third volume of H₂O and cooling slowly. It also crystallises from EtOH (yellow needles) or Me₂CO (green-yellow plates). Dry it on a steam oven. The *N-acetyl* derivative has m 180°. [*Beilstein* 12 IV 1689.]

2,6-Dinitroaniline (DNAN) [606-22-4] has m 134°(dec), 139-140°, pK²⁵ -5.37 (aqueous H₂SO₄). Purify the nitroaniline by chromatography on alumina, then crystallise it from *benzene or EtOH (pale orange needles). The *N-acetyl* derivative has m 197° (from EtOH). [Shultz *Org Synth Coll Vol* **4** 364 1963, DOI: 1.15227/orgsyn.031.0045; *Beilstein* 12 IV 1729.]

2,4-Dinitroanisole [119-27-7] C₇H₆N₂O₅, M 198.1, m 87-88°(metastable), 94.5°, 94-95°, b 206-207°/12mm. Crystallise the anisole from aqueous EtOH. The *naphthalene complex* has m 50° (from EtOH). [*Beilstein* 6 H 254, 6 III 858, 6 IV 1372.] It is a low sensitivity explosive. **3,5-Dinitroanisole** [5327-44-6] has m 105-106°. Purify the anisole by repeated crystallisation from EtOH, MeOH or H₂O and dry it in a vacuum desiccator over P₂O₅. The *naphthalene complex* has m 69° (from EtOH). [*Beilstein* 6 III 869, 6 IV 1385.]

1,2-Dinitrobenzene [528-29-0] C₆H₄N₂O₄, M 168.1, m 114-117°, 116.5°, 118°, b 319°/773mm. Steam distil it and then recrystallise it from EtOH. [*Beilstein* 5 IV 738.] **1,3-Dinitrobenzene** [99-65-0] has m 84-86°, 90.5-91°, b 297°/atm, d²⁵ 1.365. Crystallise 1,3-dinitrobenzene from alkaline EtOH solution (20g in 750ml 95%

EtOH at 40°, plus 100ml of 2M NaOH) by cooling and adding 2.5L of H₂O. The precipitate, after filtering off, is washed with H₂O, sucked dry, and crystallised from 120ml, then 80ml of absolute EtOH [Callow et al. *Biochem J* **32** 1312 1938, DOI: 10.1042/bj0321312]. Alternatively, crystallise it from MeOH, CCl₄ or EtOAc. It can be sublimed in a vacuum. [Tanner et al. *J Org Chem* **52** 2142 1987, DOI: 10.1021/jo00387a005; *Beilstein* **5** IV 739.]

1,4-Dinitrobenzene [100-25-4] C₆H₄N₂O₄, M 168.1, m 170-172°, 173°, b 183.4°/34mm, d²⁵ 1.625. Crystallise 1,4-dinitrobenzene from EtOH or EtOAc. Dry it under vacuum over P₂O₅. It can be sublimed in a vacuum. [*Beilstein* **5** IV 741.]

2,4-Dinitrobenzoic acid [610-30-3] C₇H₄N₂O₆, M 212.1, m 176° to 180°, 183°, pK²⁵ 1.42. Crystallise the acid from aqueous 20% EtOH (10ml/g) and let it dry at 100°. [*Beilstein* **9** II 279, **9** III 1776, **9** IV 1239.]

2,5-Dinitrobenzoic acid [610-28-6] has m 178° to 182°, 179.5-180°, pK²⁵ 1.62. Crystallise the acid from distilled H₂O. Dry it in a vacuum desiccator. [*Beilstein* **9** II 279, **9** III 1778, **9** IV 1241.]

2,6-Dinitrobenzoic acid [603-12-3] has m 202-203°, pK²⁵ 1.14. Crystallise the acid from water. [*Beilstein* **9** II 279, **9** III 1778, **9** IV 1242.]

3,4-Dinitrobenzoic acid [528-45-0] has m 166°, 165.5-166.5°, pK²⁵ 2.81. Crystallise the acid from EtOH by addition of water. It is soluble in hot H₂O (solubility is 0.67w/v% at 25°). It sublimes *in vacuo*. [*Beilstein* **9** III 1778, **9** IV 1242.]

3,5-Dinitrobenzoic acid [99-34-3] has m 204° to 206°, 205°, 206.5-207.2°, pK²⁵ 2.73 (2.79). Crystallise the acid from distilled H₂O (solubility is 2w/v% at 100°), or 50% EtOH (4ml/g). Dry it in a vacuum desiccator or at 70° over BaO under a vacuum for 6 hours. It sublimes *in vacuo*. It has been used for the derivatisation of modified polystyrene-divinylbenzene resins for HPLC [Bourque et al. *Anal Chem* **65** 2983 1993, DOI: 10.1021/ac00069a008]. [Brewster et al. *Org Synth Coll Vol* **3** 337 1955, DOI: 1.15227/orgsyn.022.0048; *Beilstein* **9** II 279, **9** III 1779, **9** IV 1242.] **3,5-Dinitrobenzoyl chloride** [99-33-2] C₇H₃ClN₂O₅, M 230.6, has m 68-69°, 69.5°, b 196°/11mm. Crystallise it from CCl₄ or petroleum ether (b 40-60°). It reacts readily with H₂O and should be kept in sealed ampoules. [*Beilstein* **9** IV 1350.] Used for characterisation and group protection of alcohol, amines and amino acids.

2,2'-Dinitrobiphenyl [2436-96-6] C₁₂H₈N₂O₄, M 244.2, m 123-124°, 125°, 126°, b 194°/4mm, 305°/atm. Crystallise the biphenyl from EtOH, AcOH (m 124.5°) or petroleum ether (m 125.5-126°). [*Beilstein* **5** H 538, **5** III 1759, **5** IV 1826.] **2,4'-Dinitrobiphenyl** [606-81-5] has m 92.7-93.7°. Crystallise the biphenyl from EtOH. [*Beilstein* **5** H 538, **5** I 274, **5** II 491, **5** III 1759, **5** IV 1827.] **4,4'-Dinitrobiphenyl** [1528-74-1] has m 240.9-241.8°, 242°. Crystallise the yellow biphenyl from *C₆H₆, EtOH (charcoal) or Me₂CO. Dry it under vacuum over P₂O₅. It is steam volatile. It sublimes at ~138°/2.9x10⁻³mm, and has a musky odour. [*Beilstein* **5** H 584, **5** III 1760, **5** IV 1827.]

2,6-Dinitro-*p*-cresol (2,6-dinitro-4-methylphenol) [609-93-8] C₇H₆N₂O₅, M 198.1, m 78-79°, 80.5°, 81-82°, 85°, pK_{Est}~3.7. Recrystallise the cresol from EtOH. It is slightly soluble in H₂O, but is steam volatile. The *piperidine salt* has m 195°. [*Beilstein* **6** H 414, **6** II 391, **6** III 1390, **6** IV 2152.] **TOXIC IRRITANT.**

4,6-Dinitro-*o*-cresol (4,6-dinitro-2-methylphenol) [534-52-1] has m 85-86°, 86-87°, 87°, pK²⁵ 4.70. The cresol crystallises from aqueous EtOH. [*Beilstein* **6** H 369, **6** III 1276.]

2,4-Dinitrodiphenylamine [961-68-2] C₁₂H₉N₃O₄, M 259.2, m 157°, 159°, 160°, 161°, b 413.8°/760mm, pK_{Est} <0. The amine forms red crystals from aqueous EtOH or CHCl₃/EtOH (m 158°). It dissolves in Me₂CO (25mg/ml) to give a clear orange to red solution. The UV has λ_{max} at 335nm (cyclohexane). [*Beilstein* **12** H 751, **12** III 1683, **12** IV 1692.] **IRRITANT.**

4,4'-Dinitrodiphenylurea [1,3-bis-(4-nitrophenyl)urea] [587-90-6] C₁₃H₁₄N₂O, M 302.2, m 312°(dec). Crystallise the urea from EtOH (m 364°, long heating), EtOH/Me₂CO (m 301-303°, 300-304° dec, 318-319°) or Me₂CO (m 289° dec). Its solubility in Me₂SO is 0.47w/v% at 25°. It sublimes *in vacuo*. [*Beilstein* **12** H 723, **12** II 393, **12** III 1619, **12** IV 1646.]

2,4-Dinitrofluorobenzene (FDNB, 1-fluoro-2,4-dinitrobenzene, Sanger's reagent) [70-34-8] C₆H₃FN₂O₄,

M 186.1, m 25-27°, b 133°/2mm, 137°/2mm, 140-141°/5mm, 178°/25mm, d₄²⁰ 1.483, n_D²⁰ 1.569. Crystallise the reagent from Et₂O or EtOH. Distil it in a vacuum through a Todd Column. If it is to be purified by distillation *in vacuo*, the distillation unit must be allowed to cool before air is allowed into the apparatus; otherwise the residue carbonises spontaneously and an **EXPLOSION** may occur. The material is a **skin irritant** and may cause serious dermatitis. It was developed for peptide analysis, characterisation of hydroxyl compounds, degradation of aldoses and replacement of a phenolic OH group by H. [Sanger *Biochem J* **39** 507 1945, DOI: 10.1042/bj0390507; **40** 261 1946, DOI: 10.1042/bj0400261; **45** 563 1949, DOI: 10.1042/bj0450563, Porter and Sanger *Biochem J* **42** 287 1948, DOI: 10.1042/bj0420287; Finger & Finnerty *Biochem Preparations* **3** 120 1958, See Fieser **1** 321, *Beilstein* **5** IV 742.]

1,8-Dinitronaphthalene [602-38-0] C₁₀H₆N₂O₄, M 218.2, m 161°, 170-171°, 171-172°, 171.3°. Crystallise the yellow crystalline powder from *benzene. The solubility in H₂O is 0.1w/v% at ~25°. It is a yellow colourant, and may react violently if heated or ground. [*Beilstein* **5** H 559, **5** II 455, **5** III 1608.]

2,4-Dinitro-1-naphthol (Martius Yellow) [605-69-6] C₁₀H₆N₂O₅, M 234.2, m 127-133°, 130-133°, pK_{Est} ~3.7. Crystallise the naphthol from *benzene or aqueous EtOH. Moisten with ~15% of H₂O as stabiliser. [*Beilstein* **6** IV 4240.] It is harmful if inhaled, if in contact with skin and irritates eyes, skin and respiratory system.

2,4-Dinitrophenetole [610-54-8] C₈H₈N₂O₅, M 212.2, m 85-86°, 86°, b 210-211°/15mm, 359.1°/atm, d 1.381. Crystallise it from aqueous EtOH. The 1:1 *naphthalene complex* has **m 41°** and is obtained by fusing the compound with naphthalene in various ratios, then crystallising the solidified mix from a little EtOH (Dermer & Smith *J Am Chem Soc* **61** 748 1939, DOI: 10.1021/ja01872a504). [*Beilstein* **6** H 254, **6** III 858, **6** IV 1373.] **TOXIC.**

2,4-Dinitrophenol [51-28-5] C₆H₄N₂O₅, M 184.1, m 108°, 112°, 114°, pK²⁵ 4.12. Crystallise it from *benzene, EtOH, EtOH/H₂O or H₂O acidified with dilute HCl, dry it, then recrystallise it from CCl₄. Dry it in an oven and store it in a vacuum desiccator over CaSO₄. It is sparingly soluble in cold H₂O but more soluble at ~90° (1.2w/w%), and is much more soluble in organic solvents. An aqueous solution is colourless below pH 2.2 and yellow above pH 4.4 (see pK_a, the anion is yellow in colour). It is **TOXIC** and has insecticidal properties. The *benzoate* has **m 132°** (from EtOH). [*Beilstein* **6** IV 1369.]

2,5-Dinitrophenol [329-71-5] has **m 103° to 106°, 108°, pK²⁵ 5.20.** Crystallise 2,5-dinitrophenol from H₂O with a little EtOH. It is sparingly soluble in cold H₂O but more soluble in organic solvents. It is colourless in the acid form and the anion is yellow (cf. pK_a). **EXPLOSIVE** when dry, store it with ~10% H₂O. [*Beilstein* **6** IV 1383.] **TOXIC.**

2,6-Dinitrophenol [573-56-8] has **m 63.0-63.7°, 63-64°, pK²⁵ 3.73.** Crystallise it from H₂O, aqueous EtOH, *C₆H₆/cyclohexane, or *C₆H₆/petroleum ether (b 60-80°, 1:1). Like the preceding phenols, it is colourless in the acid form and the anion is yellow (cf. pK_a). [*Beilstein* **6** III 867, **6** IV 1383.]

3,4-Dinitrophenol [577-71-9] has **m 130° to 135°, 138°, pK²⁵ 5.42.** Steam distil and crystallise it from H₂O then dry it in air. **EXPLOSIVE** when dry, store it with ~10% H₂O. Like the preceding phenols, it is colourless in the acid form and the anion is yellow (cf. pK_a). [*Beilstein* **6** III 868, **6** IV 1384.]

3,5-Dinitrophenol [586-11-8] has **m 125.1°, 126°, pK²⁵ 6.68.** Crystallise it from *C₆H₆ or CHCl₃/petroleum ether. Store it with ~10% water as it is **EXPLOSIVE** when dry. Like the preceding phenols, it is colourless in the acid form and the anion is yellow (cf. pK_a). [*Beilstein* **6** III 869, **6** IV 1383.]

2,4-Dinitrophenylacetic acid [643-43-6] C₈H₆N₂O₆, M 226.2, m 19° to 175°, 179°(dec), b 430.2°/760mm, pK²⁵ 3.50. Crystallise the acid from H₂O. [*Beilstein* **9** IV 1691.]

2,4-Dinitrophenylhydrazine (DNPH) [119-26-6] C₆H₆N₄O₄, M 198.1, m 194°, 197°, 200°(dec), 200-202°, pK_{Est} ~2.0. Crystallise the red DNPH from butan-1-ol (30ml/g), dioxane (10ml/g), EtOH, *C₆H₆ or EtOAc. It is used for characterising carbonyl compounds and for dehydrohalogenation [Fieser **1** 321]. The *hydrochloride* has **m 186° (dec)**. [*Beilstein* **15** IV 380.] **DNPH reagent** is prepared by adding a solution of it (1g) in concentrated H₂SO₄ (7ml) to EtOH (75ml) and dilute with H₂O (250ml) and filter if necessary. *Alternatively*, dissolve DNPH (0.25g or its hydrochloride) in a mixture of concentrated HCl and H₂O (50ml) by warming (at ~95°)

then diluting with H₂O (250ml). This is useful with water-soluble aldehydes and ketones. Crystalline derivatives are prepared by adding a couple of drops (0.05–0.1g) of the carbonyl compound to the DNPH reagent (3ml) and shaking. If no precipitate separates then allow to stand for 15 minutes. A crystalline precipitate separates if an aldehyde or ketone is present, and if this separates as an oil then cool and scratch until crystals are formed. Collect and recrystallise from a suitable solvent. Usually the first crystals are quite pure.

2,4-Dinitroresorcinol [519-44-8] C₆H₄N₂O₆, M 200.1, 146-148°, 147.5°, m 212.5-214.5°, b 280.9°/760mm, d₄²⁵ 1.819, pK_a²⁵ 3.05 (3.79). Crystallise the resorcinol from aqueous EtOH (yellow crystals). Its solubility at 25° is 0.25% in EtOH and 0.08g/L in H₂O. It forms a monobasic Pb salt. Store < 10°. **EXPLOSIVE**. [Beilstein 6 II 824, 6 III 4352, 6 IV 5696.]

3,5-Dinitrosalicylic acid [609-99-4] C₇H₄N₂O₇, M 228.1, m 169° to 172°, 173-174°, pK_a₁²⁵ 0.70, pK_a₂²⁵ 7.40. Crystallise the acid from H₂O (yellow crystals). Used for quantifying carbohydrate levels in blood and reducing substances in urine to form 3-amino-5-nitrosalicylic acid, which absorbs strongly at 540nm [Miller *Anal Chem*, 31 426 1959, DOI: 10.1021/ac60147a030]. [Beilstein 10 IV 270.]

2,6-Dinitrothymol [303-21-9] C₁₀H₁₂N₂O₅, M 240.2, m 54.5-55°, 80-81°, 81°. Crystallise 2,4-dinitrothymol from aqueous EtOH or petroleum ether. [Ganguly & LeFèvre *J Chem Soc* 848 1934, DOI: 10.1039/JR9340000848; Beilstein 6 H 543, 6 III 1911.]

2,3-Dinitrotoluene [602-01-7] C₇H₆N₂O₄, M 182.1, m 59.2°, 61°, 63°. Distil the toluene in steam and crystallise it from H₂O or *benzene/petroleum ether. Store it with 10% H₂O as it could be **EXPLOSIVE** when dry. [Beilstein 5 H 339, 5 III 758, 5 IV 865.]

2,4-Dinitrotoluene [121-14-2] has m 67° to 70.0°, 70.5-71.0°, 71.8-72.2°, b 102.7°/1mm. Crystallise it from Me₂CO, isopropanol or MeOH. Dry it in a vacuum over H₂SO₄. It has also been purified by zone melting. Store it with ~10% of H₂O. It could be **EXPLOSIVE** when dry. [Beilstein 5 H 339, 5 III 759, 5 IV 865.]

2,5-Dinitrotoluene [619-15-8] has m 50.3°, 51.2°, 52.5°, b 284°/atm (reported, but may be dangerous to distil at atm pressure) Crystallise it from *benzene. Store it with ~10% of H₂O. Its solubility in H₂O is very poor (0.03w/v% at 20°). **EXPLOSIVE** when dry. [Beilstein 5 III 760, 5 IV 866.] **TOXIC** to aquatic organisms, keep away from the environment.

2,6-Dinitrotoluene [606-20-2] has m 56° to 61°, 64.3°, 66.1°. Crystallise it from acetone. Store it with 10% of H₂O. **EXPLOSIVE** when dry. [Beilstein 5 III 761, 5 IV 866.]

3,4-Dinitrotoluene [610-39-9] has m 54° to 57°, 58.5°, 60°, 61°. Steam distil it and crystallise it from *C₆H₆/petroleum ether (pale yellow crystals). Store it with 10% of H₂O to avoid **EXPLOSION**. Used as standard in GS-MS. [Beilstein 5 H 341, 5 III 761, 5 IV 866.]

3,5-Dinitro-*o*-toluic acid [28169-46-2] C₈H₆N₂O₆, M 226.2, m 205°, 206°, 207.5-208°, pK_{Est} ~3.0. Crystallise the acid from H₂O or aqueous EtOH. The *ammonium salt* forms yellow crystals from EtOH with m 218-219°, and the *urea salt* has m 189-190° (prisms from EtOH). [Beilstein 9 H 474, 9 II 323, 9 III 2316.]

2,4-Dinitro-*m*-xylene [603-02-1] C₈H₈N₂O₄, M 196.2, m 83-84°. Crystallise the yellow 2,4-dinitro-*m*-xylene from EtOH. [Beilstein 5 H 379, 5 II 295, 5 III 844.]

Dinonyl phthalate (mainly 3,5,5-trimethylhexyl phthalate isomer) [84-76-4, 14103-61-8, 28553-12-0] C₂₆H₄₂O₄, M 418.6, m 26-29°, b 170°/2mm, 413°/atm, d₄²⁰ 0.9640, n_D²⁰ 1.4825. Wash the colourless ester with aqueous Na₂CO₃ then shake it with water. Ether is added to break the emulsion, and the solution is washed twice with water, and dried (CaCl₂). After evaporating the ether, the residual liquid is distilled three times under reduced pressure. It is stored in a vacuum desiccator over P₂O₅. [Beilstein 9 IV 3183.] **IRRITANT**.

4,4'-Di-*n*-pentyloxyazoxybenzene [64242-26-8] C₂₂H₃₀N₂O₃, M 370.6, m 124.5° (with phase change at 82°, becomes clear and remelts at 119°). Crystallise it from Me₂CO, and dry it by heating under vacuum. [For Raman and mesogenic transition studies see Venugopalan & Rossabi *J Chem Phys* 85 5273 1986, DOI: 10.1631/1.1451669; Beilstein 16 III 599, 16 IV 175.]

Diphenic acid (diphenyl-2,2'-dicarboxylic acid) [482-05-3] $C_{14}H_{10}O_4$, M 242.2, m 228-229°, pK²⁵ 3.46. Crystallise diphenic acid from water. Hot Ac₂O provides the *anhydride* below. The *dimethyl ester* m 74° and b 204-204°/14mm, crystallises from MeOH as colourless prismatic plates, and the *diethyl ester* m 74°, crystallises as colourless cubes from EtOH. [Beilstein 9 H 922, 9 IV 3552.] **Diphenic anhydride (diphenyl-2,2'-dicarboxylic anhydride)** [6050-13-1] $C_{14}H_8O_3$, M 466.3, has m 217°, 222-224°, 225-227°. After removing free acid by extraction with cold aqueous Na₂CO₃, the residue is crystallised from acetic anhydride and dried at 100°. Acetic anhydride converts the acid to the anhydride. It also crystallises from *C₆H₆ (m 219°) or chlorobenzene (m 224.5-225.5°). [Beilstein 17 H 526, 17 II 495, 17 III/IV 6425.]

N,N-Diphenylacetamidine [621-09-0] $C_{14}H_{14}N_2$, M 210.3, m 131°, 131-133°, pK²⁵ 8.298. Crystallise it from EtOH, then sublime it under vacuum at ca 96° onto a 'finger' cooled in solid CO₂/MeOH, with continuous pumping to free it from occluded solvent. Store in a sealed container <10°. [Schwarzenbach & Lutz *Helv Chim Acta* 23 1162 1940, DOI: 10.1002/hlca.194002301141; Beilstein 12 H 248, 12 II 144, 12 III 471, 12 IV 384.]

Diphenylacetic acid [117-34-0] $C_{14}H_{12}O_2$, M 212.3, m 147° to 149°, 147.4-148.4°, pK²⁵ 3.94. Crystallise the acid from *benzene, H₂O or aqueous 50% EtOH. [Marvel et al *Org Synth Coll Vol I* 224 1941, DOI: 10.15227/orgsyn.003.0045; Yost & Hauser *J Am Chem Soc* 69 2325 1947, DOI: 10.1021/ja01202a023; Beilstein 9 H 673, 9 IV 2492.] **Diphenylacetyl chloride** [1871-29-3] $C_{14}H_{11}ClO$, M 230.7, has m 49° to 53°, 57° (from petroleum ether), 178°/15mm; **methyl diphenylacetate** [3469-00-9] $C_{15}H_{14}O_2$, M 226.3, has m 59° to 62° (from aqueous MeOH); **diphenylacetic anhydride** [1760-46-9] $C_{28}H_{22}O_2$, M 406.5, has m 98° (96° to 100°), b 182°/3mm, 178°/15mm, and is very reactive and sensitive to moisture, and **diphenylacetamide** [519-87-9] $C_{14}H_{13}NO$, M 211.3, has m 167-168°.

Diphenylacetonitrile [86-29-3] $C_{14}H_{10}$, M 193.2, m 73-75°, 76°, b 181°/12mm. Crystallise the nitrile from EtOH or petroleum ether (b 90-100°), and/or distil it in a vacuum. It is used as a herbicide. [Beilstein 9 IV 2505.]

Diphenylacetylene (tolan) [501-65-5] $C_{14}H_{12}O_2$, M 178.2, m 59°, 61°, 62.5°, b 90-97°/0.3mm, 170°/19mm, 300°/atm. Crystallise tolan from EtOH, and/or distil it in a vacuum. [Beilstein 5 H 656, 5 IV 2276.]

Diphenylamine [122-39-4] $C_{12}H_{11}N$, M 169.2, m 50° to 53°, 62.0-62.5°, pK²⁵ 0.77 (aqueous H₂SO₄). Crystallise diphenylamine from petroleum ether, MeOH or EtOH/water. Dry it under vacuum. [Beilstein 12 H 174, 12 IV 271.] Used for its reaction with oxidising substances, e.g. NO₃, ClO₃, which when used with H₂SO₄ gives a deep blue colour. The *sulfate* [587-84-8] $C_{12}H_{11}N$. H₂SO₄, M 267.3, has m 144-146° (123-125°), b 302°/atm, and is a yellow powder which is almost insoluble in H₂O, but soluble in EtOH/H₂SO₄.

Diphenylamine-2-carboxylic acid (DPC, N-phenylanthranilic acid) [91-40-7] $C_{13}H_{11}NO_2$, M 213.2, m 182-183°, 184°, pK²⁵₁ -1.28 (aqueous H₂SO₄), pK²⁵₂ 3.86 (CO₂H). Crystallise the acid from EtOH (5ml/g, leaflets) or AcOH (2ml/g) by adding hot water (1ml/g). It is an oxidation-reduction indicator for cerimetric titrations and detection of vanadium, and also serves as a source of non-steroidal anti-inflammatory drugs. [Beilstein 14 IV 1019.]

Diphenylamine-2,2'-dicarboxylic acid (2,2'-iminodibenzoic acid) [579-92-0] $C_{14}H_{11}NO_4$, M 257.2, m 296-297°(dec), 298°(dec), 302°(dec), 320°(dec, long heating), pK³⁵₁ 6.05, pK³⁵₂ 7.02 (in 50% aqueous dioxane). Crystallise the acid from EtOH (yellow crystals). It complexes with Cu²⁺, Zn²⁺, Cd²⁺, Co²⁺ and Ni²⁺, and is an oxidation-reduction indicator in acidic solutions, see previous entry. [Beilstein 14 H 354, 14 I 545, 14 III 942, 14 IV 1058.]

9,10-Diphenylanthracene [1499-10-1] $C_{26}H_{18}$, M 330.4, m 245° to 247°. 248-249°. Crystallise the anthracene from acetic acid or xylene [Baumstark et al. *J Org Chem* 52 3308 1987, DOI: 10.1021/jo00391a024]. [Beilstein 5 IV 2807.]

N,N'-Diphenylbenzidine [531-91-9] $C_{24}H_{20}N_2$, M 336.4, m 242°, 245-247°, 248°, 251-252°, pK²⁵ 0.30. Crystallise the benzidine from toluene or ethyl acetate. Store it in the dark. [Beilstein 13 H 223, 13 IV 368.]

***trans-trans*-1,4-Diphenylbuta-1,3-diene (DPB, β,β' -bistyryl) [538-81-8] $C_{16}H_{14}$, M 206.3, m 150° to 152°. 153-153.5°, b 350°/atm.** Its solution in petroleum ether (b 60-70°) is chromatographed on an alumina-Celite column (4:1), and the column is washed with the same solvent. The main zone is cut out, eluted with ethanol and transferred to petroleum ether, which is then dried and evaporated [Pinckard et al. *J Am Chem Soc* **70** 1938 1948, DOI: 10.1021/ja01185a088]. Recrystallise it from hexane. Useful for Diels-Alder reactions. [McDonald & Campbell *Org Synth Coll Vol* 4 99 1973, DOI: 10.15227/orgsyn.040.0036; *Beilstein* **5** H 676, **9** IV 2319.]

***sym*-(1,5)-Diphenylcarbazine [140-22-7] $C_{13}H_{14}N_4O$, M 242.3, m 168-171° 170°, 172°, 175°.** A common impurity is phenyl-semicarbazide which can be removed by chromatography: ~8g in H_2O is placed on a column (polyamide 6 powder, Macherey-Nagel-GmbH-Germany, washed several times with MeOH), eluted with H_2O /MeOH/AcOH (1:3:0.04) at 7-8 drops/second, then eluted with the same solvent mixture but diluted 5 fold with H_2O . The purification is followed by UV light at 280nm. The effluent is evaporated to dryness *in vacuo* at ~28°. [For chromatography, IR & UV see Willems et al. *Anal Chim Acta* **51** 553 1970, DOI:10.1016/S0003-2670(01)95761-7]. Recrystallise it from EtOH by adding CCl_4 to induce crystallisation, or AcOH to give a white crystalline powder which turns pink in air. It is air and light sensitive and should be stored in the dark under N_2 . [*Beilstein* **15** H 292, **15** IV 182.]

1,5-Diphenylcarbazon (phenyldiazinecarboxylic acid 2-phenylhydrazide) [538-62-5] $C_{13}H_{12}N_4O$, M 240.3, m 119° to 123°, 124-127°, 156-159°(dec). It crystallises from EtOH (*ca* 5ml/g), and dry the orange-red needles at 50°. A commercial sample, nominally *sym*-diphenylcarbazon (m 154-156°) was a mixture of diphenylcarbazine and diphenylcarbazon. The former was removed by dissolving 5g of the crude material in 75ml of warm EtOH, then adding 25g Na_2CO_3 dissolved in 400ml of distilled water. The alkaline solution was cooled and extracted six times with 50ml portions of diethyl ether (discarded). Diphenylcarbazon was then precipitated by acidifying the alkaline solution with 3M HNO_3 or glacial acetic acid. It was filtered off, air dried, and stored in the dark [Gerlach & Frazier *Anal Chem* **30** 1142 1958, DOI: 10.1021/ac60138a044]. Other impurities are phenylsemicarbazide and diphenylcarbodiazone. Impurities can be detected by chromatography [Willems et al. *Anal Chim Acta* **51** 544 1970, DOI:10.1016/S0003-2670(01)95761-7]. It is used for detection and estimation of Hg, Zn, Cd, Cr, Cu, Fe, Mg and Mo. [Cheng et al. *Handbook of Organic Analytical Reagents*, Boca Baton 277 1982, *Beilstein* **16** H 24, **16** IV 17.]

Diphenyl carbonate (phenyl carbonate) [102-09-0] $C_{13}H_{10}O_3$, M 214.2, m 79°, 80°, 81-82°, 168°/15mm, 301-302°/760mm, 306°/760mm. Purify it by distillation under reduced pressure, sublimation, or by gas chromatography with 20% Apiezon on Embacel, and crystallisation from EtOH. It is soluble in organic solvents. [*Beilstein* **6** H 158, **6** IV 629.]

Diphenylcyclopropenone (Diphencyprone) [886-38-4] $C_{15}H_{10}O$, M 206.2, m 87-90°(hydrate), 118°, 119-121°, 122°(anhydrous). Crystallise it from cyclohexane. Its UV (MeCN) has λ_{max} at 226, 282, 297nm. It is an anti-baldness substance. [*Beilstein* **17** IV 1736.]

1,1-Diphenylethanol (α -methylbenzhydrol) [599-67-7] $C_{14}H_{14}O$, M 198.3, m 77° to 81°, 80-81°, 81-81.5°, 87°, 90°, b 144-145°/12mm, 260°/760mm (dec), d^{15} 1.1057. Crystallise 1,1-diphenylethanol from *n*-heptane and/or distil it under vacuum. The *benzoyl* derivative has m 115° and the *phenylurethane* has m 119°. [Bromberg et al. *J Am Chem Soc* **107** 83 1985, DOI: 10.1021/ja00287a016; *Beilstein* **6** H 685, **6** I 330, **6** II 639, **6** III 3395, **6** IV 4713.] **2,2-Diphenylethanol [1883-32-5] has m 53° to 56°, 54.5°, 57-58°, 59.1-59.5°, 64-65°, b 150-151°/3mm, 160-162°/5mm, 192-194°/20mm (dec), d^{15} 1.1057.** If too impure, dissolve in Et_2O , wash with H_2O , dry ($MgSO_4$), evaporate and distil through a short column in a vacuum, whereby a clear colourless oil is obtained which solidifies on standing to give a solid with a sharp melting point. It can be recrystallised from petroleum ether. The *tosylate*, m 117-118° (116° also reported) crystallises from Me_2CO as white needles, readily forms an anion which undergoes interesting rearrangements. [Hamrick & Hauser *J Org Chem* **26** 4199 1961, DOI: 10.1021/jo01069a005; Brown & Subba Rao *J Am Chem Soc* **78** 5694 1956, DOI: 10.1021/ja01602a063; Winstein et al. *J Am Chem Soc* **74** 1113 1952, DOI: 10.1021/ja01125a001; *Beilstein* **6** II 640.]

1,1-Diphenylethylene [530-48-3] $C_{14}H_{12}$, M 180.3, m 6°, 8.2°, b 101°/2mm, 134°/10mm, 268-270°/760mm,

d₄²⁰ 1.024, **n**_D²⁰ 1.6088. Distil it under reduced pressure from KOH. Dry it with CaH₂ and redistil it. [Allen & Converse *Org Synth Coll Vol* 1 226 1941, DOI: 10.15227/orgsyn.006.0032; *Beilstein* 5 H 639, 5 III 1975, 5 IV 2173.]

(±)-1*RS*,2*RS*-, (+)-1*R*,2*R*-, (-)-1*S*,2*S*-enantiomers of, and *meso*- **1,2-Diphenylethylene-1,2-diamine** (DPEN) see [16635-95-3], [35132-20-8], [29841-69-8] and [951-87-1] respectively in Chapter 5, Catalysts—Part 2.

***N,N'*-Diphenylethylenediamine** (Wanzlick's Reagent, **1,2-dianilinoethane**) [150-61-8] C₁₄H₁₆N₂, **M 212.3, m 65-66°, 67.5°, b 178-182°/2mm, 228-230°/12mm, pK_{Est(1)} ~0.5, pK_{Est(2)} ~3.8**. Crystallise the reagent from aqueous EtOH or MeOH, and/or distil it in a vacuum. Store <10°. It is a reagent for aldehydes which react with it to form imidazolidines, and used to prepare crystallisable derivatives. Apart from using these imidazolines to characterise the aldehydes, they can also be hydrolysed to provide the pure recovered aldehydes almost quantitatively. [Wanzlick & Löchel *Chem Ber* 86 1463 1953, DOI: 10.1002/cber.19530861116; Giannis et al. *Tetrahedron* 44 7177 1988, DOI: 10.1016/S0040-4020(01)86086-5; Fieser 1 187, 2 98; *Beilstein* 12 H 543, 12 I 282, 12 II 287, 12 III 1042, 12 IV 986.] The *dihydrochloride* [99590-70-2] M 285.2 is commercially available. Useful as a resins stabiliser.

***N,N'*-Diphenylformamidine** [622-15-1] C₁₃H₁₂N₂, **M 196.2, m 142°, 137°, 136-139°, 141°**. Crystallise it from absolute EtOH. The *hydrate* is obtained from aqueous EtOH. [Kuhn & Staab *Chem Ber* 87 272 1954, DOI: 10.1002/cber.19540870223; *Beilstein* 12 H 236, 12 IV 372.]

1,3-Diphenylguanidine [102-06-7] C₁₃H₁₃N₃, **M 211.3, m 146-147°, 148°, 150°, pK²⁵ 10.12**. Crystallise it from toluene, aqueous acetone, Et₂O or EtOH, and dry it in a vacuum. [*Beilstein* 12 H 369, 12 IV 769.]

1,6-Diphenyl-1,3,5-hexatriene (DPH, *dicinnamyl*) [1720-32-7] C₁₈H₁₆, **M 232.3, m 199°, 200-203°**. Crystallise the hexatriene from CHCl₃ or EtOH/CHCl₃ (1:1). [*Beilstein* 5 H 691, 5 IV 2425.]

1,1-Diphenylhydrazine [530-50-7] C₁₂H₁₂N₂, **M 184.2, m 34.5°, 44°, b 120°/1.1mm, 220°/40-50mm, pK_{Est} ~9.1**. Purify it via the *hydrochloride* [530-47-2] M 220.7, which has **m 165-170°(dec)** after crystallisation from aqueous EtOH (plus a few drops of HCl) and recover the free base with aqueous NaOH, extract it into Et₂O, dry it (KOH), filter, evaporate and distil the residue under a vacuum. The distillate crystallises on cooling. The *benzoyl* derivative has **m 194°** (from Me₂CO), and the *4-nitrophenyl hydrazone* (with the aldehyde) has **m 131-132°**. [Koga & Anselme *J Org Chem* 33 3963 1968, DOI: 10.1021/jo01274a068; Prep: Beringer et al. *J Am Chem Soc* 75 2705, DOI: 10.1021/ja01107a046; 7708 1953, DOI: 10.1021/ja01107a047; 84 2819 1962, DOI: 10.1021/ja00873a035; *Beilstein* 15 IV 55.] **1,2-Diphenylhydrazine (hydrazobenzene)** [122-66-7] has **m 124-126°, b 175°/10mm, 222°/40mm, pK_{Est} ~1.7**. Crystallise hydrazobenzene from hot EtOH containing a little ammonium sulfide or H₂SO₃ (to prevent atmospheric oxidation), preferably under N₂. Dry it in a vacuum desiccator, and store it in the dark or under N₂. It has been distilled in a vacuum. Alternatively, crystallise it from petroleum ether (b 60-100°) to constant absorption spectrum. It is almost colourless but in air it turns yellow, then red with the formation of azobenzene. The *hydrochloride* crystallises from EtOH and has **m 163-164°(dec)**; however, hydrazobenzene readily rearranges to *benzidine* in the presence of acid. The *picrate* crystallises from *C₆H₆ and has **m 123°(dec)**. [*Beilstein* 15 H 123, 15 IV 56.]

Diphenylmethane [101-81-5] C₁₃H₁₂, **M 168.2, m 20° to 24°, 25.4°, b 74°/0.4mm, 157°/40mm, 129°/11mm, 260.5°/764.5mm, d**₄²⁵ 1.0008 (liq), **n**_D²⁰ 1.57683, Sublime it under vacuum, or distil it at 72-75°/0.4mm. Recrystallise it from cold EtOH. It has also been purified by fractional crystallisation from the melt. Freely soluble in organic solvents but insoluble in H₂O and liquid NH₃. [Armarego *Aust J Chem* 13 95 1960, DOI: 10.1071/CH9600095.] **Potassium diphenylmethide**, PhCHKPh, is prepared by dissolving K (e.g. 0.55 mole) in liquid NH₃ (800ml) until the blue colour is discharged (forming KNH₂), then adding to diphenylmethane (0.5 mole) in dry Et₂O (800ml), whereby the orange coloured diphenylmethide anion is formed. The NH₃ is evaporated off on a steam-bath (Et₂O being added to maintain the volume at 800ml) to provide a suspension of the methide in Et₂O. The methide can then be used in a variety of carbonylation reactions, e.g. with Dry-Ice (CO₂) it provides a 90% yield of *diphenylacetic acid* [117-34-0]. [Yost & Hauser *J Am Chem Soc* 69 2325 1947, DOI: 10.1021/ja01202a023; Hamrick & Hauser *J Org Chem* 26 4199 1961, DOI: 10.1021/jo01069a005.]

[Hartman & Phillips *Org Synth Coll Vol* **2** 232 1943, DOI: 10.15227/orgsyn.014.0034; *Beilstein* **5** II 498, **5** IV 1841.]

1,1-Diphenylmethylaniline (benzhydrylaniline, aminodiphenylmethane) [91-00-9] $C_{13}H_{13}N$, *M* 183.2, *m* 12°, 34°, *b* 166°/12mm, 295°/atm, d_4^{25} 1.063, n_D^{20} 1.596, pK_{Est} ~9.1. Crystallise the amine from H_2O . The *free base* absorbs CO_2 from the atmosphere; store it accordingly. The *hydrochloride* [5267-34-5] *M* 219.7 *m* 293-295°, crystallises from H_2O . [*Beilstein* **12** H 1323, **12** IV 3282.]

§A polymer bound *benzhydrylaniline hydrochloride* is commercially available.

Diphenylmethyl chloride (benzhydryl chloride) [90-99-3] $C_{13}H_{11}Cl$, *M* 202.7, *m* 15° to 17°, *b* 140°/3mm, 167°/17mm, n_D^{20} 1.5960. Dry the chloride with Na_2SO_4 and fractionally distil it under reduced pressure. [*Beilstein* **5** H 590, **5** I 278, **5** II 500, **5** III 1790, **5** IV 1847.]

***all-trans*-1,8-Diphenyl-1,3,5,7-octatetraene** [3029-40-1] $C_{20}H_{18}$, *M* 258.4, *m* 232°, 235°, 235-237°. Crystallise the yellow octatetraene from EtOH or $CHCl_3/MeOH$. It gives a red colour with concentrated H_2SO_4 and its **1,3,5-trinitrobenzene complex**, *m* 55°, crystallises in red-brown rhombs [Kuhn & Winterstein *Helv Chim Acta* **11** 87 1928, DOI: 10.1002/hlca.19280110107.] Its UV (hexane) has one λ_{max} at 271-272nm (E_{cm}^{mol} 10.8 x 10⁴) [Zechmeister & Pinckard *J Am Chem Soc* **76** 4144 1954, DOI: 10.1021/ja01645a027]. [*Beilstein* **5** H 709, **5** II 620, **5** III 2328, **5** IV 2508.]

***N,N'*-Diphenyl-*p*-phenylenediamine** [74-31-7] $C_{18}H_{16}N_2$, *M* 260.3, *m* 143° to 145°, 148-149°, *b* 219-224°/0.7mm, pK_{Est} <0. Crystallise the diamine from EtOH, chlorobenzene/petroleum ether or *benzene. It has also been crystallised from aniline, then extracted three times with absolute EtOH. [*Beilstein* **13** H 80.]

1,1-Diphenyl-2-picrylhydrazine [1707-75-1] $C_{18}H_{13}N_5O_6$, *M* 395.3, *m* 174°(dec), 178-179.5°(dec). Crystallise the hydrazine from $CHCl_3$, or *benzene/petroleum ether (1:1), then degas it at 100° and <10⁻⁵mm Hg for *ca* 50 hours to decompose the 1:1 molar complex formed with *benzene. [*Beilstein* **15** II 221, **15** IV 1210.] The respective *hydrazyl free radical* [1898-66-4] $C_{18}H_{12}N_5O_6$, *M* 394.3, crystallises from * C_6H_6 /petroleum ether as dark violet prisms with *m* 127-129°(dec), [also ~135°(dec)], and is used as a reducing agent [Schenck et al. *Tetrahedron Lett* 193 1967, DOI: 10.1016/S0040-4039(00)90515-X].

2,2-Diphenylpropionic acid [5558-66-7] $C_{15}H_{14}O_2$, *M* 226.3, *m* 172°, 173-174°, 175°, *b* 300°/atm, pK_{Est} ~3.8. Crystallise the acid from EtOH. [*Beilstein* **9** II 474.] **3,3-Diphenylpropionic acid** [606-83-7] has *m* 151° to 154°, 155°, pK_{Est} ~4.5. Crystallise the acid from EtOH. [*Beilstein* **9** H 680, **9** IV 2532.]

1,1-Diphenylurea [603-54-3] $C_{13}H_{12}N_2O$, *M* 212.3, *m* 238-239°. Crystallise 1,1-diphenylurea from MeOH. [*Beilstein* for 1,3 **12** IV 741.]

2,6-Di-*iso*-propylaniline [24544-04-5] $C_{12}H_{19}N$, *M* 177.3, *m* -45°, -54°, *b* 120-122°/10mm, 257°/atm, d_4^{25} 0.9367, n_D^{20} 1.5330, pK_{Est} ~0. Nitration of 1,3-di-*iso*-propylbenzene in AcOH/Ac₂O (2:1 v/v) with 96% HNO_3 (1.24-2.05 equivalents) below 45-50°/24 hours gave, on fractionation through a 2.5 x 180cm column at 5-10mm, a mixture of lower boiling 4-nitro- (74%) and the higher boiling 2-nitro-1,3-di-*iso*-propylbenzene (24%). Reduction of the later in *iso*-propanol with H_2 (~1200psi, <100°) catalysed by Raney Ni (5-10% by weight), followed by filtration, evaporation of the solvent and fractionation through a 180cm column at 10mm, with a reflux ratio of 20:1 and a rate of 0.33ml/minute gave **2,6-di-*iso*-propylaniline** as a colourless oil in 94.9% yield. The *benzoyl derivative* separated as an oil which solidifies and provides a white crystalline powder, *m* 106.0-106.7°, on recrystallisation from *iso*-octane. [Newton *J Am Chem Soc* **65** 2434 1943, DOI: 10.1021/ja01252a059.] Alternatively, aniline (300g), $AlCl_3$ (18g), and Al powder (6g) are heated in a pressure vessel at 290°, and propylene is pumped into the vessel to a pressure of 250 atmospheres. From time to time more propylene is added to sustain the pressure. After *ca* 8 hours the pressure ceased to decrease, the vessel is opened and the product is shaken with dilute aqueous NaOH and fractionated under a vacuum. The three fractions that separate are 2,6-di-*iso*-propylaniline (78%), 2-*iso*-propylaniline (15%), and a small amount of 2,4,6-tri-*iso*-propylaniline [Stroh et al. *Angew Chem* **69** 124 1957, DOI: 10.1002/ange.19570690403; also report *m* 184.5-187° for the *acetyl derivative*, and *m* 254-256° for the *benzoyl derivative*.] [*Beilstein* **12** III 2764.]

It is used for making NHCs for metal catalysed α -arylation of acyclic ketones, e.g. propiophenones, and amination of haloarenes [Matsubara et al. *J Org Chem* **72** 5069 2007, DOI: 10.1021/jo070313d]. It has been sulfonated for making sulfonated anilines, and consequently sulfonated NHCs, which provide water soluble Pd-NHCs used for Suzuki coupling of arylhalides with arylboronic acids in aqueous medium [Fleckenstein et al. *Chem Commun* 2870 2007, DOI: 10.1039/B703658B]. ***N*-Trimethylsilyl 2,6-di-isopropylaniline** [78923-65-6], prepared by reaction of Me_3SiCl and the Li salt of the aniline in Et_2O , is purified by distillation in vacuum. It is used in preparing the key intermediate for the Schrock-Hoveyda molybdenum catalyst (see [205815-80-1]), and sometimes it is prepared *in situ* without isolating it while using Et_3N instead of Li as base. [Schrock et al. *J Am Chem Soc* **112** 3875 1990, DOI: 10.1021/ja00166a023; Alexander et al. *Organometallics* **19** 3700 2000, DOI: 10.1021/om000336h].

***N,N*-Di-*n*-propylaniline** [2217-07-4] $\text{C}_{12}\text{H}_{19}\text{N}$, **M 177.3**, **b 127°/10mm**, **238-241°/760mm**, **pK²³ 5.68**. Reflux the aniline for 3 hours with acetic anhydride, then fractionally distil under reduced pressure.

Di-*p*-tolyl carbonate [621-02-3] $\text{C}_{15}\text{H}_{14}\text{O}_3$, **M 242.3**, **m 115°**. Purify the carbonate by GLC with 20% Apiezon on Embacel followed by sublimation *in vacuo* and recrystallisation from EtOH (**m 114°**). [Beilstein **6** H 398, **6** I 201, **6** II 380, **6** III 1366.]

***N,N'*-Di-*o*-tolylguanidine** [97-39-2] $\text{C}_{15}\text{H}_{17}\text{N}_3$, **M 239.3**, **m 179° (175-176°)**, **pK¹⁸ 9.62**. The guanidine crystallises from aqueous EtOH. The *sulfate* has **m 253-254°**(dec, H_2O). [Beilstein **12** H 803, **12** II 445, **12** III 1871, **16** IV 1764.]

Di-*p*-tolylphenylamine (4,4'-dimethyltriphenylamine) [20440-95-3] $\text{C}_{20}\text{H}_{19}\text{N}$, **M 273.4**, **m 108.5°, 112°, b 250°/20-40mm**, **417.5°/760mm**, **d 1.079**, **pK_{Est} ~ -5.0**. It is prepared from di-*p*-tolylamine, iodobenzene, K_2CO_3 and Cu bronze in nitrobenzene, boiled for 11 hours, nitrobenzene is steam distilled off, the residue is extracted with C_6H_6 , the extract is evaporated and the residue is distilled under reduced pressure. The distillate solidified and the solid is recrystallised once from EtOAc, then twice from EtOH to give pale yellow needles, **m 109°**(corr). [Marsden et al. *J Chem Soc* 627 1937, DOI: 10.1039/JR9370000627; Beilstein **12** III 2033.]

1,2-Divinylbenzene [1321-74-0] $\text{C}_{10}\text{H}_{10}$, **M 130.2**, **b 53°/3mm**, **79.2-79.6°/15mm**, **195°/760mm**, **d²⁰ 0.919**, **n_D²⁰ 1.573**. Purify divinylbenzene by dissolving in Et_2O , shaking with H_2O , drying over CaCl_2 , filtering, evaporating and distilling *in vacuo*. It polymerises within 2-3 days unless 4-*tert*-butylcatechol (0.05%) is added as stabiliser. [Fries & Bestian *Chem Ber* **69** 715 1936, DOI: 10.1002/cber.19360690416; Beilstein **5** III 1366.]

Duroquinone (tetramethylbenzoquinone) [527-17-3] $\text{C}_{10}\text{H}_{12}\text{O}_2$, **M 164.2**, **m 110-111°, 111-112°**. Crystallise duroquinone from 95% EtOH (yellow needles). Dry it *in vacuo*. It is poorly soluble in H_2O but is steam volatile, and is soluble in organic solvents. It can be sublimed unchanged. [Beilstein **7** H 669, **7** III 3417, **7** IV 2101.] **Hydroduroquinone** [527-18-4] $\text{C}_{10}\text{H}_{14}\text{O}_2$, **M 166.2**, **m 233°**(sintering at $>200^\circ$), crystallises in almost colourless needles from EtOH, and is oxidised to duroquinone with FeCl_3 . Its **diacetyl derivative**, **m 207°**, also crystallises in colourless needles from EtOH. [Smith *Org Synth Coll Vol* **2** 254 1943, DOI: 10.15227/orgsyn.010.0040.]

Ellagic acid (4,4',5,5',6,6'-hexahydroxy-2,2'-diphenic acid dilactone) [476-66-4] $\text{C}_{14}\text{H}_6\text{O}_8$, **M 338.2**, **m $>360^\circ$** , **pK_{Est(1)} ~8**, **pK_{Est(2)} ~11**. This antioxidant, isolated from the kino (exudate) of Australian *Eucalyptus* 'gum tree' species, crystallises from pyridine. It forms a dark green solution in aqueous *N* NaOH, gives a blue FeCl_3 test, a cherry red colour with concentrated HNO_3 , and under UV light it exhibits mauve fluoresces which turns yellow when fumed with NH_3 . It is best purified by conversion into the **tetra-acetate dilactone**, **$\text{C}_{22}\text{H}_{14}\text{O}_{12}$** , by refluxing with $\text{Ac}_2\text{O}/\text{AcONa}$ /4 hours and recrystallising from Ac_2O which provides colourless needles with **m ~340°** depending on heating rate (343-346° also reported). The **tetra-acetyl derivative** (e.g. 0.3g) is then hydrolysed by refluxing with AcOH (5ml) and with H_2SO_4 (0.5ml) for 2 hours, diluted with H_2O , the solid ellagic acid is collected and recrystallised from pyridine to give cream coloured needles **m $> 360^\circ$** . [Gell et al. *Aust J Chem* **11** 372 1958, DOI: 10.1071/CH9580372.] The UV (EtOH) has λ_{max} at 255nm (log ϵ 4.60) and 366nm (log ϵ 3.93). The **tetra-carbethoxy derivative** has **m 245-246°** [Hillis & Carle *Aust J Chem*

16 147 1963, DOI: 10.1071/CH9630147]. [*Beilstein* **19** H 261, **19** III/IV 3164, **19/7** V 108.] Possible anticarcinogen.

Emodine (1,3,8-trihydroxy-6-methyl-9,10-anthracenedione, archin) [518-82-1] $C_{15}H_{10}O_5$, **M 270.2**, **m 253-257°**, **255-256°**, **256-257°**, **262°**, **264°** (phenolic pKs 7–10). Archin forms orange needles from EtOH, Et₂O, *C₆H₆, toluene or pyridine. It sublimes above 200° at 12mm. [Tutin & Clewer *J Chem Soc* **99** 946 1911, DOI:10.1039/CT9119900946; IR: Bloom et al. *J Chem Soc* 178 1959, DOI: 10.1039/JR9590000178; UV: Birkinshaw *Biochem J* **59** 485 1955, DOI: 10.1042/bj0590485; Anslow et al. DOI: 10.1042/bj0340159; *Biochem J* **34** 159 1940, DOI: 10.1042/bj0340159; *Beilstein* **8** IV 3575.] It is a purgative.

3-Ethoxy-N,N-diethylaniline [1846-92-2] $C_{12}H_{19}NO$, **M 193.3**, **b 97-98°/0.6mm**, **145°/14mm**, **268-270°/760mm**, **n_D²⁰ 1.5342**, **n_D²⁵ 1.5325**, **pK_{Est} ~6.1**. Reflux it for 3 hours with Ac₂O, then fractionally distil it, preferably under reduced pressure. The *picrate* has **m 132-133°** (from EtOH). The *N,N*-diethyl-N-methyl-m-phenetidinium iodide has **m 137-138°** (from MeOH/Et₂O) and its *picrate* has **m 101-102°** (from Me₂CO/H₂O) [Fahim & Fleifel *J Chem Soc* 2761 1951, DOI: 10.1039/JR9510002761]. [For 2-ethoxy-4-diethylaminobenzene 1-diazonium BF₄ see: Klaassens & Schoot *Recl Trav Chim Pays-Bas* **71** 1086 1952, DOI: 10.1002/recl.19520711106; Gilman & Kyle *J Am Chem Soc* **74** 3027 1952, DOI: 10.1021/ja01132a023; Brown & Mason *J Chem Soc* 1269 1933, DOI: 10.1039/JR9330001269; *Beilstein* **13** I 131, **13** III 942, **13** IV 970.]

1-Ethoxynaphthalene [5328-01-8] $C_{12}H_{12}O$, **M 172.2**, **b 136-138°/14mm**, **282°/760mm**, **d₄²⁰ 1.061**, **n_D²⁰ 1.604**. Fractionally distil it (twice) under a vacuum, then dry it with, and distil it under a vacuum from sodium. The *picrate* has **m 118.5-119°** (from EtOH). [*Beilstein* **6** H 606, **6** II 578, **6** III 2924, **6** IV 4212.]

2-Ethoxynaphthalene [93-18-5] has **m 35.6-36.0°**, **37-38°**, **b 142-143°/12mm**, **280°/760mm**. Crystallise it from petroleum ether or EtOH (**m 37-38°**). Dry it *in vacuo*, or distil it in a vacuum. The *picrate* has **m 104.5°** (from EtOH or CHCl₃). [*Beilstein* **6** H 606, **6** II 578, **6** III 2972, **6** IV 4257.] It is used in perfumery.

Ethyl p-aminobenzoate (Benzocaine) [94-09-7] $C_9H_{11}NO_2$, **M 165.2**, **m 88° to 90°**, **92°**, **pK²⁵ 2.39**. Crystallise benzocaine from EtOH/H₂O or EtOH (**m 93-94°**), and dry it in air. [Adams & Cohen *Org Synth Coll Vol I* 240 1941, DOI: 10.15227/orgsyn.008.0066; *Beilstein* **14** H 422, **14** IV 1129.] Anaesthetic.

p-Ethylaniline [589-16-2] $C_8H_{11}N$, **M 121.2**, **m -5°, -4.8°**, **b 88°/8mm**, **216°/atm**, **d₄²⁰ 0.975**, **n_D²⁰ 1.554**, **pK²⁵ 5.00**. Dissolve p-ethylaniline in *benzene, then acetylate it. Recrystallise the *acetyl* derivative (**m 93-94°**, Hickinbottom & Waite *J Chem Soc* 1565 1930, DOI: 10.1039/JR9300001558) from *C₆H₆/petroleum ether or EOH, and hydrolyse by refluxing 50g with 500ml of H₂O and 115ml of concentrated H₂SO₄ until the solution becomes clear. The amine sulfate is isolated, suspended in H₂O and solid KOH is added to generate the free base, which separates. Dry it (KOH), and distil it from zinc dust in a vacuum [Berliner & Berliner *J Am Chem Soc* **76** 6179 1954, DOI: 10.1021/ja01652a092]. The *picrate* has **m 171-172°** (from 1,2-dichloroethane). [*Beilstein* **12** H 1090, **12** III 2380, **12** IV 2419.]

trans-Ethyl cinnamate [103-36-6] $C_{11}H_{12}O_2$, **M 176.2**, **m 6.7°**, **b 127°/6mm**, **272.7°/768mm**, **d₄²⁰ 1.040**, **n_D²⁰ 1.55983**. Wash the ester with aqueous 10% Na₂CO₃, then water, dry (MgSO₄), and distil it. The purified ester is saponified with aqueous KOH, and, after acidifying the solution, cinnamic acid is isolated, washed and dried. The ester is reformed by refluxing for 15 hours the cinnamic acid (25g) with absolute EtOH (23g), conc H₂SO₄ (4g) and dry *benzene (100ml), after which it is isolated, washed, dried and distilled under reduced pressure [Jeffery & Vogel *J Chem Soc* 658 1948, DOI: 10.1039/JR9480000658]. [*Beilstein* **9** IV 2006.]

(±)-Ethyl α,β-dibromo-β-phenylpropionate [5464-70-0, erythro 30983-70-1] $C_{11}H_{12}Br_2O_2$, **M 336.0**, **m 74-75°**, **75°**. Crystallise the propionate from petroleum ether (b 60-80° or 70-90°), EtOH or aqueous EtOH (**m 78°**, and an *unstable* form **m 65°**). [*Beilstein* **9** H 512, **9** I 202, **9** III 2406, **9** IV 1772.]

Ethyl gallate (ethyl 3,4,5-trihydroxybenzoate) [831-61-8] $C_9H_{10}O_5$, **M 198.2**, **m 149°**, **150-151°**, **153°**, **163-165°**. Recrystallise the gallate from 1,2-dichloroethane. Its UV has λ_{max} (neutral species) at 275nm (ϵ 10,000), (anion) at 235nm (ϵ 10,300), 279nm (ϵ 11,400) and 324nm (ϵ 8 500) [Campbell & Coppinger *J Am Chem Soc* **73** 2708 1951, DOI: 10.1021/ja01150a079]. [*Beilstein* **10** IV 2002.] It is an antioxidant.

Ethyl *p*-nitrobenzoate [99-77-4] $C_9H_9NO_4$, **M 195.2**, **m 55°, 56°, 57°, 59°, b 314°/atm**. Dissolve it in Et_2O and wash it with aqueous alkali, then the ether is evaporated and the solid recrystallised from EtOH. Take all necessary precautions against explosion if it is to be distilled. [Beilstein 13 H 3787, 13 IV 1074.]

2-Ethylphenol [90-00-6] $C_8H_{10}O$, **M 122.2**, **m -28° and -3.4° (dimorphic), b 204.5°/760mm, 210-212°/atm, d_4^{20} 1.020, n_D^{20} 1.537, pK^{25} 10.20**. Purify as for *p*-ethylphenol below. [Beilstein 6 H 470, 6 IV 3011.]

4-Ethylphenol [123-07-9] has **m 41° to 44°, 47-48°, b 76°/1mm, 153°/100mm, 218.0°/762mm, n_D^{25} 1.5239, pK^{25} 10.21**. Non-acidic impurities are removed by passing steam through a boiling solution containing 1 mole of the phenol and 1.75 moles of NaOH (as an aqueous 10% solution). The residue is cooled and acidified with 30% (v/v) H_2SO_4 , and the free phenol is extracted into diethyl ether. The extract is washed with water, dried with $CaSO_4$ and the ether is evaporated. The phenol is distilled at 100mm pressure through a Stedman gauze-packed column. It is further purified by fractional crystallisation by partial freezing, and by zone refining, under N_2 [Biddiscombe et al. *J Chem Soc* 5764 1963, DOI: 10.1039/JR9630005764]. Alternatively, purify it via the benzoate, as for phenol. The **4-nitrophenylbenzoate** has **m 80°** (from EtOH). [Beilstein 6 H 470, 6 III 1663, 5 IV 3020.]

Ethyl phenylacetate [101-97-3] $C_{10}H_{12}O_2$, **M 164.2**, **b 99-99.3°/14mm, 133°/30mm, 229°/atm, d_4^{20} 1.030, n_D^{20} 1.499**. Shake the ester with saturated aqueous Na_2CO_3 (three times), aqueous 50% $CaCl_2$ (twice) and saturated aqueous NaCl (twice). Dry with $CaCl_2$ and distil it under reduced pressure. [Adams & Thal *Org Synth* 2 27 1922, DOI: 10.15227/orgsyn.002.0027; Beilstein 9 H 434, 9 IV 1618.] It has a pleasant odour and is used in perfumery.

Ethyl Red [2-(4-diethylaminophenylazo)benzoic acid] [76058-33-8] $C_{17}H_{19}N_3O_2$, **M 297.4**, **m 135°, 150-152°, pK_1 2.5, pK_2 9.5**. Crystallise the acid dye from EtOH/diethyl ether or toluene. It is an indicator: pH 4.4 (red) and 6.2 (yellow). Its UV has λ_{max} at 447nm. [Beilstein 16 I 316.]

Eugenol (4-allyl-2-methoxyphenol, 4-allylguaiacol) [97-53-0] $C_{10}H_{12}O_2$, **M 164.2**, **m -12° to -10°, b 253°/760mm, 255°/760mm, d_4^{20} 1.066, n_D^{20} 1.540, pK^{25} 10.19**. Fractional distillation of eugenol gives a pale yellow liquid which darkens and thickens on exposure to air. It should be stored under N_2 at -20°. [Waterman & Priedster *Recl Trav Chim Pays-Bas* 48 1272 1929, DOI: 10.1002/recl.19290481210; Beilstein 6 H 961, 6 IV 6337.] It has a spicy odour like cloves, is used in perfumery and has some analgesic properties. It is also an insect attractant. Its **benzoate** has **m 60-70°**.

Eugenol methyl ether (4-allyl-1,2-dimethoxybenzene) [93-15-2] $C_{11}H_{14}O_2$, **M 178.2**, has **m -4°, b 127-129°/11mm, 146°/30mm, 154.7°/760mm, d_4^{20} 1.0354, n_D^{20} 1.53411**. Recrystallise the ether from hexane at low temperature and redistil it (preferably *in vacuo*). [Hillmer & Schorning *Z Phys Chem [A]* 167 407 1934, Briner & Fliszár *Helv Chim Acta* 42 2063 1959, DOI: 10.1002/hlca.19590420637; Beilstein 6 IV 6337.]

Fluoranthene (benzo[*j,k*]fluorene) [206-44-0] $C_{16}H_{10}$, **M 202.3**, **m 105° to 110°, 110-111°, b 384°/760mm**. Purify it by chromatography of CCl_4 solutions on alumina, with *benzene as eluent. Crystallise it from EtOH, MeOH or *benzene. Also purify it by zone melting. [Gorman et al. *J Am Chem Soc* 107 4404 1985, DOI: 10.1021/ja00301a006; Beilstein 5 I 344, 5 IV 2463.]

Fluorene [86-73-7] $C_{13}H_{10}$, **M 166.2**, **m 111° to 114°, 114.7-115.1°, b 160°/15mm, 298°/atm**. Purify fluorene by chromatography of CCl_4 or petroleum ether (b 40-60°) solution on alumina, with *benzene as eluent. Crystallise it from 95% EtOH, 90% acetic acid and again from EtOH. Crystallisation using glacial acetic acid retains an impurity which is removed by partial mercuration and precipitation with LiBr [Brown et al. *J Am Chem Soc* 84 1229 1962, DOI: 10.1021/ja00866a032]. It has also been crystallised from hexane, or *benzene/EtOH, distilled under vacuum and purified by zone refining. [Gorman et al. *J Am Chem Soc* 107 4404 1985, DOI: 10.1021/ja00301a006; Beilstein 5 IV 2142.]

Fluorene-2,7-diamine [525-64-4] $C_{13}H_{12}N_2$, **M 196.3**, **m 160° to 162°, 165-166°**. Crystallise the diamine from hot H_2O or aqueous EtOH, dry it *in vacuo* and store it in the dark. [Beilstein 13 H 266, 13 II 123, 12 III 507, 13 IV 449.] It is a useful analytical reagent for metal dications, such as Cd, Cu, Co, Zn, as well as the

bromide, chloride, nitrite and persulfate anions.

9-Fluorenone [486-25-9] $C_{13}H_8O_2$, **M 180.2**, **m 82.5-83.0°, 85-86°, b 341°/760mm**. Crystallise 9-fluorenone from absolute EtOH, MeOH or *benzene/pentane. [Ikezawa *J Am Chem Soc* **108** 1589 1986, DOI: 10.1021/ja00267a032.] Also recrystallise it twice from toluene and sublime it in a vacuum [Saltiel *J Am Chem Soc* **108** 2674 1986, DOI: 10.1021/ja00270a028]. It can be distilled under high vacuum. The **oxime** [2157-52-0] $C_{13}H_9NO$, **M 195.2**, has **m 195.2° (194-195°, 198°)** (yellow crystals from *C₆H₆ or xylene, Anet et al. *Can J Chem* **35** 180 1957, DOI: 10.1139/v57-027; Wislicenus & Waldmuller *Chem Ber* **41** 3334 1908, DOI: 10.1002/cber.19080410309.) [Beilstein **7** H 465, **7** III 2330, **7** IV 1629.] The **hydrazone** [13629-22-6] $C_{13}H_{10}O_2$, **M 194.2**, crystallises from *C₆H₆/petroleum ether (**m 148-150°**) or MeOH (**m 152°**). It is used for the photometric determination of ketosteroids. [Beilstein **7** I 251.]

9-Fluorenylmethyl chloroformate (Fmoc-Cl) [28920-43-6] $C_{15}H_{11}ClO_2$, **M 258.7**, **m 61-63°, 61.4-63°, 62° to 64°**. If the IR contains no OH bands (at ~3000 cm⁻¹) due to the hydrolysis product 9-fluorenylmethanol, then purify it by recrystallisation from dry Et₂O. Its IR (CHCl₃) has a band at 1770 cm⁻¹ (C=O), and the ¹H NMR (CDCl₃) has δ at 4-4.6 (m 2H, CHCH₂) and 7.1-7.8 (m, 8 aromatic H). The **azide** (Fmoc-N₃) has **m 89-90°** (from hexane) and IR (CHCl₃) at 2135 (N₃) and 1730 (C=O) cm⁻¹, and the **carbazate** (Fmoc-NHNH₂) has **m 171°(dec)** (from nitromethane) with IR (KBr) at 3310, 3202 (NH) and 1686 (CONH) cm⁻¹. [Caprino & Han *J Org Chem* **37**, 3404 1972, DOI: 10.1021/jo00795a005; *J Am Chem Soc* **92** 5748 1970, DOI: 10.1021/ja00722a043; Koole et al. *J Org Chem* **54** 1657 1989, DOI: 10.1021/jo00268a030; Fürst et al. *J Chromatogr* **499** 557 1990, DOI: 10.1016/S0021-9673(00)97000-6.] The reagent is used extensively for derivatising amino acids and in solid phase peptide synthesis. It is readily used for protection and de-protection of the N-terminal amino acid of peptides and proteins [Johnon et al. *JCS Chem Commun* 369 1993, DOI: 10.1039/C39930000369], as well as in oligonucleotide synthesis [Seela & Wenzel *Helv Chim Acta* **75** 1111 1992, DOI: 10.1002/hlca.19920750414].

9-Fluorenylmethyl succinimidyl carbonate [82911-69-1] $C_{19}H_{15}NO_5$, **M 337.3**, **m 147-151°(dec), 151°(dec), 153°(dec)**. Recrystallise the carbonate from CHCl₃/Et₂O, or from petroleum ether (b 40-60°). [Paquet *Can J Chem* **60** 976 1982, DOI: 10.1139/v82-146; Lapatsanis et al. *Synthesis* 671 1983, DOI: 10.1055/s-1983-30468.] Efficient reagent for the selective N- protection.

Fluorobenzene [462-06-6] C_6H_5F , **M 96.1**, **m -42°, b 84.8°/760mm**, **d₄²⁰ 1.025**, **n_D²⁰ 1.46573**, **n_D³⁰ 1.4610**. Dry fluorobenzene for several days with P₂O₅, then fractionally distil it. [Beilstein **5** H 198, **5** IV 632.]

2-Fluorobenzoic acid [445-29-4] $C_7H_5FO_2$, **M 140.1**, **m 122° to 125°, 126°, 127°, d₂₅ 1.460**, **pK²⁵ 3.27**. Crystallise the acid from 50% aqueous EtOH, dilute HCl or *C₆H₆, then purify it by zone melting or vacuum sublimation at 130-140°. [Beilstein **9** H 333, **9** III 1324, **9** IV 950.]

3-Fluorobenzoic acid [445-38-9] has **m 122-123°, 124°, 125°, d₂₅ 1.474** **pK²⁵ 3.86**. Crystallise the acid from 50% aqueous EtOH or *C₆H₆, then sublime it *in vacuo* at 130-140°. [Beilstein **9** H 333, **9** IV 952.]

4-Fluorobenzoic acid [456-22-4] $C_7H_5FO_2$, **M 140.1**, **m 182°, 184°, b 253.7°/atm**, **d₂₅ 1.479**, **pK²⁵ 4.15**. Crystallise the acid from 50% aqueous EtOH, then purify it by zone melting or vacuum sublimation at 130-140°. [Schiemann & Winkelmüller *Org Synth Coll Vol* **2** 299 1943, DOI: 10.15227/orgsyn.013.0052; Beilstein **9** H 333, **9** III 1327, **9** IV 953.]

3-Fluoro-4-hydroxyphenylacetic acid [458-09-3] $C_8H_7FO_3$, **M 170.1**, **m 132° to 134°, pK_{Est(1)} ~4.4**, **pK_{Est(2)} ~9.4**. Crystallise the acid from water. [Beilstein **10** III 440.]

4-Fluoro-2-methylbenzaldehyde [63082-45-1] C_8H_7FO , **M 138.1**, **d₄²⁵ 1.144**, **n_D²⁵ 1.526**. The aldehyde has been purified by gas chromatography and should be kept under N₂ as it readily oxidises in air (flash point is 28°). [Burgess et al. *Aust J Chem* **30** 543 1977, DOI: 10.1071/CH9770543].

2-Fluoro-4-nitroaniline [369-35-7] $C_6H_5FN_2O_2$, **M 156.1**, **m 122° to 130°, 134-135°, 135-136°, pK²⁵ -0.44** (**aqueous HClO₄**). The aniline forms yellow crystals on recrystallisation from aqueous MeOH or EtOH. The

acetyl derivative has **m 203-204°** (from EtOH). [Wepster & Verkade *Recl Trav Chim Pays-Bas* **68** 77 1949, DOI: 10.1002/recl.19490680109; *Beilstein* **12** III 1647.]

1-Fluoro-4-nitrobenzene [350-46-9] $C_6H_4FNO_2$, **M 141.1**, **m 21.5°(unstable form), 27°(stable form), 22-24°, b 86.6°/14mm, 95-97.5°/22mm, 205.3°/735mm**. Crystallise it from EtOH or distil it in a vacuum. [*Beilstein* **5** H 241, **5** IV 719.]

1-Fluoro-4-nitronaphthalene [341-92-4] $C_{10}H_6FNO_2$, **M 191.2**, **m 80°, b 314.6°/760mm, d 1.369**. It crystallises from EtOH as yellow needles [Bunce et al. *J Org Chem* **52** 4214 1987, DOI: 10.1021/jo00228a012]. [*Beilstein* **5** III 1596, **5** IV 1675.]

4-Fluoro-3-nitrophenylazide [28166-06-5] $C_6H_3FN_4O_2$, **M 182.1**, **m 52-53°, 53-55°, 54-56°**. Dissolve the azide in Et₂O, dry it over MgSO₄, filter, evaporate and recrystallise the residue from petroleum ether (b 20-40°) to give orange needles. Store it in a stoppered container at ~0°. The ¹H NMR has δ at 7.75 (m 1H) and 7.35 (m 2H) in CDCl₃. [Hagedorn et al. *J Org Chem* **43** 2070 1978, DOI: 10.1021/jo00404a056.]

2-Fluorophenol [367-12-4] C_6H_5FO , **M 112.1**, **m 14° to 16°, 16.1°, b 53°/14mm, 171-172°/714mm, 172.06-173.06°/760mm, d₄²⁰ 1.257, n_D²⁰ 1.514, pK²⁵ 8.70**. Pass *o*-fluorophenol at least twice through a gas chromatographic column for small quantities; otherwise fractionally distil it under reduced pressure. [*Beilstein* **6** I 97, **6** IV 770.]

4-Fluorophenoxyacetic acid [405-79-8] $C_8H_7FO_3$, **M 170.1**, **m 104.2-104.6°, 106°, pK²⁵ 3.13**. Crystallise the acid from EtOH or H₂O. [Hayes & Branch *J Am Chem Soc* **65** 1555 1943, DOI: 10.1021/ja01248a031; *Beilstein* **6** III 971, **6** IV 776.]

4-Fluorophenylacetic acid [405-50-5] $C_8H_7FO_2$, **M 154.1**, **m 81° to 83°, 86°, 94°, b 164°/2mm, pK²⁵ 4.22**. Crystallise it from heptane or CHCl₃, but it is best purified by distillation at high vacuum. [Bergmann et al. *J Am Chem Soc* **78** 6037 1956, DOI: 10.1021/ja01604a023; Olah et al. *J. Org Chem* **22** 879 1957, DOI: 10.1021/jo01359a005; *Beilstein* **9** III 2261, **9** IV 1672.]

4-Fluorophenyl isocyanate [1195-45-5] C_7H_4FNO , **M 137.1**, **b 55°/8mm, 166.9°/760mm, d 1.206, n_D²⁰ 1.514**. Purify the isocyanate by repeated fractionation through an efficient column. If its IR indicates that there is too much urea (in the presence of moisture the symmetrical urea is formed), then dissolve it in dry EtOH-free CHCl₃, filter, evaporate and distil it. It hydrolyses in water. **It is a pungent LACHRYMATORY liquid, TOXIC**. [See Hardy *J Chem Soc* 2011 1934, DOI: 10.1039/JR9340002010, and Hickinbottom *Reactions of Organic Compounds* Longmans p 493 1957.]

4-Fluorophenyl 2-nitrobenzyl ether [448-37-3] $C_{13}H_{10}FNO_3$, **M 247.2**, **m 62°**. Crystallise the ether from EtOH. [Jones *J Chem Soc* 1414 1938, DOI: 10.1039/JR9380001414; *Beilstein* **6** III 1564, **6** IV 2608.]

2-Fluorotoluene [95-52-3] C_7H_7F , **M 110.1**, **m -80°, -62°, b 114.4°/760mm, d₄²⁰ 1.005, n_D²⁰ 1.475**. Dry *o*-fluorotoluene with P₂O₅ or CaSO₄ and fractionally distil it through a silvered vacuum-jacketed glass column with 1/8th-in glass helices. A high reflux ratio is necessary because of the closeness of the boiling points of the *o*-, *m*- and *p*- isomers [Potter & Saylor *J Am Chem Soc* **73** 90 1951, DOI: 10.1021/ja01145a032]. [*Beilstein* **5** H 290, **5** III 676, **5** IV 799.] **TOXIC do not breath the vapours.** **3-Fluorotoluene** [352-70-5] has **m -111°, -87°, b 115°/atm, 116.5°/760mm, d₄²⁰ 1.00, n_D²⁷ 1.46524**. Purify it as for *o*-fluorotoluene. [*Beilstein* **5** H 290, **5** III 676, **5** IV 799.] **TOXIC do not breath the vapours.** **4-Fluorotoluene** [352-32-9] has **m -56°, 116.0°/760mm, d₄²⁰ 1.00, n_D²⁰ 1.46884**. Purify it as for *o*-fluorotoluene. [*Beilstein* **5** H 290, **5** III 677, **5** IV 799.] **TOXIC do not breath the vapours.**

Formanilide (N-phenylformamide) [103-70-8] C_7H_7NO , **M 121.1**, **m 46.6-47.5°, 48°, 50°, b 166°/14mm, 216°/120mm, 271°/atm, d₄²⁰ 1.14**. Crystallise formanilide from Et₂O (**m 45.3°**), Et₂O/petroleum ether (**m 46°**), petroleum ether (**m 47.6°**), ligroin/xylene, or distil it preferably under reduced pressure. Its solubility in H₂O is 2.9w/v% at 25°. [*Beilstein* **12** H 230, **12** II 135, **12** III 453, **12** IV 368.]

Gallic acid (H₂O) (3,4,5-trihydroxybenzoic acid) [5995-86-8 (H₂O), 149-91-7 (anhydrous)] C₇H₆O₅, **M 188.1, m 251°(dec), 253°(dec), pK₁²⁵ 4.27, pK₂²⁵ 8.68.** Crystallise gallic from water, EtOH or CHCl₃. Its solubility in w/v is 1% in Et₂O, 1.2% in H₂O, 10% in glycerol, 20% in Me₂CO and 33% in EtOH. [Beilstein 10 H 470, 10 IV 1993.] The *tri-O-acetyl* derivative has **m 172°** (from MeOH), and the *anilide* has **m 207°** (from EtOH). The *methyl ester* [99-24-1] C₈H₈O₅, **M 184.1**, has **m 202°** (dry), and crystallises in prisms from MeOH. [Beilstein 10 IV 1998.] These have antioxidant, angiogenic and anticarcinogenic properties.

Galvinoxyl [2,6-di-*tert*-butyl-α-(3,5-di-*tert*-butyl-4-oxo-2,5-cyclohexadiene-1-ylidene)-*p*-tolyl-oxy] [2370-18-5] C₂₉H₄₁O₂, **M 421.65, m 153.2-153.6°, 158-159°.** It is a stable free radical inhibitor and a scavenger of short-lived free radicals with odd electrons on C or O. It is best prepared freshly by oxidation of 3,3',5,5'-tetra-*tert*-butyl-4,4'-dihydroxydiphenyl-methane [**m 154°**, 157.1-157.6°, obtained by gently heating for 10-15 minutes 2,6-di-*tert*-butylphenol, formaldehyde and NaOH in EtOH and recrystallised from EtOH (20g/100ml) as colourless plates [Karasch & Joshi *J Org Chem* 22 1435 1957, DOI: 10.1021/jo01362a033; Bartlett et al. *J Am Chem Soc* 82 1756 1960, DOI: 10.1021/ja01492a054; and 84 2596 1962, DOI: 10.1021/ja00872a026.] Oxidation is carried out under N₂ with PbO₂ in Et₂O or isooctane [Coppinger *J Am Chem Soc* 79 501 1957, DOI: 10.1021/ja01559a073; Bartlett et al. above] or with alkaline potassium ferricyanide [Karasch & Joshi, above], whereby Galvinoxyl separates as deep blue crystals and is recrystallised twice under N₂ from *C₆H₆ solution by suction evaporation at 30°. The VIS spectrum has λ_{max} at 407nm (ε 30,000), 431nm (ε 154,000) and weak absorption at 772.5nm, and its IR has ν_{max} at 1577 and 2967cm⁻¹, and is estimated by iodometric titration. It is sensitive to O₂ in the presence of OH⁻ ions and traces of strong acid in hydroxylic or hydrocarbon solvents. At 62.5° in a 0.62mM solution in *C₆H₆ the radical decays with a first order $k = 4 \times 10^{-8} \text{ sec}^{-1}$ (half life $1.7 \times 10^{17} \text{ sec}$, ~200 days) as observed by the change in OD at 550nm [see Greene & Adam *J Org Chem* 28 3550 1963, DOI: 10.1021/jo01047a505].

R(-)-Guaiaretic acid [4,4'-(2,3-dimethyl-1-butene-(1,4-diyl)-bis-(2-methoxyphenol)] [500-40-3] C₂₂H₂₄O₄, **M 352.4, m 99-100.5°, [α]_D¹⁶ -91 (c 1.1, EtOH), pK_{Est} ~10.0.** Crystallise the acid once from petroleum ether (b 60-80°), twice from 60% aqueous MeOH and finally from EtOH (**m 100-101°**). Its UV has λ_{max} (EtOH) at 207 and 260nm (log ε 4.64 and 4.28). The *dimethyl ether*, C₂₄H₂₈O₄, crystallises from MeOH in tiny needles with **m 94-94.5°, [α]_D¹⁶ -94 (c 0.6, EtOH)**, and UV has λ_{max} (EtOH) at 211 and 259nm (log ε 4.45 and 4.20). [King et al. *J Chem Soc* 4011 1964, DOI: 10.1039/JR9640004011; Schrecker & Hartwell *J Org Chem* 21 381 1956, DOI: 10.1021/jo01109a617.]

Guaiacol (2-methoxyphenol) [90-05-1] C₇H₈O₂, **M 124.1, m 26° to 29°, 32°, b 53-55°/4mm, 106°/24mm, 205°/746mm, d₂₅ 1.129g/ml, pK₂₅ 9.90.** Crystallise guaiacol from *benzene/petroleum ether or distil it in a vacuum. It has some solubility in H₂O (1.5w/v%) but is much more soluble in glycerol (1g/ml). [Beilstein 6 H 768, 6 IV 5563.] *Guaiacol carbonate* [553-17-3] C₁₅H₁₄O₅, **M 274.3**, has **m 88.1°, 89°, b 392.6°/760mm, d₂₅ 1.208g/ml**, Crystallise the carbonate from EtOH. It is estimated by bromination to the *monobromo guaiacol carbonate* **m 178°** (from EtOH) [Chernoff *J Am Chem Soc* 51 3072 1929, DOI: 10.1021/ja01385a027]. Useful in bronchitis and pneumonia [Seifert *The Lancet* 154 710 1899, DOI:10.1016/S0140-6736(01)58972-2]. [Beilstein 6 H 776, 6 I 386, 6 II 784, 6 III 4233.]

Guaiazulene (1,4-dimethyl-7-isopropylazulene, see also Chamazulene above) [489-84-9] C₁₅H₁₈, **M 198.3, m 29.5°, 30.5°, 31.5°, b 153°/7mm, d₂₅ 0.976.** This hydrocarbon is present in and was isolated from chamomile oil and guaiac wood oil. [Sorm et al. *Coll Czech Chem Commun* 16 168, DOI: org/10.1135/cccc19510168; Sorm et al. *Coll Czech Chem Commun* 16 626 1951, DOI: org/10.1135/cccc19510626]. It forms blue-violet plates from EtOH. Also redistil it *in vacuo* until distillate solidifies. Its UV has λ_{max} at 284nm (log ε 4.6, heptane) and also maxima at 285, 351, 369, 605, 660, and 735 nm. Purification can also be achieved by chromatography over Al₂O₃ (Basic I) and eluting the blue band with petroleum ether. The *trinitrobenzene complex* [4968-29-0] C₁₅H₁₈.C₆H₃N₃O₆, **M 411.4**, crystallises as black needles from EtOH with **m 151°**. The *picrate* has **m 122°** (from EtOH). A water soluble form is obtained by sulfonation to give the *sodium 3-sulfonate* (sodium gualenate, **Azulon**) [6223-35-4] C₁₅H₁₇O₃S.Na, **M 300.3**. It has anti-inflammatory and antiulcerative activity. [cf. IR, UV: Plattner, Furst & Marti *Helv Chim Acta* 32 2452 1949, DOI: 10.1002/hlca.19490320732; Mukherjee, Dunn and Houk *J Am Chem Soc* 101 251 1979, DOI: 10.1021/ja00495a058; Beilstein 5 III 1677, 5 IV 1751.]

Guanabenz (1-[2,6-dichlorobenzylideneamino]guanidine, Wyntensin, WY-8677) [5051-62-7] $\text{C}_8\text{H}_8\text{Cl}_2\text{N}_4$, **M 231.1**, **m 227-229°**(dec), **pK_{Est} ~9**. Crystallise the guanidine from MeCN and store it in a CO_2 -free atmosphere. The *monoHCl* [23256-5-0] has **m 178°**(dec) [solubility w% at 25° is 1.1(H_2O), 5(EtOH), 0.1(EtOAc), 0.06(CHCl_3)]. It is an antihypertensive drug. [Holmes et al. *Drugs* **26** 212 1983.]

Hexachlorobenzene [118-74-1] C_6Cl_6 , **M 284.8**, **m 227° to 229°, 230.2-231.0°, b 323-326°/atm.** Crystallise hexachlorobenzene repeatedly from *benzene. Dry it under vacuum over P_2O_5 . It can be sublimed. [Beilstein **5** H 205, **5** IV 670.] A former herbicide.

Hexachlorophene [2,2'-methylenebis(3,4,6-trichlorophenol)] [70-30-4] $\text{C}_{13}\text{H}_6\text{Cl}_6\text{O}_2$, **M 406.9**, **m 163-165°, 166-167°, b 471°/atm, d 1.71, pK₁²⁰ 5.41, pK₂²⁰ (30% aqueous MeOH).** It forms needles from MeOH, $\text{C}_2\text{H}_4\text{Cl}_2$, or toluene. The *diacetate* has **m 175-176°** (EtOH). This disinfectant is also available in MeOH (5mg/l). [Beilstein **6** III 5407, **6** IV 6659.]

Hexadecyl 3-hydroxynaphthalene-2-carboxylate [531-84-0] $\text{C}_{27}\text{H}_{40}\text{O}_3$, **M 412.6**, **m 72-73°, 73-74°, d¹³⁵ 0.824.** Recrystallise the ester from hot EtOH (greenish-white flakes) and sublime it in a vacuum. It is soluble in * C_6H_6 , petroleum ether and AcOH. [Oshima & Hayashi *J Soc Chem Ind Jpn* **44** 821 1941, *Chem Abs* **42** 2108 1948, Beilstein **5** H 471, **5** I 227, **5** II 358, **5** III 1106, **5** IV 1208.] Used for waterproofing synthetic cloth.

Hexaethylbenzene [604-88-6] $\text{C}_{18}\text{H}_{30}$, **M 246.3**, **m 128.7-129.5°.** Crystallise this hydrocarbon from * C_6H_6 , or * $\text{C}_6\text{H}_6/\text{EtOH}$. The *1,3,5-trinitrobenzene complex* (1:1) has **m 174°**(EtOH). [Beilstein **5** III 3038, **5** IV 1137.]

Hexafluorobenzene [392-56-3] C_6F_6 , **M 186.1**, **m -13° to -11°, 3.7° to 4.1°, 5.1°, b 79-80°/760mm, d₄²⁰ 1.61, n_D²⁰ 1.378.** Main impurities are incompletely fluorinated benzenes. Purify it by standing in contact with oleum for 4 hours at room temperature, repeating until the oleum does not become coloured. Wash with water several times, then dry with P_2O_5 . Finally purify it by repeated fractional crystallisation. [Beilstein **5** III 523, **5** IV 640.]

Hexamethylbenzene [87-85-4] $\text{C}_{12}\text{H}_{18}$, **M 162.3**, **m 164°, 165-165.5°, 165.75°, 166°, b 264°/atm.** It is prepared, in an efficient fume cupboard, by passing a solution of phenol (100g, 1.06moles) in MeOH (1L) dropwise at a rate of 110ml/hour through a column (5.08cm diameter x 40.6cm long) packed with Al_2O_3 (300g, 4-8 or 8-14 mesh) as catalyst held at 370-380°, and arranged so that the effluent is connected to a receiver that allows the exit gasses (CO , CH_4 and H_2) into the fume cupboard. The yellow product (112-115g, 65-67%, m 134-145°) from the effluent which solidifies is washed with cold MeOH, and purified by recrystallisation from EtOH (50g/ 650ml) with 65-67% recovery, or from * C_6H_6 (50g/130ml) with 60% recovery to give pure colourless *hexamethylbenzene* **m 165-166°**. [Cullinane et al. *Org Synth Coll Vol* **4** 520 1963, DOI: 10.15227/orgsyn.035.0073.] It is also formed in >80% yields when *hexamethyl-Dewar benzene* [HMDB, 7641-77-2] comes into contact with anhydrous Al_2Cl_3 in hot *benzene (highly exothermic) [Schäfer & Hellmann *Angew Chem Int Ed* **6** 518 1967, DOI: 10.1002/anie.196705181]. It has been purified by sublimation, then recrystallisation from absolute EtOH, *benzene, EtOH/*benzene or EtOH/cyclohexane. It has also been purified by zone melting. Dry it in a low vacuum over P_2O_5 . It forms 1:1 addition compounds with *2,4,7-trinitrofluorene-one* (**m 165-166.3°**), *1,3,5-trinitrobenzene* (**m 173°**), *2,4,6-trinitrotoluene* (**m 123°**), *2-chloro-1,3,5-trinitrobenzene* (**m 151°**), and 1:1 complexes with Ag^+ , $\text{Cr}(\text{CO})_3$ and $\text{W}(\text{CO})_3$. Its FT-IR (Nujol) has ν_{max} at 1300.8, 1249.1, 1180.6, 1129.2, 1057.1, 994.9, 943.5, 794.7 and 772.0 cm^{-1} ; the ^1H NMR (60MHz, CDCl_3 , TMS) has δ at 2.22 (s, 18H, Me_6); and the ^{13}C NMR (15MHz, CDCl_3 , TMS) has δ at 17.72 (6-Me C) and 132.89 (6-aromatic C). [Beilstein **5** H 450, **5** I 213, **5** II 341, **5** III 1038, **5** IV 1137.]

Homophthalic acid [89-51-0] $\text{C}_9\text{H}_8\text{O}_4$, **M 180.2**, **m 178° to 182°, 180-181°, 182-183°, 189-190°, (depends on heating rate) pK_{Est(1)} ~3.5, pK_{Est(2)} ~4.3.** Crystallise the acid from boiling H_2O (25ml/g; 63ml/g at 20°), aqueous AcOH (**m 180°**) Dry it at 100°. The *S-benzylisothiuronium salt* has **m 155-156°** (from EtOH). [Price *Org Synth* **22** 611942, Grummitt et al. *Org Synth Coll Vol* **3** 449 1955, DOI: 10.15227/orgsyn.029.0049; Beilstein **9** H 857, **9** II 617, **9** III 4266, **9** IV 3343.] The *anhydride* [703-59-3] **M 162.1** has **m 141-142°**. [Beilstein **17** IV 3343, **17/11** V 270.]

Homoveratronic nitrile (3,4-dimethoxybenzyl nitrile) [93-17-4] $C_{10}H_{11}NO_2$, M 177.2, m 62-64°, 68°, b 171-178°/10mm, 184°/20mm, 195-196°/2mm, 319.7°/760m, d^{25}_4 1.082g/ml. Its solubility is 10% in MeOH, and it has been recrystallised from EtOH or MeOH. Purify it also by distillation followed by recrystallisation. [Niederl & Ziering *J Am Chem Soc* **64** 885 1942, DOI: 10.1021/ja01256a042; Julian & Sturgis *J Am Chem Soc* **57** 1126 1935, DOI: 10.1021/ja01309a053; *Beilstein* **10** I 198.]

Homoveratrylamine (2-[3,4-dimethoxyphenyl]ethylamine) [120-20-7] $C_{10}H_{15}NO_2$, M 181.2, b 99.3-101.3°/0.5mm, 168-170°/15mm, d^{20}_4 1.091, n^{20}_D 1.5460, pK_{Est} ~9.8. Purify the amine by fractionation through an efficient column in an inert atmosphere as it is a relatively strong base. [Horner & Sturm *Justus Liebigs Ann Chem* **608** 128 1957, DOI: 10.1002/jlac.19576080112; Jung et al. *J Am Chem Soc* **75** 4664 1953, DOI: 10.1021/ja01115a012.] The *hydrochloride* has m 152°, 154°, 156° (from EtOH, Me_2CO or EtOH/ Et_2O), the *picrate* has m 165-167°(dec), and the 4-nitrobenzoyl derivative has m 147° [Buck *J Am Chem Soc* **55** 2593 1933, DOI: 10.1021/ja01333a060]. [*Beilstein* **13** H 800, **13** IV 2604.]

Hordenine {4-[(2-dimethylamino)ethyl]phenol} [539-15-1] $C_{10}H_{15}NO_2$, M 165.2, m 116-118°, 117-118°, b 173-174°/11mm, pK^{25} 9.46 (OH). Crystallise Hordenine from EtOH or H_2O and sublime it at 105°/5mm, or 140-150°/atm. The 4,4'-dichlorodiphenyl disulfimide salt has m 145-146° (from Me_2CO/H_2O) [Runge et al. *Chem Ber* **88** 533 1955, DOI: 10.1002/cber.19550880414]. [*Beilstein* **13** H 626, **13** I 236, **13** II 356, **13** III 1640, **13** IV 1790.]

4-Hydrazinobenzoic acid [619-67-0] $C_7H_8N_2O_2$, M 152.2, m 205°, 216-220°(dec), 217° (dec), pK^{25} 4.13. The acid crystallises from water. [*Beilstein* **15** H 631, **15** IV 1372.] The *hydrochloride* [24589-37-3] M 188.6, has m 253°(dec) [*Beilstein* **15** III 837].

Hydrobenzamide [1-phenyl-*N,N'*-bis(phenylmethylene)-methanediamine] [92-29-5] $C_{21}H_{18}N_2$, M 298.4, m 101-102°, 102-105°, 107-108°. Crystallise hydrobenzamide from absolute EtOH, petroleum ether (m 107-108°), C_6H_6 (m 103°), or cyclohexane/*benzene. Dry it under vacuum over P_2O_5 . [Pirrone *Gazz Chim Ital* **67** 534 1937, *Beilstein* **7** H 215, **7** I 120, **7** II 166, **7** III 838.]

dl-Hydrobenzoin (1,2-diphenyl-1,2-ethanediol, *iso*-hydrobenzoin) [492-70-6] $C_{14}H_{14}O_2$, M 214.3, m 120°, 123°. Crystallise the diol from Et_2O /petroleum ether or H_2O (m 121-122°) [*Beilstein* **6** H 1004, **6** I 490, **6** II 969, **6** III 5431.] The *R,R*-(+)- [52340-78-0] and *S,S*-(-)- [2325-10-2] *enantiomers* have m 148.5-149.5°(dec), $[\alpha]^{20}_D$ + and -94.0 (c 2, EtOH) [Eisenlohr & Hill *Chem Ber* **70** 942 1937, DOI: 10.1002/cber.19370700514]. *meso*-Hydrobenzoin [579-43-1] has m 134-136°, 139°, 139-140°. Crystallise it from EtOH or water. [*Beilstein* **6** H 1003, **6** I 490, **6** II 967, **6** III 5429, **6** IV 6682.]

Hydroquinone (1,4-dihydroxybenzene, quinol) [123-31-9] $C_6H_6O_2$, M 110.1, m 170-171°, 172-175°, 175.4°, 176.6°, b 285°/atm, d^{25} 1.332, pK^{20}_1 9.91, pK^{20}_2 11.56. Crystallise quinol from acetone, *benzene, EtOH, EtOH/*benzene, water or acetonitrile (25g in 30ml), preferably under nitrogen. Dry it under vacuum. [Wolfenden et al. *J Am Chem Soc* **109** 463 1987, DOI: 10.1021/ja00236a026; *Beilstein* **6** H 836, **6** IV 5712.] It has reducing properties and is used as a photographic developer, de-pigmentor and antioxidant.

4-Hydroxyacetophenone [99-93-4] $C_8H_6O_2$, M 136.2, m 109°, 111°, 147-148°/3mm, pK^{25} 8.01. Crystallise it from diethyl ether, aqueous EtOH or *benzene/petroleum ether. [*Beilstein* **8** H 87, **8** IV 339.]

3-Hydroxyanthranilic acid (2-amino-3-hydroxybenzoic acid) [548-93-6] $C_7H_7NO_3$, M 153.1, m >240°(dec), 245-265°(dec), 253-154°, 252-255°, λ_{max} 298nm, log ϵ 3000 (0.1M HCl), pK^{20}_1 2.7, pK^{20}_2 5.19, pK^{20}_3 10.12. Crystallise the acid from H_2O or EtOH (m 254-255°). Sublime it at 170°/0.1mm. The *hydrochloride* has m 227° (from dilute HCl). [*Beilstein* **14** H 587, **14** III 1463, **14** IV 2071.] **Possible carcinogen.**

5-Hydroxyanthranilic acid (2-amino-5-hydroxybenzoic acid) [394-31-0] has m 233-234°, 235-236°, 247°(dec), 248°(dec), b 417.8°/760mm, d 1.491, pK^{25}_1 2.72, pK^{25}_2 5.37, pK^{20}_3 10.12. Crystallise the acid from water. The *benzamide* has m 240-242° (from AcOH). It is a hypoglycemic agent. [*Beilstein* **14** H 591, **14** II 357, **14** III 1468, **14** IV 2080.]

3-Hydroxybenzaldehyde [100-83-4] $C_7H_6O_2$, M 122.1, m 100° to 103°, 108°, b 191°/50mm, pK_1^{25} 8.98, pK_2^{25} 15.81. Crystallise the aldehyde from water. [Beilstein 8 H 58, 8 IV 240.]

4-Hydroxybenzaldehyde [123-08-0] has m 112° to 115°, 115-116°, 117-119°, b 191°/50mm, 308.3°/atm, pK^{25} 7.61. Crystallise it from water (containing some H_2SO_4). Dry it over P_2O_5 under vacuum. [Beilstein 8 IV 251.]

3-Hydroxybenzoic acid [99-06-9] $C_7H_6O_3$, M 138.1, m 200° to 203°, 200° to 202°, 200.8°, b 346°/atm, pK_1^{25} 4.08, pK_2^{25} 9.98. Crystallise the hydroxyacid from absolute EtOH. [Beilstein 10 IV 315.]

4-Hydroxybenzoic acid [99-96-7] has m 213-214°, 214.5°, d 1.46, pK_1^{25} 4.50, pK_2^{25} 9.11. Crystallise the hydroxyacid from water (solubility 0.5w.v%). [Beilstein 10 IV 345.]

2-Hydroxybenzyl alcohol (saligenine) [90-01-7] $C_7H_8O_2$, M 124.1, m 83° to 85°, 86-87°, 87°, d 1.613, pK^{25} 9.92. Crystallise saligenine from H_2O (6.7w/v% at 22°) or EtOH/ $*C_6H_6$ (m 89°). [Beilstein 6 IV 5896.]

3-Hydroxybenzyl alcohol [620-24-6] has m 69-70°, 71°, 72°, pK^{25} 9.83. Crystallise the alcohol from $*C_6H_6$ or $CHCl_3$. [Beilstein 6 III 4545, 6 IV 5907.]

4-Hydroxybenzyl alcohol (gastrodigenine) [623-05-2] has m 114° to 121°, 122°, 127°, 132°, pK^{25} 9.82. Crystallise the alcohol from H_2O , $*C_6H_6$ (m 124°), $*C_6H_6$ /EtOH or $ClCH_2CH_2Cl$ (m 122°). [Beilstein 6 III 4546, 6 IV 5909.] A food flavor ingredient. It is a pleiotropic agent inhibiting development of new blood vessels — tumour inhibitor.

2-Hydroxybiphenyl [90-43-7] $C_{12}H_{10}O$, M 170.2, m 55.5° to 57.5°, 56°, 57° to 59°, b 145°/14mm, 282°/760mm, d 1.293, pK^{20} 10.01. Crystallise it from petroleum ether. [Beilstein 6 IV 4579.] It forms a *sodium tetrahydrate salt* [6152-33-6] which is soluble in H_2O (1.2g/ml), in Me_2CO (1.4g/ml) and in MeOH (1.6/ml). [Beilstein 6 III 3281.] It is a disinfectant and a fungicide.

4-Hydroxybiphenyl (4-phenylphenol) [92-69-3] has m 164-165°, 166°, b 305-308°/760mm, 321°/atm, pK^{23} 9.55. Crystallise the phenol from aqueous EtOH, $*C_6H_6$, and dry it in a vacuum over $CaCl_2$ [Buchanan et al. *J Am Chem Soc* 108 7703 1986, DOI: 10.1021/ja00284a039]. Also purified it by sublimation. [Beilstein 6 IV 4600.]

trans-4-Hydroxycinnamic acid (p-coumaric acid) [501-98-4] $C_9H_8O_3$, M 164.2, m 210-213°, 214-215°, 215°, pK_1^{25} 4.64, pK_2^{25} 9.45. Crystallise *p*-coumaric acid from H_2O (charcoal). It forms needles from concentrated aqueous solutions as the *anhydrous acid*, but from hot dilute solutions the *monohydrate acid* separates on slow cooling. The acid (33g) has been crystallised from 2.5L of H_2O (1.5g charcoal) yielding 28.4g of recrystallised acid, m 207°. It is insoluble in $*C_6H_6$ or petroleum ether. The UV in 95% EtOH has λ_{max} at 223 and 286nm (ϵ 14,450 and 19000 $M^{-1}cm^{-1}$). [UV Wheeler & Covarrubias *J Org Chem* 28 2015 1963, DOI: 10.1021/jo01043a016; Corti *Helv Chim Acta* 32 681 1949, DOI: 10.1002/hlca.19490320309; Beilstein 10 IV 1005.]

3-(4-Hydroxy-3,5-dimethoxyphenyl)acrylic acid (sinapinic acid) [530-59-6] $C_{11}H_{12}O_5$, M 224.2, m 195°(dec), 204-205°(dec), b 249.2°/atm, $pK_{Est(1)}$ ~4.6, $pK_{Est(2)}$ ~9.3. Crystallise it from water. [Beilstein 10 H 508, 10 IV 2104.]

4-Hydroxydiphenylamine [122-37-2] $C_{12}H_{11}NO$, M 185.2, m 72-73°, pK_{Est} ~10.0. Crystallise the amine from chlorobenzene/petroleum ether, pentane (m 72°) or $*C_6H_6$ /petroleum ether (m 70°). [Beilstein 13 III 1019, 13 IV 1052.] **Environmentally toxic.**

2-Hydroxy-4-(n-dodecyloxy)benzophenone [2985-59-3] $C_{25}H_{34}O_3$, M 382.5, m 50-52°, b 506.3°/760mm, d 1.029, pK_{Est} ~7.1. Recrystallise it from *n*-hexane and then 10% (v/v) EtOH in acetonitrile [Valenty et al. *J Am Chem Soc* 106 6155 1984, DOI: 10.1021/ja00333a006].

4-Hydroxyindane [1641-41-1] $C_9H_{10}O$, M 134.2, m 49-50°, b 120°/12mm, pK^{25} 10.32. Crystallise 4-hydroxyindane from petroleum ether, pentane (m 50-50.5°) or $*C_6H_6$ (m 39.5-40°). It has UV with λ_{max} at 277nm (cyclohexane). The *acetyl* derivative has m 30-32° (from EtOH), b 127°/14mm and the *3,5-dinitrobenzoyl* derivative has m 114°. [Dallacker et al. *Chem Ber* 105 2565 1972, DOI: 10.1002/cber.19721050818; Beilstein 6 III 2427, 6 IV 3827.] **5-Hydroxyindane** [1470-94-6] has m 51° to 53°,

55°, **b** 127°/14mm, **255°/760mm**, **pK²³ 10.24**. Crystallise 5-hydroxyindane from petroleum ether (**m** 56°) or pentane (**m** 59-60°). It has UV with λ_{max} at 283.5nm (cyclohexane). The **3,5-dinitrobenzoyl** derivative has **m** 156°. [*Beilstein* 6 III 2428, 6 IV 3829.]

4-Hydroxy-3-methoxyacetophenone (Apocynin, Acetovanilone) [498-02-2] **C₉H₁₀O₃**, **M 166.2**, **m** 112° to 115°, **b** 263-265°/17mm, **pK_{Est} ~7.9**. Crystallise apocynin from water, or EtOH/petroleum ether. [*Beilstein* 8 IV 1814.] It has the odour of vanilla. **2-Hydroxy-4-methoxybenzophenone** (Oxybenzone) [131-57-7] **C₁₄H₁₂O₃**, **M 228.2**, **m** 65-66°, **b** 224-227°/atm, **pK²⁵ 7.6**. Recrystallise from *iso*PrOH. It is soluble in most organic solvents. Effective UV sunscreen. [Hardy & Forster USPat 2773903 1956 to American Cyanamid Co.]

trans-4-Hydroxy-3-methoxycinnamic acid (ferulic acid) [537-98-4; 1135-24-6] **C₁₀H₁₀O₄**, **M 194.2**, **m** 168° to 172°, 174°, **pK₁²⁵ 4.58**, **pK₂²⁵ 9.39**. Crystallise ferulic acid from H₂O or EtOH/H₂O. UV has λ_{max} (EtOH) at 236nm (specific ϵ 6,000) and 322nm (specific ϵ 9,300). Its **ethyl ester**, C₁₂H₁₄O₄, provides **anhydrous** crystals **m** 38°, from petroleum ether (b 65-110°), and crystals of the **monohydrate** **m** 75.5-76.5°, from aqueous EtOH [Pearl & Beyer *J Org Chem* 16 216 1951, DOI: 10.1021/jo01142a008]. [*Beilstein* 10 H 436, 10 IV 1776.] It is used in food preservation.

4-Hydroxy-2-methylazobenzene (3-methyl-4-phenylazophenol) [1435-88-7] **C₁₃H₁₂N₂O**, **M 212.2**, **m** 100-101°, 112°, **pK_{Est} ~9.5**. Crystallise the phenol from hexane. [*Beilstein* 16 II 61, 16 IV 195.]

4-Hydroxy-3-methylazobenzene (2-methyl-4-phenylazophenol) [621-66-9] has **m** 125-126°. Crystallise the phenol from hexane or petroleum ether (**m** 130°). [*Beilstein* 16 II 59, 16 III 104, 16 IV 193.]

3-Hydroxy-4-methylbenzaldehyde [57295-30-4] **C₈H₈O₂**, **M 136.1**, **m** 73°, **b** 252.4°/760mm, **pK_{Est} ~10.2**. Crystallise it from water or *C₆H₆ (**m** 71-71°). The **O-methyl ether** has **m** 45-46° (from Et₂O/hexane). [Sidgwick & Allott *J Chem Soc* 2819 1923, DOI: 10.1039/CT9232302819; Flitsch et al. *Justus Liebigs Ann Chem* 1413 1985, DOI: 10.1002/jlac.198519850712; *Beilstein* 8 H 100, 8 II 103, 8 IV 368.]

5-Hydroxy-2-methyl-1,4-naphthaquinone (plumbagin) [481-42-5] **C₁₁H₈O₃**, **M 188.2**, **m** 78-79°, **pK_{Est(1)} ~9.5**, **pK_{Est(2)} ~11.0**. Crystallise it from aqueous EtOH (yellow needles). It is steam volatile and sublimes on heating *in vacuo*. [Fieser & Dunn *J Am Chem Soc* 58 572 1936, DOI: 10.1021/ja01295a010; *Beilstein* 8 III 2576, 8 IV 2376.] It inhibits UV-induced development of squamous carcinoma [Sand et al. *Carcinogenesis* 33 184 2012, DOI: 10.1093/carcin/bgr249, PMID: 22072620].

6-Hydroxy-2-methyl-1,4-naphthaquinone [633-71-6] has **pK_{Est} ~10.0**. Crystallise the naphthaquinone from aqueous EtOH and sublime it in a vacuum.

2-Hydroxy-1-naphthaldehyde [708-06-5] **C₁₁H₈O₂**, **M 172.2**, **m** 76° to 80°, 82°, **b** 139-142°/4mm, **192°/27mm**, **pK²⁵ 8.27** (50% aqueous EtOH). Crystallise the aldehyde from EtOH (1.5ml/g), aqueous EtOH, aqueous AcOH (**m** 84°), EtOAc or H₂O. It is air sensitive. [Russell & Lockhart *Org Synth Coll Vol* 3 463 1955, DOI: 10.15227/orgsyn.022.0063; *Beilstein* 8 H 143, 8 I 564, 8 II 171, 8 III 1108, 8 IV 1160.]

2-Hydroxy-1-naphthaleneacetic acid [10441-45-9] **C₁₂H₁₀O₃**, **M 202.2**, **m** 147-148°, **pK_{Est(1)} ~4.2**, **pK_{Est(2)} ~8.3**. Crystallise the acid from EtOH/water (1:9, v/v, activated charcoal), H₂O (**m** 157°) or xylene (**m** 147°). Dry it under vacuum, over silica gel, in the dark. Store it in the dark at -20° [Gafni et al. *J Phys Chem* 80 898 1976, DOI: 10.1021/j100549a027]. It readily forms a **cyclic lactone** (**m** 107°) [4352-63-0] **C₁₂H₈O₂**. [*Beilstein* 10 III 1102, 10 IV 1201.] **6-Hydroxy-2-naphthalenepropionic acid** [553-39-9] **C₁₃H₁₂O₃**, **M 216.2**, **m** 180-181°, **pK_{Est(1)} ~4.6**, **pK_{Est(2)} ~9.0**. Crystallise the acid from aqueous EtOH or aqueous MeOH. [Ormancey & Horeau *Bull Soc Chim Fr* 962 1955, *Beilstein* 10 III 1113.]

3-Hydroxy-2-naphthalide (Naphthol AS) [92-77-3] **C₁₇H₁₃NO₂**, **M 263.3**, **m** 246° to 248°, 248.0-248.5°, **CI 37505**. Crystallise it from xylene or AcOH which forms plates **m** 243-244° [Schnopper et al. *Anal Chem* 31 1542 1959, DOI: 10.1021/ac60153a035]. Its UV has λ_{max} (EtOH) at 394nm. [*Beilstein* 12 H 505.]

3-Hydroxy-2-naphtho-4'-chloro-o-toluidide [92-76-2] **C₁₈H₁₄ClNO₂**, **M 311.8**, **m** 243.5-244.5°. Crystallise it from xylene [Schnopper et al. *Anal Chem* 31 1542 1959, DOI: 10.1021/ac60153a035].

3-Hydroxy-2-naphthoic-1'-naphthylamide [123-68-3] $C_{21}H_{15}NO_2$, M 313.3, m 217-5-218.0°, 222-223°, b 479.3°/760mm. Crystallise the amide from xylene [Schnopper et al. *Anal Chem* **31** 1542 1959, DOI: 10.1021/ac60153a035].

3-Hydroxy-2-naphthoic-2'-naphthylamide [136-64-8] has m 243.5-244.5°, and other naphthol AS derivatives. Crystallise it from xylene [see Schnopper et al. *Anal Chem* above DOI: 10.1021/ ac60153a035].

2-Hydroxy-1,4-naphthoquinone (Lawsone B, Neutral Orange 6, hennotanic acid, tautomeric with 4-hydroxy-1,2-naphthoquinone) [83-72-7] $C_{10}H_6O_3$, M 174.2, m 191° to 193°(dec), 192° to 195°(dec), CI 75480 pK_1^{25} -5.6 (C=O protonation), pK_2^{25} 2.38, pK_3^{25} 4.00 (phenolic OH). Crystallise Lawsone B from * C_6H_6 or AcOH (m 192.5°, 195-196°). Its solubility in $CHCl_3$ is 0.1g/ml. It sublimes in a vacuum (m 194°). It has UV with λ_{max} at 455nm (aqueous NaOH). [Beilstein **8** H 300, **8** I 635, **8** II 344, **8** III 2543, **8** IV 2360.] It is an antimicrobial antioxidant dye isolated from hemp and used to dye hair and skin.

5-Hydroxy-1,4-naphthoquinone (Juglone) [481-39-0] has m 155°, 161° to 163°, 164-165°, pK^{25} 8.7. Crystallise Juglone from *benzene/petroleum ether or petroleum ether. It is steam volatile and can be sublimed. Its UV has λ_{max} (MeOH) at 420nm (ϵ 7482). It dissolves in alkalis to give a purple-red solution. [Beilstein **8** III 2558, **8** IV 2368.]

2-Hydroxy-5-nitrobenzyl bromide (Koshland's reagent) [772-33-8] $C_7H_6BrN_3$, M 232.0, m 144-146°, 147°, 148-149°, pK_{Est} ~8.0. Crystallise the bromide from * C_6H_6 or * C_6H_6 /ligroin. It is slightly soluble in EtOH, soluble in * C_6H_6 and AcOH, and very soluble in ligroin. Store at 2-8°. [Beilstein **6** H 367.] Enzyme inhibitor.

2-Hydroxyphenylacetic acid [614-75-5] $C_8H_8O_3$, M 152.2, m 145-147°, 148-149°, 152-153°, b 240-243°/760mm, $pK_{Est(1)}$ ~4.3, $pK_{Est(2)}$ ~10.1. Crystallise the acid from ether or chloroform (m 147°, m from latter solvent is always lower). [Beilstein **10** H 187, **10** I 81, **10** II 112, **10** III 422, **10** IV 536.]

3-Hydroxyphenylacetic acid [621-37-4] has m 128°, 129° to 133°, 130°, $pK_{Ext(1)}$ ~4.3, $pK_{Ext(2)}$ ~10. Crystallise the acid from * C_6H_6 /ligroin or EtOAc/cyclohexane (m 131-132°). [Beilstein **10** II 112, **10** III 428, **10** IV 541.]

4-Hydroxyphenylacetic acid [156-38-7] has m 148-150°, 150-151°, 152°, pK_1 4.28, pK_2 10.1. Crystallise the acid from water or Et₂O/petroleum ether. The *p*-bromophenacyl ester has m 117° (from EtOH). [Beilstein **10** II 112, **10** III 430, **10** IV 543.]

N-(4-Hydroxyphenyl)-3-phenylsalicylamide [550-57-2] $C_{19}H_{15}NO_3$, M 305.3, m 183-184°, pK_{Est} ~9.5. Crystallise the amide from aqueous MeOH. [Beilstein **13** IV 224.]

R(+)-2-Hydroxy-3-phenylpropionic acid (3-phenyl lactic acid) [7326-19-4] and **S(-)-2-Hydroxy-3-phenylpropionic acid** [20312-36-1] $C_9H_{10}O_3$, M 166.2, m 122° to 124°, 125-126°, $[\alpha]_D^{25}$ (+) and (-) 18.7 (EtOH), $[\alpha]_D^{20}$ (+) and (-) 22 (c 1, H₂O), for pK see below. Crystallise the acid from water, MeOH, EtOH or *benzene. [Beilstein **10** IV 653.]

dl-2-Hydroxy-3-phenylpropionic acid [828-01-3] has m 97-98°, b 148-150°/15mm, pK_{Est} ~3.7. Crystallise the propionic acid from *benzene or chloroform. [Beilstein **10** IV 653.]

3-p-Hydroxyphenylpropionic acid (phloretic acid) [501-97-3] has m 129-130°, 131°, 131-133°, $pK_{Est(1)}$ ~4.7, $pK_{Est(2)}$ ~10.1. Crystallise phloretic acid from ether or H₂O. [Beilstein **10** IV 631.] It is a fluorogenic reagent for amine oxidase assays [Matsumoto et al. *Anal Biochem* **138** 133 1984, DOI: 10.1016/0003-2697(84)90780-2].

4-Hydroxyphenylpyruvic acid [156-39-8] $C_9H_8O_4$, M 180.2, m 199-220°(dec), pK_{Est} ~2.3. Crystallise it three times from 0.1M HCl/EtOH (4:1, v/v) immediately before use [Rose & Powell *Biochem J* **87** 541 1963, DOI: 10.1042/bj0870541], or from Et₂O. The **3,4-dinitrophenylhydrazone** has m 178°. [Beilstein **10** IV 3630.]

N-Hydroxyphthalimide [524-38-9] $C_8H_5NO_3$, M 163.1, m 230°, ~235°(dec), 237-240°, pK^{30} 7.0. Dissolve the imide in H₂O by adding Et₃N to form the salt and while hot, acidify, cool and pour into a large volume of H₂O. Filter off the solid, wash it with H₂O and dry it over P₂O₅ in a vacuum. [Nefkens & Tesser *J Am Chem Soc* **83** 1263 1961, DOI: 10.1021/ja01466a068; Fieser **1** 485, Nefkens et al. *Recl Trav Chim Pays-Bas* **81** 683 1962, DOI: 10.1002/recl.19620810807.] It crystallises from H₂O in yellow needles. The **O-acetyl** derivative has m 178-180° (from EtOH). [Beilstein **21/11** V 100.] Activates terminal carboxy groups in peptide synthesis.

4'-Hydroxypropiophenone [70-70-2] $C_9H_{10}NO_2$, **M 150.2**, **m 147.5-148.5°**, **149°**, **b 140-145°/0.5mm**, **pK²⁵ 8.05**. Crystallise the phenone from H₂O (**m** 149.8-150.2°) (with solubilities of 0.03w/v% at 15° and 3.3w/v% at 100°) or EtOH (**m** 147°). It is an inhibitor of pituitary gonadotrophin. The *benzoyl derivative* has **m 117°**, and the *semicarbazone* has **m 183°** (EtOH). [Miller & Hartung *Org Synth Coll Vol* **2** 543 1943, DOI: 10.15227/orgsyn.013.0090; *Beilstein* **8** H 102, **8** II 104, **8** III 379.]

trans-4-Hydroxystilbene [6554-98-9] $C_{14}H_{12}O$, **M 196.3**, **m 185° to 188°**, **189°**. Crystallise it from *C₆H₆, MeOH (**m** 186-187°) or acetic acid. [*Beilstein* **6** H 693, **6** II 657, **6** III 3497, **6** 4855.]

6-Hydroxy-1-tetralone [3470-50-6] $C_{10}H_{10}O_2$, **M 162.2**, **m 154-157°**. It has been prepared by catalytic reduction with Pd/SrCO₃ in EtOAc [Cornforth et al. *J Chem Soc* 3348 1955, DOI: 10.1039/JR9550003348], RaNi in aqueous NaOH [Papa et al. *J Org Chem* **71** 3241 1949, DOI: 10.1021/jo01155a005] or Pt₂O/AcOH [Papa *J Am Chem Soc* **71** 3241 1949, DOI: 10.1021/ja01177a515] of 1,6-dihydroxynaphthalene [see 575-44-0 above] and purified by recrystallisation from 20% aqueous MeOH or H₂O. **NOTE:** two hydrogen atoms are introduced into the ring containing the 1-hydroxygroup of 1,6-dihydroxynaphthalene which produces 3,4-dihydro-1,6-dihydroxynaphthalene (the enol) which ketonises to 6-hydroxy-1-tetralone. The *semicarbazone*, prepared in the usual way, has **m 216.5-217.5°** after crystallisation from EtOH. Acetylation gave **6-acetoxy-1-tetralone** with **m 61-62°** (from aqueous MeOH). Methylation with Me₂SO₄/alkali produced **6-methoxy-1-tetralone** [1078-19-9] $C_{11}H_{12}O_2$, **M 176.2**, with **b 135-139°/1mm(171°/11mm)**, **m 75-77°(77-79°)** [*Beilstein* **8** IV 904.] The isomeric **5-methoxy-2-tetralone** [32940-15-1] $C_{11}H_{12}O_2$, **M 176.2**, has **b 120-122°/0.4mm**, **m 32-36°**, and was prepared by reduction of **1,6-dimethoxynaphthalene** with Na/EtOH, isolation as the bisulfite which was hydrolysed with aqueous Na₂CO₃ and extracted into EtOH [Cornforth et al. *J Chem Soc* 689 1942, DOI: 10.1039/JR9420000689].

2R(+) 2-Hydroxy-1,2,2-triphenylacetate [(R-HYTRA, 1,2-ethanediol, 1,1,2-triphenylacetate) [95061-47-5] and **2S(-) 2-Hydroxy-1,2,2-triphenylacetate** [95061-51-1, 89559-96-6] $C_{22}H_{20}O_3$, **M 332.4**, **m 237° to 240°**, **249-251°**, $[\alpha]_D^{20}$ **(+) or (-) 212 to 218 (c 1, pyridine)**. The ester is prepared by adding, under N₂ with stirring, a solution of Sc(OTf)₃ (1.23g 2.5mmol) in dry MeCN (125ml) slowly (~35min) to a solution of *R*(+)-1,1,2-triphenylethane-1,2-diol (35g, 0.121mol, [9506-46-4] see below) and Ac₂O (17.1ml, 0.181mol) and stirring is continued for 3 hours. The white solid, which started to separate after ~8 minutes, is filtered off washed with dry MeCN (2 x 25ml) and dried *in vacuo* at 40 overnight to give the pale yellow ester (35.4g 88% yield). The melting point varied with solvent of recrystallisation; and was **m 213-221°** from *C₆H₆ [Macor et al. *Org Synth* **77** 45 2000, *Org Synth Coll Vol* **10** 464 2000, DOI: 10.15227/orgsyn.077.0045], **m 249-251°** from toluene [Braun et al. *Org Synth Coll Vol* **9** 507 1998, *Org Synth* **72** 32 1995, DOI: 10.15227/orgsyn.072.0032, from CH₃COCl/pyridine/CH₂Cl₂/0° then ~25°/4hr in 92.3% yield], **m 220-221°** from Et₂O [Polansky et al. *Monatsh Chem* **87** 24 1956, DOI: 10.1007/BF00903587], **m 220-220.5°** from Et₂O/CH₂Cl₂ [Corey & Casanova *J Am Chem Soc* **85** 165 1963 DOI: DOI: 10.1021/ja00885a013] and **m 224.5-225.5°** from *C₆H₆ [Ito, Nishino & Kurosawa *Bull Chem Soc Jpn* **56** 3527 1983, DOI: 10.1246/bcsj.56.3527], and **m 239° (R) and 237° (S) after sublimation at 180/0.001mm** [Devant, Mahler and Braun *Chem Ber* **121** 397 1988, DOI: 10.1002/cber.19881210303]. **Note** that acetylation takes place with *retention* of configuration. It has IR (CHCl₃) with ν_{max} at 3064, 3024, 1737, 1495, 1372, 1239, 1168, 779 cm⁻¹ and the ¹H NMR (300MHz, CDCl₃ TMS) has δ at 1.96 (3H, s, CH₃CO), 2.78 (1H, s, OH), 6.68 (1H, s, PhCHOAc), 7.05-7.40 (13H, m, ArH) and 7.54-7.57 (2H, m, Ar-H); ¹³C NMR (125MHz, CDCl₃, TMS) has δ at 21.1 (CH₃), 78.5 (CHOAc), 80.3 (Ph₂COH), 126.2, 126.3, 127.0, 127.3, 125.5, 127.8, 127.9, 128.4, 128.5 (Ar-C), 135.9, 142.7, 144.8 (*ipso*-Ar) and 169.7 (C=O). Similar reactions with the same consequences have been performed with the *S*(-)-enantiomers [Devant, Mahler and Braun *Chem Ber* **121** 397 1988, DOI: 10.1002/cber.19881210303]. This ester undergoes stereospecific aldol condensations *via* a doubly deprotonated acetate methyl group; e.g. the *R*(+)-ester with 2-methylpropanal in the presence of lithium diisopropylamide/THF at -126° followed by hydrolysis with KOH/H₂O/MeOH/reflux provides ***R*(+)-3-hydroxy-4-methylpentanoic acid** [77981-87-4] $[\alpha]_D^{20}$ +32 to +37 (c 1.22, CHCl₃; 86-92% optical purity) in 61-78% yield. [Braun & Gräf *Org Synth Coll Vol* **9** 497 1998, DOI: 10.15227/orgsyn.072.0038; Devant et al. *Chem Ber* **121** 397 1988, DOI: 10.1002/cber.19881210303; Fieser **16** 180.]

9-Hydroxytriptycene [73597-16-7] $C_{20}H_{14}O$, **M 270.3**, **m 242-243°**, **245-246.5°**. Prepared by hydrolysis of 9-acetoxytriptycene. It was recrystallised from *benzene/petroleum ether (colourless leaves). Dried it at 100° in

a vacuum (turning into white leaves). [Bartlett & Greene *J Am Chem Soc* **76** 1088 1954, DOI: 10.1021/ja01633a047.] The ^1H NMR (400MHz, CDCl_3 , and TMS) has δ at 7.53 (1-H, 8-H, 13-H), 7.06 (2-H, 7-H, 14-H), 7.01 (3-H, 6-H, 15-H), 7.37 (4-H, 5-H, 16-H), 5.39 (10-H, bridgehead) and 3.34 (OH) with $J_{1,2} = 7.7\text{Hz}$, $J_{1,3} = 0.9\text{Hz}$, $J_{1,4} = 0.1\text{Hz}$, $J_{2,3} = 7.4\text{Hz}$, $J_{2,4} = 0.8\text{Hz}$ and $J_{3,4} = 7.5\text{Hz}$; the ^{13}C NMR (100MHz, CDCl_3 , and TMS) has δ at 118.8 (1-C, 8-C, 13-C), 124.9 (2-C, 7-C, 14-C), 125.3 (3-C, 6-C, 15-C), 123.1 (4-C, 5-C, 16-C), 143.5 (4a-C, 10a-C, 11-C), 145.7 (9a-C, 8a-C, 12-C), 80.6 (9-C) and 52.6 (10-C); and the ^{13}C NMR for the solid state has been recorded but is only slightly different from the solution spectrum [Imashiro et al. *J Am Chem Soc* **109** 729 1987, DOI: 10.1021/ja00237a016].

Hypericin (hypericum, **4,5,7,4',5',7'-hexahydroxy-2',2'-dimethylnaphthodianthrone**) [548-04-9] $\text{C}_{30}\text{H}_{16}\text{O}_6$, **M 504.4**, **m 320°(dec)**. Crystallise hypericin from pyridine by addition of methanolic HCl to give a bluish-black solid. [Beilstein **8** IV 3761.] This active ingredient of the plant called St John's Wort is an antibiotic, an antiviral and a non-specific kinase inhibitor. It is used also to treat depression, cancer and AIDS.

Ibuprofen [(*S*+) and (*R*-) 4-isobutyl- α -methylphenylacetic acid, α -methyl-4-(2-methylpropyl)phenyl acetic acid, Brufen, Nurofen, Motrui] [(*S* +) 51146-56-6, (*R* -) 51146-57-7, (*RS* -) 15687-27-1] $\text{C}_{13}\text{H}_{18}\text{O}_2$, **M 206.3**, **m 52-53°**, $[\alpha]_{\text{D}}^{20} +59$ (c 2, EtOH). Crystallise the (+) and (-) acids from EtOH or aqueous EtOH. The *racemate*, which crystallises from petroleum ether with **m 75-77°**, is sparingly soluble in H_2O and has IR (film) with ν_{max} at 1705 (C=O), 2300–3700 (OH broad) cm^{-1} . [Shiori et al. *J Org Chem* **43** 2936 1978, DOI: 10.1021/jo00408a049; Kaiser et al. *J Pharm Sci* **65** 269 1976, DOI: 10.1002/jps.2600650222; *J Pharm Sci* **81** 221 1992, DOI: 10.1002/jps.2600810306; Freer *Acta Cryst (C)* **49** 1378 1993, DOI: 10.1107/S0108270193000629 for the (*S*+) *enantiomer*; for a new synthesis using the reaction of *p*-isobutylphenylPb(OAc)₃ and the 5-methyl-Meldrum acid derivative (2,2,5-trimethyl-1,3-dioxan-4,6-dione) to give 2,2-dimethyl-5-methyl-5-isobutylphenyl-1,3-dioxan-4,6-dione quantitatively followed by hydrolysis and decarboxylation to *ibuprofen* see Pinhey & Rowe [*Tetrahedron Lett* **21** 965 1980, DOI: 10.1016/S0040-4039(00)77752-5.] It is a cyclooxygenase (COX) inhibitor, and a non-steroidal anti-inflammatory (NSAID) drug. Used with codeine as an analgesic. The physiological activity resides mainly in the *S*-*enantiomer*.

The *S*(+)-*Ibuprofen-S*-lysine salt (*Dexibuprofen lysine*, MK-223) [141505-32-0] $\text{C}_{13}\text{H}_{18}\text{O}_2 \cdot \text{C}_6\text{H}_{14}\text{N}_2\text{O}_2 \cdot \text{H}_2\text{O}$, **M 370.5**, is prepared by adding *S*(+)-ibuprofen (52.6g, 0.25mole) in EtOH (250ml) with stirring over 5 minutes to a solution of *S*-lysine (37.5, 0.25mole) in H_2O (42ml) at $\leq 35^\circ$, filtered through a medium frit glass funnel, diluted with EtOH (575ml) over 5 minutes at 20-25°, seeded with the salt (50mg), diluted further with EtOH (400ml) over 10 minutes and the suspension is stirred at 20-25° for 1 hour then cooled to 0-5° for 1 hour. The salt is filtered off, washed with cold EtOH (2 x 5ml) and dried at 50° *in vacuo* to give *pure salt* (74g, 84%) with $[\alpha]_{405}^{25} +14.9$ (c 1, MeOH). [King C. Kwan US Pat 5200558A to Merck & Co, Aug 9, 1991.]

1,3-Indandione [606-23-5] $\text{C}_9\text{H}_7\text{O}_2$, **M 146.2**, **m 129-132°**, **pK¹⁸ 7.2** (1% aqueous EtOH). Recrystallise it from EtOH or C_6H_6 . In dilute alkali it gives a deep yellow solution of the enol. [Bernasconi & Paschalis *J Am Chem Soc* **108** 2969 1986, DOI: 10.1021/ja00271a027]. [Beilstein **7** IV 2344.] Its mono-oxime **2-oximino-1-indanone** [15028-10-1] $\text{C}_9\text{H}_7\text{NO}_2$, **M 161.2**, has **m 209°(dec)** (from EtOH).

Indane (indan, hydrindene, 1,2-trimethylenbenzene) [496-11-7] C_9H_{10} , **M 118.1**, **m -51.4°**, **b 79°/29mm**, **177-179.5°/760mm**, $d_4^{20} 0.960$, $n_{\text{D}}^{20} 1.536$. Shake indane with concentrated H_2SO_4 , then water, dry (Na_2SO_4) and fractionally distil it. [Beilstein **5** H 486, **5** I 234, **5** II 376, **5** III 1200, **5** IV 1371.]

Indene [95-13-6] C_9H_8 , **M 116.2**, **f -1.5°**, **m -5° to -3°**, **b 114.5°/100mm**, **181-182°/atm**, $d_4^{20} 0.994$, $n_{\text{D}}^{20} 1.5763$. Shake indene with 6M HCl for 24 hours (to remove basic nitrogenous material), then reflux it with 40% NaOH for 2 hours (to remove benzonitrile). Fractionally distil, then fractionally crystallise it by partial freezing. The higher-melting portion is converted into its sodium salt by adding a quarter of its weight of sodamide under nitrogen and stirring for 3 hours at 120°. Unreacted organic material is distilled off at 120°/1mm. The sodium salts are hydrolysed with water, and the organic fraction is separated by steam distillation, followed by fractional distillation. Before use, the distillate is passed, under nitrogen, through a column of activated silica gel. It turns yellow in air as it readily oxidises and polymerises. Store it in the presence of *tert*-butylcatechol (50-100ppm) as antioxidant. [Russell *J Am Chem Soc* **78** 1041 1956, DOI: 10.1021/ja01586a045; Beilstein **5** IV 1532.]

2-Iodoaniline [615-43-0] C_6H_6IN , **M 219.0**, **m 55-58°, 60-61°, pK²⁵ 2.54**. Distil 2-iodoaniline with steam and crystallise it from *benzene/petroleum ether. The *N*-**acetyl** derivative has **m 110°**. [Beilstein 12 IV 1542.]

4-Iodoaniline [540-37-4] has **m 62-63°, pK²⁵ 3.81**. Crystallise it from petroleum ether (b 60-80°) by refluxing, then cool it in an ice-salt bath freezing mixture. Dry it in air. *Alternatively*, crystallise it from EtOH and dry it *in vacuo* for 6 hours at 40° [Edidin et al. *J Am Chem Soc* 109 3945 1987, DOI: 10.1021/ja00247a019]. The *N*-**acetyl** derivative has **m 184°** (from MeOH). [Beilstein 12 IV 1544.]

4-Iodoanisole [696-62-8] C_7H_7IO , **M 234.0**, **m 48° to 51°, 51-52°, b 133-133.3°/25mm, 139°/35mm, 237°/726mm**. Crystallise 4-iodoanisole from aqueous EtOH and/or distil it under vacuum. [Beilstein 6 H 208, 6 I 109, 6 II 199, 6 III 744, 6 IV 1075.]

Iodobenzene [591-50-4] C_6H_5I , **M 204.0**, **m -29°, b 63-65°/10mm, 188°/atm, d₄²⁰ 1.829, n_D²⁵ 1.6169**. Wash it with dilute aqueous Na₂S₂O₃, then water. Dry it with CaCl₂ or CaSO₄, decolourise with charcoal and distil it under reduced pressure then store it in the dark with Hg or silver powder to stabilise it. [Beilstein 5 IV 688.]

2-Iodobenzoic acid [88-67-5] $C_7H_5IO_2$, **M 248.4**, **m 160-161°, 161-162°, 161.6-162°, 162°, 162.7-163.5°, 164°, d 2.25, pK²⁰ 2.93, pK²⁵ 2.84 (H₂O), pK²⁰ 2.93 (1% EtOH), pK -7.78 (H₂SO₄)**. Crystallise the acid repeatedly from water (charcoal), EtOH, aqueous EtOH, aqueous Me₂CO, Me₂CO/Et₂O or *C₆H₆. Sublime it under vacuum at 100°. *o*-Iodobenzoic acid is prepared by the following procedure: anthranilic acid (31.9g, 233mmol, see [118-92-3]) is dissolved in dilute H₂SO₄ (50g, 27ml, 98% acid, 500mmol, in 250ml of H₂O), cooled to 0°; and while stirring NaNO₂ (20g, 290mmol) in H₂O (40ml) is added dropwise keeping the temperature below 10° (add crushed ice to the mixture if necessary). Diazotisation is complete when starch-iodide paper wetted with a drop of the mixture turns blue or brown due to the presence of excess of HNO₂. A solution of KI (60g, 360mmol) in H₂O (100ml) is gradually added with stirring, the mixture is set aside for 1 hour then heated at ~100° until effervescence (N₂ liberated) ceases. The mixture is cooled in ice, the brown iodo-acid (~92%) is filtered off, washed well with H₂O and dried *in vacuo*. The brown colour is not easily removed by recrystallisation so the acid is converted into the ethyl ester e.g. by Fischer and Speier's method [Fischer & Speier *Chem Ber* 28 3252 1895, DOI: 10.1002/cber.189502803176]. Dry HCl gas is bubbled through EtOH (~200ml) cooled in ice-water until the increase in weight is ~4g. The crude iodobenzoic acid is added to it and the mixture is boiled under reflux for 2 hours, or until an aliquot poured into H₂O does not deposit a solid. The mixture is poured into cold H₂O, the heavy oil that separated is extracted into Et₂O, the extract is washed with H₂O, saturated aqueous NaHCO₃ (care-effervescence), brine, dried (CaCl₂), filtered, evaporated and the residual oil is distilled to give a 73% yield of pure **ethyl o-iodobenzoate M 276.0** [1829-28-3], **b 151-152°/12-13mm (163-165°/23mm, 275°/atm)**. This ester is dissolved in 0.5 *N* alcoholic KOH (from 7g of KOH dissolved in 7ml of H₂O and diluted to 250ml with EtOH) in the ratio of 1g:25ml, and refluxed for 30 minutes (or until a drop added to H₂O dissolves completely), poured it into H₂O, and the clear solution of the potassium salt is acidified to pH <0 when pure iodobenzoic acid (theoretical yield) separates. It is collected, washed with cold H₂O, and dried *in vacuo*. [Baker et al. *J Chem Soc* 3721 1965, DOI: 10.1039/JR9650003721; Cohen & Raper *J Chem Soc* 1271 1904, DOI: 10.1039/CT9048501271.] The FT-IR (Nujol) has ν_{max} at 3061.7, 2646.1, 1683.9, 1582.4, 1405.0, 1269.8, 1015.8, 739.0 and 679.7 cm⁻¹; the ¹H NMR (DMSO-d₆, TMS) has δ at 7.25 (t of d, 1H, *J* = 7 Hz and 1.5 Hz, H-5), 4.85 (t, 1H, *J* = 7 Hz, H-4), 7.72 (d of d, 1H, *J* = 7 and 1.5 Hz, H-6) and 8.01 (d, 1H, *J* = 7 Hz, H-3); and the ¹³C NMR (DMSO-d₆, TMS) has δ at 167.93, 140.32, 136.72, 132.25, 129.88 127.97 and 93.92.

The **acid chloride** [609-67-6] **M 266.4**, has **m 35-40°, b 135°/19mm (159°/27mm)**, and the **amide** [3930-83-4] **M 247.0**, has **m 183°** (needles from H₂O). The **4-bromobenzyloisothiuronium salt** has **m 154°** (from EtOH), and the **4-chlorobenzyloisothiuronium salt** has **m 162°** (from dioxane). [Beilstein 9 H 363, 9 I 148, 9 II 239, 9 III 1432, 9 IV 1030.]

3-Iodobenzoic acid [618-51-9] has **m 185-186°, 186.6-186.8°, 189°, 189-189.2°, pK²⁵ 3.85, pK⁵⁰ 3.97, pK -7.64 (H₂SO₄)**. Crystallise the acid repeatedly from water, EtOH, aqueous EtOH, Me₂CO or aqueous AcOH. Sublime it under vacuum at 100°. **Ethyl m-iodobenzoate** [58313-23-8] **M 276.0b**, has **b 150.5°/15mm (165-166°/24mm)**, the **m-acid chloride** [1711-10-0] **M 266.4**, has **b 104-105°/1mm (159-150°/23mm)**, and the **m-amide** [10388-19-9] **M 247.0**, has **m 186°** (needles from H₂O). The **4-bromobenzyloisothiuronium salt** has **m 152°** (from EtOH), and the **4-chlorobenzyloisothiuronium salt** has **m 154°** (from dioxane). [Wallingford & Krueger *Org Synth Coll Vol* 2 353 1948, DOI: 10.15227/orgsyn.019.0057; Beilstein 9 III 1437, 9 IV 1033.]

4-Iodobenzoic acid [619-58-9] $C_7H_5IO_2$, M 248.4, m 269-270°, 271-272°, 272.5°, 273°, pK^{25} 4.00, pK -7.50 (H_2SO_4). Crystallise the acid repeatedly from water, aqueous EtOH, aqueous Me_2CO , aqueous AcOH. Sublime it under vacuum at 200°/0.4mm. **Ethyl p-iodobenzoate** M 276.0 [51934-41-9] has b 154°/15mm, the **p-acid chloride** has M 266.4 [1711-02-0] has m 83° (77-78°) (needles from Et_2O), b 126°/9mm (163-164°/32mm), and the **p-amide** has M 247.0 [3956-07-8], m 217.6° (209°) (needles from H_2O). The **4-bromobenzylisothiuronium salt** has m 181° (from EtOH), and the **4-chlorobenzylisothiuronium salt** has m 177° (from dioxane). [Whitmore & Woodward *Org Synth Coll Vol I* 325 1948, DOI: 10.15227/orgsyn.007.0058; *Beilstein* 9 H 366, 9 I 149, 9 II 240, 9 III 1442, 9 IV 1035.]

4-Iodobiphenyl [1591-31-7] $C_{12}H_9I$, M 280.1, m 110° to 113°, 113.7-114.3°, b 207°/28mm. Crystallise 4-iodobiphenyl from EtOH/* C_6H_6 , and dry it *in vacuo* over P_2O_5 . [*Beilstein* 5 H 581, 5 I 273, 5 II 486, 5 III 1748, 5 IV 1821.]

1-Iodo-2,4-dinitrobenzene [709-49-9] $C_6H_3IN_2O_4$, M 294.0, m 87°, 88°, 89-90°. Crystallise it from EtOAc. [*Beilstein* 5 H 270.]

1-Iodo-4-nitrobenzene [636-98-6] $C_6H_4INO_2$, M 249.0, m 171-172°, 173°, b 289°/772mm. Precipitate it from acetone by addition of water, followed by recrystallisation from EtOH. [*Beilstein* 8 H 523, 8 H 523, 8 II 191, 8 III 623, 8 IV 743.]

2-Iodophenol [533-58-4] C_6H_5IO , M 280.0, m 37° to 43°, b 186-187°/160mm, d^{25} 1.947, pK^{25} 8.51. Crystallise 2-iodophenol from $CHCl_3$ or diethyl ether. The **acetate** has m 65-66° (from MeOH). [*Beilstein* 6 H 208, 6 II 198, 6 III 774, 6 IV 1074.]

4-Iodophenol [540-38-5] has m 92° to 94°, b 138-140°/5mm, pK^{25} 9.30. Crystallise 4-iodophenol from petroleum ether (b 80-100°) or distil it *in vacuo*. If the material has a brown or violet color, then dissolve it in $CHCl_3$, shake it with 5% sodium thiosulfate solution until colourless. Dry (Na_2SO_4) the organic layer, filter, evaporate and distil the residue *in vacuo*. [Dains & Eberly *Org Synth Coll Vol 2* 355 1943, DOI: 10.15227/orgsyn.015.0039; *Beilstein* 6 IV 1074.]

5-Iodosalicylic acid (2-hydroxy-5-iodobenzoic acid) [119-30-2] $C_7H_5IO_3$, M 264.0, m 189° to 191°, 194°, 197°, 196-200°, pK_1^{25} 2.65, pK_2^{25} 13.05. Crystallise the acid from water. [*Beilstein* 10 H 112.]

2-Iodosobenzoic acid [304-91-6] $C_7H_5IO_3$, M 264.0, m >200°, 230°(dec), pK_{Est} ~2.6. Crystallise the acid from EtOH and dry it *in vacuo*. Store at -0°. [*Beilstein* 9 H 363.]

4-Iodotoluene [624-31-7] C_7H_7I , M 218.0, m 33-34°, 35°, b 211-212°/atm. Crystallise 4-iodotoluene from EtOH and/or distil it. [*Beilstein* 5 IV 840.]

Isonitrosoacetophenone (phenylglyoxaldoxime) [532-54-7] $C_8H_7NO_2$, M 149.2, m 126-128°. Crystallise it from water. It is used for detecting Fe^{2+} which gives a blue $CHCl_3$ soluble complex. [*Beilstein* 7 IV 2132.]

Isophthalic acid (benzene-1,3-dicarboxylic acid) [121-91-5] $C_8H_6O_4$, M 166.1, m 341° to 343°, 345-348°, pK_1^{25} 3.70, pK_2^{25} 4.60. Crystallise the acid from aqueous EtOH, or H_2O (with solubilities of 0.013w/v% at ~20° and 0.22w/v% at 100°). It sublimes undecomposed. [*Beilstein* 9 IV 3292.]

4,4'-Isopropylidenediphenol (Bisphenol A, 2,2-bis[4-hydroxyphenyl]propane) [80-05-7] $C_{15}H_{16}O_2$, M 228.3, m 158-159°, b 220°/4mm, pK_{Est} ~10.3. Crystallise bisphenol from acetic acid/water (1:1). It decomposes above 220° even in a vacuum. It is used for making polycarbonate bottles and leaches out slowly on heating. It is a known 'estrogenic chemical' shown to disrupt chemical signaling in the complex network of glands, hormones and cell receptors which make up the endocrine system. It causes low sperm count and damages the ecosystem by the feminisation of fish, reptiles and birds. [See **Bisphenol A** in SOURCES OF IMPURITIES in Chapter 1; *Beilstein* 6 IV 6717.]

Isopropyl p-nitrobenzoate [13756-40-6] $C_{10}H_{11}NO_4$, M 209.2, m 105-106°, 108-110°. Dissolve it in Et_2O ,

wash it with aqueous alkali, then H₂O and dry it. Evaporate the ether and recrystallise it from EtOH to give pure material. It gives yellow crystals from petroleum ether (**m** 109°, 110°). [*Beilstein* 9 H 391, 9 II 258, 9 III 1543, 9 IV 1075.]

Isovanillin (3-hydroxy-4-methoxybenzaldehyde) [621-59-0] C₈H₈O₃, **M 152.2**, **m** 113-114°, 116°, 117°, **b** 175°/14mm, 342.9°/760mm, **pK²⁵ 8.89**. Crystallise isovaniline from H₂O or *C₆H₆. The *oxime* has **m** 147°. [*Beilstein* 8 IV 1764.] It is a selective inhibitor of aldehyde oxidase.

Isoviolanthrone [128-64-3] C₃₄H₁₄O₂, **M 456.5**, **m** 510-511°(uncorrected). Dissolve isoviolanthrone in 98% H₂SO₄ and precipitate it by adding water to reduce the acid concentration to about 90%. It sublimes *in vacuo* to give dark green-violet needles [Parkyns & Ubbelohde *J Chem Soc* 4188 1960, DOI: 10.1039/JR9600004188]. [*Beilstein* 7 I 465, 7 II 815, 7 III 4538, 7 IV 2747.]

Janus Red B {3-[(2-hydroxy-1-naphthol)azo-2-methylphenylazo]N,N,N-trimethyl-benzenaminium chloride} [2636-31-9] C₂₆H₂₆ClN₅O, **M 460.0**. Crystallise the dye from EtOH/H₂O (1:1 v/v) and dry in vacuum. Store it in a dark bottle. [*Beilstein* 16 II 149.]

Ketone moschus (4-tert-butyl-2,6-dimethyl-3,5-dinitroacetophenone, Musk ketone) [81-14-1] C₁₄H₁₈N₂O₅, **M 294.3**, **m** 134-137°, 137-138°. Purify the ketone by recrystallisation from MeOH. It has a strong odour of musk and is used in perfumery. [Fuson et al. *J Org Chem* 12 587 1947, DOI: 10.1021/jo01168a016; *Beilstein* 7 IV 808.]

Lapachol [2-hydroxy-3-(3-methyl-2-butenyl)-1,4-naphthalenedione, Neutral Yellow 16] [84-79-7] C₁₅H₁₄O₃, **M 242.3**, **CI 7549**, **m** 141-143°, 140°. Crystallise Lapachol from petroleum ether/EtOH, EtOH or Et₂O. [*Beilstein* 8 H 326, 8 I 644, 8 II 365, 8 III 2720.]

Leucomalachite Green C₂₃H₂₆N₂, [129-73-7] **M 330.5**, **m** 92-93°, 100-102°, **pK²⁵ 6.90** (several pK's). Crystallise it from 95% EtOH (10ml/g), then from *benzene/EtOH, and finally from petroleum ether. Its UV-VIS has λ_{max} at 623nm. [*Beilstein* 13 H 275, 13 II 89, 13 II 135, 13 III 529, 13 IV 481.]

Malachite Green (carbinol) [510-13-4] C₂₃H₂₆N₂O, **M 346.4**, **m** 112-114°, **CI 42000**, **pK²⁴ 6.84**. The *oxalate* [2437-29-8] [*Beilstein* 8 H 326, 13 IV 2279.] is recrystallised from hot water and dried in air. The carbinol is precipitated from the oxalate (1g) in distilled water (100ml) by adding M NaOH (10ml). The precipitate is filtered off, recrystallised from 95% EtOH containing a little dissolved KOH, then washed with ether, and crystallised from petroleum ether. Dry it in a vacuum at 40°. An acid, almost colourless, solution (2 x 10⁻⁵M in 6 x 10⁻⁵M H₂SO₄) rapidly reverts to the coloured dye. [Swain & Hedberg *J Am Chem Soc* 72 3373 1950, DOI: 10.1021/ja01164a015; *Beilstein* 13 H 243, 744.]

Mandelic acid (α-hydroxyphenylacetic acid) [S-(+)- 17199-29-0, R-(-)- 611-71-2] C₈H₈O₃, **M 152.2**, **m** 130-133°, 133°, 133.1° (evacuated capillary), 133-133.5°, [α]_D²⁰ (+) and (-) 188 (c 5, H₂O), [α]_D²⁰ (+) and (-) 155 (c 5, H₂O) and (+) and (-) 158 (c 5, Me₂CO), **pK²⁵ 3.41**. Purify the mandelic acids by recrystallisation from H₂O, *C₆H₆ or CHCl₃. [Roger *J Chem Soc* 2168 1932, DOI: 10.1039/JR9320002168; Jamison & Turner *J Chem Soc* 611 1942, DOI: 10.1039/JR9420000611.] They have solubilities in H₂O of ca 11% at 25°. [Banks & Davies *J Chem Soc* 73 1938, DOI: 10.1039/JR9420000611.] The *S-benzylisothiuronium salt* has **m** 180° (from H₂O) and [α]_D²⁵ (+) and (-) 57 (c 20, EtOH) [El Masri et al. *Biochem J* 68 199 1958, DOI: 10.1042/bj0680199]. [*Beilstein* 10 IV 564.] **Methyl R-(-)-mandelate** [20698-91-3] C₉H₁₀O₃, **M 166.2**, **b** 85°/1.5mm, **m** 56-58°, 57-58°, is prepared by dissolving R-(-)-mandelic acid (10g, 0.066mol) and excess MeOH (5.3ml, 132mol) in *C₆H₆ (50ml) containing concentrated H₂SO₄ (0.1ml) and refluxed under a Dean-Stark trap while H₂O is removed azeotropically (~3 hours). The mixture is cooled, washed with aqueous saturated NaHCO₃ (3 x 50ml) and the

aqueous phase is extracted with Et₂O (3 x 50ml), the organic layers are combined, dried (MgSO₄), filtered, evaporated to dryness and the residual yellow oil is distilled in a vacuum. The distillate solidified and is recrystallised from low boiling petroleum ether to give white crystalline (-)-*ester* (9.3g, 85%) with the above physical properties. [Toniolo et al. *J Org Chem* 35 6 1970, DOI: 10.1021/jo00826a002]

RS-(±)-Mandelic acid [61-72-3] has **m 118°, 120-121°**. Purify mandelic acid by Soxhlet extraction with *C₆H₆ (about 6ml/g) and allow the extract to crystallise. It can also be recrystallised from CHCl₃. The **S-benzylisothiuronium salt** has **m 169° (166°)** (from H₂O). Dry it at room temperature *in vacuo* [Beilstein 10 IV 565.]

Mescaline sulfate [2-(3,4,5-trimethoxyphenyl)ethylamine sulfate] [642-73-9, 1152-765967-42-0] **C₁₁H₁₉NO₇S**, **M 309.3**, **m 181-184°, 183-184°, 186-189°, pK_{Est} ~9.7**. The salt crystallises from water with 2H₂O or from hot MeOH. The **acid sulfate** has **m 158°**. [Salomon & Bina *J Am Chem Soc* 68 2403 1946, DOI: 10.1021/ja01215a509; Beilstein 13 I 338, 13 II 521, 13 III 2375, 13 IV 2919.] It is a hallucinogen

Mesitylene (1,3,5-trimethylbenzene) [108-67-8] **C₉H₁₂**, **M 120.2**, **m -44.7°, b 61°/20mm, 99.0-99.8°/100mm, 166.5-167°/760mm, d₄²⁰ 0.865, n_D²⁵ 1.4967**. Dry it with CaCl₂ and distil it from Na in a glass helices-packed column. Treat it with silica gel and redistil it. *Alternative* purifications include vapour-phase chromatography, or fractional distillation followed by azeotropic distillation with 2-methoxyethanol (which is subsequently washed out with H₂O), drying and fractional distilling. More exhaustive purification uses sulfonation by dissolving in two volumes of concentrated H₂SO₄, precipitating with four volumes of concentrated HCl at 0°, washing with concentrated HCl and recrystallising from CHCl₃. The mesitylene sulfonic acid is hydrolysed with boiling 20% HCl and steam distilled. The separated mesitylene is dried (MgSO₄ or CaSO₄) and distilled. It can also be fractionally crystallised from the melt at low temperatures. [Adams & Hufferd *Org Synth Coll Vol* 1 341 1941, DOI: 10.15227/orgsyn.002.0041; Beilstein 5 IV 1016.] **IRRITANT and TOXIC.**

4'-Methoxyacetophenone [100-06-1] **C₉H₁₀O₂**, **M 150.2**, **m 36° to 38°, 39°, b 139°/15mm, 152-154°/26mm, 264°/736mm**. Crystallise the ketone from diethyl ether/petroleum ether, and/or distil it under reduced pressure. The **oxime** has **m 86°, 87°** (from petroleum ether, v. Auwers et al. *Chem Ber* 58 36 1925, DOI: 10.1002/cber.19250580109). [Beilstein 8 H 87, 8 I 536, 8 II 84, 8 III 277, 8 IV 340.]

4-Methoxyazobenzene [2396-60-3] **C₁₃H₁₂N₂O**, **M 212.3**, **m 54-56°, 340°/atm**. Crystallise 4-methoxyazobenzene from EtOH. [Beilstein 16 IV 162.]

3-Methoxybenzanthrone [3688-79-7] **C₁₈H₁₂O₂**, **M 274.3**, **m 173°**. Crystallise it from *benzene, EtOH or Me₂CO to give yellow needles. It has light fastening, photo tendering and fluorescence properties [Allen et al. *J Appl Chem & Biotechnology* 27 269 1977, DOI: 10.1002/jctb.5020270140]. [Beilstein 8 II 239, 8 III 1629, 8 IV 1476.]

3-Methoxybenzoic acid (m-anisic acid) [586-38-9] **C₈H₈O₃**, **M 152.2**, **m 105° to 107°, 110°, b 170-172°/10mm, pK₂₅ 4.09**. Crystallise *m*-anisic acid from H₂O (**m 109°, 110.5°**) or EtOH/water. The **S-benzylisothiuronium salt** has **m 176°** (from EtOH). [Beilstein 10 II 80, 10 III 244, 10 IV 316.]

4-Methoxybenzoic acid (p-anisic acid) [100-09-4] has **m 182° to 185°, 184.0-184.5°, b 276-280°/atm, pK₂₅ 4.51**. Crystallise *p*-anisic acid from EtOH, water, EtOH/water or toluene. It sublimes above its melting point. The **S-benzylisothiuronium salt** has **m 189°** (from EtOH). [Beilstein 10 II 91, 10 III 280, 10 IV 346.] It is a tyrosinase inhibitor.

4-Methoxybenzyl chloride (anisyl chloride) [824-94-2] **C₈H₉Cl₃O**, **M 156.6**, **m -1°, b 76°/0.1mm, 95°/5mm, 110°/10mm, 117-117.5°/14mm, 117°/18mm, d₄²⁰ 1.15491, n_D²⁰ 1.55478**. Purify 4-anisyl chloride by fractional distillation under vacuum, and the middle fraction is redistilled at 10⁻⁶ mm at room temperature by intermittent cooling of the receiver in liquid N₂, and the middle fraction is collected. [Kosower & Mohammed *J Am Chem Soc* 93 2709 1971, DOI: 10.1021/ja00740a021; Beilstein 6 IV 2137.] Useful O- and N- protective group reagent, may contain some K₂CO₃ as stabiliser. **LACHRYMATORY.**

'Methoxychlor' (1,1-bis[*p*-methoxyphenyl]-2,2,2-trichloroethane, DMDT) [72-43-5] **C₁₆H₁₅Cl₃O₂**, **M**

345.7, m 78-78.2°, or 86-88°. Free the insecticide from 1,1-bis(*p*-chlorophenyl)-2,2,2-trichloroethane by crystallising from EtOH. It is dimorphic and also crystallises from Et₂O/EtOH (**m 92°**). [Fritsch & Feldmann *Justus Liebigs Ann Chem* **306** 72 1899, DOI: 10.1002/jlac.18993060106; Smith et al. *Aust J Chem* **29** 743 1976, DOI: 10.1071/CH9760743; *Beilstein* **6** H 1007.]

***trans-p*-Methoxycinnamic acid** [830-09-1, 943-89-5 (*trans*)] C₁₀H₁₀O₃, **M 178.2, m 170° to 173°, 173.4-174.8°, pK²⁵ 4.54.** Crystallise the acid from MeOH to constant melting point and UV spectrum. It forms liquid crystals with a crystalline to nematic phase change at 173.5°, and a nematic to isotropic phase at 190°. [*Beilstein* **10** H 928, **10** IV 1005.]

6-Methoxy-1-indanone [13623-25-1] C₁₀H₁₀O₂, **M 162.2, m 103°, 104°, 105-109°, 107-111°, b 291.7°/760mm, d 1.166.** Crystallise the yellow powder from MeOH, then sublime it at high vacuum. [*Beilstein* **8** IV 894.]

1-Methoxy-4-nitronaphthalene [4900-63-4] C₁₁H₉NO₃, **M 203.2, m 83-84°, 85°.** Purify it by chromatography on silica gel, then recrystallise it from MeOH or EtOH (yellow needles). [Hodgson & Habeshaw *J Chem Soc* 47 1942, DOI: 10.1039/JR9420000045; Bunce et al. *J Org Chem* **52** 4214 1987, DOI: 10.1021/jo00228a012; *Beilstein* **6** H 616, **6** III 2938.]

4-Methoxyphenol [150-76-5] C₇H₈O₂, **M 124.1, m 54-55°, 55-57°, b 243°/atm, pK²⁵ 10.21.** Crystallise 4-methoxyphenol from *benzene, petroleum ether or H₂O, and dry it under vacuum over P₂O₅ at room temperature. Sublime it *in vacuo*. [Wolfenden et al. *J Am Chem Soc* **109** 463 1987, DOI: 10.1021/ja00236a026; *Beilstein* **6** IV 5717.]

α-Methoxyphenylacetic acid (*O*-methyl mandelic acid) [*R*-(-)- 3966-32-3, *S*-(+)- 26164-26-1] C₉H₁₀O₃, **M 166.2, m 62.9°, 62-65°, 65-66°, [α]_D²⁰ (-) and (+) 179 (169.8°), [α]_D²⁰ (-) and (+) 150.7 (148) (c 0.5, EtOH), pK_{Est} ~3.1.** Purify the acids by recrystallising from *C₆H₆/petroleum ether (**b 80-100°**). [Neilson & Peters *J Chem Soc* 1519 1962, Weizmann et al. *J Am Chem Soc* **70** 1153 1948, DOI: 10.1021/ja01183a082; Pirie & Smith *J Chem Soc* 338 1932, DOI: 10.1039/JR9320000337; NMR: Dale & Mosher *J Am Chem Soc* **95** 512 1973, DOI: 10.1021/ja01183a082; for resolution see Roy & Deslongchamps *Can J Chem* **63** 651 1985, DOI: 10.1139/v85-106; Trost et al. *J Am Chem Soc* **108** 4974 1986, DOI: 10.1021/ja00276a044.] The **racemic mixture** [7021-09-2] has **m 72°, b 121-122°/0.4mm, 165°/18mm** (from petroleum ether) [Braun et al. *Chem Ber* **63** 2847 1930, DOI: 10.1002/cber.19300631027]. [*Beilstein* **10** IV 566.]

2-Methoxyphenylacetic acid [93-25-4] C₉H₁₀O₃, **M 166.2, m 124-125°, pK_{Est} ~4.4.** Crystallise the acid from H₂O, EtOH or aqueous EtOH, petroleum ether/Et₂O and dry it in a vacuum desiccator over Sicapent. The **amide** has **m 131°** (from EtOH). [*Beilstein* **10** H 188, **10** III 422, **10** IV 536.]

3-Methoxyphenylacetic acid [1798-09-0] has **m 66-67°, 68-69°, 71.0-71.2°, pK_{Est} ~4.3.** Crystallise the acid from H₂O, or aqueous EtOH. The ***S*-benzylisothiuronium salt** has **m 160-161°** (from EtOH). [*Beilstein* **10** I 82, **10** III 428, **10** IV 541.]

4-Methoxyphenylacetic acid (homoanisic acid) [104-01-8] has **m 85-87°, b 138-140°/2-3mm, pK²⁵ 4.36.** Crystallise the acid from EtOH/water, EtOAc/petroleum ether (**m 87°**) or *C₆H₆/petroleum ether (**m 84-86°**). [*Beilstein* **10** III 431, **10** IV 544.] The **acid chloride** [4693-91-8] has **M 184.6, b 143°/10mm, d₄²⁵ 1.208.** [*Beilstein* **10** III 434.]

***N*-(*p*-Methoxyphenyl)-*p*-phenylenediamine** [101-64-4] C₁₃H₁₄N₂O, **M 214.3, m 102°, b 200°/2mm, 238°/12mm, pK²⁵ 6.6 (5.9).** Crystallise the diamine from ligroin or *C₆H₆/petroleum ether (**m 99-100°**). The **picrate** has **m 164°** (from EtOH). [*Beilstein* **13** III 1161, **13** IV 1243.]

α-Methoxy-α-trifluoromethylphenylacetic acid (MTPA, Mosher's acid) [*R*-(+)- 20445-31-2, *S*-(-)- 17257-71-5] C₁₀H₉F₃O₃, **M 234.2, m 43-45°, 95-97°/0.05mm, 90°/0.1mm, 105-107°/1mm, [α]_D²⁰ (+) and (-) 87, [α]_D²⁰ (+) and (-) 73 (c 2, MeOH), pK_{Est} ~2.5.** A likely impurity is phenylethylamine from the resolution. Dissolve the acid in Et₂O/*benzene (3:1), wash with 0.5N H₂SO₄, then water, dry over magnesium sulfate, filter, evaporate and distil it. Store in a sealed container under Ar or N₂ in a cool place. [Dale et al. *J Org Chem*

34 2543 1969, DOI: 10.1021/jo01261a013, *J Am Chem Soc* **95** 512 1973, DOI: 10.1021/ja00783a034.] The acid, by conversion to the acid chloride, see below) is a useful derivatising reagent for determination of enantiomeric excess of chiral alcohols and amines by NMR [George & Sudalai *Tetrahedron Asymmetry* **18** 975 2007, DOI: 10.1016/j.tetasy.2007.04.008; De Luca et al. *J Org Chem* **72** 3955 2007, DOI: 10.1021/jo070142c]. The (\pm)-acid [81655-41-6] $\text{C}_{10}\text{H}_9\text{F}_3\text{O}_3$, **M 234.2**, has **m 41-46°** (from aqueous MeOH) and **b 95-98°/0.05mm**.

α -Methoxy- α -trifluoromethylphenylacetyl chloride [*R*(-)- 39637-99-5, *S*(+)- 20445-33-4] $\text{C}_{10}\text{H}_8\text{ClF}_3\text{O}_2$, **M 252.6**, **b 54-56°/1mm**, **213-214°/760mm**, **d** $^{20}_4$ **1.353**, **n** $^{20}_D$ **1.468**, **[\alpha]** $^{20}_{546}$ **(-) and (+) 167**, **[\alpha]** $^{20}_D$ **(-) and (+) 137 (c 4, CCl₄)**, **[\alpha]** $^{24}_D$ **(-) and (+) 10.0 (neat)**. The most likely impurity is the free acid due to hydrolysis and should be checked by IR. If free from acid, then distil, taking care to keep moisture out of the apparatus. Otherwise add SOCl_2 and reflux for 5 hours then distil it. *Note* that shorter reflux times result in a higher boiling fraction (**b 130-155°/1mm**) which has been identified as the anhydride. Store under argon or N_2 in a sealed container (in aliquots) at -20° . [Dale et al. *J Org Chem* **34** 2543 1969, DOI: 10.1021/jo01261a013 for enantiomeric purity see Dale & Mosher *J Am Chem Soc* **95** 512 1973, DOI: 10.1021/ja00783a034.] See previous entry for use in determining enantiomeric excess of chiral alcohols, thiols and amines.

***N*-Methylacetanilide** [579-10-2] $\text{C}_9\text{H}_{11}\text{NO}$, **M 149.2**, **m 101°**, **102-104°**, **b 145-146°/30mm**, **225°/760mm**. Crystallise the anilide from water, ether, petroleum ether (b 80-100°) or C_6H_6 (**m 105°**). The BF_3 salt has **m 114°** (from Me_2CO /diisopropyl ether). [Beilstein **12** III 465, **12** IV 378.]

4-Methylacetophenone [122-00-9] $\text{C}_9\text{H}_{10}\text{O}$, **M 134.2**, **m 22-24°**, **b 93.5°/7mm**, **110°/14mm**, **d** $^{20}_4$ **1.000**, **n** $^{20}_D$ **1.5335**. Impurities, including the *o*- and *m*-isomers, are removed by forming the *semicarbazone* (**m 212-213.5°**) which, after repeated crystallisation, is hydrolysed to the ketone. [Brown & Marino *J Am Chem Soc* **84** 1236 1962, DOI: 10.1021/ja00866a034.] It can also be purified by distillation under reduced pressure, followed by low temperature crystallisation from isopentane. The *oxime* has **m 87-88°** (from MeOH then cyclohexane Pearson & Ball *J Org Chem* **14** 125 1949, DOI: 10.1021/jo01153a018). [Beilstein **7** H 307, **7** I 154, **7** II 238, **7** III 1060, **7** IV 701.]

1-Methylaminoanthraquinone [82-38-2] $\text{C}_{15}\text{H}_{11}\text{NO}_2$, **M 237.3**, **m 166.5°**, **pK_{Est} ~2**. Crystallise it to constant melting point from butan-1-ol, then from EtOH. It can be sublimed under vacuum. [Beilstein **7** IV 2574.]

***N*-Methyl-*o*-aminobenzoic acid** (*N*-methylantranilic acid) [119-68-6, 730910-17-5] $\text{C}_8\text{H}_9\text{NO}_2$, **M 151.2**, **m 178.5°**, **pK₁²⁵ 1.97**, **pK₂²⁵ 5.34**. Crystallise the red powder from water, MeOH (**m 177°**) or EtOH. [Cosulich & Smith *J Am Chem Soc* **70** 1923 1948, DOI: 10.1021/ja01185a081; Beilstein **14** H 323, **14** III 895, **14** IV 1015.]

3-(Methylamino)benzoic acid [51524-84-6] $\text{C}_8\text{H}_9\text{NO}_2$, **M 151.1**, **m 129°**, **pK₁²⁵ 3.08**, **pK₂²⁵ 5.10**. Crystallise the acid from H_2O (needles), petroleum ether (plates) or CHCl_3 (**m 127°**). The *hydrochloride* has **m 244°** (plates from EtOH). [Beilstein **14** H 391, **14** I 559.]

4-Methylaminophenol hemisulfate (Photol, Metol) [55-55-0] $\text{C}_7\text{H}_{11}\text{NO}_5\text{S}$, **M 344.4**, **m 260°(dec)**, **261-263°(dec)**, **pK₂₅ 5.9**. Crystallise this photographic developer from H_2O (needles **m 250-260°**) (its solubility is 5% at 20° and 17% at 100°) or MeOH. It is used for determining Ag. [Palit et al. *Indian J Chem, Sect B* **28** 64 1989.]

***N*-Methylaniline** [100-61-8] $\text{C}_7\text{H}_9\text{N}$, **M 107.2**, **b 57°/4mm**, **81-82°/14mm**, **d** $^{20}_4$ **0.985**, **n** $^{20}_D$ **1.570**, **pK₂₅ 4.56**. Dry it with KOH pellets and fractionally distil it under vacuum. Acetylate, and the *acetyl* derivative is recrystallised to constant melting point (**m 101-102°**), then hydrolysed with aqueous HCl and distilled from zinc dust under reduced pressure. [Hammond & Parks *J Am Chem Soc* **77** 340 1955, DOI: 10.1021/ja01607a029; Beilstein **12** IV 241.] ***N*-Methylaniline hydrochloride** [2739-12-0] $\text{C}_7\text{H}_9\text{N.HCl}$, **M 143.7**, has **m 123.0-123.1°**. Crystallise the salt from dry $\text{C}_6\text{H}_6/\text{CHCl}_3$ and dry under vacuum. [Beilstein **12** IV 241.]

Methyl *p*-anisate [121-98-2] $\text{C}_9\text{H}_{10}\text{O}_3$, **M 166.2**, **m 48°, 49° to 51°**, **b 123.8°/12mm**, **244-245°/760mm**. Distil the ester and/or crystallise it from EtOH. [Beilstein **10** H 159, **10** III 297, **10** IV 360.]

4-Methyl anisole [104-93-8, 3494-45-9] $C_8H_{10}O$, **M 122.2**, **b 56.2°/9mm**, **67°/20mm**, **175-176°/760mm**, **d**₁₅¹⁵ **0.9757**, **n**_D²⁰ **1.512**. Dissolve 4-methyl anisole in diethyl ether, wash it with M NaOH, water, dry (Na₂CO₃), evaporate and distil it under vacuum. The *picrate* has **m 103°** (from aqueous EtOH). [Beilstein 6 IV 2098.]

2-Methylantracene [613-12-7] $C_{15}H_{12}$, **M 192.3**, **m 204-206°**. Chromatograph it on silica gel with cyclohexane as eluent and then recrystallise it from EtOH [Werst et al. *J Am Chem Soc* **109** 32 1987, DOI: 10.1021/ja00235a005]. [Beilstein 5 IV 2311.]

9-Methylantracene [779-02-2] has **m 77-79°**, **b 196-197°/12mm**, **d**₄²⁴ **1.066**. Chromatograph it on silica gel with cyclohexane as eluent and recrystallise it from EtOH [Werst et al. *J Am Chem Soc* **109** 32 1987, DOI: 10.1021/ja00235a005]. [Beilstein 5 IV 2312.]

2-Methylantraquinone [84-54-8] $C_{15}H_{10}O_2$, **M 222.3**, **m 170° to 173°**, **176°**, **b 236-238°/10mm**. Crystallise the quinone from EtOH, then sublime it. It has λ_{max} at 257, 275 and 330nm (EtOH). [Hersberg & Fieser *J Am Chem Soc* **63** 2562 1941, DOI: 10.1021/ja01855a005; Beilstein 7 H 809, 7 III 4104, 7 IV 2574.]

Methylarenes (see also pentamethyl- and hexamethyl- benzenes). Recrystallise them from EtOH and sublime them in a vacuum [Schlesener et al. *J Am Chem Soc* **106** 7472 1984, DOI: 10.1021/ja00336a029].

Methyl benzoate [93-58-3] $C_8H_8O_2$, **M 136.2**, **m -12°**, **b 104-105°/39mm**, **199.5°/760mm**, **d**₄²⁰ **1.087**, **n**_D¹⁵ **1.52049**, **n**_D²⁴ **1.51701**, **pK**²⁰ **-8.11**, **-6.51** (H_0 scale, aqueous H₂SO₄). Wash the ester with dilute aqueous NaHCO₃, then water, dry with Na₂SO₄ and fractionally distil it in a vacuum. [Beilstein 9 IV 283.]

4-Methylbenzophenone [134-84-9] $C_{14}H_{12}O$, **M 196.3**, **m 56.-57°**, **57°**, **b 154-155°/3mm**, **183-184°/15mm**, **277-281°/3mm**, **d**₂₀²⁰ **0.9926**. Crystallise the ketone from MeOH, Et₂O (**m 58-59°**) or petroleum ether. The *cis-oxime* has **m 154°(153-156°)** (from EtOH), and the *trans-oxime* has **m 114-116°** (from petroleum ether). [Beilstein 7 H 440, 7 III 2127, 7 IV 1403.]

Methyl-1,4-benzoquinone (*p*-toluoquinone) [553-97-9] $C_7H_6O_2$, **M 122.1**, **m 66-67°**, **68-69°**. Crystallise *p*-toluoquinone from heptane or EtOH, dry rapidly (vacuum/P₂O₅) and store it in a vacuum. [Beilstein 7 IV 2088.]

Methyl benzoylformate (methyl phenylglyoxalate) [15206-55-0] $C_9H_8O_3$, **M 164.2**, **m 246-248°**, **d**_D²⁵ **1.155**, **n**_D²⁰ **1.526**. Purify the ester by radial chromatography (diethyl ether/hexane, 1:1), and dry it at 110-112°/6mm. [Meyers & Oppenlaender *J Am Chem Soc* **108** 1989 1986, DOI: 10.1021/ja00268a043; Beilstein 10 IV 2738.]

2-Methyl-3,4-benzphenanthrene (2-methylbenzo[*c*]phenanthrene) [652-04-0] $C_{19}H_{14}$, **M 242.3**, **m 70°**, **80.6-81.4°**, **b 200°/0.4mm**. Crystallise it from EtOH (**m 81-82.5°**). The *picrate* has **m 118-118.5°** (yellow needles from MeOH). [Mukerji & Rao *Nature* **168** 1041 1951, DOI:10.1038/1681041a0; Beilstein 5 III 2394, 5 IV 2570.] **POTENT CARCINOGEN**.

R-(+)- α -Methylbenzylamine [*R*(+) 3886-69-9, *RS*(\pm) 618-36-0] $C_8H_{11}N$, **M 121.2**, **b 187-188°/atm**, [α]₅₄₆²⁰ **+35** (*c* 10, EtOH), [α]_D²⁵ **+39.7** (neat), **pK 9.08** (for *RS*). Dissolve the amine in toluene, dry over NaOH and distil; fraction boiling at 187-188°/atm is collected. Store it under N₂ to avoid forming the carbamate and urea. Similarly for the *S*-(-) *enantiomer* [2627-86-3]. [For the optical resolution of phenethylamine see: Ingersoll *Org Synth Coll Vol* **2** 506 1943, DOI: 10.15227/orgsyn.017.0080; Beilstein 12 IV 2424, 2425.] The (\pm)-*racemic base* purified in the same manner has **b 185°/756mm**; and can be fractionated under reduced pressure, **b 80-81°/18mm**, **d**₄²⁵ **0.94**, **n**_D²⁰ **1.526**. [Synthesis of *dl*:- Ingersoll *Org Synth Coll Vol* **2** 503 1943, DOI: 10.15227/orgsyn.017.0076; Robinson & Snyder *Org Synth Coll Vol* **3** 717 1955, DOI: 10.15227/orgsyn.023.0068]. [Beilstein 12 IV 2452.]

4-Methylbenzyl chloride [104-82-5] C_8H_9Cl , **M 140.6**, **m 4-6°**, **b 80°/2mm**, **98-101°/27mm**, **199-201°/atm**, **d**₄²⁰ **1.085**, **n**_D²⁰ **1.543**. Dry the chloride with CaSO₄ and fractionally distil it under vacuum. [Beilstein 5 H 384, 5 III 854, 5 IV 966.] **Lachrymatory**, handle with care.

Methyl 2-bromobenzoate [610-94-6] $C_8H_7BrO_2$, **M 215.1**, **b 131.4-132°/16mm**, **234-244°/760mm**, **252°/**

atm, d_4^{24} 1.532, n_D^{20} 1.559. A solution of the ester in ether is washed with 10% aqueous Na_2CO_3 , water, then dried and distilled. [Beilstein 9 H 348, 9 III 1385, 9 IV 1012.]

Methyl 4-bromobenzoate [619-42-1] $\text{C}_8\text{H}_7\text{BrO}_2$, M 215.1, m 77° to 81°, 79.5-80.5°, d 1.689. Crystallise the ester from MeOH. EtOH (m 81°, also 80.5°, 79.5°) or C_6H_6 /petroleum ether (m 78-79°). [Beilstein 9 H 352, 9 III 1405, 9 IV 1017.]

(±)-3'-Methyl-1,2-cyclopentenophenanthrene (16,17-dihydro-17-methyl-15H-cyclopenta-[a]-phenanthrene [549-38-2] $\text{C}_{18}\text{H}_{16}$, M 232.3, m 126-127°. Crystallise it from AcOH or EtOH (plates, m 125.5-126°). The *picrate* has m 130-131° (orange-red needles from EtOH). [Tatta & Bardhan *J Chem Soc (C)* 893 1968, DOI: 10.1039/J39680000893; Beilstein 5 IV 2433.]

Methyl 2,4-dichlorophenoxyacetate [1928-38-7] $\text{C}_9\text{H}_8\text{Cl}_2\text{O}_3$, M 235.1, m 43°, b 119°/11mm. Crystallise the herbicide ester from MeOH. [Branch & Jones *J Chem Soc* 2921 1955, DOI: 10.1039/JR9550002921; Beilstein 6 III 705, 6 IV 909.]

3,4-Methylenedioxyaniline [14268-66-7] $\text{C}_7\text{H}_7\text{NO}_2$, M 137.1, m 39° to 41°, 44.5-45.5°, 45-46°, b 108°/1mm, 144°/14mm, 156°/30mm, $\text{pK}_{\text{Est}} \sim 3.8$. Crystallise the base from petroleum ether and/or distil it in a vacuum. The *hydrochloride* has m 198°(dec). [Sonn & Benirschke *Chem Ber* 54 1730 1921, DOI: 10.1002/cber.19210540808; Beilstein 19 H 328, 19 II 341, 19 III/IV 4056.]

trans-3,4-Methylenedioxycinnamic acid [2373-80-0] $\text{C}_{10}\text{H}_8\text{O}_4$, M 192.2, m 243-244°(dec), $\text{pK}_{\text{Est}} \sim 4.6$. Crystallise the acid from glacial AcOH, EtOH (m 247°) or aqueous EtOH (m 240-242°), and it has m 242° after sublimation. [Beilstein 19 H 278, 19 II 299, 19 III/IV 3548.]

5,5'-Methylenedisalicylic acid [122-25-8; 27496-82-2] $\text{C}_{15}\text{H}_{12}\text{O}_6$, M 288.2, m 238°(dec), 242°, 243.5°, b 569.6°/atm. Crystallise the acid from Me_2CO , C_6H_6 or $\text{CHCl}_3/\text{MeOH}$ (m 268-268°). It has λ_{max} at 312nm (ϵ 7530) in EtOH. [Cushman & Kanamathareddy *Tetrahedron* 46 1491 1990, DOI: 10.1016/S0040-4020(01)81957-8.]

N-Methylephedrine (2-dimethylamino-1-phenylpropanol) [1S,2R-(+)- 42151-56-4, 1R,2S-(-) 552-79-4] $\text{C}_{10}\text{H}_{15}\text{NO}_3$, M 179.3, m 85-86°, 87-87.5°, 90°, b 115°/2mm, $[\alpha]_{546}^{20}$ (+) and (-) 35, $[\alpha]_D^{20}$ (+) and (-) 30 (c 4.5, MeOH), pK^{26} 9.22. N-Methylephedrine has been recrystallised from Et_2O , petroleum ether, of aqueous EtOH or aqueous MeOH and has been distilled under reduced pressure. [Smith *J Chem Soc* 2056 1927, DOI: 10.1039/JR9270002056; Tanaka & Sugawa *Yakugaku Zasshi (J Pharm Soc Japan)* 72 1548 1952 (*Chem Abstr* 47 8682 1953), Takamatsu *Yakugaku Zasshi (J Pharm Soc Japan)* 76 1227 1956, *Chem Abstr* 51 4304 1957.] The *hydrochloride* has m 192-193° and $[\alpha]_D^{20} + 30^\circ$ (c 5, H_2O) [Prelog & Häfliger *Chim Acta* 33 2021 1950, DOI: 10.1002/hlca.19500330708]. [Beilstein 13 IV 1884.]

Methyl gallate [99-24-1] $\text{C}_8\text{H}_8\text{O}_5$, M 184.2, m 201°, 202°, 203°, 198-203°, b 450°/atm. Crystallise the gallate ester from MeOH, or aqueous MeOH, usually solvated but can be readily de-solvated. [Beilstein 10 IV 1998.]

Methyl Green [82-94-0, 7114-03-6 (ZnCl_2 salt)] $\text{C}_{27}\text{H}_{35}\text{BrClN}_3 \cdot \text{ZnCl}_2$, M 517.0 (without Zn), 653.2, m >200°(dec), >300°(dec). Crystallise the dye (violet or green crystals) from hot water. [Beilstein 13 IV 2286.] It is an indicator which is yellow at pH 0.2 and blue at 1.8. It is a biological stain, e.g. for DNA and amyloid.

Methyl 4-hydroxybenzoate (methyl paraben) [99-76-3] $\text{C}_8\text{H}_8\text{O}_3$, M 152.2, m 125°, 127.5°, 185°, b 270°/atm with dec, $\text{pK}_{\text{Est}} \sim 9.3$. Fractionally crystallise the ester from its melt, and recrystallise it from C_6H_6 , then from $\text{C}_6\text{H}_6/\text{MeOH}$ and dry it over CaCl_2 in a vacuum desiccator. Its solubility in H_2O at 25° is $\sim 0.3\text{w/v}\%$. It is a food additive. [Beilstein 10 IV 360.]

Methyl 3-hydroxy-2-naphthoate [883-99-8] $\text{C}_{12}\text{H}_{10}\text{O}_3$, M 202.2, m 73-74°, 76° to 78°, b 205-207°/atm, $\text{pK}_{\text{Est}} \sim 9.0$. Crystallise the ester from MeOH (charcoal) containing a little water. [Beilstein 10 IV 1186.]

3-Methylmercaptoaniline [1783-81-9] C_7H_9NS , M 139.2, b 101.5-102.5°/0.3mm, 163-165°/16mm, d_4^{20} 1.147, n_D^{20} 1.641, pK^{25} 4.05. Purify the aniline by fractional distillation in an inert atmosphere. It has UV max at 226 and 300nm. [Bordwell & Cooper *J Am Chem Soc* 74 1058 1952, DOI: 10.1021/ja01124a057.] The *N*-acetyl derivative has m 78-78.5° (from aqueous EtOH). The *hydrochloride* has m 260-261° (aqueous EtOH/HCl) or m 225-227° (EtOH/Et₂O). [Beilstein 13 H 533, 13 III 1221, 13 IV 1289.]

4-Methylmercaptoaniline [104-96-1] has b 140°/15mm, 151°/25mm, 155°/23mm, d_4^{20} 1.137, n_D^{20} 1.639, pK^{25} 4.40. Purify the aniline by fractional distillation in an inert atmosphere. The *hydrochloride* has m 242-246° (from aqueous EtOH/HCl). The *sulfone* has m 137° (from H₂O), pK^{25} 1.48, and the *sulfone hydrochloride* has m 260-261° (from aqueous EtOH/HCl). [Lumbroso & Passerini *Bull Soc Chim Fr* 311 1957, Mangini & Passerini *J Chem Soc* 4954 1956, DOI: 10.1039/JR9560004954; Beilstein 13 H 533, 13 II 297, 13 IV 1221.]

1-Methylnaphthalene [90-12-0] $C_{11}H_{10}$, M 142.2, m -22°, -30°, b 244.6°/atm. d_4^{20} 1.021, n_D^{20} 1.6108. Dry 1-methylnaphthalene for several days with CaCl₂ or by prolonged refluxing with BaO. Fractionally distil it through a glass helices-packed column from sodium. Purify it further by solution in MeOH and precipitation of its *picrate complex* by adding to a saturated solution of picric acid in MeOH. The picrate, after crystallisation to constant melting point (m 140-141°) from MeOH, is dissolved in *benzene and extracted with aqueous 10% LiOH until the extract is colourless. Evaporation of the *benzene solution under vacuum gives 1-methylnaphthalene [Kloetzel & Herzog *J Am Chem Soc* 72 1991 1950, DOI: 10.1021/ja01161a032]. However, neither the picrate nor the styphnate complexes satisfactorily separate 1- and 2- methylnaphthalenes. To achieve this, 2-methylnaphthalene (10.7g) in 95% EtOH (50ml) has been precipitated with 1,3,5-trinitrobenzene (7.8g) and this complex has been crystallised from MeOH to m 153-153.5° (m of the 2-methyl isomer is 124°). [Alternatively, 2,4,7-trinitrofluorenone in hot glacial acetic acid could be used, and the derivative (m 163-164°) is recrystallised from glacial acetic acid]. The 1-methylnaphthalene is regenerated by passing a solution of the complex in dry *benzene through a 15-in column of activated alumina and washing with *benzene/petroleum ether (b 35-60°) until the coloured band of the nitro compound had moved down near the end of the column. The complex can also be decomposed using tin and acetic-hydrochloric acids, followed by extraction with diethyl ether and *benzene; the extracts are washed successively with dilute HCl, strongly alkaline sodium hypophosphite, water, dilute HCl and water. [Soffer & Stewart *J Am Chem Soc* 74 567 1952, DOI: 10.1021/ja01122a518.] It can be freed from anthracene by zone melting [Beilstein 5 IV 1687.]

2-Methylnaphthalene [91-57-6] has m 34.7-34.9°, 34° to 36°, b 129-130°/25mm, 141-142°/atm, d^{25} 1.00. Fractionally crystallise repeatedly from its melt, then fractionally distil under reduced pressure. It has been crystallised from *benzene and dried under vacuum in an Abderhalden pistol. It can be purified *via its picrate* (m 114-115°) or better *via the 1,3,5-trinitrobenzene complex* as for 1-methylnaphthalene (above). [Beilstein 5 IV 1693.]

6-Methyl-2-naphthol [17579-79-2] $C_{11}H_{10}O$, M 158.2, m 128-129°, b 177.5-178°/15mm, pK_{Est} ~9.8. Crystallise the naphthol from EtOH or ligroin. Sublime it *in vacuo*. [Beilstein 6 II 618, 6 III 3028.]

7-Methyl-2-naphthol [26593-50-0] has m 118°, b 312°/760mm, d 1.1, pK_{Est} ~9.7. Crystallise the naphthol from EtOH or ligroin. It has m 118° after sublimation *in vacuo*. [Halsall & Thomas *J Chem Soc* 2431 1956, DOI: 10.1039/JR9560002431; Beilstein 6 IV 3029.]

Methyl 1-naphthyl ether (1-methoxynaphthalene) [2216-69-5] $C_{11}H_{10}O$, M 158.2, b 90-91°/2mm, 135-137°/12mm, d_4^{20} 1.095, n_D^{20} 1.6210. Steam distil the ether from alkaline solution. The distillate is extracted with Et₂O. After drying (MgSO₄) the extract and evaporating Et₂O, the methyl naphthyl ether is fractionated under reduced pressure from CaH₂. The *picrate* has m 129.5-130.5° (from EtOH). [Beilstein 6 IV 4211.]

Methyl 2-naphthyl ether (2-methoxynaphthalene, Nerolin) [93-04-9] has m 70-72°, 73.0- 73.6°, b 138°/10mm 273°/760mm. Fractionally distil the ether under vacuum. Crystallise it from absolute EtOH, aqueous EtOH, *C₆H₆, petroleum ether or *n*-heptane, and dry it under vacuum in an Abderhalden pistol or distil it *in vacuo*. The *picrate* has m 118° (from EtOH or CHCl₃). [Kikuchi et al. *J Phys Chem* 91 574 1987, DOI: 10.1021/j100287a017; Hiers & Hager *Org Synth Coll Vol I* 58 1941, DOI: 10.15227/orgsyn.009.0012; Beilstein 6 III 2969, 6 IV 4257.]

2-Methyl-3-nitroaniline [603-83-8] $C_7H_8N_2O_2$, M 152.2 m 88° to 90°, 92°, b 305°/760mm, pK_{Est} ~2.3.

Crystallise the yellow nitrotoluidine from EtOH or $\ast\text{C}_6\text{H}_6$. It is steam volatile. The *acetyl* derivative crystallises from aqueous EtOH and has **m 164°**. [*Beilstein* 12 I 395, 12 II 460, 12 III 1944, 2 IV 1811.]

2-Methyl-4-nitroaniline [99-52-5] has **m 129°, 130° to 132°, pK₂₅ 0.93**. Crystallise the nitrotoluidine from EtOH. The *acetyl* and *benzoyl* derivatives have **m 200°** and **174°** (EtOH) respectively. [*Beilstein* 12 IV 1809.]

2-Methyl-5-nitroaniline [99-55-8] has **m 103° to 106°, 109°, pK₂₅ 2.35**. Acetylate the aniline, and the acetyl derivative is crystallised to constant melting point; then hydrolyse it with 70% H_2SO_4 and the free base is regenerated by treatment with NH_3 [Bevan et al. *J Chem Soc* 4284 1956, DOI: 10.1039/JR9560004284]. [*Beilstein* 12 H 844, 12 IV 1807.]

4-Methyl-3-nitroaniline [119-32-4] has **m 74° to 77°, 81.5°, d 1.312, pK₂₅ 3.02**. Crystallise the aniline from hot water (charcoal), then ethanol and dry it in a vacuum desiccator. [*Beilstein* 12 H 966.]

N-Methyl-4-nitroaniline [100-15-2] has **m 149° to 151°, 152.2°, 152-154°, pK₂₅ 0.55**. Crystallise the aniline from aqueous EtOH. [*Beilstein* 12 H 714.]

2-Methyl-3-nitroanisole [4837-88-1] $\text{C}_8\text{H}_9\text{N}_2\text{O}_3$, **M 176.2, m 54-56°**. 2-Methyl-5-nitroanisole crystallises from MeOH (yellow needles) and sublimes *in vacuo*. [Kuffner et al. *Monatsh Chem* 91 1152 1960, DOI: 10.1007/BF00899842; *Beilstein* 6 I 178.]

Methyl 3-nitrobenzoate [618-95-1] $\text{C}_8\text{H}_7\text{NO}_4$, **M 181.2, m 76°, 78°, 79-80°, b 279°/atm**. Crystallise the benzoate from MeOH (1g/ml). [*Beilstein* 9 H 378, 9 I 153, 9 II 248, 9 III 1493, 9 IV 1056.]

Methyl 4-nitrobenzoate [619-50-1] has **m 95-95.5°, 96°, 95-97°**. Dissolve the benzoate in diethyl ether, then wash it with aqueous alkali; the ether is evaporated and the ester is recrystallised from EtOH. [*Beilstein* 9 H 390, 9 IV 1074.]

3-Methyl-2-nitrobenzoic acid [5437-38-7] $\text{C}_8\text{H}_7\text{NO}_4$, **M 181.2, m 219-222°, 220-222.5°, 223°, b 340°/atm, d₂₅ 1.393g/ml, pK₂₀ 2.91 (1% aqueous EtOH)**. Recrystallise it from EtOH or hot H_2O (solubility at 22° is 0.1w/v%). The *methyl ester* has **m 74°** (from MeOH), and the *amide* [60310-07-8] **M 180.1, has m 192°** (needles from H_2O , prisms from EtOH). [*Beilstein* 9 H 480, 9 IV 1722.]

4-Methyl-3-nitrobenzoic acid (3-nitro-*p*-toluic acid) [96-98-0] has **m 187-190°, 190-191°, pK₂₀ 3.62 (1% aqueous EtOH)**. Recrystallise the acid from EtOH. The *S-benzylisothiuronium salt* has **m 167-168°** (EtOH). The *acid chloride* [10397-30-5] has **m 20-21°, b 185°/36mm**, and the *methyl ester* [7356-11-8] crystallises as pale yellow needles from MeOH with **m 51°**. [*Beilstein* 9 H 502, 9 II 334, 9 III 2359, 9 IV 1746.]

N-Methyl-4-nitrosoaniline [10595-51-4] $\text{C}_7\text{H}_8\text{N}_2\text{O}$, **M 136.2, m 114-115°, 118°, pK_{Est} ~1.0**. Crystallise it from $\ast\text{C}_6\text{H}_6$. The *picrate* has **m 166°(dec)** (from MeOH or CHCl_3). [*Beilstein* 7 III 3370, 12 IV 1228.]

4-Methylphenylacetic acid (*p*-tolylacetic acid) [622-47-9] $\text{C}_9\text{H}_{10}\text{O}_2$, **M 150.2, m 88° to 92°, 90-93°, 94°, b 265-267°/atm, pK₂₅ 4.37**. Crystallise the acid from heptane or water. [*Beilstein* 9 IV 1795.]

1-Methyl-1-phenylhydrazine sulfate [33008-18-3] $\text{C}_7\text{H}_{12}\text{N}_2\text{O}_4\text{S}$, **M 218.2, pK₂₅ 4.98 (free base)**. Crystallise the sulfate from hot H_2O by addition of hot EtOH. [*Beilstein* 15 IV 53 for free base.]

N-Methylphthalimide [550-44-7] $\text{C}_9\text{H}_7\text{NO}_2$, **M 161.1, m 129° to 132°, 133.8°**. Recrystallise the imide from absolute EtOH or AcOH (**m 134°**). The IR has ν_{max} at 1780 and 1380 cm^{-1} . [*Beilstein* 21 III/IV 5030.]

Methyl Red (4-dimethylaminoazobenzene-2'-carboxylic acid) [493-52-7] $\text{C}_{15}\text{H}_{15}\text{N}_3\text{O}_2$, **M 269.3, m 179-180°, 181-182°, CI 13020, pK₁²⁵ 2.30, pK₂²⁵ 4.82, pK₃²⁵ 9.5**. The acid is extracted with boiling toluene using a Soxhlet apparatus. The crystals which separate on slow cooling to room temperature are filtered off, washed with a little toluene and recrystallised from glacial acetic acid, \ast benzene or toluene followed by pyridine/water. Alternatively, dissolve it in aqueous 5% NaHCO_3 solution, and precipitate it from a hot solution by dropwise addition of aqueous HCl. Repeat this until the extinction coefficients do not increase at λ_{max} at 410nm (H_2O). It is a titration indicator which is red at pH 4.4 and below, and yellow at pH 6.2 and above (see pK_a values). It is easily reduced to a colourless product. [Clarke & Kirner *Org Synth Coll Vol* 1 374 1941, DOI: 10.15227/orgsyn.002.0047; *Beilstein* 16 IV 504.]

Methyl salicylate (methyl 2-hydroxybenzoate, oil of winter green) [119-36-8] $C_8H_8O_3$, M 152.2, m -8.6° , b $79^\circ/6\text{mm}$, $104\text{--}105^\circ/14\text{mm}$, $223.3^\circ/\text{atm}$, d_4^{20} 1.1149, n_D^{20} 1.5380, pK^{25} 10.19. Dilute the ester with Et_2O , wash with saturated $NaHCO_3$ (it may effervesce due to the presence of free acid), brine, dry $MgSO_4$, filter, evaporate and distil it. Its solubility is 1g/1.5L of H_2O . The *benzoyl* derivative has m 92° (b $270\text{--}280^\circ/120\text{mm}$), and the *3,5-dinitrobenzoate* has m 107.5° , and the *3,5-dinitrocarbamoyl* derivative has m $180\text{--}181^\circ$. [Hallas *J Chem Soc* 5737 p5770 1965, DOI: 10.1039/JR9650005737; *Beilstein* 10 IV 143.] It has a fragrant odour and is used in foods and beverages, and in liniments.

α -Methylstyrene (monomer, 2-phenylpropene) [98-83-9] C_9H_{10} , M 118.2, m -24° , b $57^\circ/15\text{mm}$, $165\text{--}169^\circ/\text{atm}$, $172\text{--}173^\circ/\text{atm}$, d_4^{20} 0.910, n_D^{20} 1.5368. Wash the monomer three times with aqueous 10% $NaOH$ (to remove inhibitors such as quinol), then six times with distilled water, dry with $CaCl_2$ and distil it under vacuum. The distillate is kept under nitrogen, in the cold, and redistilled if kept for more than 48 hours before use. It can also be dried with CaH_2 . Add stabiliser (e.g. 4-*tert*-butylcatechol) if it is to be stored for long periods. [*Beilstein* 5 IV 1364.]

***trans*- β -Methylstyrene (1-phenylpropene)** [873-66-5] C_9H_{10} , M 118.2, b $64\text{--}65^\circ/10\text{mm}$, $176^\circ/760\text{mm}$, d_4^{20} 0.910, n_D^{20} 1.5496. Distil it under N_2 from powdered $NaOH$ through a Vigreux column, and pass it through activated neutral alumina before use [Wong et al. *J Am Chem Soc* 109 3428 1987, DOI: 10.1021/ja00245a039]. Store at $\sim 2\text{--}8^\circ$. [*Beilstein* 5 III 1184, 5 IV 1359.]

4-Methylstyrene [622-97-9] C_9H_{10} , M 118.2, b $60^\circ/12\text{mm}$, $106^\circ/10\text{mm}$, d_4^{20} 0.9173, n_D^{20} 1.542. Purify it as the above styrenes and add a small amount of antioxidant if it is to be stored. It has UV in $EtOH$ at λ_{max} at 285nm (log ϵ 3.07), and in $EtOH + HCl$ 295nm (log ϵ 2.84) and 252nm (log ϵ 4.23). [Schwartzman & Corson *J Am Chem Soc* 78 322 1956, DOI: 10.1021/ja01583a020; Joy & Orchin *J Am Chem Soc* 81 305 1959, DOI: 10.1021/ja01511a012; Buck et al. *J Chem Soc* 2377 1949, DOI: 10.1039/JR9490002377; *Beilstein* 5 IV 1369.]

Methyl Violet 2B [4,4'-bis-(diethylamino)-4"-methyliminotriphenylmethyl hydrochloride] [8004-87-3] $C_{24}H_{28}ClN_3$, M 394.0, m $137^\circ(\text{dec})$, CI 42535, $\lambda_{\text{max}} \sim 580\text{nm}$. Crystallise the dye from $EtOH$ by precipitation with Et_2O during cooling in an ice-bath. Filter it off and dry it at 105° . [*Beilstein* 13 H 755, 13 IV 2283.] It is a biological stain which, with iodine, is used for identifying chromosomes, chromatin, nucleoli and amyloid.

Michler's ketone [4,4'-bis(dimethylamino)benzophenone] [90-94-8] $C_{17}H_{20}N_2O$, M 268.4, m 173° , $174\text{--}176^\circ$, 179° , pK^{25} 9.84. Dissolve the ketone in dilute HCl , filter and precipitate it by adding ammonia (to remove water-insoluble impurities such as benzophenone). Then crystallise it from $EtOH$ or petroleum ether. [Suppan *JCS Faraday Trans 1* 71 539 1975, DOI: 10.1039/F19757100539.] It is also purified by dissolving in *benzene, then washing with water until the aqueous phase is colourless. The *benzene is evaporated off, and the residue is recrystallised three times from *benzene and $EtOH$ [Hoshino & Kogure *J Phys Chem* 92 417 1988, DOI: 10.1021/j100313a035]. [*Beilstein* 14 IV 255.]

Naphthacene (benz[b]anthracene, 2,3-benzanthracene, rubene) [92-24-0] $C_{18}H_{12}$, M 228.3, m $>300^\circ$, 341° (open capillary), 349° , 357° . Naphthacene crystallises in orange needles from $EtOH$, $*C_6H_6$ or toluene. Dissolve it in sodium-dried *benzene and pass it through a column of alumina. The eluent is evaporated under vacuum, and the chromatography is repeated using fresh *benzene. Finally, the naphthacene is sublimed *in vacuo* at 186° . [Martin & Ubbelohde *J Chem Soc* 4948 1961, DOI: 10.1039/JR9610004948; UV: Clar *Chem Ber* 65 503 1932, DOI: 10.1002/cber.19320650402; Clar *Chem Ber* 69 607 1936, DOI: 10.1002/cber.19360690327; IR: Cannon & Sutherland *Spectrochim Acta* 4 373 1951, *Beilstein* 5 IV 2545.]

1-Naphthaldehyde [66-77-3] $C_{11}H_8O$, M 156.2, m $1\text{--}2^\circ$, b $160\text{--}161^\circ/15\text{mm}$, $292^\circ/760\text{mm}$, d 1.155, pK^{20} -7.04 (aqueous H_2SO_4). Distil 1-naphthaldehyde with steam, extract the distillate into Et_2O , dry (Na_2SO_4), filter, evaporate the filtrate and distil the residue in a vacuum. [*Beilstein* 7 IV 1286.]

2-Naphthaldehyde [66-99-9] has m 59° , 60° , 61° , $59\text{--}62^\circ$, b $160^\circ/19\text{mm}$, $308.4^\circ/760\text{mm}$, d^{25} 1.6211g/ml, pK^{20} -7.04 (aqueous H_2SO_4). Distil 2-naphthaldehyde with steam, then crystallise it from water or $EtOH$. [*Beilstein* 7 IV 1288.]

Naphthalene [91-20-3] $C_{10}H_8$, **M 128.2**, **m 80.3°**, **80° to 82°**, **b 87.5°/10mm**, **218.0°/atm**, d_4^{20} **1.0253**, d_4^{100} **0.9625**, n_D^{85} **1.5590**. Crystallise naphthalene once or more times from the following solvents: EtOH, MeOH, CCl_4 , $*C_6H_6$, glacial acetic acid, acetone or diethyl ether, followed by drying at 60° in an Abderhalden drying apparatus. It has also been purified by vacuum sublimation and by fractional crystallisation from its melt. Other purification procedures include refluxing in EtOH over Raney Ni and chromatography of a CCl_4 solution on alumina with $*benzene$ as eluting solvent. Baly and Tuck [*J Chem Soc* 1902 1908, DOI: 10.1039/CT9089301902,] purified naphthalene for spectroscopy by heating with concentrate H_2SO_4 and MnO_2 , followed by steam distillation (repeating the process), and formation of the *picrate* which, after recrystallisation (**m 150°**) is decomposed with base and the naphthalene is steam distilled. It is then crystallised from dilute EtOH. It can be dried over P_2O_5 under vacuum (take care not to make it sublime). Also purify it by sublimation and subsequent crystallisation from cyclohexane. Alternatively, it has been washed at 85° with 10% NaOH to remove phenols, with 50% NaOH to remove nitriles, with 10% H_2SO_4 to remove organic bases, and with 0.8g $AlCl_3$ to remove thianaphthalenes and various alkyl derivatives. Then it is treated with 20% H_2SO_4 , 15% Na_2CO_3 and finally distilled. [Gorman et al. *J Am Chem Soc* **107** 4404 1985, DOI: 10.1021/ja00301a006.] Zone refining purified naphthalene from anthracene, 2,4-dinitrophenylhydrazine, methyl violet, benzoic acid, methyl red, chrysene, pentacene and indoline. [Beilstein **5** IV 1640.]

1,8-Naphthalic acid (naphthalene-1,8-dicarboxylic acid) [518-05-8] $C_{12}H_8O_4$, **M 216.9**, **m 270°**, $pK_{Est(1)} \sim 2.1$, $pK_{Est(2)} \sim 4.5$. Crystallise the acid from EtOH or aqueous EtOH. [Raecke & Schrip *Org Synth* **40** 71 1960, DOI: 10.15227/orgsyn.040.0071; Beilstein **9** II 651, **9** III 4466.] **1,8-Naphthalic anhydride** [81-84-5] $C_{12}H_6O_3$, **M 198.2**, **m 267° to 296°, 274°, 274-275°**. Extract it with cold aqueous Na_2CO_3 to remove free acid, then crystallise from acetic anhydride. [Beilstein **17** III/IV 6392, **17/11** V 492.]

2-Naphthamide (2-naphthoic acid amide) [2243-82-5] $C_{11}H_9NO_2$, **M 171.2**, **m 195°**, pK^{20} **-2.30** (H_0 scale, aqueous H_2SO_4). Crystallise it from EtOH (197°). [Clemo & Spence *J Chem Soc* 2818 1928, DOI: 10.1039/JR9280002811; Beilstein **9** H 657, **9** II 45, **9** IV 2417.] Useful for preparing antiallergic agents [Ikawa et al. US Patent 5,714,613, February 3, 1998].

Naphthazarin (5,8-dihydroxy-1,4-naphthoquinone) [475-38-7] $C_{10}H_6O_4$, **M 190.2**, **m ~ 220-230°(dec)**, **m 225-230°**, pK_1^{20} **8.2**, pK_2^{20} **10.2**. It crystallises in red-brown needles with a green shine from EtOH or dioxane (m 228-232°). It also crystallises from hexane and is further purified by sublimation at 2-10mm. [Huppert et al. *J Phys Chem* **89** 5811 1985, DOI: 10.1021/j100272a047.] It is sparingly soluble in H_2O but soluble in alkalis. The *diacetate* forms golden yellow prisms from $CHCl_3$, **m 192-193°** and the *5,8-dimethoxy* derivative has **m 157° (155°)** (from petroleum ether) [Bruce & Thompson *J Chem Soc* 1089 1955, DOI: 10.1039/JR9550001089; IR: Schmand & Boldt *J Am Chem Soc* **97** 447 1975, DOI: 10.1021/ja00835a052; NMR: Brockmann & Zeeck *Chem Ber* **101** 4221 1968, DOI: 10.1002/cber.19681011222]. The *monothiosemicarbazone* has **m 168°(dec)** from EtOH [Gardner et al. *J Am Chem Soc* **74** 2106 1952, DOI: 10.1021/ja01128a509]. [Beilstein **8** H 412, **8** III 3600.]

1-Naphthoic acid [86-55-5] $C_{11}H_8O_2$, **M 172.2**, **m 157° to 160°, 160.5-162.0°, 162.5-163.0°, >300°/atm**, pK^{25} **3.60**. Crystallise the acid from toluene (3ml/g) (charcoal), petroleum ether (b 80-100°), or aqueous 50% EtOH. The *amide* has **m 202°** (from EtOH). [Gilman et al. *Org Synth Coll Vol* **2** 425 1943, DOI: 10.15227/orgsyn.011.0080; Beilstein **9** IV 2402.] Mild irritant.

2-Naphthoic acid [93-09-4] has **m 180-184°, 184-185°, 186-187°, >300°/atm**, d **1.08**, pK^{25} **4.14**. Crystallise the acid from 95% EtOH (4ml/g), aqueous 50% EtOH or Me_2CO (**m 185-186°**). Dry it at 100°. The *acid chloride* [2243-83-6] has **m 52-52°** (from $*C_6H_6$ /petroleum ether) and **b 160-162°/11mm** [Hersberg & Carson *Org Synth Coll Vol* **3** 629 1955], the *amide* has **m 192°** (from EtOH), and the *N-Methyl amide* has **m 109-109.5°** (form $*C_6H_6$). [Newman & Holmes *Org Synth Coll Vol* **2** 428 1943, DOI: 10.15227/orgsyn.017.0065; Beilstein **9** H 656, **9** III 3174, **9** IV 2414.] Mild irritant.

1-Naphthol [90-15-3] $C_{10}H_8O$, **M 144.2**, **m 94-96°, 95.5-96°, 278-280°/atm**, $d_4^{98.7}$ **1.0954**, pK^{25} **9.34**. Sublime 1-naphthol, then crystallise it from aqueous MeOH (charcoal), aqueous 25% or 50% EtOH, $*C_6H_6$, cyclohexane, heptane, CCl_4 or H_2O . Dry it over P_2O_5 *in vacuo*. The *4-nitrobenzoate* has **m 143°** (from EtOH). [Shizuka et al. *J Am Chem Soc* **107** 7816 1985, DOI: 10.1021/ja00312a003; Beilstein **8** H 596, **6** IV 4208.]

2-Naphthol [135-19-3] $C_{10}H_8O$, **M 144.2**, **m** 121°, 122.5-123.5°, **b** 285-286°/atm, **d** 1.22, **pK²⁵** 9.57. Crystallise 2-naphthol from aqueous 25% EtOH (charcoal), H₂O (solubility is 0.1 w/v% at ~20° and ~1.3w.v% at 100°), *benzene, toluene or CCl₄. Alternatively, extract it repeatedly with small amounts of EtOH (solubility is 125w/v% at ~20°), followed by dissolution in a minimum volume of EtOH and precipitation with distilled water, then drying over P₂O₅ under vacuum. It has also been dissolved in aqueous NaOH and precipitated by adding acid (repeat several times), then precipitated from *benzene by addition of heptane. Final purification can be by zone melting or sublimation *in vacuo*. Its solubility at ~20° in CHCl₃ is 5.9w/v%, and in Et₂O it is 77w.v%. The **4-nitrobenzoate** has **m** 104° (from EtOH). [Bardez et al. *J Phys Chem* **89** 5031 1985, DOI: 10.1021/j100269a029; Kikuchi et al. *J Phys Chem* **91** 574 1987, DOI: 10.1021/j100287a017; *Beilstein* **6** IV 4253.] Used in making antioxidants, is an antiseptic and an anthelmintic.

Naphthol AS-D (3-hydroxy-2-naphthoic-o-toluide) [135-61-5] $C_{18}H_{15}NO_2$, **M 277.3**, **m** 196-198°, **CI 37520**, **pK_{Est}** ~8.5. Purify it by recrystallisation from xylene. This gives yellow-green fluorescent solutions at pH 8.2-9.5. [IR: Schnopper et al. *Anal Chem* **31** 1542 1959, DOI:10.1021/ac60153a035.] The **naphthol AS-D acetate** is obtained with AcCl, **m** 168-169°, and with chloroacetyl chloride **naphthol AS-D-chloroacetate** is obtained [Moloney et al. *J Histochem Cytochem* **8** 200 1960, DOI:10.1177/8.3.200; Burstone *Arch Pathology* **63** 164 1957, PMID: 13393921]. [*Beilstein* **12** H 505.]

α -Naphtholbenzein [bis-(α -{4-hydroxynaphth-1-yl})-benzyl alcohol] [145-50-6 (anhydrous), 6948-88-5 (covalent hydrate)] $C_{27}H_{18}O_2$, $C_{27}H_{18}O_2 \cdot H_2O$, **M 374.5** (anhydrous), **392.5** (covalent hydrate), **m** 122-125°, 230-235°, **pK_{Est}** ~9.3. Crystallise the alcohol from EtOH, aqueous EtOH or glacial acetic acid. It is an indicator in titrations of weak bases and amino acids, also in non-aqueous media, e.g. glacial AcOH, for the pH range (pH 0.0-0.8/pH 8.2-10.0) 8.2 to 10 with a visual transition from yellow to blue. It is soluble in EtOH, Me₂CO and AcOH, and almost insoluble in H₂O. [Nabeau & Branchen *J Am Chem Soc* **57** 1363 1935, DOI: 10.1021/ja01310a056; *Beilstein* **6** H 1150.]

1-Naphthol-2-carboxylic acid (1-hydroxy-2-naphthoic acid) [86-48-6] $C_{11}H_8O_3$, **M 188.2**, **m** 191-192°, 195° to 200°, 203-204°, **pK_{Est(1)}** ~2.5, **pK_{Est(2)}** ~12. Successively crystallise the acid (red crystals) from EtOH/water, diethyl ether and acetonitrile, with filtration through a column of charcoal and Celite. Its solubility in H₂O is low but it is soluble in organic solvents and aqueous alkalies where it forms salts. [Tong & Glesmann *J Am Chem Soc* **79** 583 1957, DOI: 10.1021/ja01560a023; *Beilstein* **10** H 331, **10** IV 1194.]

3-Naphthol-2-carboxylic acid (3-hydroxy-2-naphthoic acid) [92-70-6] has **m** 218-221°, 222-223°, **pK₁²⁵** 2.79, **pK₂²⁵** 12.84. Crystallise it from water or acetic acid. It is soluble in organic solvents and aqueous alkalies. The **S-benzisothiuronium salt** has **m** 216-217° (from EtOH). It forms many metal complex salts. [*Beilstein* **10** H 333, **10** III 1084.]

1,2-Naphthaquinone (o-, β -naphthoquinone) [524-42-5] $C_{10}H_6O_2$, **M 158.2**, **m** 115-120°(dec), 139-140°(dec), 140-142°(dec), 145-147°(softening at ~140° dec). Crystallise the quinone from ether (red needles) or *benzene (orange leaflets). [Note the variations in **m**.] Unlike the α -isomer below it is non-steam volatile. It has been prepared by the oxidation of 1-amino-2-naphthol hydrochloride of reasonably high purity in H₂O, to which FeCl₃ 6H₂O in concentrated HCl and diluted with H₂O was added all at once with stirring, whereby the golden yellow microcrystalline quinone separated. This was filtered off then washed well with H₂O; the solid was sliced and dried on a filter paper at ~25° in an acid-free atmosphere. Recrystallisation from the above stated solvents provides nice crystals but is wasteful and the solid does not keep well. The crude dry quinone [145-147°(softening at ~140° dec)] is essentially pure and will store, as it is, indefinitely without serious deterioration, preferably in the absence of light. It can become very *electrified* if pulverised, and it is advised not to do so. [Fieser *Org Synth Coll Vol* **2** 430 1943, *Org Synth* **17** 68 1937, DOI: 10.15227/orgsyn.017.0068.] [*Beilstein* **7** IV 2417.] The **dioxime** [1Z,2E: 6436-72-1] crystallises in yellow needles from EtOH with **m** 169° (also reported m 142-144°, 149°, isomers?). The oxime complexes with Ni, Co and Fe²⁺ [Kuse et al. *Anal Chim Acta* **70** 75 1974, DOI:10.1016/S0003-2670(01)82911-1] and its derivatives can be used as analytical reagents for the spectrophotometric determination of Ni [Töel et al. *Anal Chim Acta* **75** 323 1975, DOI:10.1016/S0003-2670(01)85357-5]. The **2-semicarbazone** (Haemostop, Naftazone) [31853-38-0] $C_{11}H_9N_3O_2$, **M 215.2** is haemostatic.

1,4-Naphthaquinone (p-naphthoquinone, α -naphthoquinone) [130-15-4] $C_{10}H_6O_2$, **M 158.2**, **m 119-122°**, **125-125.5°**, **pKa -7.5 (in H_2SO_4)**. Crystallise the quinone from diethyl ether (charcoal). It distils in steam. It also crystallises from *benzene, petroleum ether or aqueous EtOH (yellow needles) and sublimes in a vacuum. It has been prepared by the oxidation of 1-amino-4-naphthol hydrochloride with $K_2Cr_2O_7/H_2SO_4/H_2O$ in 78-81% yield [Fieser *Org Synth Coll Vol 1* 383 1941, DOI: 10.15227/orgsyn/005.0079], or by direct oxidation of naphthalene with $CrO_3/AcOH/H_2O$ (2-3 hours during naphthalene/AcOH addition to oxidant at 10-15°, warmed overnight, then the dark green solution is set aside with occasional stirring for 3 days at ~25°) in consistent 32-35% yields (50% yields have also been reported [Chawla et al. *Tetrahedron* **51** 2709 1995, DOI: 10.1016/0040-4020(95)00019-5]). Purification by crystallisation from petroleum ether (b 80-100°) is more convenient than by steam distillation, and one crystallisation gives a product of high purity. Its IR (KBr) has a band at ν_{max} 1644 cm^{-1} . [Braude & Fawcett *Org Synth Coll Vol 4* 698 1963, DOI: 10.15227/orgsyn.033.0050.] The quinone is an **IRRITANT** and a **VESICANT**. [For the fluorescence detection of cyanide by quinone derivatives see Guilbault & Kramer *Anal Chem* **37** 1395 1965, DOI: 10.1021/ac60230a027; for UV spectra of naphthoquinones see Singh et al. *Tetrahedron* **24** 6053 1968, DOI:10.1016/S0040-4020(01)90989-5; *Beilstein* **7 IV** 2422.] The *dioxime* [14140-02-4] $C_{10}H_8N_2O_2$, **M 188.2**, crystallises in needles with **m 207°(dec)** from EtOH.

1,5-Naphthaquinone (1,5-naphthoquinone) [51583-62-1] $C_{10}H_6O_2$, **M 158.2**. Theoretical studies by Boldt and coworkers [Schmand & Boldt *J Am Chem Soc* **97** 447 1975, DOI: 10.1021/ja00835a052; and Schmand et al. *Justus Liebigs Ann Chem* **1976** 1560 1976, DOI: 10.1002/jlac.197619760905; see also 1,7- and 2,3-naphthaquinone below] showed that this quinone should not be *isolable* unless stabilised by inserting shielding alkyl groups. Attempted synthesis of 2,3,4,6,7,8-tetramethyl-1,5-naphthoquinone failed. However, they succeeded in preparing **3,5-di-tert-butyl-1,5-naphthoquinone** by oxidation of the respective 1,5-diol, with 2,3-dichloro-5,6-dicyanobenzoquinone (DDQ) in $CHCl_3$ under N_2 , as red crystals in nearly quantitative yield. The quinone was fully characterised and was stable under N_2 ; thus the TLC (silica gel, CH_2Cl_2) under N_2 and absence of moisture gave a spot with $R_F = 0.2$. However, on exposure to H_2O it was slowly converted to **5-hydroxy-3,5-di-tert-butyl-1,4-naphthoquinone** which crystallised from MeOH with **m 113°**, and was characterised fully (IR, NMR and MS). They also showed from SCF-LCAO-MO calculations and spectral comparisons that **naphthazarin** (5,8-dihydroxy-1,4-naphthoquinone, see [475-38-7]) is a 1,4- and not a 1,5-naphthoquinone.

1,7-Naphthaquinone (1,7-naphthoquinone) [46001-16-5] $C_{10}H_6O_2$, **M 158.2**. Theoretical calculations by Bolt and coworkers [Menting et al. *J Org Chem* **48** 2814 1983, DOI: 10.1021/jo00165a006] on the naphthaquinones showed the reasons why some quinones are unstable; 1,7-naphthaquinone being one of them (the 1,5-naphthaquinone above was another unstable isomer as was the 2,3-isomer below). Their instability at room temperature being possibly due to their reactivity with H_2O and polymerisation which they argued could be avoided by heavy substitution with alkyl groups. In an attempt to improve their stability, they synthesised **3,6-di-tert-butyl-8-methyl-1,7-naphthoquinone** [83021-64-1] (in 13 steps), successfully by final DDQ oxidation of **3,6-di-tert-butyl-8-methyl-naphthalene 1,7-diol** ([83021-63-0], sublimes at 160-200/vacuum; **m 167°**, 1H NMR, MS and analyses) as a bright red residue which they characterised fully (IR, UV/vis. NMR and MS) as well as reductive acetylation to **1,7-diacetoxy-3,6-di-tert-butyl-8-methyl-naphthalene** ([86392-51-0], **m 60-61°**, 1H NMR, MS and analyses). However, it proved to be unexpectedly unstable and the authors concluded from MNDO calculations that the alkyl groups caused high steric strain, making it non-planar and thus unstable.

2,3-Naphthaquinone (2,3-naphthoquinone) [4939-92-8] $C_{10}H_6O_2$, **M 158.2**. All attempts to prepare this quinone by oxidation of 2,3-dihydroxynaphthalene failed because of the expected instability of the quinone. Apparently it possibly reacts readily with H_2O and polymerises. However, its transient existence was demonstrated by Horak and coworkers by oxidation of 2,3-dihydroxynaphthalene with aqueous KIO_3 in the presence of a large excess of cyclopentadiene, used also as solvent, that provided a 56% yield of the **1,4-(1:1)-Diels-Alder adduct**, **m 170°**, which crystallised from cyclohexane and sublimed *in vacuo*. Its IR, 1H NMR and MS are consistent with its structure, and on reduction with $NaBH_4$ it yielded 1,2-dihydroxynaphthalene, **m 160°** (from cyclohexane) see [92-44-4] above. [Horak et al. *Tetrahedron Lett* **22** 3577 1981, DOI:10.1016/S0040-4039(01)81962-6.]

2,6-Naphthaquinone (amphi-naphthaquinone, 2,6-naphthoquinone) [613-20-7] $C_{10}H_6O_2$, **M 158.2**, **m 130-**

135°, 135°(dec). It is formed by the oxidation of 2,6-dihydroxynaphthalene in C_6H_6 with suspended active lead dioxide (prepared from lead tetraacetate) by shaking for 1-2 minutes, filtering and evaporating [Kuhn & Hammer *Chem Ber* **83** 413 1950, DOI: 10.1002/cber.19500830415; Willstätter & Parnas *Chem Ber* **40** 1406 1907, DOI: 10.1002/cber.19070400224]. Alternatively, 2,6-naphthaquinone was prepared by dissolving 2-aminonaphthalene (0.1g) in a mixture of CHCl_3 and MeOH (1:1 v/v, 100ml) and a dye sensitizer (5mg, methylene blue) and illuminated with a tungsten lamp (2 x 200Watt) through a $\text{K}_2\text{Cr}_2\text{O}_7$ screen as O_2 was bubbled through the solution for 18 hours, and the 2,6-naphthaquinone, **m 132-134°**, was isolated after filtration and evaporation [Chawla et al. *Tetrahedron* **51** 2709 1995, DOI: 0.1016/0040-4020(95)00019-5; Chawla et al. *Indian J Chem* **32B** 733 1993.] The quinone crystallises from C_6H_6 , C_6H_6 /light petroleum or petroleum ether in yellowish-red to brick-red prisms, and forms yellow needles from ligroin. It is odourless, non-volatile and stable, and unchanged for a long time. It is soluble in EtOH and MeOH but not in H_2O . When heated to 130-135°, it turns grey and the product then dissolves in alkali. It turns a solution of hematoxylin red in colour, and a solution of $\text{K}_3[\text{Fe}(\text{CN})_6]$ blue in colour. It shows the characteristic oxidising power of quinones and is a better oxidant than 1,2- and 1,4-naphthaquinone. [Beilstein **7** 733, **7** II 656, **7** III 3706, **7** IV 2417.]

β -Naphthoxyacetic acid (BNOA) [120-23-0] $\text{C}_{12}\text{H}_{10}\text{O}_3$, **M 202.2**, **m 151° to 154°, 155°, 156°, $\text{pK}_{\text{Est}} \sim 3.0$** . Crystallise the acid from hot water or C_6H_6 . The *ethyl ester*, **m 49°**, crystallises from EtOH. [Beilstein **6** IV 4274.] It has herbicide activity.

β -Naphthoyltrifluoroacetone (4,4,4-trifluoro-2-naphthylbutan-1,3-dione) [893-33-4] $\text{C}_{14}\text{H}_9\text{F}_3\text{O}_2$, **M 266.2**, **m 70-71°, 72°, 74-76°, $\text{pK}^{20} 6.35$** . Crystallise the dione from EtOH. It forms complexes with metals, and the lanthanide complexes are particularly useful. The *mono-oxime* crystallises from H_2O or aqueous EtOH and has **m 137-138°**. [Reid & Calvin *J Am Chem Soc* **72** 2948 1950, DOI: 10.1021/ja01163a038; Beilstein **7** IV 2478.]

Naphthvalene (2,3-dihydro-1,2,3-metheno-1H-indene) [34305-47-0] C_{10}H_8 , **M 128.1**, **m dec at 175° to benzvalene**. Purify it by chromatography on alumina and eluting with pentane, also by reverse phase (ODS) HPLC using MeCN as solvent. It is stable at room temperature [Kjell et al. *J Am Chem Soc* **108** 4111 1986, DOI: 10.1021/ja00274a042; Abelt et al. *J Am Chem Soc* **107** 4148 1985, DOI: 10.1021/ja00300a010]. The ^1H NMR in CCl_4 has τ 3.18 (4H), 6.17 (t J 1.5Hz, 2H), 7.60 (t J 1.5Hz 2H).

1-Naphthyl acetate [830-81-9] $\text{C}_{12}\text{H}_{10}\text{O}_2$, **M 186.2**, **m 43° to 46°, 45-46°, 48-49°, b 114-115°/1mm**. Chromatograph the acetate on silica gel and crystallise as the 2-isomer below. The **2,4,7-trinitrofluoren-9-one complex** has **m 120°** (from EtOH). [Beilstein **6** H 608, **6** III 2928, **6** IV 4217.]

2-Naphthyl acetate [1523-11-1] has **m 68° to 70°, 71°, 71-72°, b 133-134°/2mm**. Distil the acetate in a high vacuum and/or crystallise it from petroleum ether (b 60-80°) or dilute aqueous EtOH. The *picrate* has **m 80°** (from EtOH). [Beilstein **6** H 644, **6** I 313, **6** II 600, **5** III 2982, **6** IV 4267.]

1-Naphthylacetic acid (NAA) [86-87-3] $\text{C}_{12}\text{H}_{10}\text{O}_2$, **M 186.2**, **m 129° to 131.5°, 132°, $\text{pK}^{25} 4.23$** . Crystallise the acid from EtOH or water. It has plant growth substance activity. [Beilstein **9** H 666, **9** IV 2424.]

2-Naphthylacetic acid [581-96-4] has **m 143.1-143.4°, $\text{pK}^{25} 4.30$** . Crystallise the acid from water or C_6H_6 . [Beilstein **9** H 666, **9** IV 2431.]

1-Naphthylamine [134-32-7] $\text{C}_{10}\text{H}_9\text{N}$, **M 143.2**, **m 47° to 50°, 50.8-51.2°, b 301°/atm, d 1.114, $\text{pK}^{25} 3.94$** . Sublime the amine at 120° in a stream of nitrogen, then crystallise it from petroleum ether (b 60-80°), or absolute EtOH then diethyl ether. Its solubility in H_2O is 0.002w.v% at 20°, but it is steam volatile. Dry it *in vacuo* in an Abderhalden pistol. It has also been purified by crystallisation of its *hydrochloride* (see below) from water, followed by liberation of the free base and distillation; it is finally purified by zone melting. The *styphnate* has **m 181-182°** (from EtOH). [Beilstein **12** III 2846, **12** IV 3009.] **CARCINOGEN.**

1-Naphthylamine hydrochloride [552-46-5] $\text{C}_{10}\text{H}_9\text{N.HCl}$, **M 179.7**, has **m 240-250°** sublimes on heating. Crystallise the salt from water (charcoal). [Beilstein **12** III 2849, **12** IV 3009.]

2-Naphthylamine [91-59-8] $\text{C}_{10}\text{H}_9\text{N}$, **M 143.2**, **m 111-112°, 113°, b 306°/atm, d 1.061, $\text{pK}^{25} 4.20$** . Sublime the amine at 180° in a stream of nitrogen. Crystallise it from hot water (charcoal) or C_6H_6 . Dry it under vacuum in a drying pistol. The *styphnate* has **m 194-195°** (from EtOH). [Beilstein **12** H 1265, **12** III 2989, **12**

IV 3122.] **CARCINOGEN.**

1-(1-Naphthyl) ethanol [*R*-(+)- 42177-25-3, *S*-(-)- 15914-84-8] $C_{12}H_{12}O$, **M 172.2, m 46°, 45-47.5°, 48°, $[\alpha]_{D}^{20}$ (+) and (-) 94, $[\alpha]_{D}^{20}$ (+) and (-) 78 (c 1, MeOH).** Purify the alcohol by recrystallisation from Et_2O /petroleum ether, Et_2O , hexane [Balfe et al. *J Chem Soc* 797 1946, DOI: 10.1039/JR9460000797; IR, NMR: Theisen & Heathcock *J Org Chem* **53** 2374 1988, DOI: 10.1021/jo00245a051; see also Fredga et al. *Acta Chem Scand* **11** 1609 1957, DOI: 10.3891/acta.chem.scand.11-1609]. The *RS-alcohol* [57605-95-5] has **m 63-65°, 65-66°** from hexane. [Beilstein **6** III 3034, **6** IV 4346.]

1-(1-Naphthyl)ethylamine [*R*-(+)- 3886-70-2, *S*-(-)- 10420-89-0] $C_{12}H_{13}N$, **M 171.2, b 153°/11mm, 178-181°/20mm, d_4^{20} 1.067, n_D^{20} 1.624, $[\alpha]_{D}^{20}$ (+) and (-) 65, $[\alpha]_{D}^{20}$ (+) and (-) 55 (c 2, MeOH), $[\alpha]_{D}^{17}$ (+) and (-) 82.8 (neat), $pK_{Est} \sim 9.3$.** Purify the amine by distillation in a good vacuum. [Mori et al. *Tetrahedron* **37** 1343 1981, DOI: 10.1016/S0040-4020(01)92450-0, cf. Wilson in *Topics Stereochem* (Allinger and Eliel eds) **vol 6** 135 1971, Fredga et al. *Acta Chem Scand* **11** 1609 1957, DOI: 10.3891/acta.chem.scand.11-1609]. The *hydrochlorides* crystallise from H_2O [$\alpha]_{D}^{18} \pm 3.9$ (c 3, H_2O), and the *sulfates* recrystallise from H_2O as *tetrahydrates* **m 230-232°**. The *RS-amine* [42882-31-5] has **b 153°/11mm, 156°/15mm, 183.5°/41mm** [Blicke & Maxwell *J Am Chem Soc* **61** 1780 1939, DOI: 10.1021/ja01876a039]. [Beilstein **12** III 3111.]

***N*-(α -Naphthyl)ethylenediamine dihydrochloride** $C_{12}H_{16}Cl_2N_2$, [1465-25-4] **M 291.2, m 188-190°, 200°, $pK_{Est(1)} \sim 3.8$, $pK_{Est(2)} \sim 9.4$.** Crystallise the salt from water. It has applications in qualitative inorganic analysis of nitrates, nitrites, sulfonamides and serum free fatty acids in blood. [Beilstein **12** II 699.]

1-Naphthyl isocyanate [86-84-0] $C_{10}H_7NCO$, **M 169.2, m 3-5°, b 140-142°/12mm, 269-270°/760mm, d_4^{20} 1.1774.** Distil the isocyanate at atmospheric pressure or in a vacuum. It can be crystallised from petroleum ether (b 60-70°) at low temperature. Used for derivatising alcohols, acids, amines and compounds with related functional groups. *It has a pungent odour, is TOXIC, and is absorbed through the skin.* [Beilstein **12** H 1244, **12** III 2948.]

2-(2-Naphthyloxy)ethanol [93-20-9] $C_{12}H_{12}O_2$, **M 188.2, m 72-74°, 76.7°.** Crystallise it from *benzene/petroleum ether. Its solubility in 95%EtOH is 25w/w%, in Me_2CO is 50w/w% and is also freely soluble in Et_2O and $CHCl_3$. [Yoshino et al. *Bull Chem Soc Jpn* **46** 553 1973, DOI: 10.1246/bcsj.46.553.]

***N*-1-Naphthylphthalamic acid (Naptalam)** [132-66-1] $C_{18}H_{13}NO_3$, **M 291.3, m 189° (dec), 203°.** Crystallise the herbicide from EtOH (**m 183-185°**). The *Na salt* has **m 185°**. [Beilstein **12** H 1236, **12** I 525, **12** III 2876.]

2-Naphthyl salicylate (Betol) [613-78-5] $C_{17}H_{12}O_3$, **M 264.3, m 94.4°, 95°, b 423.2°/atm, d 1.286, $pK_{Est} \sim 10.0$.** Crystallise Betol from EtOH. It is *dimorphic* with *m*'s at $\sim 55^\circ$ and $\sim 96^\circ$. [Beilstein **10** H 80, **10** II 53, **10** III 136, **10** IV 158.]

1-Naphthyl urea [6950-84-1] $C_{11}H_{10}N_2O$, **M 186.2, m 215-220°.** Crystallise the urea from EtOH (**m 213-214°** also **215°**). [Beilstein **12** H 1238, **12** IV 3076.]

2-Naphthyl urea [13114-62-0] has **m 212°, 219-220°.** Crystallise the urea from EtOH. [Beilstein **12** H 1292, **12** III 3029, **12** IV 3149.]

Narcein {6-[6-(2-dimethylaminoethyl)]-2-methoxy-3,4-(methylenedioxy)phenylacetyl]-2,3-dimethoxybenzoic acid} [131-28-2] $C_{23}H_{27}NO_8$, **M 445.4, m 176-177° (145° anhydrous), pK_1^{15} 3.5, pK_2^{15} 9.3.** Recrystallise this alkaloid, which is present in opium and used as a morphine substitute from water (as fine white crystals of the trihydrate). The *styphnate* has **m 185-189°** (from EtOH), and the *picrate* has **m 200°** (from EtOH). [Beilstein **19** H 370, **19** I 797, **19** II 386, **19** IV 4382.]

Neostigmine [(3-dimethylcarbamoylphenyl)trimethylammonium] bromide [114-80-7] $C_{12}H_{19}BrN_2O_2$, **M 303.2, m 176°(dec), 181°(dec).** Crystallise neostigmine bromide from EtOH/diethyl ether. Its solubility in H_2O is $\sim 50\%$. [Beilstein **13** III 939.] (It is cholinergic and parasympathomimetic, and *highly TOXIC*.) The starting material **3-dimethylcarbamoyl-*N,N*-dimethylaniline** [59-99-4] has **b 195°/20mm** [Beilstein **13** III 936], and its

picrate has **m 138°** (from EtOH).

Neostigmine methyl sulfate (Prostigmine B) [51-60-5] $\text{C}_{13}\text{H}_{22}\text{N}_2\text{O}_6\text{S}$ **M 334.4, m 142-145°**. Crystallise the sulfate from EtOH or Me_2CO (**m 143-144°**). Its solubility in H_2O is ~10%. [*Beilstein* 13 III 939.] (It is cholinergic and parasympathomimetic, and **highly TOXIC**.)

Ninhydrin (1,2,3-triketohydrindene hydrate) [485-47-2] $\text{C}_9\text{H}_6\text{O}_4$ (hydrate), **M 178.1, m 241-243°(dec), 250°(dec), d 0.86, pK³⁰ 8.82**. Crystallise ninhydrin from hot water (charcoal). Dry it under vacuum and store it in sealed brown containers. It becomes red on heating above 100°. It is a yellow toxic powder, soluble in EtOH, and a useful reagent for amines, amino acids and related compounds giving a deep blue or purple colour (Ruhemann's purple) and used in detection and analysis.[*Beilstein* 7 IV 2786.]

2-Nitroacetanilide [552-32-9] $\text{C}_8\text{H}_8\text{N}_2\text{O}_3$, **M 180.2, m 90-92°, 93-94°, pK_{Est} <0**. Crystallise the anilide from H_2O , aqueous EtOH (**m 92-93°**) or EtOH (**m 92.5-93.5°**). [*Beilstein* 12 II 371, 12 III 1523, 12 IV 1574.]

4-Nitroacetanilide [104-04-1] has **m 213-215°, 217°, pK_{Est} <0**. Precipitate the anilide from 80% H_2SO_4 by adding ice, then wash with water, and crystallise from aqueous EtOH. Dry it in air. [*Beilstein* 12 IV 1632.]

3-Nitroacetophenone [121-89-1] $\text{C}_8\text{H}_7\text{NO}_3$, **M 165.2, m 76-80°, 79°, 81°, b 167°/18mm, 202°/760mm**. Distil the ketone in steam and crystallise it from EtOH. [*Beilstein* 7 IV 656.]

4-Nitroacetophenone [100-19-6] has **m 77-81°, 79°, 80-81°, 81.8°, b 202°/760mm**. Crystallise the ketone from EtOH or aqueous EtOH. [*Beilstein* 7 IV 657.]

3-Nitroalizarin (1,2-dihydroxy-3-nitro-9,10-anthraquinone, Alizarin Orange) [568-93-4] $\text{C}_{14}\text{H}_7\text{NO}_6$, **M 285.2, m 244° (dec), pK_{Est(1)} ~4.6, pK_{Est(2)} ~9.6**. Crystallise the dye from AcOH (**m 244-245°**). It has λ_{max} at 494nm (H_2SO_4) and forms Cu salts. [*Beilstein* 8 H 447, 8 II 491, 8 III 3774, 8 IV 2359.]

2-Nitroaniline [88-74-4] $\text{C}_6\text{H}_6\text{N}_2\text{O}_2$, **M 138.1, m 70-73°, 72.5-73.0°, b 284°/atm, pK²⁵ -0.25 (-0.31)**. Crystallise the aniline from hot water (charcoal), then from aqueous 50% EtOH, or EtOH, and dry it in a vacuum desiccator. It has also been chromatographed on alumina, then recrystallised from *benzene. [Ehrenfeld & Puterbaugh *Org Synth Coll Vol I* 388 1941, DOI: 10.15227/orgsyn.009.0064; *Beilstein* 12 IV 1563.]

3-Nitroaniline [99-09-2] has **m 114°, pK²⁵ 2.46**. Purify it as for 2-nitroaniline. **Warning: it is absorbed through the skin.** [*Beilstein* 12 IV 1589.]

4-Nitroaniline [100-01-6] has **m 148-148.5°, pK²⁵ 1.02**. Purify it as for 2-nitroaniline. It also crystallises from acetone. It is freed from *o*- and *m*-isomers by zone melting and sublimation. [*Beilstein* 12 IV 1613.]

2-Nitroanisole (2-methoxynitrobenzene) [91-23-6] $\text{C}_7\text{H}_7\text{NO}_2$, **M 153.1, m 9.4°, 11-12°, b 265°/737mm, 273°/atm, d₄²⁰ 1.251, n_D²⁰ 1.563**. Purify it by repeated vacuum distillation in the absence of oxygen. It is volatile in steam. [*Beilstein* 6 IV 1249.] Possible carcinogen.

4-Nitroanisole (4-methoxynitrobenzene) [100-17-4] has **m 54°, b 260°/atm, b 274°/atm, d₄²⁵ 1.233**. Crystallise it from petroleum ether or hexane and dry it *in vacuo*. [*Beilstein* 6 IV 1282.]

9-Nitroanthracene [602-60-8] $\text{C}_{14}\text{H}_9\text{NO}_2$, **M 223.2, m 141° to 144°, 142-143°**. Purify it by recrystallisation from EtOH or MeOH. Further purify it also by sublimation or TLC. [*Beilstein* 5 H 666, 5 II 578, 5 III 2136, 5 IV 2296.]

2-Nitrobenzaldehyde [552-89-6] $\text{C}_7\text{H}_5\text{NO}_3$, **M 151.1, m 42-43°, 44-45°, b 120-144°/3-6mm, 153°/23mm**. Crystallise the aldehyde from toluene (2-2.5ml/g) by addition of 7ml petroleum ether (b 40-60°) for 1ml of solution. It can also be distilled under reduced pressures. It is steam volatile. [*Beilstein* 7 IV 584.]

3-Nitrobenzaldehyde [99-61-6] has **m 55-56°, 58°, b 164°/23mm**. Crystallise the aldehyde from water or EtOH/water, then sublime it twice at 2mm pressure at a temperature slightly above its melting point. It is steam volatile. [*Beilstein* 7 IV 591.]

4-Nitrobenzaldehyde [555-16-8] has **m 106°, 106-107°**. Purify it as for 3-nitrobenzaldehyde above. It is steam volatile and can be sublimed. [*Beilstein* 7 H 256, 7 I 141, 7 II 196, 7 III 907, 7 IV 589.]

Nitrobenzene [98-95-3] $C_6H_5NO_2$, M 123.1, m 5.8°, 6°, b 84-86.5°/6.5-8mm, 210.8°/760mm, d_4^{20} 1.206, n_D^{15} 1.55457, n_D^{20} 1.55257, pK^{18} -11.26 (aqueous H_2SO_4). Common impurities include nitrotoluene, dinitrothiophene, dinitrobenzene and aniline. Most impurities can be removed by steam distillation in the presence of dilute H_2SO_4 , followed by drying with $CaCl_2$, and shaking with, then distilling at low pressure from BaO , P_2O_5 , $AlCl_3$ or activated alumina. It can also be purified by fractional crystallisation from absolute EtOH (by refrigeration). Another purification process includes extraction with aqueous 2M NaOH, then water, dilute HCl, and water, followed by drying ($CaCl_2$, $MgSO_4$ or $CaSO_4$) and fractional distillation under reduced pressure. The pure material is stored in a brown bottle, in contact with silica gel or CaH_2 . It is very hygroscopic. [Beilstein 5 H 233, 5 I 124, 5 II 171, 5 III 591, 5 IV 708.]

4-Nitrobenzene-azo-resorcinol (magneson-I) [74-39-5] $C_{12}H_9N_3O_4$, M 259.2, m 185°(dec), 195-198°(dec), 199-200°(dec). Crystallise the dye from EtOH. It has λ_{max} at 432nm. It is insoluble in H_2O but soluble in aqueous alkalis and gives a red-violet complex with molybdenum. [Beilstein 16 H 181, 16 IV 1266.]

4-Nitrobenzenediazonium fluoroborate [456-27-9] $C_6H_4BF_4N_3O_2$, M 236.9, m 144-148°(dec). It crystallises from water. Drain it well and use it as soon as possible. Store at 2-8°. [Whetsel et al. *J Am Chem Soc* 78 3360 1956, DOI: 10.1021/ja01595a027; Beilstein 16 IV 816.] It can be EXPLOSIVE when dry.

4-Nitrobenzhydrazide (4-Nitrobenzoic hydrazide) [636-97-5] $C_7H_7N_3O_3$, M 181.1, m 124°, 124°, 210-214°, 216-218°, 218°(dec). Crystallise the hydrazide from EtOH or EtOAc (m 120°). [Beilstein 9 H 375, 9 I 152, 9 II 246, 9 III 1481, 9 IV 1052.]

2-Nitrobenzoic acid [552-16-9] $C_7H_5NO_4$, M 167.1, m 141° to 148°, 147.5°, 146-148°, 146-149°, pK^{25} 2.21. Crystallise the acid from *benzene (twice), *n*-butyl ether (twice), then water (twice). Dry and store it in a vacuum desiccator. [Le Noble & Wheland *J Am Chem Soc* 80 5397 1958, DOI: 10.1021/ja01553a020.] It has also been crystallised from EtOH/ H_2O . The *amide* [610-15-1] $C_7H_6N_2O_3$, M 166.1, has m 176.5° (from H_2O , also m 174-178°). [Beilstein 9 III 1466, 9 IV 1046.] It is used as an NH protecting group.

3-Nitrobenzoic acid [121-92-6] has m 139-141°, 143-143.5°, pK^{25} 3.46. Crystallise the acid from *benzene, H_2O , EtOH (charcoal), glacial acetic acid or MeOH/ H_2O . Dry and store it in a vacuum desiccator. The *amide* [645-09-0] $C_7H_6N_2O_3$, M 166.1, has m 143° (from H_2O or * C_6H_6 , also m 143° and 140-143°). [Beilstein 9 III 1489, 9 IV 1055.]

4-Nitrobenzoic acid [62-23-7] has m 137-240°, 241-242°, pK^{25} 3.43. Purify it as for 3-nitrobenzoic acid above. The *amide* [619-80-7] $C_7H_6N_2O_3$, M 166.1, has m 201.6° (from H_2O , also m 202.5° and 201-204°). [Beilstein 9 III 1537, 9 IV 1072.] **4-Nitrobenzoic anhydride** [902-47-6] $C_{14}H_8N_2O_7$, M 316.2, has m 192°, 195-195.5°. Crystallise the anhydride from Me_2CO , toluene, * C_6H_6 /EtOAc or EtOAc. [Beilstein 9 II 268, 9 III 1684.]

4-Nitrobenzoyl chloride [122-04-3] $C_7H_4ClNO_3$, M 185.6, m 71° to 74°, 75°, b 155°/20mm, 202-205°/105mm. Crystallise the acid chloride from dry petroleum ether (b 60-80°), * C_6H_6 or CCl_4 (yellow crystals). Distil it under vacuum. IRRITANT. [Adams & Jenkins *Org Synth Coll Vol* 1 394 1941, DOI: 10.15227/orgsyn.003.0075; Beilstein 9 III 1709, 9 IV 1191.]

3-Nitrobenzyl alcohol (3-NBA, NOBA) [619-25-0] $C_7H_7NO_3$, M 153.1, m 30° to 32°, 94°, b 178-180°/3mm, d_4^{20} 1.29. If the IR contains OH bands, then heat with oxalyl chloride until evolution of gasses ceases and then evaporate excess of oxalyl chloride and distil the residue in a vacuum. [Beilstein 6 IV 2609.] It is a very useful material as a matrix in mass spectrometry [Meili & Seibl *Org Mass Spectrom* 19 581 1984, DOI: 10.1002/oms.1210191111; Zhao et al. *Anal Chem* 63 450 1991, DOI: 10.1021/ac00005a012; Chan et al. *Org Mass Spectrom* 27 53 1992, DOI: 10.1002/oms.1210270114; Iavarone & Williams *J Am Chem Soc* 125 2319 2003, DOI: 10.1021/ja021202t].

4-Nitrobenzyl alcohol [619-73-8] has m 92°, 93°, 94°, b 185°/12mm. Crystallise the alcohol from EtOH and sublime it *in vacuo*. Purity should be at least 99.5%. Sublimed samples should be stored in the dark over anhydrous $CaSO_4$ (Drierite). If the IR contains OH bands, then the sample should be resublimed before use. [Kosower & Mohammed *J Am Chem Soc* 93 2709 1971, DOI: 10.1021/ja00740a021; Beilstein 6 IV 2611.]

4-Nitrobenzyl bromide [100-11-8] $C_7H_6BrNO_2$, M 216.0, m 96-98°, 98.5-99.0°, 99-101°. Recrystallise the

bromide four times from absolute EtOH, then twice from cyclohexane/hexane/*benzene (1:1:1), followed by sublimation at 0.1mm and final recrystallisation from the same solvent mixture. [Lichtin & Rao *J Am Chem Soc* **83** 2417 1961, DOI: 10.1021/ja01472a002.] It has also been crystallised from petroleum ether (b 80-100°, 10ml/g, charcoal). It slowly decomposes even when stored in a desiccator in the dark. **IRRITANT**. [Beilstein **5** IV 861.]

3-Nitrobenzyl chloride [619-23-8] $C_7H_6ClNO_2$, **M 171.6**, **m 41-44°, 45°, 46°, b 85-87°/5mm**. Crystallise the chloride from petroleum ether (b 90-120°), and/or distil in a vacuum. **IRRITANT**. [Beilstein **5** IV 855.]

4-Nitrobenzyl chloride [100-14-1] has **m 70-73°, 72.5-73°, 70-74°, b 112°/0.6mm**. Crystallise the chloride from CCl_4 , dry diethyl ether, or *n*-heptane, and dry it under vacuum. **IRRITANT**. [Beilstein **5** IV 856.]

4-Nitrobenzyl cyanide [555-21-5] $C_8H_6N_2O_2$, **M 162.2**, **m 117°, 115-118°, b 197°/12mm, 336.2°/760mm, d 1.272**. Crystallise the nitrile from EtOH, and/or distil it under vacuum. **TOXIC**, can liberate HCN. [Beilstein **9** H 456, **9** I 183, **9** II 313, **9** III 2291.]

2-Nitrobiphenyl [86-00-0] $C_{12}H_9NO_2$, **M 199.2**, **m 36.7°, 37°, 37.5°, 36-38°, b 165-170°/13mm, 185-195°/20-30mm**. *o*-Nitrobiphenyl has been prepared by a Gomberg reaction between diazotised 2-nitroaniline and $*C_6H_6$ [Elks et al. *J Chem Soc* 1284 1940, DOI: 10.1039/JR9400001284] or by the nitration of biphenyl [Morgan & Walls, **49** 15T 1930]. Although the respective yields are ~45% and ~25%, the latter method is by far more convenient to perform experimentally. Crystallise it from EtOH (seeding required) or petroleum ether (b 40-60°). Sublime it under vacuum. [Beilstein **5** H 582, **5** I 273, **5** II 487, **5** III 1750.]

trans-3-Nitrocinnamic acid [555-68-0] $C_9H_7NO_4$, **M 193.2**, **m 200-201°, 200-202°, pK²⁵ 2.58 (trans)**. Crystallise the acid from $*C_6H_6$ or EtOH. The *p*-bromophenacyl ester has **m 178° (from AcOH)**. [Thayer *Org Synth Coll Vol* **1** 398 1941, DOI: 10.15227/orgsyn.005.0083; Beilstein **9** III 271, **9** IV 2043.]

trans-4-Nitrocinnamic acid [882-06-4] has **m 143° (cis), 286°(trans), pK_{Est} ~2.6 (trans)**. Crystallise the acid from H_2O . The *p*-bromophenacyl ester has **m 191° (from AcOH)**. [Beilstein **9** H 606, **9** III 2744, **9** IV 2045.]

4-Nitrodiphenylamine [836-30-6] $C_{12}H_{10}N_2O_2$, **M 214.2**, **m 132-135°, 133-134°, 135.3°, b 211°/30mm, 355.2°/760mm, pK²⁵ -2.5**. Crystallise the amine from EtOH or aqueous EtOH (**m 135-136°**) and has **m 131.5-132°** after sublimation *in vacuo*. [Beilstein **12** H 715, **12** III 1586.]

2-Nitrodiphenyl ether [2216-12-8] $C_{12}H_9NO_3$, **M 215.2**, **b 106-108°/0.01mm, 137-138°/0.5mm, 161-162°/4mm, 188-189°/12mm, 195-200°/25mm, d₄²⁰ 1.241, n_D²⁵ 1.600**. Purify the ether by fractional distillation. Its UV (EtOH) has λ_{max} at: 255, 315nm (ϵ 6200 and 2800), and the IR (CS_2) has bands at 1350 (NO_2) and 1245, 1265 (COC) cm^{-1} [UV, IR: Dahlgard & Brewster *J Am Chem Soc* **80** 5861 1958, DOI: 10.1021/ja01554a070; Tomita & Takase *Yakugaku Zasshi (J Pharm Soc Japan)* **75** 1077 1955, Fox & Turner *J Chem Soc* 1115 1930, DOI: 10.1039/JR9300001115; Henley *J Chem Soc* 1222 1930, DOI: 10.1039/JR9300001222]. [Beilstein **6** H 218, **6** II 210, **6** III 801.]

Nitrodurene (1,2,4,5-tetramethyl-3-nitrobenzene) [3463-36-3] $C_{10}H_{13}NO_2$, **M 179.2**, **m 113-114°, b 143-144°/10mm**. Distil nitrodurene in a vacuum and/or crystallise it from EtOH (yellow prisms, **m 113-113.5°**), MeOH, acetic acid, petroleum ether or chloroform. It has been crystallised by dissolving in hexane and kept at -20° for crystals to form (**m 111-112°**). Its UV has λ_{max} at 238 and 400nm (iso-octane). [Masnovi et al. *J Am Chem Soc* **111** 2263 1989, DOI: 10.1021/ja00188a048; cf. Powell & Johnson *Org Synth Coll Vol* **2** 449 1943, DOI: 10.15227/orgsyn.014.0068; Beilstein **5** H 432, **5** III 982, **5** IV 1080.]

3-Nitrofluoranthene [892-21-7] $C_{16}H_9NO_2$, **M 247.3**, **m 157-159°, 159-160°**. Recrystallise it from AcOH or EtOAc (yellow crystals) and/or sublime it at high vacuum. It is soluble in CH_2Cl_2 , Me_2CO or $*C_6H_6$, soluble in warm EtOH and very soluble in Et_2O . [Kloetzel et al. *J Am Chem Soc* **78** 1165 1956, DOI: 10.1021/ja01587a020; Beilstein **5** III 2279.] It elicits apoptosis and necrosis [Asare et al. *Toxicology* **255** 140 2009, DOI: 10.1016/j.tox.2008.10.021].

2-Nitrofluorene [607-57-8] $C_{13}H_9NO_2$, **M 211.2**, **m 156°, 156-158°**. Crystallise 2-nitrofluorene from aqueous

acetic acid or Me₂CO (**m** 158-158.5°, also 160-160.5°). [*Beilstein* **5** H 628, **5** III 1948.] Possible carcinogen.

Nitromesitylene (2-nitro-1,3,5-trimethylbenzene) [603-71-4] C₉H₁₁NO₂, **M** 165.2, **m** 41-43°, 44°, **b** 255°/760mm. Crystallise it from EtOH, or a small volume of MeOH and cool in an ice-salt bath (**m** 43.5°). [Powell & Johnson *Org Synth Coll* Vol **2** 449 1943, DOI: 10.15227/orgsyn.014.0068; *Beilstein* **5** H 410, **5** III 923, **5** IV 1028.]

Nitromethylphenylsulfone [21272-85-5] C₇H₇NO₄S, **M** 201.2, **m** 78-79°. Recrystallise the sulfone from CHCl₃, or 95% EtOH, then sublime it at 100-120°/0.1 Torr and recrystallise again. It is an acidic analogue of nitromethane [Wade et al. *J Org Chem* **46** 765 1981, DOI: 10.1021/jo00317a023], used making phenylsulfonyl nitrile oxide for syn-cyanohydroxylations of olefins for the general synthesis of aliphatic acids and nitriles [Wade & Pillay *J Org Chem* **46** 5425 1981, DOI: 10.1021/jo00339a041; Dubey & Knaus *J Org Chem* **49** 123 1984, DOI: 10.1021/jo00175a025], and in cycloaddition reactions with olefins to form heterocycles [Wade et al. *J Org Chem* **49** 4595 1984, DOI: 10.1021/jo00198a005]. [*Beilstein* **6** II 292.]

1-Nitronaphthalene [86-57-7] C₁₀H₇NO₂, **M** 173.2, **m** 53-57°, 57.3-58.3°, **b** 30-40°/0.01mm, 304°/atm, **d**₄²⁵ 1.223. Fractionally distil 1-nitronaphthalene under reduced pressure, then crystallise it from EtOH, aqueous EtOH or heptane. Chromatograph it on alumina with *benzene/petroleum ether as eluent. It sublimes *in vacuo*. The **1:1 picrate complex** has **m** 72° (from EtOH). [*Beilstein* **5** H 553, **5** III 1593, **5** IV 1673.]

2-Nitronaphthalene [581-89-5] has **m** 79°, **b** 165°/15mm. Distil it in a vacuum and/or crystallise it from aqueous EtOH and sublime in a vacuum. The **1:1 1,3,5-trinitrobenzene complex** has **m** 75.5° (from EtOH). [*Beilstein* **5** H 555, **5** III 1596, **5** IV 1675.]

1-Nitro-2-naphthol [550-60-7] C₁₀H₇NO₃, **M** 189.2, **m** 103°, **b** 115°/0.05mm, **pK**²⁵ 5.93. Distil it under high vacuum and/or crystallise the naphthol (repeatedly) from *benzene/petroleum ether (**b** 60-80°)(1:1), or aqueous EtOH (yellow needles or plates). It forms an insoluble complex with Co. [*Beilstein* **6** III 3002, **6** IV 4370.]

2-Nitro-1-naphthol [607-24-9] has **m** 123-125°, 127-128°, **pK**²⁵ 5.89. Crystallise the naphthol (repeatedly) from EtOH. [*Beilstein* **6** H 615, **6** III 2938, **6** IV 4236.]

2-Nitrophenol [88-75-5] C₆H₅NO₃, **M** 139.1, **m** 44.8°, 44.5-45.5°, 46°, 44-48°, **b** 214-215°/760mm, **pK**²⁵ 7.23. Crystallise 2-nitrophenol from EtOH/water, water, EtOH, *benzene or MeOH/petroleum ether (**b** 70-90°). It can be steam distilled. Petrucci and Weygandt [*Anal Chem* **33** 275 1961, DOI: 10.1021/ac60170a038] crystallised it from hot water (twice), then EtOH (twice), followed by fractional crystallisation from the melt (twice), drying over CaCl₂ in a vacuum desiccator and then in a drying pistol. The **4-nitrobenzoate** had **m** 141° (from EtOH). [*Beilstein* **6** IV 1246.]

3-Nitrophenol [554-84-7] has **m** 96°, **b** 160-165°/12mm, 214-216°(dec)/atm, **pK**²⁵ 8.36. Crystallise 3-nitrophenol from water, CHCl₃, CS₂, EtOH or petroleum ether (**b** 80-100°), and dry it under vacuum over P₂O₅ at room temperature. It can also be distilled at low pressure (it decomposes at its bp at 760mm). The **4-nitrobenzoate** has **m** 174° (from EtOH). [*Beilstein* **6** IV 1269.]

4-Nitrophenol [100-02-7] has **m** 110-115°, 113-114°, **b** 279°/atm, **pK**²⁵ 7.16. Crystallise 4-nitrophenol from water (which may be acidified, e.g. with *N* H₂SO₄ or 0.5*N* HCl), EtOH, aqueous MeOH, CHCl₃, *benzene or petroleum ether, then dry it *in vacuo* over P₂O₅ at 25°. It can be sublimed at 60°/10⁻⁴mm. A 0.1% aqueous EtOH has been used as an indicator giving a colourless solution of the neutral species below pH 5.6, and a yellow solution of the anionic species at pH values above 7.6 (see pK_a). The **4-nitrobenzoate** has **m** 159° (from EtOH). [*Beilstein* **6** IV 1279.]

2-Nitrophenoxyacetic acid [1878-87-1] C₈H₇NO₅, **M** 197.2, **m** 150-159°, 156-159°, 158.2-158.5°, **pK**²⁵ 2.90. Crystallise the acid from water, and dry it over P₂O₅ *in vacuo*. The ***S*-benzylisothiuronium salt** has **m** 155-156° (from EtOH). [Hayes & Branch *J Am Chem Soc* **65** 1555 1943, DOI: 10.1021/ja01248a031; *Beilstein* **6** H 220, **6** II 211, **6** III 804, **6** IV 1261.]

4-Nitrophenyl acetate [830-03-5] C₈H₇NO₄, **M** 181.2, **m** 75-77°, 77-79°, 78-79°. Recrystallise the ester from absolute EtOH [Moss et al. *J Am Chem Soc* **108** 5520 1986, DOI: 10.1021/ja00278a025]. [*Beilstein* **6** IV 1298.] It is a chromogenic esterase substrate since the released 4-nitrophenol formed provides a solution of the

yellow anion.

2-Nitrophenylacetic acid [3740-52-1] $\text{C}_8\text{H}_7\text{NO}_4$, M 181.2, m 137-140°, 141-142.5°, pK^{25} 3.95. The acid crystallises as yellow needles from EtOH, EtOH/water and dry it over P_2O_5 under vacuum. The *amide* has m 160-161° (from $^*\text{C}_6\text{H}_6$ plates or EtOH, needles). [Beilstein 9 III 2282, 9 IV 1687.]

4-Nitrophenylacetic acid [104-03-0] has m 150-155°, 153.4-154.6°, pK^{25} 3.92. Crystallise the acid from EtOH/water (1:1), then from sodium-dried diethyl ether and dry it over P_2O_5 *in vacuo*. The *styphnate* has m 196.5-197° (prisms from EtOH). The *amide* has m 191° (from EtOH). [Beilstein 9 III 2284, 9 IV 1698.]

4-Nitro-1,2-phenylenediamine (1,2-diamino-4-nitrobenzene) [99-56-9] $\text{C}_6\text{H}_7\text{N}_3\text{O}_2$, M 153.1, m 199-201°, 201°, pK_1^{25} 1.39 (1-NH₂), pK_2^{25} 2.61 (2-NH₂). Crystallise the diamine from water (solubility is 1.2g/L at 20°). [Beilstein 13 IV 75.]

1-(4-Nitrophenyl)ethylamine hydrochloride [*R*-(+)- 57233-86-0, *S*-(-)- 132873-57-5] $\text{C}_8\text{H}_{11}\text{ClN}_2\text{O}_2$, M 202.6, m 225°, 240-242°(dec), 243-245°(dec), 248-250°, $[\alpha]_D^{20}$ (+) and (-) 72 (c 1, 0.05 M NaOH), (+) and (-) 0.3 (H₂O), $\text{pK}_{\text{Est}} \sim 8.6$. To ensure dryness, the hydrochloride (*ca* 175 g) is extracted with EtOH (3x100ml) and evaporated to dryness (any residual H₂O increases the solubility in EtOH and lowers the yield). The hydrochloride residue is triturated with absolute EtOH and dried *in vacuo*. The product is further purified by refluxing with absolute EtOH (200 ml for 83g) for 1 hour, and cooling to 10° to give 76.6g of the hydrochloride m 243-245°(dec). The *free base* [*R*, 22038-87-5; *S*, 4187-53-5] $\text{C}_8\text{H}_{10}\text{N}_2\text{O}_2$, M 166.2, is prepared by dissolving in N NaOH, extracting with CH_2Cl_2 (3 x 500ml), drying (Na_2CO_3), filtering, evaporating and distilling it. It has m 27°, b 119-120°/0.5mm [105-107°/0.5mm, 157-159°/9mm, d_4^{20} 1.1764, n_D^{20} 1.5688, $[\alpha]_D^{24}$ (+) and (-) 17.7 (neat)] [Perry et al. *Synthesis* 492 1977, DOI: 10.1055/s-1977-24459; ORD: Nerdel & Liebig *Justus Liebigs Ann Chem* 621 42 1959, DOI: 10.1002/jlac.19596210107]. [Beilstein 12 IV 2451.]

4-Nitrophenylhydrazine [100-16-3] $\text{C}_6\text{H}_7\text{N}_3\text{O}_2$, M 153.1, m 156°(dec), 158°(dec), pK_1^{25} -9.2 (aqueous H_2SO_4), pK_2^{25} 3.70. Crystallise the hydrazine from EtOH. Add water to stabilise it. The *hydrochloride* has m 212°(dec) (from EtOH, also m 202-203° dec). [Beilstein 15 III 331, 15 IV 317.] Reagent for making hydrazones for characterisation of carbonyl compounds.

3-Nitrophenyl isocyanate [3320-87-4] $\text{C}_7\text{H}_4\text{N}_2\text{O}_3$, M 164.1, m 51-52°, 52-54°, b 130-131°/11mm. Distil the isocyanate in a vacuum, and/or recrystallise it from petroleum ether (b 28-38°) or toluene/petroleum ether (m 52°). Store at 2-8°. [Beilstein 12 H 708, 12 III 1573.]

4-Nitrophenyl isocyanate [100-28-7] has m 56-59°, b 137-138°/11mm, 162-164°/20mm. Distil the isocyanate in a vacuum, and/or recrystallise it from petroleum ether (b 28-38°), $^*\text{C}_6\text{H}_6$ /petroleum ether (m 58°) or CCl_4 (m 56-57°). Store at 2-8°. [Beilstein 12 H 725, 12 III 1630.]

2-Nitrophenylpropionic acid [530-85-8, 92358-21-4] $\text{C}_9\text{H}_5\text{NO}_4$, M 191.1, m 157°(dec), 160.5-161°, 166-167°(dec), pK^{25} 2.83 (50% aqueous dioxane), 3.39 (50% aqueous EtOH). Digest the acid with boiling CHCl_3 , then crystallise it from H₂O. It can explode (store cold). The *amide* has m 159° (plates, from H₂O). [Schofield & Simpson *J Chem Soc* 512 1945, DOI: 10.1039/JR9450000512; Beilstein 9 H 636, 9 I 267, 9 II 438, 9 III 3067, 9 IV 2330.]

4-Nitrophenyl trifluoroacetate [658-78-6] $\text{C}_8\text{H}_4\text{F}_3\text{NO}_4$, M 235.1, m 35-37°, 37-39°, b 120°/12mm. Recrystallise the ester from CHCl_3 /hexane [Margolis et al. *J Biol Chem* 253 7891 1978, <http://www.jbc.org/content/253/21/7891.full.html#ref-list-1>]. It sublimes *in vacuo* (m 36-38°). It is *moisture sensitive*. Store at 2-8°. [Sakakibara & Inukai *Bull Chem Soc Jpn* 37 1231 1964, DOI: 10.1246/bcsj.37.1231.]

4-Nitrophenyl urea [556-10-5] $\text{C}_7\text{H}_7\text{N}_3\text{O}_3$, M 181.2, m 232°(dec), 238°, 242°. Crystallise the urea from EtOH or hot water. Its UV has λ_{max} at 322nm (EtOH). [Beilstein 12 II 392, 12 III 1617, 12 IV 1645.]

3-Nitrophthalic acid [603-11-2] $\text{C}_8\text{H}_5\text{NO}_6$, M 211.1, m 210°, 216-218°, b 441.3°/atm, d 1.671, pK^{25} 3.93. Crystallise 3-nitrophthalic acid from hot water (1.5ml/g). Dry it in air. The *amide* has m 201° (from EtOH). [Beilstein 9 H 823, 9 IV 3275.]

4-Nitrophthalic acid [610-27-5] $\text{C}_8\text{H}_5\text{NO}_6$, M 211.1, m 165°, pK²⁵ 4.12. Crystallise 4-nitrophthalic acid from Et₂O, EtOAc or *C₆H₆ (m 166°). The *amide* has m 200° (from EtOH). [Huntress et al. *Org Synth Coll Vol* 2 457 1943, DOI: 10.15227/orgsyn.016.0056; *Beilstein* 9 H 828, 9 IV 4234.]

3-Nitrophthalic anhydride [641-70-3] $\text{C}_8\text{H}_3\text{NO}_5$, M 193.1, m 163°, 164°, 165°. Crystallise it from *C₆H₆, *C₆H₆/petroleum ether, Me₂CO, AcOH, or Ac₂O (m 164-165°). Dry it at 100°. [*Beilstein* 17/11 V 266.]

4-Nitrophthalic anhydride [5466-84-2] has m 116-120°, 120-121.5°, b 196-197°/8mm. Distil the anhydride in a vacuum and/or recrystallise it from *C₆H₆ or Et₂O/petroleum ether. Dry it *in vacuo*. It forms addition compounds with anthracene (m 118°), and phenanthrene (m 96°). [*Beilstein* 17 III/IV 6150, 17/11 V 267.]

5-Nitro-2-*n*-propoxyaniline [553-79-7] $\text{C}_9\text{H}_{12}\text{N}_2\text{O}_3$, M 196.2, m 47.5-48.5°, 48°, 49°. pK_{Est} ~2.32. Crystallise the aniline from *n*-propyl alcohol/petroleum ether or H₂O (136mg/L at 20°). It is an orange solid that is 4,000 the sweetness of sucrose, but was banned in the US due to its possible toxicity. [*Beilstein* 13 III 878, 13 IV 897.]

2-Nitroresorcinol [601-89-8] $\text{C}_6\text{H}_5\text{NO}_4$, M 155.1, m 81-82°, 83°, 84-85°, pK₁²⁰ 6.37, pK₂²⁰ 9.46. Recrystallise 2-nitroresorcinol from aqueous EtOH. [*Beilstein* 6 H 823.]

4-Nitrosalicylic acid [619-19-2] $\text{C}_7\text{H}_5\text{NO}_5$, M 183.1, m 234°, 235-238°(dec), 237-240°, pK²⁵ 2.23. Crystallise the acid from H₂O (m 236-238°) or aqueous EtOH (m 235-236° dec). [*Beilstein* 10 III 194, 10 IV 231.]

5-Nitrosalicylic acid [96-97-9] has m 233°, b 1.730, pK₁²⁵ 2.32, pK₂²⁵ 10.34. Crystallise the acid from Me₂CO (charcoal), then twice more from Me₂CO alone, aqueous EtOH (m 234-236°) or H₂O (m 232-233°). [*Beilstein* 10 III 197, 10 IV 255.]

Nitrosobenzene (NOB) [586-96-9] $\text{C}_6\text{H}_5\text{NO}$, M 107.1, m 67.5-68°, 65-69°, b 57-59°/18mm. Steam distil nitrosobenzene, then crystallise it from a small volume of EtOH with cooling below 0°, dry it over CaCl₂ in a desiccator at atmospheric pressure, and store it under N₂ at 0°. *Alternatively*, it can be distilled onto a cold finger cooled with brine at ~-10° in a vacuum at 17mm (water pump), while heating in a water bath at 65-70° [Robertson & Vaughan *J Chem Educ* 27 605 1950, DOI: 10.1021/ed027p605]. [*Beilstein* 5 IV 702.] It is a spin trap reagent and used in the study of oxidative DNA damage [Ohkuma & Kawanishi *Biochem Biophys Res Commun* 257 555 1999, <http://www.idealibrary.com/bbrc.1999.0525>], and nitroso-induced respiratory burst in neutrophils [Nakata *J Biochem (Tokyo)* 122 188 1997, PMID: 9276687].

4-Nitrosodiphenylamine (tautomer of benzoquinone-1,4-phenylimine oxime) [156-10-5] $\text{C}_{12}\text{H}_{10}\text{N}_2\text{O}$, M 198.2, m 144°(dec), 144-145°(dec), 145.4-146.6°, 144-148°. The amine forms dark green crystals from EtOH or *C₆H₆ (m 143°). It has UV with λ_{max} at 421nm (EtOH), and it is used for detecting Pd and Rh. *It is highly toxic and a possible carcinogen*. [*Beilstein* 12 H 207, 12 II 122, 12 III 347, 12 IV 1860B.]

1-Nitroso-2-naphthol [131-91-9] $\text{C}_{10}\text{H}_7\text{NO}_2$, M 173.2, m 106-108°, 110.4-110.8°, pK²⁵ 7.63. Crystallise the naphthol from petroleum ether (b 60-80°, 7.5ml/g). Its solubility in cold H₂O is 0.1w/w%, and in cold EtOH it is 2.9w/w%, but more soluble on heating. It forms a Cobalt complex and is used to separate it from Ni. [Marvel & Porter *Org Synth Coll Vol* 1 411 1941, DOI: 10.15227/orgsyn.002.0061; *Beilstein* 7 H 712, 7 IV 2419.]

2-Nitroso-1-naphthol [132-53-6] has m 150-155°, 162-164°, pK²⁵ 7.24. Purify the naphthol by recrystallisation from petroleum ether (b 60-80°) or by dissolving it in hot EtOH, followed by successive addition of small volumes of water (m 158° dec). It also crystallises from *C₆H₆ or H₂O. Crystallisation from *C₆H₆/petroleum ether gives m (106-109 °, also 109.5°). It has λ_{max} at 274.5 and 382nm (CHCl₃). It complexes with metals. [*Beilstein* 7 H 712, 7 I 385, 7 II 647, 7 III 3688, 7 IV 2419.]

4-Nitroso-1-naphthol [605-60-7] has m 198°(dec), pK²⁵ 8.18. Crystallise the naphthol from *C₆H₆ or H₂O (m 197°, dec). and sublime it at 75-80°/0.2mm. It has λ_{max} at 336nm (EtOH). [*Beilstein* 7 H 715, 7 II 653, 7 III 3700, 7 IV 2424.]

4-Nitrosophenol (benzoquinone mono oxime) [104-91-6] $\text{C}_6\text{H}_5\text{NO}_2$, M 123.1, m >124°(dec), 132-144°(dec), pK²⁵ 6.36. 4-Nitrosophenol forms yellow crystals from xylene, *C₆H₆ (m ~144°) or Et₂O (m 128-129°, dec). The solubility in H₂O is ~0.01w.v% at 21°. [*Beilstein* 7 H 622, 7 II 574, 7 III 3367.]

***N*-Nitroso-*N*-phenylbenzylamine** [612-98-6] $C_{13}H_{12}N_2O$, M 212.2, m 58°. Crystallise the amine from absolute EtOH (yellow needles) and dry it in air. It is slightly soluble in organic solvents. [Beilstein 12 H 1071, 12 II 335, 12 III 2335.]

***trans*- β -Nitrostyrene** [5153-67-3] $C_8H_7NO_2$, M 149.2, m 55-58°, 58°, 60°, b 250-260°/atm. Crystallise the styrene from absolute EtOH, or three times from *benzene/petroleum ether (b 60-80°) (1:1). [Beilstein 5 III 1180, 5 IV 1352.]

4-Nitrostyrene [100-13-0] $C_8H_7NO_2$, M 149.2, m 20.5-21°, 25-28°, 29°, b 93-96°/3mm, 120°/10mm. Crystallise it from $CHCl_3$ /hexane. Purify it by addition of MeOH to precipitate the polymer, then crystallise it at -40° from MeOH. It has also been crystallised from EtOH. [Bernasconi et al. *J Am Chem Soc* 108 4541 1986 DOI: 10.1021/ja00275a047; Beilstein 5 H 478, 5 III 1180, 5 IV 1351.]

2-Nitrotoluene [88-72-2] $C_7H_7NO_2$, M 137.1, m -9.55° (α -form), -3.85° (β -form), b 118°/16mm, 222.3°/760mm, d_4^{20} 1.163, n_D^{20} 1.545. Crystallise 2-nitrotoluene (repeatedly) from absolute EtOH by cooling in a Dry-ice/alcohol mixture. Further purify it by passing an alcoholic solution through a column of alumina. [Beilstein 5 IV 845.]

3-Nitrotoluene [99-08-1] has m 14°, 16°, b 113-114°/15mm, 232.6°/atm, d_4^{20} 1.156, n_D^{20} 1.544. Dry 3-nitrotoluene over P_2O_5 for 24 hours, then fractionally distil it under reduced pressure. [Clarke & Taylor *Org Synth Coll Vol* 1 415 1941, DOI: 10.1522/orgsyn.003.0091; Beilstein 5 IV 847.]

4-Nitrotoluene [99-99-0] has m 51°, 51.6°, 52°, 54°, b 238°/atm, d_4^{25} 1.392. Crystallise 4-nitrotoluene from EtOH, MeOH/water, EtOH/water (1:1) or MeOH. Dry it in air, then dry it in a vacuum desiccator over H_2SO_4 . [Wright & Gilliom *J Am Chem Soc* 108 2340 1986, DOI: 10.1021/ja00269a033; Beilstein 5 IV 848.]

5-Nitrovanillin (nitroveratric aldehyde) [6635-20-7] $C_8H_7NO_5$, M 197.2, m 172-175°, 176°, 178°. It forms yellow plates from AcOH and needles from EtOH [Slota & Szyszka *Chem Ber* 68 184 1935, DOI: 10.1002/cber.19350680140]. With diazomethane, 5-nitro-3,4-dimethoxyacetophenone is formed [Brady & Manjunath *J Chem Soc* 125 1060 1924, DOI: 10.1039/CT9242501060]. The *methyl ether* crystallises from EtOAc or AcOH, m 88°, 90-91°, and the *phenylhydrazone* has m 108-110° (from aqueous EtOH). [Finger & Schott *J Prakt Chem* [2] 115 281 1927, DOI: 10.1002/prac.19271150122.] The *oxime* has m 216° (from EtOH or AcOH), and the *oxime acetate* has m 147° (from aqueous EtOH) [Vogl *Monatsh Chem* 20 383 1899, DOI: 10.1007/BF01524832; Brady & Dunn *J Chem Soc* 107 1858 1915, DOI: 10.1039/CT9150701858]. [Beilstein 8 III 2064.]

Nordihydroguaiaretic [1,4-bis(3,4-dihydroxyphenyl)-2,3-dimethylbutane] acid [500-38-9] $C_{18}H_{22}O_4$, M 302.4, m 184-185°, $pK_{Est(1)} \sim 9.7$, $pK_{Est(2)} \sim 12$. Crystallise the acid from dilute acetic acid. It is an antioxidant and was banned as a nutritional supplement because it has renal and hepato toxicity. [Beilstein 6 IV 7771.]

1,2,3,4,6,7,8-Octahydroanthracene [1079-71-6] $C_{14}H_{18}$, M 186.3, m 73°, 73.5°, 78°, d_4^{80} 0.9703, n_D^{80} 1.5372. Crystallise the compound from EtOH, then purify it by zone melting. [Beilstein 5 III 1400.]

***n*-Octylammonium 9-anthranilate** [88020-99-9] M 351.5, m 134-135°, pK^{25} 10.65 (for octylamine). Recrystallise the ester several times from ethyl acetate.

4-Octylbenzoic acid [3575-31-3] $C_{15}H_{22}O_2$, M 234.3, m 97-98°, 98°, 99-100°, pK^{25} 6.5 (80% aqueous EtOH), $pK_{Est} \sim 4.5$ (H_2O). When crystallised from EtOH it has m 139°, but when crystallised from aqueous EtOH it has m 99-100°. It forms liquid crystals. [Beilstein 9 H 571, 9 III 2611.]

4-(*tert*-Octyl)phenol [140-66-9] $C_{14}H_{22}O$, M 206.3, m 79-82°, 85-86°, b 166°/20mm, 175°/30mm, $pK_{Est} \sim 10.4$. Crystallise the phenol from *n*-hexane and/or distil it in a vacuum. [Beilstein 6 III 2051, 6 IV 3484.]

Opianic acid (2-formyl-4,5-dimethylbenzoic acid) [519-05-1] $C_{10}H_{10}O_5$, M 210.2, m 145-147°, 148°, 150°,

pK²⁵ 3.07. Crystallise the acid from water. [*Beilstein* 10 H 990, 10 I 484, 10 II 719, 10 III 4511, 10 IV 3863.]

Orcine monohydrate (orcinol hydrate, 3,5-dihydroxytoluene) [6153-39-5] $C_7H_{10}O_3$, **M 142.2**, **m 56°, 56-58°, 58°, 56-61°, b 147°/5 mm, 290°/atm, pK₁²⁰ 9.48 (9.26), pK₂²⁰ 11.20 (11.66).** Purify orcine by recrystallisation from H₂O as the *monohydrate*. It sublimes *in vacuo*, and the *anhydrous* compound has **m 106.5-108° (110°, 108°)**. It can be recrystallised from CHCl₃ (plates) or *C₆H₆ (needles or prisms). [UV: Kiss et al. *Bull Soc Chim Fr* 275 1949, Adams et al. *J Am Chem Soc* 62 732 1940, DOI: 10.1021/ja01861a010.]

Orcinol (5-methylresorcinol) [504-15-4; 6153-39-5 H₂O] $C_7H_8O_2$, **M 124.2**, **m 107.5°, 106-108°, 110°, 106-13138112°, m 59-61° (hydrate cf. orcine), pK₁²⁰ 9.36 (9.48), pK₂²⁰ 11.6 (11.20).** Crystallise orcinol from CHCl₃/*benzene (2:3). See *hydrate* in previous entry. [*Beilstein* 6 H 882, 6 IV 5892.]

[2.2]-Paracyclophane (tricyclo[8.2.2.2^{4,7}]hexadeca-4,6,10,12,13,15-hexaene)[1633-22-3] $C_{16}H_{16}$, **M 208.3**, **m 284°, 285-287°, 286-288°, 288-290°.** Purify it by recrystallisation from AcOH. ¹H-NMR δ: 1.62 (Ar-H) and 1.71 (CH₂) [Waugh & Fessenden *J Am Chem Soc* 79 846 1957, DOI: 10.1021/ja01561a017; IR and UV: Cram et al. *J Am Chem Soc* 76 6132 1954, DOI: 10.1021/ja01652a081; Cram & Steinberg *J Am Chem Soc* 73 5691 1951, DOI: 10.1021/ja01156a059. It complexes with unsaturated compounds: Cram & Bauer *J Am Chem Soc* 81 5971 1959, DOI: 10.1021/ja01531a031; Syntheses: Brink DOI: 10.1055/s-1975-23940; 807 1975, Givens et al. *J Org Chem* 44 1608 1979, DOI: 10.1021/jo01324a005; Kaplan & Truesdale *Tetrahedron Lett* 3665 1976, DOI: 10.1016/S0040-4039(00)93075-2]. [*Beilstein* 5 IV 2223.]

Para Red [1-(4-nitrophenylazo)-2-naphthol] [6410-10-2] $C_{16}H_{11}N_3O_3$, **M 293.3**, **m 250-251°, 248-252°, CI 12070, pK_{est} ~ 9.3.** Crystallise this red dye from AcOH or xylene and dry it *in vacuo*. It has λ_{max} at 488nm. [*Beilstein* 16 II 70.]

Pargyline hydrochloride (Eutonyl, N-methyl-n-propargylbenzylamine hydrochloride) [306-07-0] $C_{11}H_{14}ClN$, **M 195.7**, **m 154-155°, 155°, pK²⁵ 6.9.** Recrystallise the salt from EtOH/Et₂O and dry it *in vacuo*. It is very soluble in H₂O, in which it is unstable. The *free base* has **b 101-103°/11mm**. It is a glucuronyl transferase inducer and a monoamine oxidase inhibitor. [von Braun et al. *Justus Liebigs Ann Chem* 445 201 1925, DOI: 10.1002/jlac.19254450114; Langston et al. *Science* 225 1480 1984, DOI: 10.1126/science.6332378; *Beilstein* 12 II 548.] It is an irreversible MAO inhibitor and an antihypertensive drug.

Pavatrine hydrochloride [fluorine-9-carboxylic acid, 2-(diethylamino)ethyl ester hydrochloride] [548-65-2] $C_{20}H_{24}ClNO_2$, **M 333.7**, **m 143-144°, pK_{Est} ~9.0** Recrystallise the salt from isopropanol, EtOAc/isoPrOH and dry it over P₂O₅ *in vacuo*. The *metho-bromide* has **m 111-117°** (from butanone). [*Beilstein* 9 III 3412, 9 IV 2596.]

Pentabromophenol [608-71-9] C_6HBr_5O , **M 488.7**, **m 190-191°, 223-226, 229-230°, d₄²² 1.978, pK_{Est} ~ 4.5.** Purify it by crystallisation (charcoal) from toluene then from CCl₄ (white crystals). Dry it for 2 weeks at ca 75°. Its solubility in H₂O is 0.002w/v% at 30°. The *diethylammonium salt* has **m 191-193°** (from MeOH). [*Beilstein* 6 H 206, 6 I 108, 6 II 197, 6 III 766, 6 IV 1069.]

1-Pentacene [135-48-8] $C_{22}H_{14}$, **M 278.4**, **m 300°, 372-374° (sublimes).** It forms blue crystals from *benzene or nitrobenzene and sublimes in a vacuum. [*Beilstein* 5 IV 2721; Clar et al. *Chem Ber* 62 940 1929, DOI: 10.1002/cber.19290620426; Clar et al. *Chem Ber* 64 981 1931, DOI: 10.1002/cber.19310640507.]

Pentachloronitrobenzene (Quintozene) [82-68-8] $C_6Cl_5NO_2$, **M 295.3**, **m 140-143°, 144-145°, 146°, b 328°/atm (some decomposition), d₄²⁵ 1.718.** Crystallise it from EtOH (needles) and CS₂ (plates). [*Beilstein* 5 H 247, 5 II 188, 5 III 618.] It has fungicidal properties.

Pentachlorophenol [87-86-5] C_6HCl_5O , **M 266.3**, **m 165-180°, 190-191°, b 310°/atm, d₄²⁵ 1.978, pK²⁵ 4.8.** Crystallise it twice from toluene/EtOH. Sublime it *in vacuo*. [*Beilstein* 6 IV 1025.] The *Na salt* [131-52-2] **M**

288.3, has **m** ~300°. [*Beilstein* 6 H 194.]

Pentachlorothiophenol [133-49-3] C_6HCl_5S , **M** 282.4, **m** between 223-227°, 225°, 231.5°, 228° and 235°, **pK_{Est}** ~1.1. Crystallise from *benzene, toluene (**m** 243°) or AcOH (**m** 240° also 242-244°). [*Beilstein* 6 IV 1642.]

Pentafluorobenzene [363-72-4] C_6HF_5 , **M** 168.1, **m** -47.3°, -48°, **b** 85°/atm, 85-86°/atm, 88-89°/atm, **d**₄²⁰ 1.524, **n**_D²⁰ 1.3931. Purify it by distillation and by gas chromatography. Its IR (film) has bands at 1535 and 1512 cm⁻¹ (*C₆H₆ ring). [Stephen & Tatlow *Chem Ind (London)* 821 1957, Nield et al. *J Chem Soc* 166 1959, DOI: 10.1039/JR9590000166; *Beilstein* 5 IV 639.]

2,3,4,5,6-Pentafluorobenzoic acid [602-94-8] $C_7HF_5O_2$, **M** 212.1, **m** 101-102°, 101-103°, 104-105°, 106-107°, **d** 1.942, **pK_a** 1.48, **pK₂₅** 1.75. Dissolve the acid in Et₂O, treat it with charcoal, filter, dry (CaSO₄), filter again, evaporate and recrystallise the residue from petroleum ether (b 90-100°) after adding a little toluene, to give large colourless plates. Its UV (H₂O) has λ_{max} at 265nm (ε 761) (H₂O). The *S*-benzylisothiuronium salt has **m** 187° (from H₂O). [McBee & Rapkin *J Am Chem Soc* 73 1366 1951, DOI: 10.1021/ja01147a520; Nield et al. *J Chem Soc* 166 1959, DOI: 10.1039/JR9590000166; *Beilstein* 9 IV 956.]

O-(2,3,4,5,6-Pentafluorobenzyl)hydroxylamine hydrochloride (PFBHA.HCl) [57981-02-9] $C_7H_5ClF_5NO$, **M** 249.6, **m** 215°, 215-216°, 227°, **b** 214.5°/760mm, **pK_{Est}** ~1.1. Recrystallise the salt from EtOH to form colourless leaflets. Drying the compound at high vacuum and elevated temperature will result in losses by sublimation. [Youngdale *J Pharm Sci* 65 625 1976, DOI: 10.1002/jps.2600650443; Wehner & Handke *J Chromatogr* 177 237 1979, DOI: 10.1016/S0021-9673(01)96319-8; Nambara et al. give incorrect **m** as 115-116° *J Chromatogr* 114 81 1975, DOI: 10.1016/S0021-9673(00)85244-9.]

2,3,4,5,6-Pentafluorophenol [771-61-9] C_6HF_5O , **M** 184.1, **m** 33-35°, 34-36°, 38.5-39.5°, **b** 72-74°/48mm, 142-144°/atm, 143°/atm, **n**_D²⁰ 1.4270 (liquid prep), **pK₂₅** 5.53. It is a *hygroscopic* low melting solid not freely soluble in H₂O. Purify it by distillation, preferably in a vacuum [Forbes et al. *J Chem Soc* 2019 1959, DOI: 10.1039/JR95900002019; IR and pK_a: Birchall & Haszeldine *J Chem Soc* 13 1959, DOI: 10.1039/JR9590000013]. IR of a film has ν_{max} 3600 (OH) and 1575 (fluoroaromatic breathing) cm⁻¹. The *benzoyl* derivative has **m** 74-75°, *3,4-dinitrobenzoyl* derivative has **m** 107°, the *tosylate* has **m** 64-65° (from EtOH) and the *K salt* crystallises from Me₂CO, **m** 242°(dec), with *1H₂O-salt* the **m** is 248°(dec), and the *2H₂O-salt* has **m** 245°(dec). [*Beilstein* 6 IV 782.]

1-(Pentafluorophenyl)ethanol [*R*-(+)- 104371-21-3, *S*-(-)- 104371-20-2] $C_8H_5F_5O$, **M** 212.1, **m** 41-42°, 42°, 42.5-43°, [α]₅₄₆²⁰ (+) and (-) 9, [α]_D²⁰ (+) and (-) 7.5 (c 1, *n*-pentane). Recrystallise the ethanol from *n*-pentane at -40° and sublime it at 25°/0.3mm (use ice-cooled cold finger). It has also been purified by column chromatography through Kieselgel 60 (0.063-0.2mm mesh, Merck) and eluted with EtOAc/*n*-hexane (1:5), then recrystallised from *n*-pentane and sublimed in a vacuum. It has R_F 0.28, on Kieselgel 60 F₂₅₄ TLC foil and eluting with EtOAc/*n*-hexane (1:5). [Meese *Justus Liebigs Ann Chem* 2004 1986, DOI: 10.1002/jlac.198619861120.]

The *racemate* [75853-08-6] has **m** 32-34°, **b** 77-79°/8mm, 80-82°/37mm, **n**_D²⁰ 1.4426; the *3,4-dinitrobenzoate* has **m** 83° [Nield et al. *J Chem Soc* 166 1959, DOI: 10.1039/JR9590000166]. [*Beilstein* 6 IV 3044.]

Pentamethylbenzene [700-12-9] $C_{11}H_{16}$, **M** 148.3, **m** 50-51°, 53°, 54.5°, 53.5-55.1°, **b** 231-232°/atm, **d**₄²⁵ 0.917. Successively crystallise it from absolute EtOH, aqueous EtOH, MeOH, toluene *C₆H₆, and dry it under vacuum. [Rader & Smith *J Am Chem Soc* 84 1443 1962, DOI: 10.1021/ja00867a021.] It has also been sublimed. The *1,3,5-trinitrobenzene complex (1:1)* has **m** 121° (EtOH). Its FT-IR (neat) has ν_{max} at 2939.4, 2728.0, 1568.1, 1475.5, 1381.8, 1067.6, 1013.9, 860.7 and 524.6 cm⁻¹; [*Beilstein* 5 H 443, 5 III 1010, 5 IV 1109.] It is a very electron rich hydrocarbon which readily undergoes electrophilic reactions.

Perbenzoic acid [93-59-4] $C_7H_6O_3$, **M** 138.1, **m** 42°, 41-43°, **b** 97-110°/13-15mm, **pK_{Est}** ~7.7. Crystallise the peracid from *C₆H₆, or petroleum ether. It sublimes readily and is steam volatile. It is soluble in CHCl₃, CCl₄

and Et₂O. It is advisable **NOT** to distil it unless extreme precautions are taken. [Braun *Org Synth Coll Vol 1* 431 1941, DOI: 10.15227/orgsyn.008.0030.] **EXPLOSIVE.**

Perylene [198-55-0] C₂₀H₁₂, M 252.3, m 273-274°, 276-279°. Purify perylene by silica-gel chromatography of its recrystallised picrate. [Ware *J Am Chem Soc* **83** 4374 1961, DOI: 10.1021/ja01482a019.] Crystallise it from *benzene, toluene or EtOH and sublime it at 142° in a flow of oxygen-free nitrogen. It forms a **1:1 *benzene-complex** (m 223-224.5° needles from *C₆H₆), and a **1:2 *benzene-complex** (m 154-155° from *C₆H₆ or H₂O). The **2,4,7-trinitrofluoren-9-one** has m 270-271° (from EtOH/*C₆H₆). [Gorman et al. *J Am Chem Soc* **107** 4404 1985, DOI: 10.1021/ja00301a006; Johansson et al. *J Am Chem Soc* **109** 7374 1987, DOI: 10.1021/ja00258a021; *Beilstein* **5** III 2521, **5** IV 2689.]

Phenanthrene [85-01-8] C₁₄H₁₀, M 178.2, m 98°, 98.7-99°, 99.15°, 100.8-101.3°, b 148-149°/1mm, b 340°/760mm, d₂₅ 1.175. Likely contaminants include anthracene, carbazole, fluorene and other polycyclic hydrocarbons. Purify it by distillation from sodium under vacuum, boiling with maleic anhydride in xylene, crystallisation from acetic acid, sublimation and zone melting. It has also been recrystallised repeatedly from EtOH, *benzene or petroleum ether (b 60-70°), with subsequent drying under vacuum over P₂O₅ in an Abderhalden pistol. Feldman, Pantages and Orchin [*J Am Chem Soc* **73** 4341 1951, DOI: 10.1021/ja01153a091] separated most of the anthracene impurity by refluxing phenanthrene (671g) with maleic anhydride (194g) in xylene (1.25L) under nitrogen for 22 hours, then filtered. The filtrate was extracted with aqueous 10% NaOH, the organic phase was separated, and the solvent was evaporated. The residue, after stirring for 2 hours with 7g of sodium, was distilled in a vacuum, then recrystallised twice from 30% *benzene in EtOH. It was then dissolved in hot acetic acid (2.2ml/g), and to it was slowly added an aqueous solution of CrO₃ (60g in 72ml H₂O plus 2.2L of acetic acid), followed by slow addition of concentrated H₂SO₄ (30ml). The mixture was refluxed for 15 minutes, diluted with an equal volume of water and cooled. The precipitate was filtered off, washed with water, dried and distilled, then recrystallised twice from EtOH. Further purification is possible by chromatography from a CHCl₃ solution on activated alumina, with *benzene as eluent, and by zone refining. The **picrate (1:1)** forms golden yellow needles with m 146°, and the **styphnate (1:1)** has m 138-139° (plates or needles from EtOH or EtOH/H₂O respectively). [Dornfeld et al. *Org Synth Coll Vol 3* 134 1955, DOI: 10.15227/orgsyn.028.0019; *Beilstein* **5** H 667, **5** I 327, **5** II 579, **5** III 2136, **5** IV 2297.]

Phenanthrene-9-aldehyde [4707-71-5] C₁₅H₁₀O, M 206.3, m 102-103°, pK₂₅ -6.39 (aqueous H₂SO₄). Recrystallise the aldehyde from EtOH and sublime it at 95-98°/0.07mm. It is a white powder with blue fluorescence, and its solubility in CHCl₃ is 0.04M. The **2,4-dinitrophenylhydrazone** has m 272-273°. [*Beilstein* **7** III 2532, **7** IV 1740.]

9,10-Phenanthrenequinone [84-11-7] C₁₄H₈O₂, M 208.2, m 206°, 208°, 210°, 209-212°, b 360°/760mm, d 1.405, pK₂₅ -7.1 (aqueous H₂SO₄). Crystallise the quinone from dioxane or 95% EtOH and dry it under vacuum. It gives a green colour in H₂SO₄. [*Beilstein* **7** IV 2565.]

Phenethylamine [64-04-0] C₈H₁₁N, M 121.2, m -60°, b 87°/13mm, 194-195°/atm, 200-202°/atm, d₄²⁰ 0.962, n_D²⁰ 1.535, pK₂₅ 9.88. Distil the amine from CaH₂, under reduced pressure, just before use. It is a strong base, store under N₂. [*Beilstein* **12** H 1096, **12** IV 2453.]

Phenethyl bromide [103-63-9] C₈H₉Br, M 185.1, m -65.7°, -56°, b 92°/11mm, 220-221°/atm, d₄²⁰ 1.368, n_D²⁰ 1.557. Wash the bromide with concentrated H₂SO₄, water, aqueous 10% Na₂CO₃ and water again, then dry it with CaCl₂ and fractionally distil it just before use. [*Beilstein* **5** IV 907.]

(±)-N-1-Phenethyl urea (N-α-phenethyl urea) [60295-51-4] C₉H₁₂N₂O, M 164.2, m 137°. Crystallise the urea from H₂O, EtOAc or *C₆H₆. [Buck *J Am Chem Soc* **56** 1607 1934, DOI: 10.1021/ja01322a047; *Beilstein* **12** I 1096, **12** IV 1440.]

(+)-R-N-1-Phenethyl urea (R-N-α-phenethyl urea) [16849-91-5] has m 121-122°, [α]_D²⁵ +48.8 (c 2, EtOH), [α]_D²⁵ +46.2 (c 4.0, EtOH). Crystallise the (+)-urea from H₂O or EtOH (m 122-123°). [Marcwald & Meth *Chem Ber* **38** 801 1905, DOI: 10.1002/cber.190503801136; Cairns *J Am Chem Soc* **63** 871 1941, DOI: 10.1021/

ja01848a504; *Beilstein* **12** I 1092, **12** III 2398.] This (+)-*enantiomer*, but not the (-)-*enantiomer*, stereospecifically inhibits Ca^{2+} - but not ADP- stimulated mitochondrial respiration [Davidoff et al. *Science* **193** 66 1976, DOI:10.1126/science.935857].

(-)-*S-N-I-Phenethyl urea* (*S-N- α -phenethyl urea*) [25144-64-3] has **m** 121-122°, [α]_D²⁵ -43.6° (c 14, EtOH), [α]_D²⁵ -52.1° (c 3.6, EtOH). Crystallise the (-)-urea from H₂O or EtOH. [Lovén *J Prakt Chem* [2] **72** 307 1905, DOI: 10.1002/prac.19050720123; *Beilstein* **12** I 1094, **12** III 2398.]

N-2-Phenethyl urea [2158-04-5] **C₉H₁₂N₂O**, **M** 164.2, **m** 112°. Crystallise the (±)-urea from H₂O (**m** 112-113°) or EtOH. The *picrate* has **m** 113-115° (from H₂O) [Spica *Gazzetta* **9** 567 1879, Shapiro et al. *J Am Chem Soc* **81** 2220 1959, DOI: 10.1021/ja01518a052]. [*Beilstein* **12** 1099, **12** III 2423, **12** IV 2470.]

Phenetole [103-73-1] **C₈H₁₀O**, **M** 122.2, **m** -30°, **b** 60°/9mm, 77.5°/31mm, 170.0°/760mm, **d**₄²⁰ 0.967, **n**_D²⁰ 1.50735, **n**_D²⁵ 1.50485. Small quantities of phenol can be removed by shaking with NaOH, but this is not a very likely contaminant of commercial material. Fractional distillation from sodium, at low pressures, probably gives adequate purification. It can be dissolved in diethyl ether and washed with 10% NaOH (to remove phenols), then water. The ethereal solution is evaporated, and phenetole is fractionally distilled under vacuum. [*Beilstein* **6** H 140, **6** I 80, **6** II 142, **6** III 545.]

Phenocoll hydrochloride (4-ethoxyaniline HCl, *p*-phenetidine HCl) [637-56-9] **C₈H₁₁NO.HCl**, **M** 230.7, **m** 234°, **pK**²⁸ 5.20. Crystallise the salt from water then sublime it *in vacuo*. [*Beilstein* **13** IV 1017.]

Phenol (carbolic acid) [108-95-2] **C₆H₆O**, **M** 94.1, **m** 40.5°, 40.9°, **b** 85.5-86.0°/20mm, 180.8°/760mm, 181.70°/760mm, **d**₄²⁰ 1.06, **n**_D⁴¹ 1.54178, **n**_D⁴⁰ 1.53957, **pK**²⁵ 9.86 (in H₂O, 9.95, 10.02 also reported), 29.1 (in MeCN). Steam is passed through a boiling solution containing 1mole of phenol and 1.5-2.0moles of NaOH in 5L of H₂O until all non-acidic material has distilled. The residue is cooled, acidified with 20% (v/v) H₂SO₄, and the phenol is separated, dried with CaSO₄ and fractionally distilled under reduced pressure. It is then fractionally crystallised several times from its melt [Andon et al. *J Chem Soc* 5246 1960, DOI: 10.1039/JR9600005246]. Purification *via* the benzoate has been used by Berliner, Berliner and Nelidow [*J Am Chem Soc* **76** 507 1954, DOI: 10.1021/ja01631a052]. The *benzoate*, (**m** 70°, **b** 314°/760mm), is crystallised from 95% EtOH, then hydrolysed to the free phenol by refluxing with two equivalents of KOH in aqueous EtOH until the solution becomes homogeneous. It is acidified with HCl and extracted with diethyl ether. The ether layer is freed from benzoic acid by thorough extraction with aqueous NaHCO₃, and, after drying and removing the ether, the phenol is distilled. Phenol has also been crystallised from a 75% w/w solution in water by cooling to 11° and seeding with a crystal of the hydrate. The crystals are centrifuged off, rinsed with cold water (0-2°), saturated with phenol, and dried. It can be crystallised from petroleum ether [Bernasconi & Paschalis *J Am Chem Soc* **108** 2969 1986, DOI: 10.1021/ja00271a027]. Its solubility in H₂O is 8.3w/v% at 20°.

Draper and Pollard [*Science* **109** 448 1949, DOI: 10.1126/science.109.2835.448] added 12% water, 0.1% aluminium (can also use zinc) and 0.05% NaHCO₃ to phenol, and distilled it at atmospheric pressure until the azeotrope was removed. The phenol was then distilled at 25mm. Phenol has also been dried by distillation from the *benzene solution to remove the water/*benzene azeotrope and the excess *benzene, followed by distillation of the phenol at reduced pressure under nitrogen. Processes such as this are probably adequate for analytical grade phenol which has as its main impurity water. Phenol has also been crystallised from petroleum ether/*benzene or petroleum ether (b 40-60°). The purified material is stored in a vacuum desiccator over P₂O₅ or CaSO₄. [*Beilstein* **6** IV 531.] *It has a sweet tarry odour. An aqueous solution is used as an antiseptic, should be used with gloves as it causes severe skin burns and can be lethal if ingested.*

Phenolphthalein [77-09-8] **C₂₀H₁₄O₄**, **M** 318.2, **m** 252°, 263°, **d**₄²⁵ 0.918, **pK**_{Est(1)}~ 4.2, **pK**_{Est(2)}~ 9.8. Dissolve it in EtOH (7ml/g), then dilute it with eight volumes of cold water, filter and heat on a water-bath to remove most of the alcohol and the phenolphthalein that precipitates is filtered off and dried *in vacuo* to give white or pale yellow microcrystals. A 0.5% solution in EtOH/H₂O (1:1) is used as a titration indicator in which it is colourless below pH 8.0 and red at pH values above 10 (see **pK_a** ~9.8). [*Beilstein* **18** II 119, **18** III/IV 1945, **18/4** V 188.] It has laxative properties and is used as a purgative.

Phenolphthalol {2-[bis(4-hydroxyphenyl)methyl]benzyl alcohol, Normolax, Egamol, Velaxin} [81-92-5]

C₂₀H₁₈O₃, M 306.3, m 201-202°, pK_{Est} ~ 9.8. Crystallise it from aqueous EtOH. [*Beilstein* 6 H 1146, 6 II 1110, 6 IV 7623.] Laxative.

Phenoxyacetic acid [122-59-8] **C₈H₈O₃, M 152.2, m 98-99°, 98-100°, 285°/atm (dec), pK²⁵ 3.18.** Crystallise the acid from water (solubility at ~20° is 1.33w/v%), or aqueous EtOH. [*Beilstein* 6 IV 634.] It is an irritant fungicide which can soften hard callus skin. **Phenoxyacetyl chloride** [701-99-5] **C₈H₇ClO₂, M 170.6,** has **b 112°/10mm, 102°/16mm, 225-226°/atm, d₄²⁰ 1.235, n_D²⁰ 1.534.** If it has no OH band in the IR then distil it in a vacuum, taking precautions for the moisture-sensitive compound. If it contains free acid (due to hydrolysis, OH bands in the IR), then add an equal volume of redistilled SOCl₂, reflux for 2-3 hours, evaporate and distil the residue in a vacuum as before. It is a useful reagent for generating phenyloxyketene that undergoes cycloaddition reactions to imines leading to β-lactams [Shaikh et al. *Tetrahedron* 63 3380 2007, DOI: 10.1016/j.tet.2007.02.022; Huang & Calter *Tetrahedron Lett* 48 1657 2007, DOI: 10.1016/j.tetlet.2006.12.091]. The **amide** [621-88-5] has **m 101°**. [McElvain & Carney *J Am Chem Soc* 68 2592 1946, DOI: 10.1021/ja01216a051; *Beilstein* 6 III 613.]

4-Phenoxyaniline [139-59-3, 73166-61-7] **C₁₂H₁₁NO, M 185.2, m 82-84°, 95°, b 140°/2.3mm, pK²⁰ 4.44 (50% aqueous EtOH).** Crystallise 4-phenoxyaniline from water (solubility at 20° is 0.1w/v%). [*Beilstein* 13 IV 1020.]

Phenoxybenzamine [*N*-(2-chloroethyl)-*N*-(1-methyl-2-phenoxyethyl)benzylamine] [59-96-1] **M 303.5, m 38-40°, hydrochloride** [63-92-3] **C₁₈H₂₂ClNO, M 340.0, m 137.5-140°, pK_{Est} ~4.2.** The free base is crystallised from petroleum ether, and the **hydrochloride** is recrystallised from EtOH/diethyl ether. [*Beilstein* 12 IV 2204.] Antihypertensive.

2-Phenoxybenzoic acid [2243-42-7, 36361-03-2] **C₁₃H₁₀O₃, M 214.2, m 110-112°, 113°, b 355°/760mm, pK^{15/25} 3.53.** Crystallise the acid from aqueous EtOH or H₂O (m 114°). [*Beilstein* 10 H 65, 10 I 28, 10 II 40, 10 III 99, 10 IV 132.]

3-Phenoxybenzoic acid [61-72-3, 3739-38-6] has **m 145°, 147-149°, 147-150°, pK²⁵ 3.95.** Crystallise the acid from aqueous EtOH. [*Beilstein* 10 H 138, 10 III 247, 10 IV 316.]

4-Phenoxybutyric acid [6303-58-8] has **m 64°, 63-65°, 65-66°, b 170°/7mm, 180-185°/12mm, pK 3.17.** It has been purified by recrystallisation from petroleum ether, *C₆H₆, Et₂O/petroleum ether, EtOH and from H₂O. It can be steam distilled or distilled in a good vacuum. [UV: Ramart-Lucas & Hoch *Bull Soc Chim Fr* [4] 51 824 1932, Dann & Arndt *Justus Liebigs Ann Chem* 587 38 1954, DOI: 10.1002/jlac.19545870104.] The **acid chloride** has **b 154-156°/20mm** [Hanford & Adams *J Am Chem Soc* 57 921 1935, DOI: 10.1021/ja01308a044], and the **amide** crystallises from *C₆H₆ as needles with **m 113°**. [*Beilstein* 6 IV 645.]

2-Phenoxypropionic acid (lactic acid *O*-phenylether) [940-31-8] **C₉H₁₀O₃, M 166.2, m 112-115°, 115-116°, 116-119°, b 105-106°/5mm, 265-266°/758mm, pK²⁵ 3.11.** Crystallise the acid from water. [*Beilstein* 6 H 163, 6 II 158, 6 III 614.]

Phensuximide (*N*-methyl-2-phenylsuccinimide) [86-34-0] **C₁₁H₁₁NO₂, M 189.2, m 71-73°.** Crystallise phensuximide from hot 95% EtOH (m 72-73°). At 25° 1g of the imide dissolves in 1g of *C₆H₆, 18g of Et₂O, 9.5g of EtOH, 5.1g of MeOH and 235g of H₂O. [*Beilstein* 21 II 300, 21 III/IV 5465.] It is an anticonvulsant and an antiepileptic drug.

Phenylacetic acid [103-82-2] **C₈H₈O₂, M 136.2, m 76-77°, 77.5°, 76-79°, b 140-150°/20mm, 265.5°/760mm, d 1.1, pK₁ -7.59 (aqueous H₂SO₄), pK₂ 4.31.** Crystallise the acid from petroleum ether (b 40-60°), isopropyl alcohol, 50% aqueous EtOH or hot water (m 77.8-78.2°). Dry it *in vacuo*. It can be distilled under a vacuum. [*Beilstein* 9 II 294, 9 III 2169.] Present in faint odour of flowers as well as horse urine. **Phenyl acetate** [122-79-2] **C₈H₈O₂, M 136.2,** has **m -30°, b 78°/10mm, 195-196°/atm, d₄²⁰ 1.079, n_D²² 1.5039.** Phenyl acetate acid is freed from phenol and acetic acid by washing (either directly or as a solution in pentane) with aqueous 5% Na₂CO₃, then with saturated aqueous CaCl₂, drying with CaSO₄ or Na₂SO₄, and fractionally distilling under reduced pressure. [*Beilstein* 6 II 153, 6 III 595, 6 IV 611.] **Phenylacetamide** [103-81-1] **C₁₈H₉NO, M 135.2,** has **m 155°, 156°, 157°, 158.5°.** Crystallise the acetamide repeatedly from absolute EtOH, EtOAc (m 160-161°)

or H₂O (m 159-160°). Dry it *in vacuo* over P₂O₅. [Beilstein 9 H 347, 9 III 2193, 9 IV 1632.]

Phenylacetone (1-phenylpropan-2-one) [103-79-9, 136675-26-8] C₉H₁₀O, M 134.2, m -15°, b 69-71°/3mm, 214-216°/760mm, d₄²⁰ 1.006, n_D²⁰ 1.516. Convert the ketone to the *semicarbazone* and crystallise it three times from EtOH (m 186-187°). The semicarbazone is then hydrolysed with 10% phosphoric acid, and the ketone is recovered by distillation. [Kumler et al. *J Am Chem Soc* 72 1463 1950, DOI: 10.1021/ja01160a010; Beilstein 7 H 303, 7 I 161, 7 II 223, 7 III 1037, 7 IV 687.]

4'-Phenylacetophenone (4-acetylbiphenyl) [92-91-1] C₁₄H₁₂O, M 196.3, m 117-123°, 118-121°, 120.3-121.2°, b 168°/8mm, 196-210°/18mm, 325-327°/760mm. Crystallise it from EtOH or acetone. It can also be distilled under reduced or atmospheric pressure. The *semicarbazone* has m 131-132° (aqueous EtOH). [Beilstein 7 H 443, 7 III 2134, 7 IV 1407.]

Phenylacetylene [536-74-3] C₈H₆, M 102.1, m -45°, -44.8°, b 75°/80mm, 142-144°/760mm, d₄²⁰ 0.930, n_D²⁵ 1.5463, pK ~19. Distil phenylacetylene through a spinning band column. It should be filtered through a short column of alumina before use [Campbell & Campbell *Org Synth Coll Vol* 4 763 1963, DOI: 10.15227/orgsyn.030.0072; for pK see Brandsma *Preparative Acetylenic Chemistry*, 1st Edn Elsevier 1971, p. 15, ISBN 0444409475]. [Beilstein 5 IV 1525.]

Phenylalaninol (2-amino-3-phenylpropan-1-ol) [*R*-(+)- 5267-64-1, *S*-(-)- 3182-95-4] C₉H₁₃NO, M 151.2, m 91-92°, 91.5°, 92-94°, b 80°/11mm (Kügelrohr), [α]_D²⁰ (+) and (-) 28, [α]_D²⁰⁻²⁵ (+) and (-) 23-28.7 (c 1-5, EtOH), pK_{Est} ~9.3. It has been recrystallised from Et₂O, *C₆H₆/petroleum ether (b 40-60°) or toluene and distilled in a vacuum. It has been purified by dissolving in Et₂O, drying over K₂CO₃, filtering, evaporating to a small volume, cooling in ice and collecting the plates. Store them in the presence of KOH (i.e. CO₂—free atm). [Karrer & Ehrhardt *Helv Chim Acta* 34 2202 1951, DOI: 10.1002/hlca.19510340714; Oeda *Bull Chem Soc Jpn* 13 465 1938, DOI:10.1246/bcsj.13.465.] The *picrate* has m 141-141.5° (from EtOH/petroleum ether). The *hydrogen oxalate* has m 177°, 161-162° [Hunt & McHale *J Chem Soc* 2073 1957, DOI: 10.1039/JR9570002073]. The *racemate* has m 87-88° from *C₆H₆/petroleum ether (75-77° from Et₂O), and the *hydrochloride* has m 139-141° [Fodor et al. *J Chem Soc* 1858 1951, DOI: 10.1039/JR9510001858]. [Beilstein 13 IV 1920.]

3-Phenylallyl chloride (cinnamyl chloride) [*E*: 18685-01-3, *Z*: 39199-93-4] C₉H₉Cl, M 152.6, *trans*: m -19° 7-8°, b 92-93°/3mm, 108°/12mm, d₄²⁵ 1.086, n_D²⁵ 1.5802, *cis*: 85°/3mm, d₄²⁵ 1.0891, n_D²⁵ 1.5746. Distil the chloride under vacuum three times from K₂CO₃. [Hatch & Alexander *J Am Chem Soc* 72 5643 1950, DOI: 10.1021/ja01168a078; Beilstein 5 III 1186.] **LACHRYMATORY.**

Phenyl 4-aminosalicylate (Phenamisal) [133-11-9] C₁₃H₁₁NO₃, M 229.2, m 150-152°, 153°, b 406.4°/760mm, d 1.32, pK_{Est(1)}~2.0 (NH₂), pK_{Est(2)}~9.7 (OH). Crystallise the ester from EtOH (m 155°, also 149-150.5°), aqueous EtOH (m 147-149°), or isopropanol. It is an antibiotic and is **tuberculostatic**. [Beilstein 13 IV 1979.]

4-Phenylanisole (4-methoxybiphenyl) [361-37-6] C₁₃H₁₂O, M 184.2, m 89-90°, 89.9-90.1°, 157°/10mm, 329.5°/760mm. Crystallise the biphenyl from *benzene/petroleum ether. Dry it under vacuum in an Abderhalden pistol. It has λ_{max} at 259.5nm (hexane). [Beilstein 6 H 674, 6 II 625, 6 III 3321, 6 IV 4600.]

9-Phenylanthracene [602-55-1] C₂₀H₁₄, M 254.3, m 153-154°, 156°, b 417°/760mm. Chromatograph it on alumina in *C₆H₆ and recrystallise it from AcOH or toluene. [Beilstein 5 H 725, 5 II 639, 5 III 2462.]

4-Phenylazobenzoyl chloride [104-24-5] C₁₃H₉ClN₂O, M 244.7, m 93°, 95°. Crystallise the acid chloride from petroleum ether (b 60-80°). It is moisture sensitive, store accordingly. [Beilstein 16 III 224.]

4-Phenylazo-1-naphthylamine [131-22-6] C₁₆H₁₃N₃, M 247.3, m 125-125.5°. Crystallise the dye from cyclohexane or aqueous EtOH. [Brode et al. *J Am Chem Soc* 74 4641 1952, DOI: 10.1021/ja01138a059; Beilstein 16 H 361, 16 III 406, 16 IV 546.]

4-Phenylazophenacyl bromide [62625-24-5] $C_{14}H_{11}BrN_2O$, M 317.3, 303.2, m 103-104°, 103-105°, 115°. Purify the bromide on a column silica gel, using petroleum ether/ Et₂O (9:1 v/v) as solvent. It forms orange yellow crystals from *C₆H₆ (m 113-114°) or petroleum ether/Et₂O (m 114.5-115°). [Beilstein 16 II 22, 16 III 53, 16 IV 79.]

4-Phenylazophenol (4-hydroxyazobenzene) [1689-82-3] $C_{12}H_{10}N_2O$, M 198.2, m 152-155°, 153°, 154°, 155°, pK₁²⁵ -0.93, pK₂²⁵ 8.2. Crystallise the dye from *benzene or 95% EtOH. [Beilstein 16 II 38, 16 III 86, 16 IV 159.]

Phenyl benzenethiosulfonate (diphenyldisulfoxide) [1212-08-4] $C_{12}H_{10}O_2S_2$, M 250.3, m 36-37°, 45-46°, 45-47°, 36-53°. Recrystallise the disulfoxide from EtOH or MeOH. It has also been purified from phenylsulfide impurities by dissolving in CHCl₃, washing with aqueous saturated NaHCO₃, drying (Na₂SO₄), filtering, evaporating the filtrate, and the residual oil is passed through a silica gel column (600g) and eluted with hexane/*C₆H₆ (1L, 4:1, eluting PhSSPh) then *C₆H₆ (1L) which elutes PhSSO₂Ph. [Trost & Massiot *J Am Chem Soc* 99 4405 1977, DOI: 10.1021/ja00455a032; Knoevenagel & Römer *Chem Ber* 56 215 1923, DOI: 10.1002/cber.19230560130; Beilstein 11 IV 220.]

2-Phenyl benzoate [93-99-2] $C_{13}H_{10}O_2$, M 198.2, m 68-70°, 69°, 69.5°, 70°, 71°, b 298-299°/atm. Crystallise the ester from EtOH using *ca* twice the volume needed for complete dissolution at 69°. [Beilstein 9 IV 303.]

Phenyl-1,4-benzoquinone [363-03-1] $C_{12}H_8O_2$, M 184.2, m 114-115°, 115°. Crystallise the quinone from heptane, petroleum ether (b 60-70°), *C₆H₆ (m 113.5-114.5°) or EtOH (m 112-113°) and sublime it *in vacuo*. [Carlson & Miller *J Am Chem Soc* 107 479 1985, DOI: 10.1021/ja00288a035; Beilstein 7 H 740, 7 III 3764.]

1-Phenylbiguanide [102-02-3] $C_8H_{11}N_5$, M 177.2, m 135-142°, 144-146°, d 1.33, pK₁³² 2.16, pK₂³² 10.74. Crystallise the biguanide from water or toluene. [Beilstein 12 H 370, 12 I 236, 12 III 807.] The *hydrochloride* [55-57-2] has m 244-247°.

S-(-)-1-Phenylbutanol [22135-49-5] $C_8H_{10}O$, M 150.2, m 46-47°, 46-48°, 49°, b 90-92°/2mm. [α]_D¹⁸ -51.4 (c 5, CHCl₃), -44.7 (c 5.13, *C₆H₆). Purify the alcohol by distillation, and the distillate crystallises on cooling. The *hydrochloride* has [α]_D²⁰ +45.1 (c 4.8, *C₆H₆). The (-)-*hydroperoxide* has b 58°/0.005mm, n_D²⁰ 1.5123, α _D¹⁸ -2.14, (l = 0.5dcm, neat). [Holding & Ross *J Chem Soc* 145 1954, DOI: 10.1039/JR9540000145; Davies & Feld *J Chem Soc* 4637 1958, DOI: 10.1039/JR9580004637.] The (\pm)-*racemate* has b 73°/0.05mm, and its 4-nitrophenylhydrazone has m 58°. [Beilstein 6 IV 3272.]

2-Phenylbutyramide [90-26-6] $C_{10}H_{13}NO$, M 163.2, m 86°, 87°. Crystallise the amide from H₂O, EtOH, Et₂O/petroleum ether or *C₆H₆. [Beilstein 9 I 212, 9 II 356, 9 III 2466.]

2-Phenylbutyric acid [R-(-)- 938-79-4, S-(+)- 4286-15-1] $C_{10}H_{12}O_2$, M 164.2, b 102-104°/760mm, d₄²⁰ 1.056, n_D²⁰ 1.521, [α]_D²⁰ (+) and (-) 96 (c 2.5, *C₆H₆), [α]_D²³ (-) and (+) 5.8 (neat), pK_{Est} ~4.3. Purify the acids by distillation at atmospheric pressure using an efficient column. The *acid chlorides* have b 106-107°/20mm, [α]_D¹⁸ (-) and (+) 108 (c 2, *C₆H₆). [Levene et al. *J Biol Chem* 100 589 1933, Gold-Aubert DOI: 10.1002/hlca.19580410609; *Chim Acta* 41 1512 1958, ORD in heptane: Rothen & Levene *J Chem Phys* 7 975 1939, DOI:org/10.1063/1.175037; Beilstein 9 III 2461.]

3-Phenylbutyric acid [R-(-)- 772-14-5, S-(+)- 772-15-6] $C_{10}H_{12}O_2$, M 164.2, b 94-95°/3mm, 134°/4mm, d₄²⁰ 1.066, n_D²⁵ 1.5167, [α]_D²⁰ (-) and (+) 57 (c 1, *C₆H₆), pK₂₅ 4.40. Purify the acids as the 2-isomer above, i.e. by distillation, but under a good vacuum. [Prelog & Scherrer *Helv Chim Acta* 42 2227 1959, DOI: 10.1002/hlca.19590420651; Levene & Marker *J Biol Chem* 93 761 1932, 100 685 1933, Cram *J Am Chem Soc* 74 2137 1952, DOI: 10.1021/ja01129a002.] The *R-amide* crystallises from H₂O, with m 101.5-102°, and [α]_D²⁰ -16.5 (c 1.2, EtOH). The *racemic acid* has m 39-40°, b 134-136°/6mm, 158°/12mm [Marvel et al. *J Am Chem Soc* 62 3499 1940, DOI: 10.1021/ja01869a059]. [Beilstein 9 IV 1813.]

4-Phenylbutyric acid [1821-12-1] $C_{10}H_{12}O_2$, M 164.2, m 48°, 50°, 52°, 49-53°, b 165°/10mm, 340.4°/atm, pK^{25} 4.76. Crystallise the acid from petroleum ether (b 40-60°). [Beilstein 9 IV 1811.]

O-Phenyl chlorothionoformate [1005-56-7] C_7H_5ClOS , M 172.6, b 81-83°/6mm, 91°/10mm, d_4^{20} 1.276, n_D^{20} 1.585. Purify it by dissolving in $CHCl_3$, washing with H_2O , drying ($CaCl_2$), filtering, evaporating and distilling twice under a vacuum to give a clear yellow liquid. **It is reactive and POISONOUS—work in a fume cupboard.** Store it in sealed ampoules under N_2 . A possible impurity is **O,O'-diphenyl thiocarbonate** which has m 106° and remains behind in the distilling flask. [Bögemann et al. in *Methoden Der Organischen Chemie (Houben-Weyl)* 4th edn (E. Müller Ed.) Vol 9 Schwefel-Selen-Tellur Verbindungen pp. 807-808 1955, Rivier & Schalch *Helv Chim Acta* 6 605 1923, DOI: 10.1002/hlca.19230060163; Kalson *Chem Ber* 20, 2384 1887, DOI: 10.1002/cber.18870200256; Rivier & Richard *Helv Chim Acta* 8 490 1925, DOI: 10.1002/hlca.19250080174; Schönberg & Varga *Justus Liebigs Ann Chem* 483 176 1930, DOI: 10.1002/jlac.19304830117; Schönberg & Vargha *Chem Ber* 64 1390 1931, DOI: 10.1002/cber.19310640628; Beilstein 6 III 609.] Reagent for thionocarbonylation, and to form phenoxythiocarbonyl esters.

Phenyl cinnamate [2757-04-2] $C_{15}H_{12}O_2$, M 224.3, m 75-76°, b 205-207°/15mm. Crystallise the cinnamate from EtOH (2ml/g). It can also be distilled under reduced pressure. [Womack & McWhirter *Org Synth Coll Vol* 3 714 1955, DOI: 10.15227/orgsyn.020.0077; Beilstein 9 H 583, 9 II 387, 9 III 2689, 9 IV 2011.]

α -Phenylcinnamic acid (2,3-diphenylprop-2-enoic acid) [*cis-E* 91-48-5, *trans-Z* 91-47-4] $C_{15}H_{12}O_2$, M 224.3, m 174°(*cis*), m 138-139°(*trans*), pK^{25} 4.8 (60% aqueous EtOH). Recrystallise the acid from Et_2O /petroleum ether. Crystallise the *cis-isomer* from petroleum ether or EtOH (m 174°) and has a pK^{25} of 4.44, and the *cis-amide* from aqueous Me_2CO (m 174°). Crystallise the *trans-isomer* from Et_2O /petroleum ether or EtOH (m 140°), and the *trans-amide* from $CHCl_3$ /petroleum ether (m 167-168°). [Beilstein 6 H 691, 9 III 3414.]

***o*-Phenylenediamine** [95-54-5] $C_8H_8N_2$, M 108.1, m 100-101°, 102-104°, 103-104°, 252°/atm, d 1.03, pK_1^{25} 0.67 (aqueous H_2SO_4), pK_2^{25} 4.47 (4.85). Crystallise the diamine from aqueous 1% sodium hydrosulfite (charcoal), wash it with ice-water and dry it in a vacuum desiccator, or sublime it *in vacuo*. It has been purified by recrystallisation from toluene and zone refined [Anson et al. *J Am Chem Soc* 108 6593 1986, DOI: 10.1021/ja00281a024]. Purification by refluxing a CH_2Cl_2 solution containing charcoal is also carried out followed by evaporation and recrystallisation [Koola & Kochi *J Org Chem* 52 4545 1987, DOI: 10.1021/jo00229a022], protect from light. The *acetate* has m 186°. [Beilstein 13 IV 38.]

***m*-Phenylenediamine** [108-45-2] $C_8H_8N_2$, M 108.1, m 61-63°, 62-63°, 62.5°, 63-64°, b 146°/22mm, 282-284°/760mm, 284-287°/atm, d_{10}^{10} 1.1422, $n_D^{57.7}$ 1.6340, pK_1^{25} 2.41, pK_2^{25} 4.98. Purify the diamine by distillation under a vacuum followed by recrystallisation from EtOH (rhombs) and if necessary redistillation. It should be protected from light; otherwise it darkens rapidly. [Neilson et al. *J Chem Soc* 361 p371 1962, DOI: 10.1039/JR9620000361; IR: Katritzky & Jones *J Chem Soc* 3674 1959, DOI: 10.1039/JR9590003674; 2058 1959, DOI: 10.1039/JR9590002058; UV: Forbes & Leckie *Can J Chem* 36 1371 1958, DOI: 10.1139/v58-202.] The *hydrochloride* has m 277-278°, and the *bis-4-chlorobenzenesulfonyl* derivative has m 220-221° from H_2O (214-215°, from MeOH/ H_2O) [Runge & Pfeiffer *Chem Ber* 90 1757 1957, DOI: 10.1002/cber.19570900910]. The *acetate* has m 191°. [Beilstein 13 IV 79.]

***p*-Phenylenediamine** [106-50-3] $C_8H_8N_2$, M 108.1, m 140°, 145-147°, b 267°/atm, pK_1^{25} 2.89, pK_2^{25} 6.16. Crystallise the diamine from EtOH or *benzene, and sublime it *in vacuo*; protect it from light. The *acetate* has m 304°. [Beilstein 13 IV 104.] Useful for making polymers and composites.

***o*-Phenylenediamine dihydrochloride** [615-28-1] $C_8H_8N_2 \cdot 2HCl$, M 181.1, m 258°(dec). Crystallise the salt from dilute HCl (60ml conc HCl, 40ml water, with 2g stannous chloride), after treatment of the hot solution with charcoal by adding an equal volume of concentrated HCl and cooling in an ice-salt mixture. The crystals are washed with a small amount of concentrated HCl and dried in a vacuum desiccator over NaOH. [Beilstein 13 IV 38.] It is a peroxidase substrate useful in ELISA procedures.

1-Phenyl-1,2-ethanediol [*R*-($-$)- 16355-00-3, *S*-($+$)- 25779-13-9] $\text{C}_8\text{H}_{10}\text{O}_2$, **M 138.2**, **m 64-67°**, **65-66°**, $[\alpha]_{\text{D}}^{24}$ ($-$) and ($+$) **40.5** (c **2.8**, H_2O), $[\alpha]_{\text{D}}^{20}$ ($-$) and ($+$) **39** (c **3**, EtOH). Purify the diol by recrystallisation from $^*\text{C}_6\text{H}_6$ /ligroin and sublime it at 1-2mm. [Arpesella et al. *Gazzetta* **85** 1354 1955, Prelog et al. *Helv Chim Acta* **37** 221 1954, DOI: 10.1002/hlca.19540370127; *Beilstein* **6** IV 5939.]

***dl*-1-Phenylethanol** (*dl*- α -methylbenzyl alcohol) [98-85-1; 13323-81-4] $\text{C}_8\text{H}_{10}\text{O}$, **M 122.2**, **m 19-20°**, **20°**, **b 60.5-61.0°/3mm**, **106-107°/22-23mm**, **203-205°/atm**, d_4^{20} **1.01**, n_{D}^{20} **1.5254**. Purify the alcohol *via* its hydrogen phthalate. [See Houssa & Kenyon *J Chem Soc* 2260 1930, DOI: 10.1039/JR9300002260.] Shake it with a solution of ferrous sulfate, and the alcohol layer is washed with distilled H_2O , dried (MgSO_4) and fractionally distilled. [*Beilstein* **6** II 444.]

2-Phenylethanol [60-12-8] $\text{C}_8\text{H}_{10}\text{O}$, **M 122.2**, **m -27°**, **b 215-217°/atm**, **219-221°/atm**, d_4^{20} **1.020**. Purify the ethanol by shaking it with a solution of ferrous sulfate, and the alcohol layer is washed with distilled water and fractionally distilled. [*Beilstein* **6** IV 3067.] It is an antimicrobial antiseptic.

Phenyl ether (diphenyl ether) [101-84-8] $\text{C}_{10}\text{H}_{12}\text{O}$, **M 170.2**, **m 25.0° to 27.0°**, **28°**, **b 83-84°/1mm**, **138°/21mm**, **257°/760mm**, d_4^{20} **1.074**, $n_{\text{D}}^{30.7}$ **1.57596**. Crystallise the ether from 90% EtOH. Melt it, wash it with 3M NaOH and water, dry it with CaCl_2 and fractionally distil it under reduced pressure. Fractionally recrystallise it from its melt and store over P_2O_5 . [*Beilstein* **6** IV 562.] IRRITANT solvent with a fruity odour used to perfume soaps.

1-Phenylethyl isocyanate (α -methylphenyl isocyanate) [*R*-($+$)- 33375-06-3, *S*-($-$)- 14649-03-7] $\text{C}_9\text{H}_9\text{NO}$, **M 147.2**, **b 82-83°/12-14mm**, d_4^{20} **1.045**, n_{D}^{20} **1.513**, $[\alpha]_{\text{D}}^{24}$ ($+$) and ($-$) **2** (c **3.5**, $^*\text{C}_6\text{H}_6$), ($+$) and ($-$) **10.5** (neat). Purify the isocyanates by fractional distillation under a vacuum. With ammonia they give the *ureido* derivatives which crystallise from H_2O with **m 121-122°**, $[\alpha]_{\text{D}}^{25}$ ($+$) and ($-$) **48.8**. [For the optical resolution of chiral amines, e.g. 2,3-dimethylethyleneimine see Cairns *J Am Chem Soc* **63** 871 1941, DOI: 10.1021/ja01848a504.] The *racemate* has **b 90-94°/3mm**, **96°/18mm** [Seifkan *Justus Liebigs Ann Chem* **562** 75 1949, DOI: 10.1002/jlac.19495620202]. [*Beilstein* **12** IV 2443.]

(\pm)-*p*- α -Phenylethylphenol [1988-89-2] $\text{C}_{14}\text{H}_{14}\text{O}$, **M 198.3**, **m 56.0-56.3°**, **64°**, **b 126°/0.4mm**, **165-170°/5mm**, **315-316°/742.2mm**, pK_{Est} **~10.3**. Crystallise the phenol from petroleum ether. *S*-($+$)-*enantiomer* has $[\alpha]_{\text{D}}$ **+10.3** ($^*\text{C}_6\text{H}_6$). [Okamoto et al. *Bull Chem Soc Jpn* **39** 299 1966, DOI:10.1246/bcsj.39.299.]

5-(α -Phenylethyl)semioxamazide [93-95-8] $\text{C}_{10}\text{H}_{13}\text{N}_3\text{O}_2$, **M 207.1**, **m 167-168°** (*L*-), **157°** (*dl*-). Crystallise it from EtOH. The *S*-($-$)-*enantiomer* [6152-25-6] has $[\alpha]_{\text{D}}^{25}$ **-102.5** (c **1**, CHCl_3). [Leonard & Boyer *J Org Chem* **15** 42 1950, DOI: 10.1021/jo01147a007.]

Phenylhydrazine [100-63-0] $\text{C}_6\text{H}_8\text{N}_2$, **M 108.1**, **m 19.5°**, **23°**, **b 71.8°/1mm**, **137-138°/18mm**, **241-242°/760mm**, d_{D}^{20} **1.10**, n_{D}^{20} **1.607**, pK_1^{20} **-5.2** (aqueous H_2SO_4), pK_2^{20} **5.27**. Purify phenylhydrazine by chromatography, then crystallise it from petroleum ether (b 60-80°)/ $^*\text{benzene}$. Store it in the dark under N_2 as it turns yellow, then red, on exposure to air. It is best stored as the *hydrochloride salt*; see below. [Coleman *Org Synth Coll Vol I* 442 1941, DOI: 10.15227/orgsyn/02.0071; Shaw & Stratton *J Chem Soc* 5004 1962, DOI: 10.1039/JR9620005004; *Beilstein* **15** IV 50.]

Phenylhydrazine hydrochloride [59-88-1] $\text{C}_6\text{H}_8\text{N}_2 \cdot \text{HCl}$, **M 144.5**, **m 244°**, **250-254°**(dec). Dissolve 100g of phenylhydrazine hydrochloride in 200ml of warm H_2O (60-70°) during 1-3 hours, then add 1L of boiling EtOH. The solution is filtered, while still hot, through Whatman No 2 filter paper and cooled in a refrigerator. The precipitate is collected on a medium sintered-glass filter and recrystallised twice this way, then washed with cold EtOH, dried thoroughly and stored in a stoppered brown bottle. [Peterson et al. *Anal Chem* **29** 144 1957, DOI: 10.1021/ac60121a042.] Hough, Powell and Woods [*J Chem Soc* 4799 1956, DOI: 10.1039/JR9560004799] boiled the hydrochloride with three times its weight of water, filtered hot (charcoal), added one-third volume of concentrated HCl and cooled to 0°. The crystals were washed with acetone, and dried over P_2O_5 under vacuum. The salt has also been crystallised from 95% EtOH, and it can be sublimed. [Coleman *Org Synth Coll Vol I* 442 1941, DOI: 10.15227/orgsyn/02.0071; *Beilstein* **15** III 71.]

Phenylhydroxylamine (*N*-hydroxyaniline) [100-65-2] C_6H_7NO , M 109.1, m 81°, 82°, 83-84°, pK^{25} 3.2. Impure base deteriorates rapidly. Crystallise it from H_2O , $*C_6H_6$ or $*C_6H_6$ /petroleum ether (40-60°). The *picrate* has m 186° (from EtOH), and the *benzenesulfonate salt* has m 70° (dec) (EtOH/ $*C_6H_6$). [Beilstein 15 H 2, 15 I 3, 15, II 4, 15 III 5. 15 IV 4.]

2-Phenyl-1,3-indandione [83-12-5] $C_{15}H_{10}O_2$, M 222.2, m 144-148°, 149-151°, pK^{20} 4.12 (1% aqueous MeOH). Crystallise the dione from EtOH (m 156°) or $CHCl_3$ (m 148-150°). [Beilstein 7 H 808, 7 III 4100, 7 IV 2570.]

Phenylisocyanate [103-71-9] C_7H_5NO , M 119.1, m -30°, b 45-47°/10mm, 162-163°/atm, d_4^{20} 1.093, n_D^{24} 1.536. Distil phenylisocyanate under reduced pressure from P_2O_5 . [Beilstein 12 IV 864.] Used for derivatisation.

8-Phenylmenthol [1*R*,2*S*,5*R*-(*-*)- 65253-04-5, 1*S*,2*R*,5*S*-(*+*)- 57707-91-2] $C_{16}H_{24}O$, M 232.4, $[\alpha]_D^{20}$ (–) and (+) 26 (c 2, EtOH). Dissolve the menthol in toluene, dry (Na_2SO_4), evaporate and chromatograph it on a silica gel column and eluting with 5% Et_2O in petroleum ether to give an oil with the desired rotation. Its IR has ν_{max} at $3420cm^{-1}$ (OH) with consistent 1H NMR [Ensley et al. *J Org Chem* 43 1610 1978, DOI: 10.1021/jo00402a037; Whitesell et al. *Tetrahedron* 42 2993 1986, DOI: 10.1016/S0040-4020(01)90590-3; Bednarski & Danishefsky *J Am Chem Soc* 108 7060 1986, DOI: 10.1021/ja00282a035].

2-Phenylnaphthalene [612-94-2] $C_{16}H_{12}$, M 204.3, m 102-103°, 103.5°, 103-104°, 105°, b 185-190°/5mm, 357-358°/760mm. Chromatograph it on alumina in $*benzene$ and crystallise it from aqueous EtOH or MeOH/EtOH. It has been sublimed. It has λ_{max} at 248 and 286nm (methylcyclohexane). The *2,4,7-trinitrofluoren-9-one* has m 169.5-170.5° (from EtOH/ $*C_6H_6$). [Beilstein 5 H 687, 5 II 603, 5 III 2231, 5 IV 2412.]

***N*-Phenyl-1-naphthylamine** [90-30-2] $C_{16}H_{13}N$, M 219.3, m 60-62°, 63.7-64.0°, b 226°/15mm, pK_{Est} ~0.1. Crystallise it from EtOH, petroleum ether or $*C_6H_6$ /EtOH. Dry it under vacuum in an Abderhalden pistol. It has λ_{max} at 252nm [Beilstein 12 H 1224.]

***N*-Phenyl-2-naphthylamine** [135-88-6] has m 107.5-108.5°, 110°, b 395°/atm, pK_{Est} ~0.5. Crystallise it from EtOH, MeOH, glacial acetic acid or $*benzene$ /hexane. [Beilstein 12 H 1275, 12 I 535, 12 II 716, 12 III 2991.]

4-Phenylphenacyl bromide [135-73-9] $C_{14}H_{11}BrO$, M 275.2, m 126°, b 370°/atm, d 1.379. Crystallise (charcoal) the bromide from EtOH (15ml/g), or ethyl acetate/petroleum ether (b 90-100°). [Beilstein 7 III 2137.] **IRRITANT.**

(±)-2-Phenylpropanal [93-53-8] $C_9H_{10}O$, M 134.2, b 92-94°/12mm, 206°/760mm, d_4^{20} 1.001, n_D^{20} 1.5183. It may contain up to 15% of acetophenone. Purify it *via* the bisulfite addition compound [Lodge & Heathcock *J Am Chem Soc* 109 3353 1987, DOI: 10.1021/ja00245a027] and see Chapter 2 for preparation and decomposition of bisulfite adducts. [Beilstein 7 IV 695.]

Phenylpropionic acid [637-44-5] $C_9H_8O_2$, M 146.2, m 136°, 137°, 137.8-138.4°, pK^{25} 2.23. Crystallise the acid from $*benzene$, CCl_4 (m 136°) or aqueous EtOH. The *S-benzylisothiuronium salt* has m 184-186° (from EtOH). [Beilstein 9 II 436, 9 III 3061, 9 IV 2327.]

***RS*-2-Phenylpropionic acid** [492-37-5] $C_9H_{10}O_2$, M 150.2, m 16-16.5°, b 153-155°/20mm, 189°/48mm, 260-262°/760mm, d_4^{20} 1.10, n_D^{20} 1.522, pK^{25} 4.3. Fractionally distil the acid, or recrystallise it from petroleum ether (b 40-60°) with strong cooling (see references below). [Beilstein 9 II 348.]

2-Phenylpropionic acid [*R*-(*-*)- 7782-26-5, *S*-(*+*)- 7782-24-3] has m 30.3-31°, 30-32°, b 115°/1-2mm, 142°/12mm, $[\alpha]_D^{20}$ (–) and (+) 99.7 (l = 1 dcm, neat), (–) and (+) 89.1 (c 1.7, EtOH), (–) and (+) 75 (c 1.6, $CHCl_3$). Purify the acids by vacuum distillation and by recrystallisation from petroleum ether. The *S-anilide* has m 103-104° (from H_2O or $CHCl_3$ / $*C_6H_6$), $[\alpha]_D^{25}$ +47 (c 9, Me_3CO) [Argus & Kenyon *J Chem Soc* 916 1939, DOI: 10.1039/JR9390000916; Campbell & Kenyon *J Chem Soc* 25 1946, DOI: 10.1039/JR9460000025;

Levene et al. *J Biol Chem* **88** 27, 34 1930]. [*Beilstein* **9** III 2417, **9** IV 1779.]

3-Phenylpropionic acid (hydrocinnamic acid) [501-52-0] has **m 45-48°, 48-48.5°, pK²⁵ 4.56**. Crystallise the acid from *benzene, CHCl₃ or petroleum ether (b 40-60°). Dry it in a vacuum. [*Beilstein* **9** H 508.] Food flavour and fragrance agent.

3-Phenylpropyl bromide [637-59-2] **C₉H₁₁Br, M 199.1, b 110°/12mm, 128-129°/29mm, d₄²⁰ 1.31**. Wash the bromide successively with concentrated H₂SO₄, water, 10% aqueous Na₂CO₃ and again with water, then dry it with CaCl₂ and fractionally distil it just before use. [*Beilstein* **5** IV 982.]

Phenylpyruvic acid [156-06-9] **C₉H₈O₃, M 164.2, m 152°, 150-154°, 155°, 158-159°, pK_{Est} ~2.1**. Recrystallise the acid from *C₆H₆. The **phenylhydrazone** has **m 173°** [Zeller *Helv Chim Acta* **26** 1614 1943, DOI: 10.1002/hlca.19430260524; Hopkins & Chisholm *Can J Research* [B] **24** 89 1946, DOI: 10.1139/cjr46b-016]. The **2,4-dinitrophenylhydrazone** has **m 162-164° (189°, 192-194°)** [Fones *J Org Chem* **17** 1534 1952, DOI: 10.1021/jo50011a022]. [*Beilstein* **10** IV 2760.]

Phenyl salicylate (Salol) [118-55-8] **C₁₃H₁₀O₃, M 214.2, m 41.5°, 41.8-42.6°, 43°, 173°/atm, d 1.25, pK_{Est} ~9.9**. Fractionally crystallise salol from its melt, then crystallise it from *benzene. [*Beilstein* **10** IV 154.]

3-Phenylsalicylic acid [304-06-3] **C₁₃H₁₀O₃, M 214.3, m 186°, 186-187.5°, pK_{Est(1)}~2.8 (CO₂H), pK_{Est(2)}~11.0 (OH)**. Dissolve the acid in *ca* 1 equivalent of saturated aqueous Na₂CO₃, filter and precipitate it by adding 0.8 equivalents of M HCl. Crystallise it from ethylene dichloride (charcoal), and sublime it at 0.1mm. [Brooks et al. *J Chem Soc* 661 1961, DOI: 10.1039/JR9610000661.]

1-Phenylsemicarbazide [103-03-7] **C₇H₉N₃O, M 151.2, m 171-174°, 172°, 173.5°, 174-176°**. Crystallise it from water and dry it in a vacuum over KOH. Store at -20°. [*Beilstein* **15** H 287, **15** II 106, **15** III 184, **15** IV 180.]

4-Phenylsemicarbazide [537-47-3] **C₇H₉N₃O, M 151.2, m 122°, 122-125°**. Crystallise it from water and dry it in a vacuum over KOH. Store at -20°. [*Beilstein* **12** II 221, **12** III 822.]

Phenylsuccinic acid [*R*(-)- 46292-93-7, *S*(+)- 4036-30-1] **C₁₀H₁₀O₄, M 194.2, m 173-176°, 178.5-179°, 179-180°, [α]_D²⁵ (-) and (+) 171 (c 2, Me₂CO), [α]_D²⁶⁻³⁰ (-) and (+) 148 (c 0.27-5, EtOH), pK₁²⁵ 3.78, pK₂²⁵ 5.55**. Purify the acids by re-precipitation from alkali and recrystallisation from H₂O. [Naps & Johns *J Am Chem Soc* **62** 2450 1940, DOI: 10.1021/ja01866a053; Fredga & Matell *Bull Soc Chim Belg* **62** 47 1953, DOI: 10.1002/bscb.19530620108; Wren & Williams *J Chem Soc* **109** 572 1916, DOI: 10.1039/CT9160900572.] The **racemate** [635-51-8] has **m 166-168°, 168°** after recrystallisation from H₂O or MeCN. Its ***S*-benzylisothiuronium salt** has **m 164-165°** (from EtOH) [Friediger & Pedersen *Acta Chem Scand* **9** 1425 1955, DOI: 10.3891/acta.chem.scand.09-1425]. [*Beilstein* **9** IV 3351.]

Phenyltoloxamine [2-(2-dimethylaminoethoxy)-diphenylmethane] hydrochloride [6152-43-8] **C₁₇H₂₁NO·HCl, M 291.8, m 119-120°, pK²⁵ 9.3 (free base)**. Crystallise the salt from isobutyl methyl ketone. The **free base** [92-12-6] **M 255.4, has b 144°/1mm**. [*Beilstein* **6** III 3351, **6** IV 4630.] It is an antihistamine with sedative and analgesic properties.

Phenyl 4-tolylcarbonate [13183-20-5] **C₁₄H₁₂O₃, M 228.2, m 94°**. Purify the carbonate by preparative GLC with 20% Apiezon on Embacel, and sublime it *in vacuo*. [*Beilstein* **6** H 398.]

1-Phenyl-2,2,2-trifluoroethanol [*R*(-)- 10531-50-7, *S*(+)- 340-06-7] **C₈H₇F₃O, M 176.1, b 74-76°/10mm, 125-127°/760mm, d₄²⁰ 1.301, n_D²⁰ 1.4632, [α]_D²⁰ (-) and (+) 31 (neat)**. Purify the chiral alcohols by fractional distillation preferably in a vacuum. [Morrison & Ridgeway *Tetrahedron Lett* 573 1969, DOI: 10.1016/S0040-4039(01)87751-0; NMR: Pirkle & Beare *J Am Chem Soc* **90** 6250 1968, DOI: 10.1021/ja01024a073.] The **racemate** [340-05-6] has **b 52-54°/2mm, 57-59°/2mm, 64-65°/5mm, d₄²⁰ 1.293, n_D²⁰ 1.457**, and the **2-carbobenzoyl** derivative has **m 137-138°** [Mosher et al. *J Am Chem Soc* **78** 4374 1956, DOI: 10.1021/ja01598a049]. [*Beilstein* **6** IV 3043.]

Phenylurea [64-10-8] $C_7H_8N_2O$, M 136.2, m 145-147°, 147°, 148°, b 238°/atm, $pK_{25}^{-1.45}$ (aqueous H_2SO_4). Crystallise the urea from boiling water (10ml/g) or amyl alcohol (m 149°). Dry it in a steam oven at 100°. The 1:1 *resorcinol complex* has m 115° (from EtOAc/* C_6H_6). [Beilstein 12 H 346, 12 II 204, 12 III 760, 12 IV 734.]

Phloretin [2',4',6'-trihydroxy-3-(*p*-hydroxyphenyl)propiophenone] [60-82-2] $C_{15}H_{14}O_5$, M 274.3, m 260°(dec), 264-271°(dec), $pK_{Est(1)} \sim 7.5$, $pK_{Est(2)} \sim 8.0$, $pK_{Est(3)} \sim 10$, $pK_{Est(4)} \sim 12$ (phenolic OH's). Crystallise phloretin from aqueous EtOH. [Zemplén & Bognár *Chem Ber* 75 1040 1942, DOI: 10.1002/cber.19420750903; Zemplén et al. *Chem Ber* 76 386 1943, DOI: 10.1002/cber.19430760412; Beilstein 8 IV 3518.] Inhibits active transport of glucose into certain cells. It is an antioxidant which inhibits elastase activity, and affect Ca^{2+} channels.

Phloroacetophenone ($2H_2O$) (2',4',6'-trihydroxyacetophenone) [480-66-0] $C_8H_8O_4$, M 186.2, m 218-219°, 219-221°, 218-223°, $pK_{Est(1)} \sim 7.9$, $pK_{Est(2)} \sim 12.0$. Crystallise the ketone from hot H_2O /charcoal (35ml/g). Dry the colorless to pale yellow needles in an oven at 130°, but they pick up water readily on exposure to air. They give a wine-red colour with $FeCl_3$, compare with violet colour with phloroglucinol (below) [Gulati *et al. Org Synth* 15 70 1935, DOI: 10.15227/orgsyn.015.0070]. [Beilstein 8 IV 2729.] It lowers plasma cholesterol in animals, and is used as a matrix for laser desorption/ionisation in acidic glycans and glycopeptide negative ion analysis.

Phloroglucinol ($2H_2O$) (benzene-1,3,5-triol) [6099-90-7 ($2H_2O$), 108-73-6 (anhydrous)] $C_6H_6O_3$, M 126.1, m 217-219°, 219°, 117° (anhydrous), $pK_1^{25} -7.74$ ($HClO_4$), $pK_2^{20} 7.97$ (8.43), $pK_3^{20} 9.23$. Crystallise the triol from water, and store it in the dark under nitrogen. [Clarke & Hartmann *Org Synth Coll Vol* 1 455 1941, DOI: 10.15227/orgsyn.009.0074; Beilstein 6 IV 7361.] It has antispasmodic properties.

o-Phthalic acid [88-99-3] $C_8H_6O_4$, M 166.1, m 205°(dec), 210-211°(dec), 211-211.5°(dec), d 1.593, $pK_1^{25} 2.76$ (3.05), $pK_2^{25} 4.92$ (4.73). Crystallise phthalic acid from water. [Beilstein 9 IV 3167.]

Phthalic anhydride [85-44-9] $C_8H_4O_3$, M 148.1, m 130.8°, 131°, 132°, 130-134°, b 295°/atm, d 1.53. Distil the anhydride under reduced pressure. Purify it from the acid by extracting with hot $CHCl_3$, filtering and evaporating. The residue is crystallised from $CHCl_3$, CCl_4 or *benzene, or sublimed (295 °/atm). Fractionally crystallise it from its melt. Dry it under vacuum at 100°. [Saltiel et al. *J Am Chem Soc* 108 2674 1986, DOI: 10.1021/ja00270a028; Beilstein 17/11 V 253.]

Phthalide [87-41-2] $C_8H_6O_2$, M 134.1, m 72-73°, 75-77°, b 290°/atm, $pK -7.98$ (aqueous H_2SO_4). Crystallise phthalide from water (75ml/g) and dry it in air on filter paper. [Synthesis see Gardner & Naylor *Org Synth Coll Vol* 2 526 1943, DOI: 10.15227/orgsyn.016.0071; Beilstein 17/10 V 7.]

Phthalimide [85-41-6] $C_8H_5NO_2$, M 147.1, m 234°, 235°, 238°, b 336°/atm, $pK 8.30$. Crystallise the imide from EtOH (20ml/g) (charcoal), or sublime it. For potassium phthalimide see entry in 'Metal-organic Compounds', Chapter 4. [Beilstein 21/10 V 270.]

Phthalonitrile (1,2-dicyanobenzene) [91-15-6] $C_8H_4N_2$, M 128.1, m 138°, 140°, 141°, b 304°/atm. Crystallise the nitrile from EtOH, toluene or *benzene. It has also been distilled under high vacuum. It is steam volatile. [Beilstein 9 H 815, 9 II 602, 9 III 4199, 9 IV 3268.]

Phthalylsulfacetamide [131-69-1] $C_{16}H_{14}N_2O_6S$, M 362.3, m 196°, 304°. It is prepared by hydrolysis of N-acetamidophenylphthalimide by boiling for 3 hours in 15% aqueous KOH followed by treatment with Norit, filtration and acidification with AcOH. Crystallise the phthalamide from hot H_2O , aqueous EtOH, or EtOH. [Jain et al. *J Indian Chem Soc* 24 174 1947, Basu *J Indian Chem Soc* 26 130 1949, and several patents.] It is antibacterial. [Beilstein 14 III 2073.]

Phthiocol (2-hydroxy-3-methylnaphtha-1,4-quinone) [483-55-6] $C_{11}H_8O_3$, M 188.1, m 173-174°, 173.5°, $pK_{Est} \sim 4.2$. Crystallise the quinone from diethyl ether/petroleum ether. [Beilstein 8 III 2568, 8 IV 2375.] It is

hemostatic.

Physodic acid [4,4',6'-trihydroxy-6-(2-oxoheptyl)-2'-pentyl-2,3'-oxydibenzoic acid 1,5-lactone] [84-24-2] $C_{26}H_{30}O_8$, **M 470.5**, **m 205°**, $pK_{Est(1)} \sim 3.0$, $pK_{Est(2)} \sim 10$, $pK_{Est(3)} \sim 13$. Crystallise the acid from MeOH. The *methyl ester*, **m 156-157°**, crystallises from EtOH/H₂O (4:1) in prisms. The *diacetate* has **m 155-156°** (from Me₂CO/CS₂). [Beilstein 19 II 329, 19 III/IV 3988.]

Picene (dibenzo[*a,i*]phenanthrene, 3,4-benzchrysene) [213-14-3, 213-46-7] $C_{22}H_{14}$, **M 278.3**, **m 364°**, **366-367°**, **367-369°**. Crystallise picene from isopropylbenzene/xylene. After sublimation at 300°/2mm followed by crystallisation from xylene (charcoal), it gives white glistening leaflets with **m 366-366.5°** [Newman *J Org Chem* 09 518 1944, DOI: 10.1021/jo01188a005]. The *2,4,7-trinitrofluoren-9-one* has **m 257-257.8°** (from *C₆H₆). [Beilstein 5 H 735, 5 I 369, 5 III 2555, 5 IV 2724.]

Picric acid [88-89-1] $C_6H_3N_3O_7$, **M 229.1**, **m 122-123°** (dried material), **d 1.76**, $pK^{25} 0.33$ (0.37). Crystallise the acid first from acetic acid, then acetone, toluene, CHCl₃, aqueous 30% EtOH, 95% EtOH, MeOH or H₂O. Dry it in a vacuum for 2 hours. Alternatively, dry it over Mg(ClO₄)₂ or fuse (**CARE**) and allow it to solidify under a vacuum three times. Because it is **EXPLOSIVE**, picric acid should be stored moistened with H₂O, and only small portions should be dried at any one time. The dry acid should **NOT** be heated. [Beilstein 6 IV 1388.]

Picryl chloride (2-chloro-1,3,7-trinitrobenzene) [88-88-0] $C_6H_2ClN_3O_6$, **M 247.5**, **m 83°**, **d 1.797**. Crystallise the chloride from CHCl₃ or EtOH (**m 83-84°**). The 2:1 *C₆H₆-complex has **m 39°** (from *C₆H₆). It forms a 1:1 *hexamethylbenzene complex* which has orthorhombic orange-yellow crystals [Ross et al. *J Am Chem Soc* 76 69 1954, DOI: 10.1021/ja01630a018]. [Beilstein 5 II 205, 5 III 645, 5 IV 757.]

Picryl iodide (2-iodo-1,3,7-trinitrobenzene) [4436-27-5] $C_6H_2IN_3O_6$, **M 339.0**, **m 164-165°**. Crystallise the iodide from *benzene. [Beilstein 5 III 647, 5 IV 758.]

Piperic acid [*trans,trans*-5-(3,4-methylenedioxyphenyl)-2,4-pentadieneoic acid] [136-72-1, 5285-18-7] $C_{12}H_{10}O_4$, **M 218.2**, **m 215°**, **217°**, **220-221°**, $pK_{Est} \sim 4.7$. Crystallise the acid from EtOH. It turns yellow in light. It sublimes with partial decomposition. It has UV with λ_{max} at 340nm (MeOH). [Beilstein 19 H 281, 19 II 30, 19 III/IV 3565.]

Piperonal [120-57-0] $C_8H_6O_3$, **M 150.1**, **m 37°**, **b 140°/15mm**, **263°/760mm**. Crystallise piperonal from aqueous 70% EtOH or EtOH/water. [Beilstein 19/4 V 225.] It is a food-grade flavour ingredient.

Piperonylic acid [94-53-1] $C_8H_6O_4$, **M 166.1**, **m 229°**, $pK^{25} 4.50$. Crystallise the acid from EtOH or water. [Synthesis see Shriner & Kleiderer *Org Synth* 10 82 1930, DOI: 10.15227/orgsyn.010.0082; Beilstein 19/7 V 300.]

Polystyrene (PS) [9003-53-6] (C₈H₈)_n, **m ~240°**. Precipitate polystyrene repeatedly from CHCl₃ or toluene solution by addition of MeOH. Dry it *in vacuo*. [Miyasaka et al. *J Phys Chem* 92 249 1988, DOI: 10.1021/j100313a001.]

Procaine [4-(2-diethylaminomethoxycarbonyl)aniline] [59-46-1] $C_{13}H_{20}N_2O_2$, **M 236.3**, **m 51°** (dihydrate), **61°** (anhydrous), $pK_1 2.45$, $pK_2 8.91$. Procaine crystallises as the *dihydrate* from aqueous EtOH and as the *anhydrous* material from petroleum ether or diethyl ether. The latter is *hygroscopic*. [Beilstein 14 IV 1138.] It is a sympathomimetic anti-inflammatory drug. See *hydrochloride* in 'Physiologically Active....' In Chapter 6.

p-(1-Propenyl)phenol [*cis/trans* 6380-21-8, *trans* 85960-8-2, 539-12-8] $C_9H_{10}O$, **M 134.2**, **m 93-94°**, **b (trans) 229.6°/760mm**, $pK_{Est} \sim 10.2$. Crystallise the phenol from water. [Beilstein 6 III 2394, 6 IV 3796.]

n-Propyl gallate [121-79-9] $C_{10}H_{12}O_5$, **M 212.2**, **m 147°**, **148°**, **150°**, **d 1.21**. Crystallise the ester from aqueous EtOH or *C₆H₆ (**m 146-146.5°**). [Beilstein 10 III 2078, 10 IV 2003.]

Protocatechualdehyde (3,4-dihydroxybenzaldehyde) [139-85-5] $C_7H_6O_3$, M 138.1, m 153°, 150-157°. Crystallise the aldehyde from water or toluene and dry it in a vacuum desiccator over KOH pellets or shredded wax respectively. [Beilstein 8 IV 1762.]

1*S*,2*S*-Pseudoephedrine (1-hydroxy-1-phenyl-2-methylaminopropane) [90-82-4] $C_{10}H_{15}NO$, M 165.2, m 118-119°, $[\alpha]_D^{20}$ +53.0 (EtOH), +40.0 (H₂O), pK_{25}^{25} 9.71. Crystallise the amine from dry diethyl ether, or from water and dry it in a vacuum desiccator. [Beilstein 13 IV 1878.]

1*S*,2*S*-Pseudoephedrine hydrochloride [345-78-8] $C_{10}H_{15}NO \cdot HCl$, M 210.7, m 181-182°, 185-188°, $[\alpha]_D^{20}$ +61 (c 1 H₂O). Crystallise the salt from EtOH. [Beilstein 13 IV 1878.]

Purpurin (1,2,4-trihydroxy-5,10-anthraquinone) [81-54-9] $C_{14}H_8O_5$, M 256.2, m 253-256°, $pK_{Est(1)} \sim 7.0$ (2-OH), $pK_{Est(2)} \sim 9.0$ (4-OH), $pK_{Est(3)} \sim 11.1$ (1-OH). Crystallise purpurin from aqueous EtOH (orange needles with 1 H₂O), dry it at 100°. Sublimation at ~150°/2mm provides red needles of the anhydrous dye. It has λ_{max} at 515nm and 521nm; and complexes with metal salts to produce coloured complexes (e.g. 'lakes'). It stains nuclear material in histology. [Beilstein 8 IV 3568.]

Pyrene (benzo[def]phenanthrene) [129-00-0] $C_{16}H_{10}$, M 202.3, m 145-148°, 149-150°. Crystallise pyrene from EtOH, glacial acetic acid, *benzene or toluene. Purify it also by chromatography of CCl₄ solutions on alumina, with *benzene or *n*-hexane as eluent. [Backer & Whitten *J Phys Chem* **91** 865 1987, DOI: 10.1021/j100288a021.] It can also be zone refined and purified by sublimation. Marvel and Anderson [*J Am Chem Soc* **76** 5434 1954, DOI: 10.1021/ja01650a051] refluxed pyrene (35g) in toluene (400ml) with maleic anhydride (5g) for 4 days, then added 150ml of aqueous 5% KOH and refluxed for 5 hours with occasional shaking. The toluene layer was separated, washed thoroughly with H₂O, concentrated to about 100ml and allowed to cool. Crystalline pyrene was filtered off and recrystallised three times from EtOH or acetonitrile. [Chu & Thomas *J Am Chem Soc* **108** 6270 1986, DOI: 10.1021/ja00280a026; Russell et al. *Anal Chem* **58** 2961 1986, DOI: 10.1021/ac00127a014.] The material is free from anthracene derivatives. Another purification step involves passage of pyrene in cyclohexane through a column of silica gel. It can be sublimed in a vacuum and zone refined. The *picate* has m 224°. [Kano et al. *J Phys Chem* **89** 3748 1985, DOI: 10.1021/j100263a032; Beilstein 5 IV 2467.]

Pyrene-1-aldehyde [3029-19-4] $C_{17}H_{10}O$, M 230.3, m 123-126°, 125-126°. Recrystallise the aldehyde three times from aqueous EtOH. [Beilstein 7 IV 1821.]

1-Pyrenebutyric acid [3443-45-6] $C_{20}H_{16}O_2$, M 288.4, m 184-186°, $pK_{Est} \sim 4.1$. Crystallise the butyric acid from *benzene, EtOH, EtOH/water (7:3 v/v) or *C₆H₆/AcOH. Dry it over P₂O₅. [Chu & Thomas *J Am Chem Soc* **108** 6270 1986, DOI: 10.1021/ja00280a026; Beilstein 9 IV 2731.]

1-Pyrenecarboxylic acid [19694-02-1] $C_{17}H_{10}O_2$, M 230.3, m 270-272°, 273-274°, $pK_{Est} \sim 3.2$. Crystallise the acid from *C₆H₆, chlorobenzene, nitrobenzene or 95% EtOH. [Beilstein 9 H 712, 9 III 3575.]

***p*-Quaterphenyl** [135-70-6] $C_{24}H_{18}$, M 306.4, m 312-314°, 316-318°, 317°, 320°, b 428°/18mm. Recrystallise *p*-quaterphenyl from dimethyl sulfoxide at ca 50°. [Beilstein 5 II 669.]

Quinalizarin (1,2,5,8-tetrahydroxy-9,10-anthraquinone) [81-61-8] $C_{14}H_8O_6$, M 272.2, m 275°, ~300°, $pK_{Est(1)} \sim 7.1$ (1-OH), $pK_{Est(2)} \sim 9.9$ (8-OH), $pK_{Est(3)} \sim 11.1$ (5-OH), $pK_{Est(4)} \sim 11.8$ (2-OH). Crystallise the red quinone from acetic acid or nitrobenzene. It can be sublimed *in vacuo*. The Cu salt forms red crystals. [Beilstein 8 H 549, 8 I 755, 8 II 584.]

Quinhydrone (1:1 complex of hydroquinone and benzoquinone) [106-34-3] $C_{12}H_{10}O_4$, M 218.2, m 168°, 167-172°, 168-171°, 172°, 173-174°. Crystallise quinhydrone from H₂O at 65°, then dry it *in vacuo*. [Beilstein 7 H 617, 7 IV 2069.]

Quinizarin (1,4-dihydroxy-9,10-anthraquinone, Solvent orange 86) [81-64-1] $C_{14}H_8O_4$, M 240.2, m 198-199°, 200-202°, b 450°/atm, pK_1^{25} 9.90 (9.5), pK_2^{25} 11.18. Crystallise quinizarin (orange or red-brown crystalline powder) from glacial acetic acid. It forms 'lake' pigments with Ca, Ba and Pb salts. [Beilstein 8 H 450, 8 IV 3260.]

p-Quinquephenyl (p-pentaphenyl) [3073-05-0, 61537-20-0] $C_{30}H_{22}$, M 382.5, m 388.5°, 385-390°. Recrystallise p-pentaphenyl from boiling dimethyl sulfoxide (b 189°, lowered to 110°). The solid obtained on cooling is filtered off and washed repeatedly with toluene, then with concentrated HCl. The final material is washed repeatedly with hot EtOH. It is also recrystallised from pyridine or Me_2SO (well defined leaflets), then sublimed *in vacuo*. [Campbell & McDonald *Org Synth Coll Vol* 5 985 1973, DOI: 10.15227/orgsyn.040.0085; Beilstein 5 II 709, 5 III 2680, 5 IV 2874.]

Resorcinol [108-46-3] $C_6H_6O_2$, M 110.1, m 110°, 111.2-111.6°, b 277°/atm, d 1.28, pK_1^{25} 9.23, pK_2^{25} 13.05. Recrystallise resorcinol from *benzene, toluene or *benzene/diethyl ether. The *benzoate* has m 117°. [Beilstein 6 IV 2069.]

Retene (1-methyl-7-isopropylphenanthrene) [483-65-8] $C_{18}H_{18}$, M 234.3, m 89°, 90°, 98.5°, 99°, 100.5-101°, b 208°/10mm, 390°/atm. Crystallise retene from EtOH. The *picrate* has m 126° (from EtOH). The 1:1 *1,3,5-trinitrobenzene-complex* has m 144° (from EtOH). [Beilstein 5 H 683, 5 II 598, 5 III 2199.]

ParaRosaniline (4,4',4"-triaminotriptyllium [triphenylmethane] carbonium ion, para-fuschin, paramagenta) [467-62-9] $C_{19}H_{19}N_3O$, M 305.4, m 205°, pK^{25} 7.57 and free base has $pK > 13$. Dissolve the dye in EtOH (1.16g in 30ml), filter and add aqueous NH_3 till neutral (colourless) and precipitate it by adding H_2O giving 0.8g, m 247°(dec, sintering at 230°). Dissolve it in EtOH, neutralise with NH_3 till colourless, add 0.1g of charcoal, filter, and repeat, then add H_2O (100ml) to precipitate the colourless *carbinol* (pseudo-base) and dry it *in vacuo*, m 257°(dec, also 205°, sintering at 232°). [Weissberger & Theile *J Chem Soc* 148 1934, DOI: 10.1039/JR9340000148.] The *carbinol* is slightly soluble in H_2O but is soluble in acids (e.g. HCl to give the coloured *chloride* [569-61-9]) and EtOH [pK: Goldacre & Phillips *J Chem Soc* 1724 1949, DOI: 10.1039/JR9490001724]. The *perchlorate* (dark red with a green shine) has m 300° and explodes at 317° [Dilthey & Dinklage *J Prakt Chem* [2] 129 24 1931, DOI: 10.1002/prac.19311290102]. [Beilstein 13 IV 2283.] **Rosaniline HCl (Magenta I, Fuschin)** [632-99-5] $C_{19}H_{18}ClN_3$, M 337.9, has m >200°(dec). Purify the dye by dissolving it in EtOH (1mg/ml), filtering and adding H_2O . Filter or centrifuge it and wash the precipitate with Et_2O and dry it in air. It has also been recrystallised from water and dried *in vacuo* at 40°. The crystals have a metallic green lustre. It has UV with λ_{max} in EtOH at 543nm (ϵ 93,000). Its solubility in H_2O is 0.26%. A carmine red colour is obtained in EtOH. It is paraRosaniline with a methyl group. [Scalan *J Am Chem Soc* 57 887 1937, DOI: 10.1021/ja01308a029.]

p-Rosolic acid (4-[bis-{4-hydroxyphenyl}methylene]-2,5-cyclohexadien-one, 4',4"-di-hydroxy-fuschson, aurin, corallin) [603-45-2] $C_{19}H_{14}O_3$, M 290.3, m 292°, 295-300° (dec with liberation of phenol), 308-310°(dec), pK_1 3.11, pK_2 8.62. It forms green crystals with a metallic luster, but the colour depends on the solvent used. When recrystallised from brine (saturated aqueous NaCl) acidified with HCl, it forms red needles, but when recrystallised from EtO/AcOH, the crystals have a beetle iridescent green colour. It has been recrystallised from Me_2CO (although it dissolves slowly), methyl ethyl ketone, 80-95% AcOH and from AcOH/* C_6H_6 . An aqueous KOH solution is golden yellow, and a 70% H_2SO_4 solution is deep red in colour. An *alternative* purification is to dissolve this triphenylmethane dye in 1.5% of aqueous NH_3 , filter, and heat to 70-80°, then acidify with dilute AcOH by adding it slowly with vigorous stirring, whereby the aurin separates as a brick-red powder or as purplish crystals depending on the temperature and period of heating. Filter off the solid, wash it with H_2O and a little dilute AcOH, then H_2O again. Stir this solid with Et_2O to remove any ketones and allow it to stand overnight in the Et_2O , then filter and dry it in air then in a vacuum. [Gomberg & Snow *J Am Chem Soc* 47 298 1925, DOI: 10.1021/ja01678a029; Baines & Driver *J Chem Soc* 123 1214 1923, DOI: 10.1002/prac.19311290102, UV: Burawoy *Chem Ber* 64 462 1931, DOI: 10.1002/cber.19310640243; Beilstein 8 IV 2646.] It is an indicator whose aqueous solutions are pink-red below pH 5.0 and yellow above pH 6.8 (see pK_a and above).

Salicylaldehyde (*o*-hydroxybenzaldehyde) [90-02-8] $C_7H_6O_2$, M 122.1, m -7° , 1-2 $^\circ$, b 93 $^\circ$ /25mm, 195-197 $^\circ$ /760mm, d_4^{20} 1.167, n_D^{20} 1.574, pK^{25} 8.37. It is precipitated as the bisulfite addition compound by pouring the aldehyde slowly and with stirring into a 25% solution of $NaHSO_3$ in 30% EtOH, then standing for 30 minutes. The precipitate, after filtering at the pump, and washing with EtOH, is decomposed with aqueous 10% $NaHCO_3$, and the aldehyde is extracted into diethyl ether, dried with Na_2SO_4 or $MgSO_4$, and distilled, under reduced pressure. Alternatively, salicylaldehyde is precipitated as its Cu complex by adding it to warm, saturated aqueous $Cu(OAc)_2$, shaking and standing in ice. The precipitate is filtered off, washed with EtOH, then Et_2O , and decomposed with 10% H_2SO_4 ; the aldehyde is extracted into Et_2O , dried and vacuum distilled. It was also purified by dry column chromatography on Kieselgel G [Nishiya et al. *J Am Chem Soc* **108** 3880 1986, DOI: 10.1021/ja00274a003]. The *acetyl* derivative has m 38-39 $^\circ$ (from petroleum ether or EtOH) and b 142 $^\circ$ /18mm, 253 $^\circ$ /atm. The *oxime*, [94-67-7] M 137.1, crystallises $CHCl_3$ /petroleum ether (b 40-60 $^\circ$) with m 57 $^\circ$. [Beilstein 8 IV 176, 203.]

Salicylamide [65-45-2] $C_7H_7NO_2$, M 137.1, m 140-144 $^\circ$, 142-144 $^\circ$, pK^{20} 8.37. Crystallise the amide from water or repeatedly from $CHCl_3$ [Nishiya et al. *J Am Chem Soc* **108** 3880 1986, DOI: 10.1021/ja00274a003]. [Beilstein 10 IV 169.] The *anilide* [87-17-2] M 213.2, m 135 $^\circ$ crystallises from H_2O . [Beilstein 12 H 500, 12 I 268, 12 II 256, 12 944.] Analgesic.

Salicylhydroxamic acid [89-73-6] $C_7H_7NO_3$, M 153.1, m 177 $^\circ$ (dec), 179-180 $^\circ$ (dec), pK_1^{30} 2.15, pK_2^{30} 7.46, pK_3^{30} 9.72. Crystallise the hydroxamic acid from acetic acid. [Beilstein 10 H 98.] Irreversible inhibitor of bacterial (and plant) urease and useful in urinary tract infections. [Opperdoes et al. *Exptl Parasitol* **40** 198 1976, DOI: 10.1016/0014-4894(76)90082-5].

Salicylic acid (2-hydroxybenzoic acid) [69-72-7] $C_7H_8O_3$, M 138.1, m 157-159 $^\circ$, 158-160 $^\circ$, 158.6 $^\circ$, 159 $^\circ$, 159.5 $^\circ$, 159-160 $^\circ$, 162 $^\circ$, b 211 $^\circ$ /20mm, pK_1^{25} 3.01, pK_2^{25} 13.43 (13.01). It has been purified by steam distillation, by recrystallisation from H_2O (solubility is 0.22% at room temperature and 6.7% at 100 $^\circ$), absolute MeOH, or cyclohexane and by sublimation in a vacuum at 76 $^\circ$. The *acid chloride* (needles) has m 19-19.5 $^\circ$, b 92 $^\circ$ /15mm, the *O-acetyl* derivative has m 135 $^\circ$ (rapid heating and the liquid resolidifies at 118 $^\circ$), and the *O-benzoyl* derivative has m 132 $^\circ$ (aqueous EtOH). [IR: Hales et al. *J Chem Soc* 3145 1954, DOI: 10.1039/JR9540003145; Bergmann et al. *J Chem Soc* 2351 1950, DOI: 10.1039/JR9500002351]. [Beilstein 10 IV 125.]

cis-Stilbene (Z-1,2-diphenylethylene) [645-49-8] $C_{14}H_{12}$, M 180.3, m 1-2 $^\circ$, b 145 $^\circ$ /12mm, 307 $^\circ$ /atm. Purify it by chromatography on alumina using hexane and distil it under vacuum. (The final product contains *ca* 0.1% of the *trans*-isomer.) [Lewis et al. *J Am Chem Soc* **107** 203 1985, DOI: 10.1021/ja00287a037; Saltiel et al. *J Phys Chem* **91** 2755 1987, DOI: 10.1021/j100295a022; Beilstein 5 H 630.]

trans-Stilbene (E-1,2-diphenylethylene) [103-30-0] has m 122-124 $^\circ$, 125.9 $^\circ$, 126 $^\circ$, b 305-307 $^\circ$ /744mm, d_4^{20} 0.970. Purify it by vacuum distillation. (The final product contains about 1% of the *cis* isomer.) Crystallise it from EtOH. It has also been purified by zone melting. The *styphnate* (see next entry) has m 142 $^\circ$. [Lewis et al. *J Am Chem Soc* **107** 203 1985, DOI: 10.1021/ja00287a037; Bellucci et al. *J Am Chem Soc* **109** 515 1987, DOI: 10.1021/ja00236a032; Saltiel *J Phys Chem* **91** 2755 1987, DOI: 10.1021/j100295a022; Beilstein 5 IV 2156.]

Styphnic acid (2,4,6-trinitroresorcinol) [82-71-3] $C_6H_3N_3O_8$, M 245.1, m 177-178 $^\circ$, 179-180 $^\circ$, 180 $^\circ$, d 1.83, pK_1^{25} 0.06 (1.74), pK_2^{25} 4.23 (4.86). Crystallise the phenol from ethyl acetate or water containing HCl [EXPLODES violently on rapid heating.] Its solubility in H_2O is 0.7% at 20 $^\circ$ and 3% at 100 $^\circ$. It forms *addition compounds* with aromatic hydrocarbons, e.g. naphthalene (m 168 $^\circ$), anthracene (m 180 $^\circ$), phenanthrene (m 142 $^\circ$), fluorene (m 134 $^\circ$) and retene (m 141 $^\circ$). [Beilstein 6 H 830, 6 III 4354, 6 IV 5699.]

Styrene (vinylbenzene) [100-42-5] C_8H_8 , M 104.2, m -30 $^\circ$, b 41-42 $^\circ$ /18mm, 145.2 $^\circ$ /760mm, d_4^{20} 0.907, n_D^{20} 1.5469, n_D^{25} 1.5441. Styrene is difficult to purify and keep pure. Usually it contains added inhibitors (such as a trace of hydroquinone). Wash it with aqueous NaOH to remove inhibitors (e.g. *tert*-butanol), then with water, dry it for several hours with $MgSO_4$ and distil it at 25 $^\circ$ under reduced pressure in the presence of an inhibitor (such as 0.005% *p-tert*-butylcatechol). It can be stored at -78 $^\circ$. It can also be stored and kept anhydrous with Linde type 5A molecular sieves, CaH_2 , $CaSO_4$, BaO or sodium, being fractionally distilled, and distilled in a

vacuum line just before use. *Alternatively*, styrene (and its deuterated derivative) are passed through a neutral alumina column before use [Woon et al. *J Am Chem Soc* **108** 7990 1986, DOI: 10.1021/ja00285a018; Collman *J Am Chem Soc* **108** 2588 1986, DOI: 10.1021/ja00270a016]. [*Beilstein* **5** IV 1334.]

(±)-Styrene glycol (±-1-phenyl-1,2-ethanediol) [93-56-1] $C_8H_{10}O_2$, **M 138.2**, **m 67-68°**, **67.5°**, **68°**, **272-274°/755mm**. Crystallise the diol from petroleum ether, Et_2O , $Et_2O/*C_6H_6$ (**m 69-70°**) or $*C_6H_6$. The *dibenzoyl derivative* has **m 96-97°**. [*Beilstein* **6** H 907, **6** I 444, **6** II 887, **6** III 4572, **6** IV 5939.]

Sudan II [Solvent Orange 7, 1-(2,4-xylylazo)-2-naphthol] [3118-97-6] $C_{18}H_{16}N_2O$, **M 276.3**, **m 156-158°**, **CI 12140**, λ_{max} (420nm sh) **493, 604 nm**, **pK_{Est} ~9.0**. Crystallise the red dye from EtOH, EtOH/water or $*benzene/absolute EtOH$ (1:1). This fat-soluble dye stains triglycerides and protein bound lipids in frozen paraffin sections. [*Beilstein* **16** H 168.]

Sudan III [Solvent Red 23, 1-(*p*-phenylazo-phenylazo)-2-naphthol] [85-86-9] $C_{22}H_{16}N_4O$, **M 352.4**, **m 199°(dec)**, **CI 26100**, λ_{max} **354, 508 nm**, **pK_{Est} ~9.0**. Crystallise this red lysochrome dye from EtOH, EtOH/water or $*benzene/absolute EtOH$ (1:1). It stains fatty substances, e.g. oils, fats, waxes, greases etc. [*Beilstein* **16** II 75, **16** III 148, **16** IV 248.]

Sudan IV [Solvent Red 24, 1-(4-*o*-tolylazo-*o*-tolylazo)-2-naphthol] [85-83-6] $C_{24}H_{20}N_4O$, **M 380.5**, **m ~184°(dec)**, **199°(dec)**, **CI 26105**, λ_{max} (357nm sh) **520nm**, **pK_{Est} ~9.0**. Crystallise this red-brown lysochrome dye from EtOH/water or acetone/water. [*Beilstein* **16** IV 249.]

Sudan Blue II (Solvent Blue 35, 1,4-bis-(butylamino)-9,10-anthraquinone) [17354-14-2] $C_{22}H_{26}N_2O_2$, **M 350.5**, **m 121-122°**, **122°**, λ_{max} **604, 652nm**, **pK_{Est} ~9.5 (OH)**. It is formed from quinizarin (2g see [81-64-1]), 33% EtOH/*n*-BuNH₂ (20ml) and Na₂S₂O₄ (2g) at 140°/8 hours, evaporate, extract with toluene, chromatograph (Al₂O₃), the intense blue band in toluene is evaporated, and the residue gave purple needles (Cu lustre) from petroleum ether (b 60-80°) (1.1g, 38%) [Peters & Walker *J Chem Soc* 1429 1956, DOI: 10.1039/JR9560001429; *Beilstein* **14** IV 460]. It forms Cu and Ni salts.

Syringaldehyde (3,5-dimethoxy-4-hydroxybenzaldehyde) [134-96-3] $C_9H_{10}O_4$, **M 182.2**, **m 110-113°**, **112°**, **113°**, **b 192°/14mm**, **d 1.01**, **pK_{Est} ~8**. Crystallise syringaldehyde from petroleum ether. [*Beilstein* **8** H 391, **8** IV 2718.]

Syringic acid (3,5-dimethoxy-4-hydroxybenzoic acid) [530-57-4] $C_9H_{10}O_5$, **M 198.2**, **m 204-205°**, **206.5°**, **206-209°**, **209-210°**, **b 192-193°/14mm**, **pK₁²⁵ 4.34**, **pK₂²⁵ 9.49**. Recrystallise syringic acid from H₂O using charcoal [Bogert & Coyne *J Am Chem Soc* **51** 571 1929, DOI: 10.1021/ja01377a030; Anderson & Nabenhauer *J Am Chem Soc* **48** 3001 1926 DOI: 10.1021/ja01690a037.] The *methyl ester* has **m 107°** (from MeOH), the *4-acetyl* derivative has **m 190°** and the *4-benzoyl* derivative has **m 229-232°**. [Hahn & Wassmuth *Chem Ber* **67** 696 1934, DOI: 10.1002/cber.19340670430; UV: Lemon *J Am Chem Soc* **69** 2998 1947, DOI: 10.1021/ja01204a018 and Pearl & Beyer *J Am Chem Soc* **72** 1743 1950, DOI: 10.1021/ja01160a092; *Beilstein* **10** IV 1995.]

Terephthalaldehyde [623-27-8] $C_8H_6O_2$, **M 134.1**, **m 112-115°**, **114-117°**, **116°**, **117°**, **b 245-248°/771mm**, **d 1.06**. Crystallise terephthalaldehyde from water. [*Beilstein* **7** IV 2140.]

Terephthalic acid (benzene-1,4-dicarboxylic acid) [100-21-0] $C_8H_6O_4$, **M 166.1**, **m sublimes >300° without melting**, **d 1.52**, **pK₁²⁰ 3.4**, **pK₂²⁰ 4.34**. Purify the acid *via* the sodium salt which, after crystallisation from water, is re-converted to the acid by acidification with mineral acid. Filter off the solid, wash it with H₂O and dry it in a vacuum. The *S-benzylisothiuronium salt* has **m 204°** (from aqueous EtOH). [*Beilstein* **9** IV 3301.] Used for making PET polyesters, e.g. for plastic bottles, clothing and related materials.

Terephthaloyl chloride (benzene-1,4-dicarbonyl chloride) [100-20-9] $C_8H_4Cl_2O_2$, **M 203.0**, has **m 79-83°**, **81°**, **80-82°**, **83.5°**, **b 265°/atm**, **d 1.34**. Crystallise the acid chloride from dry hexane. The *diamide* [3010-82-0] $C_8H_8N_2O_2$, **M 164.1**, has **m 332°** (from aqueous NH₃). [*Beilstein* **9** IV 3318.]

***o*-Terphenyl (1,2-diphenylbenzene)** [84-15-1] $C_{18}H_{14}$, M 230.3, m 54-57°, 56.2°, 57°, 58-59°, b 332°/atm, 337°/atm, d 1.24. Crystallise *o*-terphenyl from EtOH. Also purify it by chromatography of CCl_4 solution on alumina, with petroleum ether as eluent, followed by crystallisation from petroleum ether (b 40-60°) or petroleum ether/* C_6H_6 . It also distils under vacuum. [Beilstein 5 III 2292, 5 IV 2478.]

***m*-Terphenyl (1,3-diphenylbenzene)** [92-06-8] $C_{18}H_{14}$, M 230.3, has m 86°, 87°, 88-89°, b 363°/atm, 379°/atm, d ²⁵ 1.24g/ml. Purify it as for *o*-terphenyl above. [Beilstein 5 IV 2480.]

***p*-Terphenyl (1,4-diphenylbenzene)** [92-94-4] $C_{18}H_{14}$, M 230.3, has m 212.7°, b 389°/atm. Crystallise *p*-terphenyl from nitrobenzene or trichlorobenzene. It is also purified by chromatography on alumina in a darkened room, using petroleum ether, and then crystallising from petroleum ether (b 40-60°) or petroleum ether/*benzene. It is a fluorophore for scintillation counting and has λ_{ex} 286nm : λ_{em} 343nm in DMF, and λ_{max} at 277nm (log ϵ 4.50). [Beilstein 5 IV 2483.]

3',3'',5',5''-Tetrabromophenolphthalein ethyl ester [1176-74-5] $C_{22}H_{14}Br_4O_4$, M 662.0, m 208-211°, 212-214°. Crystallise the yellow to red ester from *benzene, dry at 120°, and keep it under vacuum or store at -20°. [Beilstein 10 III 4490.] Useful stain in microscopy.

2,3,4,5-Tetrachloroaniline [634-83-3] $C_6H_3Cl_4N$, M 230.9, m 117-118°, 119-120°, $pK_{Est} \sim 0.26$. Crystallise it from EtOH. Analytical standard available in MeCN (100mg/ml), and it store at +4°. The *acetyl* derivative has m 165-166° (from EtOH). [Beilstein 12 H 630, 12 I 313, 12 II 340, 12 IV 1286.]

2,3,5,6-Tetrachloroaniline [3481-20-7] has m 106°, 106-108°, 107-108°, $pK_{Est} \sim 1.8$. Crystallise it from EtOH. The *acetyl* derivative has m 213-214° (from EtOH). [Beilstein 12 II 340, 12 III 1414, 12 IV 1287.]

1,2,3,4-Tetrachlorobenzene [634-66-2] $C_6H_2Cl_4$, M 215.9, m 42-45°, 45-46°, 47°, 47.5°, b 254°/761mm. Crystallise it from EtOH. [Beilstein 5 H 204, 5 II 156, 5 III 550, 5 IV 667.]

1,2,3,5-Tetrachlorobenzene [634-90-2] has m 50-51°, 51°, 54-55°, b 246°/760mm, d¹⁰ 1.7344. Crystallise it from EtOH. [Beilstein 5 II 157, 5 III 551, 5 IV 668.]

1,2,4,5-Tetrachlorobenzene [95-94-3] has m 139°, 139.5°, 139.5-140.5°, 139-142°, b 240°/760mm, 245°/760mm, d 1.86. Crystallise it from EtOH, ether, *benzene, *benzene/EtOH or carbon disulfide. [Beilstein 5 IV 668.]

3,4,5,6-Tetrachloro-1,2-benzoquinone (*o*-chloranil) [2435-53-2] $C_6Cl_4O_2$, M 245.9, m 126° to 129°, 130°, 133°. Crystallise *o*-chloranil from AcOH. Dry it in vacuum desiccator over KOH. A useful dehydrogenating and oxidising agent [Fieser 1 128]. [Brook *J Chem Soc* 5035, Note p 5040, 1952, DOI: 10.1039/JR9520005035; Beilstein 7 IV 2065.]

3,4,5,6-Tetrachloro-*N*-methylphthalimide [14737-80-5] $C_9H_3Cl_4NO_2$, M 298.9, m 209.7°. Crystallise the imide from absolute EtOH. [Beilstein 21 H 505, 17/11 V 260.]

2,3,4,6-Tetrachloronitrobenzene (1,2,3,5-tetrachloro-4-nitrobenzene) [879-39-0, 3714-62-3] $C_6HCl_4NO_2$, M 260.9, m 41-42°, 42°. Crystallise it from aqueous EtOH. [Beilstein 5 II 187, 5 III 617, 5 IV 728.]

2,3,5,6-Tetrachloronitrobenzene (1,2,4,5-tetrachloro-3-nitrobenzene) [117-18-0, 28804-67-3] has m 98-101°, 99-100°, b 304°/760mm, Crystallise it from aqueous EtOH or H_2O . [Beilstein 5 III 617, 5 IV 728.]

2,3,4,5-Tetrachlorophenol [4901-51-3] $C_6H_2Cl_4O$, M 231.9, m 116.5°, 116-117°, pK^{25} 6.95 (6.35). Crystallise the phenol from petroleum ether. The *benzoate* has m 110° (from EtOH). [Beilstein 6 II 182, 6 III 729, 6 IV 1020.]

2,3,4,6-Tetrachlorophenol [58-90-2] $C_6H_2Cl_4O$, M 231.9, m 63-67°, 70°, b 150°/15mm, 164°/23mm, d 1.839, pK^{25} 5.38. Crystallise the phenol from petroleum ether and/or distil it under vacuum. The *benzoate* has m 116° (from EtOH). [Beilstein 6 H 193, 6 III 729, 6 IV 1021.] Pesticide against insects, fungi and bacteria, but is a potential carcinogen.

2,3,5,6-Tetrachlorophenol [935-95-5] has m 114-116°, 115°, pK^{25} 5.48 (5.09). Recrystallise the phenol from petroleum ethers. It is soluble in * C_6H_6 (very), H_2O (slightly), and ligroin. It irritates eyes, skin, mucous membranes and respiratory tract. The *benzoate* has m 136° (from EtOH). [Beilstein 6 III 730, 6 IV 1025.]

Tetrachlorophthalic acid [632-58-6] $\text{C}_8\text{H}_2\text{Cl}_4\text{O}_4$, **M 303.9**, **m 98°** (anhydrous), **439°/760mm**. Crystallises from hot water as the *hemihydrate* in colourless thin monoclinic plates. [For crystal structure of hemihydrate see Ito et al. *Bull Chem Soc Jpn* **48** 3078 1975, DOI: 10.1246/bcsj.48.3078.]

Tetrachlorophthalic anhydride [117-08-8] $\text{C}_8\text{Cl}_4\text{O}_3$, **M 285.9**, **m 253-256°, 254.5°, 255-256°, 255-257°, 254-258°, b 371°/760mm**. Crystallise the anhydride from chloroform or *benzene, then sublime it in a vacuum. It forms a phenanthrene adduct [3178-32-3] [*Beilstein* **17/11** V 260, Fieser **15** 300] **Tetrachlorophthalimide** [1571-13-7] $\text{C}_8\text{HCl}_4\text{NO}_2$, **M 284.9**, crystallises as colourless blades **m 338-339°** (from hot AcOH, solubility 1g/100ml); **m 336-337°** (from nitrobenzene); and yellow needles **m 345-347°** (from DMF) and is a **teratogen**. [Al-Mughaid & Grindley *e-EROS Encyclopedia of Reagents for Organic Synthesis* DOI: 10.1002/047084289X.rn00333; *Beilstein* **21** 505.]

1,2,4,5-Tetracyanobenzene [712-74-3] $\text{C}_{10}\text{H}_2\text{N}_4$, **M 178.1**, **m 270-272°, 275-280° (280°)**. The tetranitrile can be prepared by dehydration of **1,2,4,5-benzenetetracarboxamide (pyromellitimide)** [6183-35-5] $\text{C}_{10}\text{H}_{10}\text{N}_4\text{O}_4$, **M 250.2**, **m >450-454°**, with SOCl_2/DMF . Crystallise the tetra-nitrile from EtOH (fine needles) and sublime it *in vacuo*. It crystallises from AcOH in colourless plates (**m 267-269°**), and recrystallisation from $\text{MeOCH}_2\text{CH}_2\text{OH}$, EtCOMe and again from AcOH provided crystals with **m 270-272°**. Its IR (KBr pellet) has ν_{max} at 2,240 (4.5 μ , CN), 3,125 and 3,030 (3.2 μ and 3.3 μ , CH), and 926 (10.8 μ , 1,2,4,5-benzene substitution) cm^{-1} . It forms distinct molecular complexes with a variety of aromatic 'donor' molecules such as durene (**m 267-270°**), anthracene (**m 277-280°**), pyrene (**m 268-270°**), 2-naphthol (**m 265-270°**), 2,7-dimethoxynaphthalene (**m 267-270°**), dimethylaniline (**m 266-268°**), 1,5-diaminonaphthalene (**m >360°**), quinolone (**m 262-265°**) and isoquinoline (**m 194-196°**). [Lawton & McRitchie *J Org Chem* **24** 26 1959, DOI: 10.1021/jo01083a008; Bailey et al. DOI: 10.1016/0040-4020(63)80018-6; **19** 161 1963, *Beilstein* **9** IV 3804.]

7,7,8,8-Tetracyanoquinodimethane (TCNQ) [1518-16-7] $\text{C}_{12}\text{H}_4\text{N}_4$, **M 204.2**, **m 287-289°(dec), 289-291°, 293.5-296°**. It is prepared by condensation of cyclohexane-1,4-dione and malononitrile, bromination followed by dehydrobromination with pyridine. It is purified by sublimation above 250° at atmospheric pressure or at ~200° under vacuum giving rust coloured crystals. It also crystallises nicely from EtOAc, tetrahydrofuran or MeCN in rust-coloured crystals. When the crystals are crushed between soft glass melting point cover glasses and heated, a beautiful blue film forms on the glass plates at ~200° due to the reaction of TCNQ with the bases in the glass to give the TCNQ anion-radical. Its has IR bands at ν_{max} 1540 cm^{-1} (conjugated C=C for a cycloolefine) and 2220 cm^{-1} (conjugated nitrile). The UV in EtOH is complicated by the presence of the TCNQ radical anion (among other products), and the absorption from TCNQ is best determined by using a KBr wafer or measuring it in MeCN solution which has λ_{max} at 395nm (ϵ 63,600). Polarographic reduction of TCNQ in 0.2M NaOAc is at +0.15 to +0.16V reversibly to form TCNQH₂. TCNQ is a strong Lewis acid which forms π -complexes; the charge-transfer complex with anthracene is black. The association constants of the radical anions in CH_2Cl_2 were determined with durene (5.6), hexamethylbenzene (14.5) and pyrene (78.4) which compare with those of TCNE (tetracyanoethylene) that are 54.2, 263 and 29.5 respectively. [Acker & Hertler *J Am Chem Soc* **84** 3370 1962, DOI: 10.1021/ja00876a028; Melby et al. *J Am Chem Soc* **84** 3374 1962, DOI: 10.1021/ja00876a029; Fieser **1** 1136, **12** 464.]

Tetrahydroxy-p-benzoquinone (2H₂O) [5676-48-2; 123334-16-7 2H₂O] $\text{C}_6\text{H}_4\text{O}_6 \cdot 2\text{H}_2\text{O}$, **M 172.1 + xH₂O**, **m ~300°, pK₁³⁰ 4.80, pK₂³⁰ 6.8**. Crystallise the quinone from water. [*Beilstein* **8** H 534, **8** II 572, **8** III 4204, **8** IV 3604.] It is an indicator for sulfate titrations.

Tetralin (1,2,3,4-tetrahydronaphthalene) [119-64-2] $\text{C}_{10}\text{H}_{12}$, **M 132.2**, **m -35.79° (from CF₂Cl₂), b 65-66°/5mm, 207.6°/760mm, d₄²⁰ 0.968, n_D²⁰ 1.5413**. Wash tetralin with successive portions of concentrated H₂SO₄ until the acid layer is no longer coloured, then wash it with aqueous 10% Na₂CO₃, and then distilled water. Dry (CaSO₄ or Na₂SO₄), filter, reflux and fractionally distil it under reduced pressure from sodium or BaO. It can also be purified by repeated fractional freezing. Bass [*J Chem Soc* 3475(3498) 1964, DOI: 10.1039/JR9640003475 (note)] freed tetralin, purified as above, from naphthalene and other impurities by conversion to ammonium tetralin-6-sulfonate. Concentrated H₂SO₄ (150ml) is added slowly to stirred tetralin (272ml) which is then heated on a water bath for about 2 hours for complete solution. The warm mixture, when poured into aqueous NH₄Cl solution (120g in 400ml water), gives a white precipitate which, after filtering off, is crystallised from boiling water, washed with 50% aqueous EtOH and dried at 100°. Evaporation of its boiling

aqueous solution on a steam bath removes traces of naphthalene. The pure salt (229g) is mixed with concentrated H_2SO_4 (266ml) and steam distilled from an oil bath at 165–170°. An ether extract of the distillate is washed with aqueous Na_2SO_4 , and the ether is evaporated, prior to distilling the tetralin from sodium. Tetralin has also been purified *via* barium tetralin-6-sulfonate, converted to the sodium salt and decomposed in 60% H_2SO_4 using superheated steam. [*Beilstein* 5 H 491, 5 III 1219, 5 IV 1388.]

Tetralin hydroperoxide [771-29-9] $\text{C}_{10}\text{H}_{12}\text{O}_2$, M 164.2, m 55.7–56°, 56°. Crystallise the tetralin hydroperoxide from hexane, toluene at -30° (m 54.0–54.5°). The oxygen content should be ~9.70%. [Knight & Swern *Org Synth Coll Vol* 4 895 1963, DOI: 10.15227/orgsyn.034.0090.]

α -Tetralone (1,2,3,4-tetrahydro-1-oxonaphthalene) [529-34-0] $\text{C}_{10}\text{H}_{10}\text{O}$, M 146.2, m 2-7°, 7.8–8.0°, b 75–85°/0.3mm, 89°/0.5mm, 94–95°/2mm, 132–134°/15mm, 143–145°/20mm, d_4^{20} 1.0695, n_D^{20} 1.5665. Check the IR first. Purify α -tetralone by dissolving 20ml in Et_2O (200ml), washing with H_2O (100ml), 5% aqueous NaOH (100ml), H_2O (100ml), 3% aqueous AcOH (100ml), 5% NaHCO_3 (100ml) then H_2O (100ml) and dry the ethereal layer over MgSO_4 . Filter, evaporate and fractionate the residue through a 6in Vigreux column under reduced pressure to give a colourless oil (~17g) with b 90–91°/0.5–0.7mm. [Snyder & Werber *Org Synth Coll Vol* 3 798 1955, DOI: 10.15227/orgsyn.020.0094.] It has also been fractionated through a 0.5metre packed column with a heated jacket under reflux using a partial take-off head. It has λ_{max} at 247.5 and 290nm (hexane). The *phenylhydrazone* has m 83°. The *2,4,6-trinitrophenylhydrazone* has m 247.5–248° (from EtOH). [Olson et al. *Org Synth Coll Vol* 4 898 1963, DOI: 10.15227/orgsyn.035.0095; *Beilstein* 7 III 1416, 7 IV 1015.]

β -Tetralone (1,2,3,4-tetrahydro-2-oxonaphthalene) [530-93-8] has m 17–18°, ~18°, b 93–95°/2mm, 104–105°/4mm, 114–115°/4–5mm, 140°/18mm, d_4^{20} 1.1000, n_D^{20} 1.5598. If reasonably pure, then fractionate it through an efficient column. Otherwise purify it *via* the *bisulfite adduct*. To a solution of NaHSO_3 (32.5g, 0.31mol) in H_2O (57ml) is added 95% EtOH (18ml) and set aside overnight. Any bisulfite-sulfate that separated is removed by filtration, and the filtrate is added to the tetralone (14.6g, 0.1mol) and shaken vigorously. The adduct separates in a few minutes as a white precipitate and is kept on ice for ~3.5 hours with occasional shaking. The precipitate is collected, washed with 95% EtOH (13ml), then with Et_2O (4 x 15ml, by stirring the suspension in the solvent, filtering and repeating the process). The colourless product is dried in air and stored in air tight containers in which it is stable for extended periods (yield is ~17g). This bisulfite (5g) is suspended in H_2O (25ml), and $\text{Na}_2\text{CO}_3 \cdot \text{H}_2\text{O}$ (7.5g) is added (pH of solution is ~10). The mixture is then extracted with Et_2O (5 x 10ml, i.e. until the aqueous phase does not test for tetralone — see below). Wash the combined extracts with 10% aqueous HCl (10ml), H_2O (10ml, i.e. until the washings are neutral), dry (MgSO_4), filter, evaporate and distil the residual oil using a Claisen flask under reduced pressure and in a N_2 atmosphere. The pure tetralone is a colourless liquid b 70–71°/0.25mm (see also above). The yield is ~2g. **Tetralone test:** Dissolve a few drops of the tetralone solution (ethereal or aqueous) in 95% EtOH in a test tube and add 10 drops of 25% NaOH down the side of the tube. A deep blue colour develops at the interface with air. [Soffer et al. *Org Synth Coll Vol* 4 903 1963, DOI: 10.15227/orgsyn.032.0097; Cornforth et al. *J Chem Soc* 689 1942, DOI: 10.1039/JR9420000689; UV: Soffer et al. *J Am Chem Soc* 74 1556 1952, DOI: 10.1021/ja01126a063.] The *phenylhydrazone* has m 108° [Crowley & Robinson *J Chem Soc* 2001 1938, DOI: 10.1039/JR9380002001]. [*Beilstein* 7 H 370, 7 II 295, 7 III 1422, 7 IV 1018.]

1,2,3,4-Tetramethylbenzene (prehnitine) [488-23-3] $\text{C}_{10}\text{H}_{14}$, M 134.2, m -6.3°, b 79.4°/10mm, 204–205°/760mm, d_4^{20} 0.905, n_D^{20} 1.5203. Dry it over sodium and distil under reduced pressure. The *picrate* has m 92–95° (EtOH). [*Beilstein* 5 H 430, 5 I 206, 5 II 329, 5 III 974, 5 IV 1072.]

1,2,3,5-Tetramethylbenzene (isodurene) [527-53-7] has m -23.7°, -24°, b 74.4°/10mm, 198°/760mm, d_4^{20} 0.890, n_D^{20} 1.5130. Reflux isodurene over sodium and distil it under reduced pressure. [Smith *Org Synth Coll Vol* 2 248 1943, DOI: 10.15227/orgsyn.010.0032; *Beilstein* 5 H 430, 5 II 329, 5 III 976, 5 IV 1073.]

1,2,4,5-tetramethylbenzene (durene) [95-93-2] has m 77° to 80°, 79.5–80.5°, 191–192°/760mm, d_4^{20} 0.838. Chromatograph durene on alumina, and recrystallise it from aqueous EtOH or *benzene. Dry it under vacuum. Zone-refining removes duroaldehydes. [Yamauchi et al. *J Phys Chem* 89 4804 1985, DOI: 10.1021/j100268a029.] It is been prepared from xylene/ $\text{MeCl}/\text{AlCl}_3$ under conditions which produce a high proportion of durene which is separated from other methylbenzenes by fractional distillation using a good column, then steam distilled, the durene is collected from the distillate, dried (CaCl_2), filtered, distilled through a good column, then dried over Na and redistilled. It is poorly soluble in H_2O but is steam volatile, and soluble in

organic solvents. It can be sublimed unchanged, but it is best to sublime it *in vacuo*. [Johnston et al. *J Am Chem Soc* **109** 1291 1987, DOI: 10.1021/ja00239a001; Smith *Org Synth Coll Vol* **2** 248 1943, DOI: 10.15227/orgsyn.010.0032; *Beilstein* **5** H 431, **5** I 207, **5** II 329, **5** III 979, **5** IV 1076.]

***N,N,N',N'*-Tetramethylbenzidine** [366-29-0] $C_{16}H_{20}N_2$, **M 240.4**, **m 193-195°**, **195.4-195.6°**, **pK_{Est(1)}~3.4**, **pK_{Est(2)}~4.5**. Crystallise the benzidine from EtOH or petroleum ether, then from petroleum ether/*benzene, and sublime it in a vacuum. [Guarr et al. *J Am Chem Soc* **107** 5104 1985, DOI: 10.1021/ja00304a015.] Dry it *in vacuo* in a drying pistol, or a vacuum line. It has **m 195-196°** after sublimation. [*Beilstein* **13** H 221, **13** I 61, **13** II 97, **13** III 429, **13** IV 368.]

***p,p'*-Tetramethyldiaminodiphenylmethane** [bis(*p*-dimethylaminophenyl)methane, Michler's base, (*p,p'*-methylene-bis-(*N,N*-dimethylaniline)] [101-61-1, 30135-64-9] $C_{17}H_{22}N_2$, **M 254.4**, **m 89-90°**, **b 155-157°/0.1mm**, **pK_{Est(1)}~5.8**, **pK_{Est(2)}~5.1**. Crystallise the base from EtOH (2ml/g) or 95% EtOH (*ca* 12ml/g). It sublimes on heating. [*Beilstein* **13** IV 390.] Used in fixing latent fingerprints developed with iodine [Trowell *J Forensic Sci Soc* **15** 189 1975, DOI:10.1016/S0015-7368(75)70984-2].

***N,N,N',N'*-Tetramethyl-1,8-naphthalenediamine** [Proton sponge, 1,8-bis-(dimethylamino)-naphthalene [20734-58-1] $C_{14}H_{18}N_2$, **M 214.3**, **m 45-48°**, **47-48°**, **47-51°**, **b 144-145°/4mm**, **pK₁ -10.5 (from half protonation in 86% aqueous H₂SO₄, diprotonation)**, **pK₂ 12.34 (monoprotonation)**. It is prepared by methylating 1,8-diaminonaphthalene, and likely impurities are methylated products. The tetramethyl compound is a stronger base than the unmethylated, di and trimethylated derivatives. The pK_a values are: 1,8-(NH₂)₂ = 4.61, 1,8-(NHMe)₂ = 5.61, 1-NHMe-8-NHMe₂ = 6.43 and 1,8-(NMe₂)₂ = 12.34. The mixture is then treated with H₂O at pH 8 (where all but the required base are protonated) and extracted with Et₂O or CHCl₃. The dried extract (K₂CO₃) yields the tetramethyldiamine on evaporation which can be distilled. It is a strong base with weak nucleophilic properties, e.g. it could not be alkylated by refluxing with EtI in MeCN for 4 days; and on treatment with methyl fluorosulfonate only the fluorosulfonate salt of the base is obtained. [NMR: Adler et al. *Chem Commun* (London) 723 1968, DOI: 10.1039/C19680000723; Brown & Letang *J Am Chem Soc* **63** 358 1941, DOI: 10.1021/ja01847a009; Brzezinski et al. *JCS Perkin Trans 2* 857 1991, DOI: 10.1039/P29910000857.] Alternatively, crystallise *proton sponge* from EtOH and dry it in a vacuum oven. Store it in the dark in a CO₂-free atmosphere. [Benoit et al. *Can J Chem* **65** 996 1987, DOI: 10.1139/v87-170; *Beilstein* **13** IV 344.] Also used as a matrix for MALDI-MS, and traces of the cations of Ba(at ≈5mg/kg), Cd(at ≈5mg/kg), Co(at ≈5mg/kg), Cr(at ≈5mg/kg), Cu(at ≈5mg/kg), Fe(at ≈100mg/kg), K(at ≈20mg/kg), Mg(at ≈5mg/kg), Mn(at ≈5mg/kg), Na(at ≈50mg/kg), Ni(at ≈5mg/kg), Pb(at ≈5mg/kg), Zn(at ≈5mg/kg) can be detected, as well as lipids and metabolites.

***N,N,N',N'*-Tetramethyl-1,4-phenylenediamine (TMPD, Wurster's blue)** [100-22-1] $C_{10}H_{16}N_2$, **M 164.3**, **m 51°**, **b 135°/14mm**, **260°/760mm**, **pK₁²⁰ 2.29**, **pK₂²⁰ 6.35**. Crystallise the amine from petroleum ether or water. It can be sublimed or dried carefully in a vacuum line, and stored in the dark under nitrogen. It has been recrystallised from its melt. It loses two electrons stepwise upon oxidation; the radical-cation has a characteristic blue-violet colour, and is used in studies of electron transport for oxidase tests of biological electron transport systems [Michaelis et al. *J Am Chem Soc* **61** 1981 1939, DOI: 10.1021/ja01877a013]. [*Beilstein* **13** H 74, **13** I 22, **13** II 40, **13** III 111, IV 107.]

***N,N,N',N'*-Tetramethyl-1,4-phenylenediamine dihydrochloride (Wurster's Reagent; see preceding entry)** [637-01-4] $C_{10}H_{16}N_2 \cdot 2HCl$, **M 237.2**, has **m 222-224°**, **224°(dec)**. Crystallise the salt from isopropyl or *n*-butyl alcohols, saturated with HCl. Treat it with aqueous NaOH to give the *free base* (see previous entry) which is filtered, dried and sublimed in a vacuum. [Guarr et al. *J Am Chem Soc* **107** 5104 1985, DOI: 10.1021/ja00304a015; *Beilstein* **13** H 74.] Oxidase reagent (1% aqueous solution) used for testing cytochrome +ve aerobic microorganisms.

Tetra(4-nitrophenyl)ethylene [47797-98-8] $C_{26}H_{16}N_4O_8$, **M 512.4**, **m 298-299°**, **306-307°(corr)**. Crystallise it from dioxane or AcOH (**m 292°**, yellow needles), and dry it at 150°/0.1mm. [Gorvin *J Chem Soc* 678 1959, DOI: 10.1039/JR9590000678, Schlenk *Justus Liebigs Ann Chem* **394** 178 1912, DOI: 10.1002/jlac.19123940203; *Beilstein* **5** H 744, **5** III 2600, **5** IV 2782.]

Tetraphenylethylene [632-51-9] $C_{26}H_{20}$, M 332.4, m 223-224°, 224°, 225°, b 415-425°/760mm. Crystallise the ethylene from dioxane or from EtOH/*C₆H₆. Sublime it under high vacuum. [Beilstein 5 IV 2780.]

Tetraphenylhydrazine [632-52-0] $C_{24}H_{20}N_2$, M 336.4, m 147°, pK_{Est} ~0. Crystallise the hydrazine from 1:1 CHCl₃/toluene, 1:5 CHCl₃/EtOH (m 149°), *C₆H₆ or *C₆H₆/petroleum ether. Store it in a refrigerator, in the dark. [Beilstein 15 H 125, 15 I 29, 15 III 77, 15 IV 59.]

trans-1,1,4,4-Tetraphenyl-2-methylbutadiene [20411-57-8] $C_{29}H_{24}$, M 372.5. Crystallise it from EtOH or AcOH. [Wittig & Obermann *Chem Ber* 68 2214 1935, DOI: 10.1002/cber.19350681210; Beilstein 5 IV 2816.]

5,6,11,12-Tetraphenylnaphthacene (Rubrene) [517-51-1] $C_{42}H_{28}$, M 532.7, m>315°, 322°, 330-335°, d₄²⁰ 1.255. Rubrene forms orange crystals on sublimation at 250-260°/3-4mm [UV Badger & Pearce *Spectrochim Acta* 4 280 1950, DOI: 10.1016/0371-1951(51)80073-0]. It has also been recrystallised from *benzene under red light because it is chemiluminescent and light sensitive. [Beilstein 5 IV 2968.]

1,2,3,4-Tetraphenylnaphthalene [751-38-2] $C_{34}H_{24}$, M 432.6, m 199-201°, 204-204.5°. Crystallise the naphthalene from MeOH or EtOH. [Fieser & Haddadin *Org Synth* 46 107 1966, DOI: 10.15227/orgsyn.046.0107; Beilstein 5 IV 2918.]

Thymol (2-isopropyl-5-methylphenol) [89-83-8] $C_{10}H_{14}O_2S$, M 150.2, m 49-51°, 51.5°, b 232°/atm, d₄²⁵ 0.965, n_D²⁵ 1.5204, pK₂₀ 10.62. It occurs in the volatile oils of *Thymus vulgaris* and *Moranda punctata* L. from which it was first isolated [Arppe *Justus Liebigs Ann Chem* 58 41 1846, DOI: 10.1002/jlac.18460580107]. It is quite volatile at 100°, and is steam volatile, separating as white crystals which have a characteristic pungent odour, with a caustic taste. It should be stored in the dark, preferably under N₂ as it oxidises slowly in air. It possesses antimould properties, is an internal and external antiseptic, and has nematode anthelmintic activity. The *acetate* [528-93-2] is a yellow irritating oil, b 243.5-245.5°/atm, d⁰ 1.009, with the odour of thymol, and is soluble in most organic solvents, but almost insoluble in H₂O. [Beilstein 6 IV 3334.] Antimicrobial.

Thymolphthalein complexone {TPC, 3,3'-bis[N,N-di(carboxymethyl)aminomethyl]thymolphthalein} [1913-93-5] $C_{38}H_{44}N_2O_{12}$, M 720.8, m 190°(dec), 191°, pK₁^{18.2} 7.35, pK₂^{18.2} 12.25. Purify it as for phthalein complexone except that it is synthesised from thymolphthalein instead of cresolphthalein. It has absorption maxima at 600-605nm (A_{1cm}^{1%} 350-345) in aqueous 0.1NaOH. [Beilstein 18/4 V 194.]

o-Tolidine (3,3'-dimethylbenzidine) [119-93-7] $C_{14}H_{16}N_2$, M 212.3, m 129°, 131-132°, b 300.5°/atm, d 1.23, pK₂₅ 4.45. Dissolve the tolidine in *benzene by percolation through a column of activated alumina and crystallise it from *benzene/petroleum ether. [Beilstein 13 IV 410.]

p-Tolualdehyde [104-87-0] C_8H_8O , M 120.2, m -6°, b 83-85°/0.1mm, 199-200°/atm, 205°/760mm, d₄²⁰ 1.018, n_D²⁰ 1.5479. Steam distil the aldehyde, dry it with CaSO₄, then fractionally distil it. [Beilstein 7 IV 672.]

o-Toluamide [527-85-5] C_8H_9O , M 135.2, m 141°, 142°, 144-145°, 147°. Crystallise o-toluamide from hot water (10ml/g) and dry in air. [Noller *Org Synth Coll Vol* 2 586 1943, DOI: 10.15227/orgsyn.013.0094; Beilstein 9 H 465, 9 II 319, 9 III 2304.]

Toluene [108-88-3] C_7H_8 , M 92.1, m -93°, -94.9°, -95°, b 110.6°/760mm, 111°/760mm, d₄¹⁰ 0.87615, d₄²⁵ 0.86231, n_D²⁰ 1.49693, n_D²⁵ 1.49413. Dry toluene with CaCl₂, CaH₂ or CaSO₄, and dry further by standing with sodium, P₂O₅ or CaH₂. It can be fractionally distilled from sodium or P₂O₅. Unless specially purified, toluene is likely to be contaminated with methylthiophenes and other sulfur-containing impurities. These can be removed by shaking with concentrated H₂SO₄, but the temperature must be kept below 30° if sulfonation of toluene is to be avoided. A typical procedure consists of shaking toluene twice with cold concentrated H₂SO₄ (100ml of acid per L), once with water, once with aqueous 5% NaHCO₃ or NaOH, again with H₂O, then drying successively with CaSO₄ and P₂O₅, with final distillation from P₂O₅ or over LiAlH₄ after refluxing for 30 minutes. *Alternatively*, NaHCO₃ can be replaced by boiling under reflux with 1% sodium amalgam. Sulfur compounds can also be removed by prolonged shaking of the toluene with mercury, or by two distillations from

AlCl_3 , the distillate then being washed with water, dried with K_2CO_3 and stored with sodium wire. Other purification procedures include refluxing and distillation of sodium dried toluene from diphenylpicrylhydrazyl, and from SnCl_2 (to ensure freedom from peroxides). It has also been co-distilled with 10% by volume of ethyl methyl ketone, and again fractionally distilled. [Brown & Pearsall *J Am Chem Soc* **74** 191 1952, DOI: 10.1021/ja01121a049.] For removal of carbonyl impurities see **benzene*. Toluene has been purified by distillation under nitrogen in the presence of sodium benzophenone ketyl. Toluene has also been dried with MgSO_4 , after the sulfur impurities have been removed, and then fractionally distilled from P_2O_5 and stored in the dark [Tabushi et al. *J Am Chem Soc* **107** 4466 1985, DOI: 10.1021/ja00301a016]. Toluene can be purified by passage through a tightly packed column of Fuller's earth.

Rapid purification: Alumina, CaH_2 and 4A molecular sieves (3% w/v) may be used to dry toluene (6 hours stirring and standing). Then the toluene is distilled, discarding the first 5% of distillate, and is stored over molecular sieves (3A, 4A) or Na wire. [Beilstein **5** H 280, **5** I 144, **5** II 209, **5** III 651, **5** IV 766.]

Toluene-2,4-diamine (4-methyl-*m*-phenylenediamine) [95-80-7] $\text{C}_7\text{H}_{10}\text{N}_2$, **M 122.2**, **m 99°**, **b 148-150°/8mm**, **292°/760mm**, **pK_{Est(1)}~2.5**, **pK_{Est(2)}~4.4**. Recrystallise the diamine from water (solubility is 0.75w/v%) containing a very small amount of sodium dithionite (to prevent air oxidation), and dry it under vacuum. It also crystallises from **benzene*. [Beilstein **13** IV 235.]

Tolhydroquinone (1,4-dihydroxy-2-methylbenzene) [95-71-6, 96937-50-7] $\text{C}_7\text{H}_8\text{O}_2$, **M 124.1**, **m 125°**, **126°**, **128-129°**, **d 1.34**, **pK₁²⁰ 10.15**, **pK₂²⁰ 11.75**. Crystallise the quinone from EtOH. [Beilstein **6** IV 5866.]

***o*-Toluic acid** [118-90-1] $\text{C}_8\text{H}_8\text{O}_2$, **M 136.2**, **m 102-103°**, **104-105°**, **b 258-259°/760mm**, **d 1.06**, **pK²⁵ 3.91**. Crystallise the acid from **benzene* (2.5ml/g) and dry in air. The *S-benzylisothiuronium salt* has **m 146°** (from aqueous EtOH). [Beilstein **9** IV 1697.]

***m*-Toluic acid** [99-04-7] has **m 108.7°**, **108-110°**, **112°**, **111-113°**, **b 263°/760mm**, **d 1.05**, **pK²⁵ 4.27**. Crystallise the acid from water. [Beilstein **9** IV 1712.] Aromatic acid impurities (to <0.05%) can be removed *via* the (±)-α-methylbenzylamine salt as described for 2,4-dichlorobenzoic acid [Ley & Yates *Organic Process Research & Development* **12** 120 2008, DOI: 10.1021/op7001547]. The *S-benzylisothiuronium salt* has **m 140°** (from aqueous EtOH).

***p*-Toluic acid** [99-94-5] has **m 178.5-179.5°**, **180°**, **181°**, **178-182°**, **d 1.06**, **b 274-275°/760mm**, **pK²⁵ 4.37**. Crystallise the acid from water, water/EtOH (1:1), MeOH/water or **benzene*. [Beilstein **9** IV 1724.] Aromatic acid impurities (to <0.05%) can be removed *via* the (±)-α-methylbenzylamine salt as described for 2,4-dichlorobenzoic acid [Ley & Yates *Organic Process Research & Development* **12** 120 2008, DOI: 10.1021/op7001547]. The *S-benzylisothiuronium salt* has **m 164°** (from aqueous EtOH).

***o*-Toluidine (2-methylaniline)** [95-53-4, 162607-18-3] $\text{C}_7\text{H}_9\text{N}$, **M 107.2**, **m -24°**, **-16.3°**, **-14.4°**, **b 80.1°/10mm**, **200.3°/760mm**, **d₄²⁰ 0.999**, **n_D²⁰ 1.57246**, **n_D²⁵ 1.56987**, **pK²⁵ 4.45**. In general, methods similar to those for purifying aniline can be used, e.g. distillation from zinc dust, at reduced pressure, under nitrogen. Berliner and May [*J Am Chem Soc* **49** 1007 1927, DOI: 10.1021/ja01403a017] purified it *via* the oxalate. Twice-distilled *o*-toluidine is dissolved in four times its volume of diethyl ether, and the equivalent amount of oxalic acid needed to form the di-oxalate is added as its solution in diethyl ether. (If *p*-toluidine is present, its oxalate precipitates and can be removed by filtration.) Evaporation of the ethereal solution gives crystals of ***o*-toluidine di-oxalate** [Beilstein **12** III 1494, **12** IV 1817]. These are filtered off, recrystallised five times from water containing a small amount of oxalic acid (to prevent hydrolysis), then treated with dilute aqueous Na_2CO_3 to liberate the amine which is separated, dried (CaCl_2) and distilled under reduced pressure. The *benzoyl* derivative has **m 144°** (from EtOH). [Beilstein **12** H 772, **12** I 372, **12** II 429, **12** III 1837, **12** IV 1744.]

***m*-Toluidine (3-methylaniline)** [108-44-1] has **m -30.4°**, **-31.3°**, **b 82.3°/10mm**, **203.4°/760mm**, **d₄²⁰ 0.989**, **n_D²⁰ 1.56811**, **n_D²⁵ 1.56570**, **pK²⁵ 4.71**. It can be purified as for aniline. Twice-distilled, *m*-toluidine is converted to the hydrochloride using a slight excess of HCl, and the salt is fractionally crystallised from 25% EtOH (five times), and from distilled water (twice), rejecting, in each case, the first material that crystallised out. The amine is regenerated and distilled as for *o*-toluidine. The *benzoyl* derivative has **m 125°** (from EtOH). [Berliner & May *J Am Chem Soc* **49** 1007 1927, DOI: 10.1021/ja01403a017; Beilstein **12** II 463, **12** III 1949, **12** IV 1813.]

***p*-Toluidine (4-methylaniline)** [106-49-0] has **m 42-46°, 43°, 43.6°, 44°, 44.8°, b 79.6°/10mm, 200.5°/760mm, d_4^{20} 0.962, n_D^{20} 1.5636, $n_D^{59.1}$ 1.5534, pK^{25} 5.08**. In general, methods similar to those for purifying aniline can be used. It can be separated from the *o*- and *m*-isomers by fractional crystallisation from its melt. *p*-Toluidine has been crystallised from hot water (charcoal), EtOH, *benzene, petroleum ether or EtOH/water (1:4), and dried in a vacuum desiccator. It can also be sublimed at 30° under vacuum. For further purification, use has been made of the oxalate, the sulfate and acetylation. The **oxalate**, formed as described for *o*-toluidine, is filtered, washed and recrystallised three times from hot distilled water. The base is regenerated with aqueous Na₂CO₃ and recrystallised three times from distilled water. [Berliner & May *J Am Chem Soc* **49** 1007 1927, DOI: 10.1021/ja01403a017.] Alternatively, *p*-toluidine is converted to its **acetyl** derivative which, after repeated crystallisation from EtOH, is hydrolysed by refluxing (50g) in a mixture of 500ml of water and 115ml of concentrated H₂SO₄ until a clear solution is obtained. The amine sulfate is isolated, suspended in water, and NaOH is added. The free base is distilled twice from zinc dust under vacuum. The *p*-toluidine is then recrystallised from petroleum ether and dried in a vacuum desiccator or in a vacuum for 6 hours at 40°. The **benzoyl** derivative has **m 158°** (from EtOH). [Berliner & Berliner *J Am Chem Soc* **76** 6179 1954, DOI: 10.1021/ja01652a092; Moore et al. *J Am Chem Soc* **108** 2257 1986, DOI: 10.1021/ja00269a022; *Beilstein* **12** H 880, **12** I 140, **12** II 482, **12** III 2017, **12** IV 1866.] ***p*-Toluidine hydrochloride** [540-23-8] **C₇H₉N.HCl, M 143.6**, has **m 243-245°, 245.9-246.1°**. Crystallise the salt from MeOH containing a few drops of concentrated HCl or aqueous EtOH. Dry it under vacuum over paraffin chips. [*Beilstein* **12** II 587, **12** III 2021, **12** IV 1869.]

***o*-Tolunitrile** [529-19-1, 25550-22-5] **C₈H₇N, M 117.2, m -14°, -13.5°, -13°, b 205.2°/atm, d_4^{20} 0.992, n_D^{20} 1.5279**. Fractionally distil the nitrile, wash it with concentrated HCl or 50% H₂SO₄ at 60° until the smell of isonitrile has gone (this also removes any amines), then wash it with saturated NaHCO₃ and dilute NaCl solutions, then dry it with K₂CO₃ and redistil it. [*Beilstein* **9** IV 1703.]

***m*-Tolunitrile** [620-22-4] has **m -23°, b 99-101°/20mm, 209.5-210°/773mm, d_4^{20} 0.986, n_D^{20} 1.5250**. Dry the nitrile with MgSO₄, fractionally distil it, then wash it with aqueous acid to remove possible traces of amines, dry and redistil it. [*Beilstein* **9** H 477, **9** I 191, **9** II 325, **9** III 2324, **9** IV 1717.]

***p*-Tolunitrile** [104-85-8] has **m 26-29°, 27°, 28°, 29.5°, b 104-106°/20mm, 217°/atm, 218°/atm, d_4^{20} 0.981**. Melt the nitrile, dry it with MgSO₄, fractionally crystallise it from its melt, then fractionally distil it under reduced pressure in a 6-in spinning band column. [Brown *J Am Chem Soc* **81** 3232 1959, DOI: 10.1021/ja01522a018.] It can also be crystallised from *benzene/petroleum ether (b 40-60°). [*Beilstein* **9** H 489, **9** I 194, **9** II 330, **9** III 2348, **9** IV 1738.]

4-Tolyl-2-benzoic acid (4'-methylbiphenyl-2-carboxylic acid) [7148-03-0] **C₁₄H₁₂O₂, M 212.2, m 138-139°, 146-148°, 148-152°, pK^{25} 3.64**. Crystallise the acid from toluene or *C₆H₆ (m 147-148°). [*Beilstein* **9** H 677, **9** IV 2523.]

***p*-Tolyl carbinol (4-methylbenzyl alcohol)** [589-18-4] **C₈H₁₀O, M 122.2, m 58-61°, 59.5-60°, 61°, b 116-118°/20mm, 217°/760mm**. Recrystallise the alcohol from petroleum ether (b 80-100°, 1g/ml), Et₂O, pentane or H₂O (m 61-62.1°). It can also be distilled in a vacuum. [*Beilstein* **6** H 498, **6** I 248, **6** II 469, **6** III 1779.]

Tolylene-2,4-diisocyanate (toluene-2,4-diisocyanate) [584-84-9] **C₉H₆N₂O₂, M 174.2, m 19.5-21.5°, 21.8°, 20-22°, 28°, b 126°/11mm, 124-126°/18mm, 251°/760mm, d^{25} 1.214g/ml**. It is purified by fractionation in a vacuum and should be stored in a dry atmosphere. It is soluble in organic solvents but reacts with H₂O, alcohols (slowly) and amines, all of which could cause explosive polymerisation. It darkens on exposure to light. It has a sharp pungent odour, is **TOXIC** and is **IRRITATING TO THE EYES**. [Siefken *Justus Liebigs Ann Chem* **562** 75, 96, 127 1949, DOI: 10.1002/jlac.19495620202; Bayer *Angew Chem* **59** 257 1947, DOI: 10.1002/ange.19470590901] It is a reagent for covalent crosslinking of proteins [Wold *Methods Enzymol* **25** 623 1972, DOI: 10.1016/S0076-6879(72)25061-3.] [*Beilstein* **13** IV 243.]

Tolylene-2,6-diisocyanate (2-methyl-*m*-phenylenediisocyanate) [91-08-7] **C₉H₆N₂O₂, M 174.2, m 13°, b 129-133°/18mm, 246-247°/760mm, d_4^{20} 1.225, n_D^{20} 1.5715**. It is purified by fractional distillation in a vacuum. Store it under N₂ in sealed dark ampoules as it is water and light sensitive. Like the preceding 2,4-isomer, it has a sharp pungent odour, is **TOXIC** and is **IRRITATING TO THE EYES**. [*Beilstein* **13** IV 259.]

p-Tolyl urea [622-51-5] $C_8H_{10}N_2O$, M 150.2, m 177-184°, 181°, 182°, 182-83°. Crystallise the urea from H_2O (m 186°), EtOH/water (1:1) or aqueous AcOH (m 184°). [Beilstein 12 H 941, 12 I 425, 12 II 512, 12 III 2084, 12 IV 1923.]

Tribenzylamine [620-40-6] $C_{21}H_{21}N$, M 287.4, m 91.5°, 93°, 93-94°, b 230°/13mm, 417.2°/760mm, $pK_{Est} < 0$. Crystallise the amine from absolute EtOH or petroleum ether. Dry it in a vacuum over P_2O_5 at room temperature. The *hydrochloride* has m 226-228° (from EtOH) and the *picrate* has m 191° (from H_2O or aqueous EtOH). [Beilstein 12 IV 2183.]

2,4,6-Tribromoaniline [147-82-0] $C_6H_3Br_3N$, M 329.8, m 120°, 121°, 122°, b 300°/atm, $d^{25} 2.35g/ml$. $pK_{Est} \sim -0.5$ (aqueous H_2SO_4). Crystallise the aniline from MeOH. The *benzenesulfonamide derivative* has m 198°. [Beilstein 12 H 663, 12 I 329, 12 II 358, 12 III 1477, 12 IV 1538.] 2,4,6-Tribromoacetanilide [607-93-2] $C_8H_6Br_3NO$, M 451.8, has m 232°, 238-240°, 240°. Crystallise the anilide from EtOH. [Beilstein 12 II 359, 12 III 1478.]

sym-Tribromobenzene (1,3,5-tribromobenzene) [626-39-1] $C_6H_3Br_3$, M 314.8, m 119-123°, 122°, 122°, 122.8°, 124°, b 271°/atm. Crystallise it from glacial acetic acid/water (4:1), then wash with chilled EtOH and dry in air. [Beilstein 5 H 213, 5 IV 685.]

2,4,6-Tribromophenol [118-79-6] $C_6H_3Br_3O$, M 330.8, m 86-92°, 89°, 94°, 95.5°, b 244°/atm (282-289° sublimes), $d^{25} 2.55g/ml$, $pK^{25} 6.00$. Crystallise the phenol from EtOH or petroleum ether. Dry it under vacuum over P_2O_5 at room temperature. [Beilstein 6 IV 1067.]

sym-Tri-*tert*-butylbenzene (1,3,5-tri-*tert*-butylbenzene) [1460-02-2] $C_{18}H_{30}$, M 246.4, m 67° to 72°, 73.4-73.9°, b 121-122°/12mm. Crystallise it from EtOH. [Beilstein 5 IV 1206.]

2,4,6-Tri-*tert*-butylphenol [732-26-3] $C_{18}H_{30}O$, M 262.4, m 125-130°, 129-132°, 131°, 131-131.2°, b 131°/1mm, 147°/10mm, 278°/760mm, $pK^{25} 12.19$. Distil the phenol under reduced pressure and/or recrystallise it from *n*-hexane or several times from 95% EtOH until the EtOH solution is colourless [Balasubramanian & Bruce *J Am Chem Soc* 108 5495 1986, DOI: 10.1021/ja00278a021]. It has also been purified by sublimation [Yuan & Bruce *J Am Chem Soc* 108 1643 1986, DOI: 10.1021/ja00267a039; Wong et al. *J Am Chem Soc* 109 3428 1987, DOI: 10.1021/ja00245a039]. Purification has also been achieved by passage through a silica gel column followed by recrystallisation from *n*-hexane [Kajii et al. *J Phys Chem* 91 2791 1987, DOI: 10.1021/j100295a029]. [Beilstein 6 III 2094, 6 IV 3539.]

2',2',2'-Trichloroacetanilide [2563-97-5] $C_8H_6Cl_3NO$, M 238.5, m 93.5-94°, 95°. Crystallise the anilide from *benzene or 90% EtOH (m 93.5-95.5°). [Sukornick *Org Synth Coll Vol* 5 1074 1973, DOI: 10.15227/orgsyn.040.0103; Beilstein 12 H 224, 12 I 193, 12 II 142, 12 III 464, 12 IV 377.]

2,3,4-Trichloroaniline [634-67-3] $C_6H_4Cl_3N$, M 196.5, m 65-67°, 67.5°, b 292°/774mm, $pK_{Est} \sim 1.3$. Recrystallise it from ligroin. The *acetanilide* has m 120-122° (from EtOH or * C_6H_6). [Beilstein 12 H 626.]

2,4,5-Trichloroaniline [636-30-6] $C_6H_4Cl_3N$, M 196.5, has m 93-95°, 96.5°, b 270°/760mm, $pK 1.09$. Crystallise the aniline from ligroin. [Beilstein 12 H 627, 12 IV 1277.]

2,4,6-Trichloroaniline [634-93-5] $C_6H_4Cl_3N$, M 196.5, has m 73-75°, 75-79°, 74°, 77°, 78°, 78.5°, b 127°/14mm, 262°/746mm, $pK^{25} 0.03$. Crystallise the aniline from ligroin. The *benzoyl* derivative has m 174° (from EtOH). [Beilstein 12 H 627, 12 IV 1281.]

1,2,3-Trichlorobenzene [87-61-6] $C_6H_3Cl_3$, M 181.5, m 51-55°, 52.6°, 53°, 54°, 55°, 218-219°/740mm, d 1.69. Crystallise it from EtOH. [Beilstein 5 IV 664.]

1,2,4-Trichlorobenzene [120-82-1] $C_6H_3Cl_3$, M 181.5, has m 16.9°, 17°, b 210°/atm, 214.6°/760mm, d 1.454. Separate it from isomers by washing with fuming H_2SO_4 , then water, drying with $CaSO_4$ and slowly fractionally distilling. [Jensen et al. *J Am Chem Soc* 81 3303 1959, DOI: 10.1021/ja01522a038; Beilstein 5 IV 664.]

1,3,5-Trichlorobenzene [108-70-3] $C_6H_3Cl_3$, M 181.5, m 62-65°, 63°, 64°, 64-65°, 208°/740mm. Recrystallise it from dry *benzene or toluene. [Beilstein 5 IV 666.]

3,4,5-Trichloro-*o*-cresol (3,4,5-trichloro-2-methylphenol) [608-92-4] $C_7H_5Cl_3O$, M 211.5, m 77°, pK_{Est} ~7.6. Crystallise the cresol from petroleum ether.

2,3,5-Trichloro-*p*-cresol (2,3,5-trichloro-4-methylphenol) [608-91-3] has m 66-67°, pK_{Est} ~6.9. Crystallise the cresol from petroleum ether. [Datta & Mitter *J Am Chem Soc* 41 2028 1919, DOI: 10.1021/ja02233a020; Beilstein 6 I 204.]

2,4,5-Trichloro-1-nitrobenzene (1,2,4-trichloro-5-nitrobenzene) [89-69-0] $C_6H_2Cl_3NO_2$, M 226.5, m 52-57°, 57°, 288°/740mm, d^{25} 1.79g/ml. Crystallise it from EtOH. Its solubility in H_2O is 0.27w/v% at 20°. [Beilstein 5 IV 728.]

3,4,6-Trichloro-2-nitrophenol [82-62-2, 4524-78-1] $C_6H_2Cl_3NO_3$, M 226.5, m 92-93°, pK_{Est} ~4.1. Crystallise the nitro-phenol from petroleum ether or EtOH. [Beilstein 6 III 842.]

2,4,5-Trichlorophenol [95-95-4] $C_6H_3Cl_3O$, M 197.5, m 67°, 67-69°, b 72°/1mm, 248°/740mm, d^{25} 1.79g/ml, pK^{25} 7.0. Crystallise the phenol from EtOH or petroleum ether. [Beilstein 6 IV 962.]

2,4,6-Trichlorophenol [88-06-2] has m 67-68°, 69°, b 246°/760mm, d^{25} 1.675g/ml, pK^{25} 6.23. Crystallise the phenol from *benzene, EtOH or EtOH/water. [Beilstein 6 IV 1005.] It is an environmental pollutant.

3,4,5-Trichlorophenol [609-19-8] has m 100°, d^{25} 1.80g/ml, pK^{25} 7.84. Crystallise the phenol from petroleum ether/*benzene mixture. [Beilstein 6 III 729.]

2,4,5-Trichlorophenoxyacetic acid (2,4,5-T) [93-76-5] $C_8H_5Cl_3O_3$, M 255.5, m 153°, 155-158°, pK^{25} 2.83. Crystallise this phenoxyacid from *benzene. [Beilstein 6 III 721.] It is a herbicide, and CANCER SUSPECT.

1,2,4-Triethylbenzene [877-44-1] $C_{12}H_{18}$, M 162.3, b 96.8-97.1°/12.8mm, 215°/atm, d_4^{20} 0.8738, n_D^{20} 1.5015. For separation from a commercial mixture see Dillingham and Reid [*J Am Chem Soc* 60 2606 1938, DOI: 10.1021/ja01278a014]. [Beilstein 5 IV 1133.]

1,3,5-Triethylbenzene [102-25-0, 102-49-0, 25340-18-5] has m -67.1°, -66°, -65.5°, b 136.85-137.15°, 215.9°/atm, 217.5°/760mm, d_4^{20} 0.8631, n_D^{20} 1.4957. For separation from a commercial mixture see Dillingham and Reid [*J Am Chem Soc* 60 2606 1938, DOI: 10.1021/ja01278a014]. The separation was done by fractionation at atmospheric pressure in a 125-plate miniature Hyper-Cal column and the middle fraction containing about 80% of the distillate was collected and found to be free from isomeric and other impurities (purity is 99.0-99.5 mole% from freezing point data). The IR (film) has ν_{max} 865 cm^{-1} . [McCaulay et al. *J Am Chem Soc* 76 2354 1954, DOI: 10.1021/ja01638a020; Beilstein 5 IV 1133.]

4-(Trifluoromethyl)acetophenone [709-63-7] $C_9H_7F_3O$, M 188.2, m 29-35°, 31°, 32°, 31-33°, b 79-81°/9mm, 81-84°/8-9mm, d^{25} 0.924g/ml. Purify the ketone by distillation or sublimation *in vacuo*. [Beilstein 7 IV 1404.]

3-Trifluoromethyl-4-nitrophenol (TFN) [88-30-2] $C_7H_4F_3NO_3$, M 162.1, m 81°, b 135-138°/0.01mm, pK_{Est} ~6.1. Crystallise the nitrophenol from *benzene or from petroleum ether/*benzene mixture. [Beilstein 6 III 1328.] It is a lampricide and is poisonous to amphibians— an environmental hazard.

α,α,α -Trifluorotoluene (benzotrifluoride) [98-08-8] $C_7H_5F_3$, M 146.1, m -29°, b 102.5°/atm, d_4^{20} 1.190, n_D^{30} 1.4100. Purify benzotrifluoride by repeated treatment with boiling aqueous Na_2CO_3 (until no test for chloride ion is obtained), dry it with K_2CO_3 , then with P_2O_5 , and fractionally distil it. [Beilstein 5 IV 802.]

2,3,4-Trihydroxybenzoic acid [610-02-6] $C_7H_6O_5$, M 170.1, m 207-208°, 210°(dec), 221°, $pK_{Est(1)}$ ~3.4, $pK_{Est(2)}$ ~7.8, $pK_{Est(3)}$ >12. Crystallise the acid from water. [Beilstein 10 IV 1971.]

2,4,6-Trihydroxybenzoic acid [83-30-7] $C_7H_6O_5$, M 170.1, m 205-212°(dec), $pK_{Est(1)}$ ~1.5, $pK_{Est(2)}$ ~8.0, $pK_{Est(3)}$ >12. Crystallise the acid from water which provides crystals of a *monohydrate* [Beilstein 10 IV 1987.]

3,4,5-Triiodobenzoic acid [2338-20-7] $C_7H_3I_3O_2$, M 499.8, m 289-290°, 292.5°, 293°, pK^{25} 0.65. Crystallise the acid from aqueous EtOH or water. [Beilstein 9 H 367, 9 III 1475.]

3,4,5-Triiodobenzyl chloride [52273-54-8] $C_7H_4Cl_3$, M 504.4, m 138°. Crystallise the chloride from CCl_4 /petroleum ether (charcoal).

Trimellitic (benzene-1,2,4-tricarboxylic) acid [528-44-9] $C_9H_6O_6$, M 210.1, m 218-220°, pK_1^{25} 2.42, pK_2^{25} 3.71, pK_3^{25} 5.01. Crystallise the acid from acetic acid or aqueous EtOH. [Beilstein 9 IV 3746.]

1,2,3-Trimethoxybenzene [634-36-6] $C_9H_{12}O_3$, M 168.2, m 43-47°, 45-46°, b 241°/atm, d^{25} 1.112. Sublime it under vacuum. Used for studies of the solvent effects on photoinduced electron-transfer reactions [Niwa et al. *J Phys Chem* 97 11960 1993, DOI: 10.1021/j100148a019]. [Beilstein 6 I 540, 6 II 1066, 6 III 6265, 6 IV 7329.] **1,3,5-Trimethoxybenzene** [621-23-8] has m 50-53°, 53°, b 255°/atm, Sublime it under vacuum. Its solubility in MeOH is ~0.1g/ml at ~20°. [Beilstein 6 III 635, 6 IV 7362.]

3,4,5-Trimethoxyphenol (Antiarol) [642-71-7] $C_9H_{12}O_4$, M 184.2, m 146°, 145-149°, pK_{Est} ~9.4. Recrystallise the phenol from 10 times its weight of H_2O (white needles, m 148°). The *acetyl* derivative crystallises as elongated prisms from EtOH with m 74°. [Chapman et al. *J Chem Soc* 3028 1927, DOI: 10.1039/JR9270003015; Shriner et al. *J Am Chem Soc* 61 2322 1939, DOI: 10.1021/ja01878a017; Beilstein 6 H 1154, 6 II 1118, 6 III 6656.]

1,2,4-Trimethylbenzene (pseudocumene) [95-63-6] C_9H_{12} , M 120.2, m -43.78°, b 51.6°/10mm, 167-168°/760mm, d_4^{20} 0.889, n_D^{20} 1.5048. Reflux pseudocumene over sodium and distil it under reduced pressure. [Beilstein 6 H 1088, 6 I 542, 6 II 1072, 6 III 6278, 6 IV 7339.]

2,4,6-Trimethylbenzoic acid (mesitoic acid) [480-63-7] $C_9H_{12}O_2$, M 164.2, m 152°, 153-155°, 154°, 155°, pK^{25} 3.45. Crystallise mesitoic acid from water, ligroin or carbon tetrachloride [Ohwada et al. *J Am Chem Soc* 108 3029 1986, DOI: 10.1021/ja00271a036]. [Beilstein 9 H 553, 9 I 214, 9 II 360, 9 III 2489, 9 IV 1854.]

Trimethyl-1,4-benzoquinone [935-92-2] $C_9H_{10}O_2$, M 150.1, m 29-30°, 36°, b 98°/10mm, 108°/18mm, 215.3°/760mm, d 1.1, n 1.501. Distil the quinone in a vacuum or sublime it *in vacuo* before use. It is prepared in 96-100% yield by oxidation of 2,3,6-trimethylphenol with H_2O_2 over Ti(I) grafted on commercial mesoporous silica [Kholdeeva et al. *Green Chem* 9 731 2007, DOI: 10.1039/B617162A]. A quantitative structure-activity relationship of trimethylbenzoquinone, among many other quinones, in the two-electron reduction by *Enterobacter cloacae* NAD(P)H:nitroreductase was evaluated by Nivinskas et al. [*Arch Biochem Biophys* 403 249 2002, DOI: 10.1016/S0003-9861(02)00228-X]. [Beilstein 7 H 161, 7 III 3407, 7 IV 2098.]

Trimethyl-1,4-hydroquinone (2,3,5-trimethylbenzene-1,4-diol) [700-13-0] $C_9H_{12}O_2$, M 152.2, m 169-174°, 173-174°, b 298.3°/760mm, d^{25} 1.126g/ml, $pK_{Est(1)} \sim 11.1$, $pK_{Est(2)} \sim 12.7$. Recrystallise the hydroquinone from water, under anaerobic conditions. [Beilstein 6 H 931, 6 IV 5997.]

2,3,5-Trimethylphenol [697-82-5] $C_9H_{12}O$, M 136.2, m 92-95°, 95-96°, b 233°/760mm, d^{25} 0.763g/ml, pK^{25} 10.67. Crystallise the phenol from water or petroleum ether. [Beilstein 6 IV 3248.]

2,4,5-Trimethylphenol [496-78-6] has m 70.5-71.5°, 72°, b 232°/760mm, d^{25} 1.00g/ml, n_D^{20} 1.55, pK^{25} 10.57. Crystallise the phenol from water. [Beilstein 6 H 509, 6 I 255, 6 II 482, 6 III 1831, 6 IV 3247.]

2,4,6-Trimethylphenol [527-60-6] has m 69°, 70-74°, 72°, 73°, b 220-221°/760mm, pK^{25} 10.86. Crystallise the phenol from water and sublime it *in vacuo*. [Beilstein 6 IV 3253.]

3,4,5-Trimethylphenol [527-54-8] has m 104-108°, 107°, b 248-249°/760mm, pK^{25} 10.25. Crystallise the phenol from petroleum ether. [Beilstein 6 IV 3245.]

Trimethylphenylammonium benzenesulfonate [3426-74-2, 16093-66-6] $C_{15}H_{19}NO_3S$, M 293.3. Crystallise it repeatedly from MeOH (charcoal). [Beilstein 12 IV 249.]

2,4,6-Trinitroanisole [606-35-9] $C_7H_5N_3O_7$, M 243.1, m 68°, d^{25} 1.632g/ml. Recrystallise it from EtOH or

MeOH. Dry it *in vacuo*. [Beilstein 6 H 288, 6 I 140, 6 II 280, 6 III 968, 6 IV 1456.]

1,3,5-Trinitrobenzene [99-35-4] $C_6H_3N_3O_6$, M 213.1, m 122°, 122-123°. Crystallise it from glacial acetic acid, $CHCl_3$, CCl_4 , EtOH aqueous EtOH or EtOH/*benzene, after (optionally) heating with dilute HNO_3 . Dry it in air. Fuse it and crystallise it under vacuum. [Beilstein 5 H 271, 5 I 140, 5 II 203, 5 III 643, 5 IV 754.]

2,4,6-Trinitrobenzoic acid [129-66-8] $C_7H_3N_3O_8$, M 257.1, m 227-228°, 229°, pK^{25} 0.65. Crystallise the acid from distilled H_2O . Dry in a vacuum desiccator. The *amide* has m 264° (from EtOH). [Beilstein 9 H 417, 9 I 168, 9 II 285, 9 III 1956, 9 IV 1362.]

2,4,6-Trinitro-*m*-cresol [602-99-3] $C_7H_5N_3O_7$, M 243.1, m 107.0-107.5°, b 320.9°/760mm, d^{25} 1.74g/ml, pK^{25} 2.8. Crystallise the yellow cresol successively from H_2O , aqueous EtOH and *benzene/cyclohexane, then dry at 80° for 2 hours. [Davis & Paabo *J Res Nat Bur Stand* 64A 533 1960, DOI:org/10.6028/jres.064A.055; Beilstein 6 H 387, 6 I 194, 6 II 363, 6 III 1331, 6 IV 2079.]

2,4,7-Trinitro-9-fluorenone [129-79-3] $C_{13}H_5N_3O_7$, M 315.2, m 176°, 176-178°. Crystallise it from nitric acid/water (3:1), wash it with water and dry it under vacuum over P_2O_5 , or recrystallise it from dry *benzene. [Beilstein 7 II 410, 7 III 2348, 7 IV 1638.]

2,4,6-Trinitrotoluene (TNT) [118-96-7] $C_7H_5N_3O_6$, M 227.1, m 80°, 80.35°, 81.0-81.5°, b 295°/760mm (decomposes, avoid distilling due to possible explosive nature), d^{25} 1.65g/ml. Crystallise TNT from * C_6H_6 or EtOH. Then fuse (CARE) and allow to crystallise under vacuum. Gey, Dalbey and Van Dolah [*J Am Chem Soc* 78 1803 1956, DOI: 10.1021/ja01590a008] dissolved TNT in acetone and added cold water (1:2:15), the precipitate was filtered off, washed free from solvent and stirred with five parts of aqueous 8% Na_2SO_3 at 50-60° for 10 minutes. This was filtered, washed with cold water until the effluent was colourless, and air dried. The product was dissolved in five parts of hot CCl_4 , washed with warm water until the washings were colourless and TNT was recovered by cooling and filtering. It was recrystallised from 95% EtOH and carefully dried over H_2SO_4 . The dry solid should not be heated without taking precautions for a possible **EXPLOSION**. Work with small quantities. Irritates skin, mucous membrane, causes liver damage, jaundice; cyanosis, sneezing, cough, sore throat and muscle pain — **do not inhale it**. [Beilstein 5 H 347, 5 I 172, 5 II 268, 5 III 767, 5 IV 873.]

2,4,6-Trinitro-*m*-xylene [632-92-8] $C_8H_7N_3O_6$, M 241.2, m 182.2°. Crystallise the xylene from ethyl methyl ketone. [Beilstein 5 H 381, 5 I 185, 5 II 295, 5 III 845, 5 IV 950.]

Triphenylamine [603-34-9] $C_{18}H_{15}N$, M 245.3, m 123-126°, 125-129°, 125-127°, 127°, 127.3-127.9°, b 347-348°/atm, 365°/atm, d^{25} 0.77g/ml, pK^{25} -5.0 (in fluorosulfuric acid). Crystallise the amine from EtOH or from *benzene/absolute EtOH, diethyl ether and petroleum ether. It is sublimed under vacuum and carefully dried in a vacuum line. Store it in the dark under nitrogen. [Beilstein 12 IV 276.]

1,3,5-Triphenylbenzene [612-71-5] $C_{24}H_{18}$, M 306.4, m 171-172°, 173-175°, 176°, 460°/atm, d_4^{20} 1.205. Purify it by chromatography on alumina using *benzene or petroleum ether as eluents. Crystallise the triphenylbenzene from EtOH (m 174°). [Beilstein 5 H 737, 5 I 370, 5 II 670, 5 III 2563, 5 IV 2732.]

Triphenylene (9,10-benzophenanthrene) [217-59-4] $C_{18}H_{12}$, M 228.3, m 195-198°, 198°, b 425°/atm, 438°/atm. Purify triphenylene by zone refining or crystallisation from EtOH or $CHCl_3$ and sublimation. Its fluorescence has λ_{ex} at 265nm and λ_{em} at 372nm in cyclohexane. [Beilstein 5 IV 2556.]

***RS*-(±)-1,1,2-Triphenyl-1,2-ethanediol** [6296-95-3], ***R*-(+)-1,1,2-triphenyl-1,2-ethanediol** [9506-46-4], and ***S*-(-)-1,1,2-triphenyl-1,2-ethanediol** [108998-83-0] $C_{20}H_{18}O_2$, M 290.4, m 167-167.5°, 167-168° for *RS*, m 126°, 128-129° for *R*-enantiomer, $[\alpha]_D^{25}$ +213.8 to 214.6 (c 1, EtOH), $[\alpha]_D^{25}$ +220 (c 1, 95% EtOH), m 128-129° for *S*-enantiomer and $[\alpha]_D^{25}$ -214.6 (EtOH), -217(c 1, EtOH). The racemate is prepared by condensation of benzophenone, thus benzaldehyde (10.6g, 0.1 mole) in dry Et_2O (50ml) is added to a stirred solution of disodio-benzophenone (0.1 mole prepared from benzophenone and Na in liquid NH_3) in liquid NH_3 (250ml), whereby the black solution turned to light blue. The NH_3 is evaporated off and replaced by Et_2O , and the susp-

ension is then poured into Et₂O (100ml) containing glacial acetic acid (30ml) to which is added H₂O and the two layers are separated. The organic layer is washed with saturated NaHCO₃ and dried (Drierite), then evaporated to give a white solid which is dissolved in hot *C₆H₆ (25ml) and diluted with 4 volumes of petroleum ether (b 30-60°) whereby white crystals deposited (26.5g, 91%) with **m 167-168°**. [Hamrick & Hauser *J Am Chem Soc* **81** 493 1959, DOI: 10.1021/ja01511a058] The **RS-diol**, prepared by a crossed pinacol reaction, has **m 167-168°** (from hexane/EtOAc 6:1) and IR (CH₂Cl₂) with ν_{\max} at 3550, 3440 (OH), 3050, 3015, 1590 and 1480 (HC=C) cm⁻¹, and ¹H NMR with δ at 1.45 (1H, d, J = 3.2Hz, CHOH), 3.14 (1H, s, COH), 5.60 (1H, d, J = 3.2Hz, CHO), 7.02-7.17, 7.24-7.32, 7.36-7.41 (15H 4m, ArH); ¹³C NMR with δ at 77.95 (CHO), 80.75 (CO), 126.15, 126.7, 127.0, 127.35, 127.45, 127.6, 127.7, 128.05, 128.45, 138.75, 143.35 and 145.05 (ArC); and MS with m/z 183 (M⁺ -107, 15%), 105 (61%), 78 (28%) and 77 (100%). [Guijarro et al. *Tetrahedron* **49** 1327 1993, DOI: 10.1016/S0040-4020(01)85822-1]

The optical isomers, e.g. **R-(+)-1,1,2-triphenyl-1,2-ethanediol**, are prepared from their respective chiral methyl mandelates {see [*R*-(-)-methyl mandelate [20698-91-3] above} where the configuration at the chiral center is retained. Thus *R*-(-)-methyl mandelate (15.0g 500mmol) in dry THF (100ml) is added, slowly during 20-30 minutes, to a stirred solution under N₂ of phenylmagnesium bromide [prepared from Mg powder (12.15g, 500mmol) in THF (100ml) and bromobenzene (3ml) to start the reaction, followed by more bromobenzene (39ml) in THF (200ml) and boiled under reflux for 3 hours, then cooled to 5-10°], stirred for 2 hours below 10° then refluxed for 3 to 5 hours. The stirred mixture is cooled in an ice bath, treated with saturated aqueous NH₄Cl (200ml) and neutralised with 5% aqueous HCl. The organic layer is collected, the aqueous phase is extracted with CHCl₃, the combined organic phases and washed with H₂O, dried (Na₂SO₄), filtered and the filtrate is evaporated to dryness under vacuum. The residue is recrystallised from CH₂Cl₂ (50ml) to give the **R-diol** (17.0-18.6g, 65-71% yield) with **m 126°**, [α]_D²⁵ **+213.8 to +214.6 (c 1 EtOH)** and IR (KBr) with ν_{\max} at 3500—3300, 1495, 1450, 1170 1060, 1005, 770, 750, 735 and 695 cm⁻¹, and ¹H NMR (90MHz, CDCl₃, TMS) has δ at 2.42 (1H, d, J = 4Hz, PhCHOH), 3.11 (1H, s, J = 4Hz, Ph₂COH), 5.64 (1H, d, J = 4Hz, PhCHOH), 6.97—7.48 (13H, m, Ar-H), 7.60-7.77 (2H, m, Ar-H); and MS (70eV) has m/z (%): 272 (1), 256 (3), 243 (6), 183 (100), 165 (10), 105 (70), 77 (36). [For the synthesis of *R*(+)- and *S*(-)- diols and acetates see Devant et al. *Chem Ber* **121** 397 1988, DOI: 10.1002/cber.19881210303; and Braun et al. *Org Synth* **72** 32 1995, DOI: 10.15227/orgsyn.072.0032; Braun e-EUROS 15 April 2001, DOI: 10.1002/047084289X.rt363; *Beilstein* **6** I 510.] The optically active diols form 2:1 inclusion compounds with racemic compounds to form diastereoisomeric complexes which have very different solubilities and therefore effect clean optical resolutions, see for example *Lansoprazole* obtained in 99.9%ee see in 'Physiologically Active Compounds', Chapter 6.

For the 2-acetates and their stereospecific aldol condensations *via* the doubly deprotonated acetate methyl group see **2-Hydroxy-1,2,2-triphenylacetate** above.

1,2,3-Triphenylguanidine [101-01-9] C₁₉H₁₇N₃, **M 287.3**, **m 144°**, **144-147°**, **d²⁵ 1.16g/ml**, **pK²⁵ 9.10**. Crystallise the guanidine from EtOH or EtOH/water. Dry it *in vacuo*. It is a fairly strong base, store in the absence of CO₂. [*Beilstein* **12** H 451, **12** I 261, **12** II 246, **12** III 907, **12** IV 866.]

Triphenylmethane [519-73-3] C₁₉H₁₆, **M 244.3**, **m 90-92°**, **92-93°**, **93°**, **94°**, **358-359°/atm**, **d²⁵ 1.014g/ml**. Crystallise triphenylmethane from EtOH or *benzene (with one molecule of *benzene of crystallisation which is lost on exposure to air or by heating on a water bath). It can also be sublimed under vacuum. It has been given a preliminary purification by refluxing with tin and glacial acetic acid, then filtered hot through a glass sinter disc, and precipitated by addition of cold water. [*Beilstein* **5** H 698, **5** IV 2495.]

Triphenylmethanol (triphenylcarbinol) [76-84-6] C₁₉H₁₆O, **M 260.3**, **m 161-164°**, **162°**, **163°**, **164°**, **164.2°**, **b 360-380°/atm (without dec)**, **d²⁵ 1.189g/ml**, **pK²⁵ -6.63 (aqueous H₂SO₄)**. Crystallise the carbinol from EtOH, MeOH, CCl₄ (4ml/g), *benzene, hexane or petroleum ether (b 60-70°). Dry it at 90°. [Ohwada et al. *J Am Chem Soc* **108** 3029 1986, DOI: 10.1021/ja00271a036; *Beilstein* **6** IV 5014.]

Triphenylmethyl chloride (trityl chloride) [76-83-5] C₁₉H₁₅Cl, **M 278.8**, **m 109-112°**, **110-112°**, **111-112°**, **b 230-235°/20mm**, **d²⁵ 1.141g/ml**. Crystallise trityl chloride from iso-octane. Also crystallise it from 5 parts of petroleum ether (b 90-100°) and 1 part of acetyl chloride using 1.8g of solvent per g of chloride. Dry it in a desiccator over soda lime and paraffin wax, and store it under N₂ in aliquots. Its solubility in CHCl₃ is 0.1g/ml at ~20°. [Bachman *Org Synth Coll Vol* **3** 841 1955, DOI: 10.15227/orgsyn.023.0100; Thomas & Rochow *J Am*

Chem Soc **79** 1843 1957, DOI: 10.1021/ja01565a021; Meisel et al. *J Am Chem Soc* **108** 4706 1986, DOI: 10.1021/ja00276a004; *Beilstein* **5** H 750, **5** I 346, **5** II 615, **5** III 2315, **5** IV 2497.] It is moisture sensitive, store in a well sealed container. It is used to introduce a trityl protecting group. **LACHRYMATORY**.

***dl*-Tropic [(3-hydroxy-2-phenylpropionic) acid, Tropaic acid]** [529-64-6, 552-63-6, 28845-94-5] $C_9H_{10}O_3$, **M 166.2, m 116-117°, 118°, pK^{25} 4.12**. Tropic acid crystallises from H_2O or $*C_6H_6$ in needles or plates. Its solubility in MeOH is 0.1g/ml at $\sim 20^\circ$. It is the (\pm)-acid component (*dl*-tropyl ester) of atropine [51-55-8]. It is best prepared from acetophenone cyanohydrin via atrolactic acid [515-30-0] **m 94.5°** (anhydr), atropic acid [492-38-6] **m 106-107°, chlorohydratropic acid** followed by hydrolysis of the latter in boiling aqueous 1.73N Na_2CO_3 in high yielding steps. No racemisation occurred when chiral intermediates were used (see next entry). [McKenzie & Wood *J Chem Soc* **115** 828 1919, DOI: 10.1039/CT9191500828; McKenzie & Strathern *J Chem Soc* **127** 82 1925, DOI: 10.1039/CT9252700082.] [*Beilstein* **10** IV 664.]

***S*(-)-Tropic acid [(-)3-hydroxy-2-phenylpropionic acid]** [16202-15-6] has **m 128-129°, $[\alpha]_D^{13} -72.5$ (c 2.6, EtOH), $[\alpha]_D^{15} -79.0$ (c 1.5, H_2O), $[\alpha]_D^{15} -83.3$ (c 1.8, Me_2CO)**. It is the (-)-acid component of *l*-hyoscyamine (i.e *l*-tropyl atropine, [101-31-5]). It has been prepared by optical resolution of (\pm)-tropic acid using the quinine diastereomeric salt for obtaining ***R*(+)-tropic acid** [529-64-6], and then with the morphine diastereomeric salt for obtaining the **natural (-)-tropic acid**. The salts were hydrolysed with dilute H_2SO_4 , extracted thoroughly into Et_2O , dried (Na_2SO_4), evaporated, and the residue was recrystallised from $*C_6H_6$ (in which it is sparingly soluble) to give lustrous needles **m 128-129°**. It also crystallises from H_2O in glassy needles and plates grouped in rosettes. The enantiomer, obtained in the same way, has identical properties except for opposite optical rotations. [McKenzie & Wood *J Chem Soc* **115** 828 1919, DOI: 10.1039/CT9191500828; McKenzie & Strathern *J Chem Soc* **127** 86 1925, DOI: 10.1039/CT9252700082]. The absolute configuration of the (-)-tropic acid was deduced as *S*- by conversion into (+)-alanine. [Fedor & Csepregy *J Chem Soc* 3222 1961, DOI: 10.1039/JR9610003222; for enzymic kinetic resolution see Klomp et al. *Tetrahedron Asymmetry* **16** 3892 2005, DOI: 10.1016/j.tetasy.2005.10.032.]

Tropolone (2-hydroxycyclohepta-2,4,6-trien-1-one) [533-75-5] $C_7H_6O_2$, **M 122.1, m 49-50°, 50-52°, b 81-84°/0.1mm, $pK_1^{25} -0.53$ (protonation of CO, aqueous H_2SO_4), $pK_2^{25} 6.67$ (acidic OH)**. Crystallise tropolone from hexane or petroleum ether and sublime it at $40^\circ/4mm$. Also distil it at high vacuum. Store it at $2-8^\circ$. [Pauson *Chem Rev* **55** 9 1955, DOI: 10.1021/cr50001a002; *Beilstein* **8** IV 159.] It complexes with Cu^{2+} , Ga^{3+} , In^{3+} among other metals.

Tropone (2,4,6-cycloheptatrien-1-one) [539-80-0] C_7H_6O , **M 106.1, m -7°, b 92-95°/0.25mm, 91-92°/4mm, 113°/15mm, $d^{25}_D 1.094$, $n^{20}_D 1.615$** . Among the various syntheses, tropone was prepared by the oxidation of cycloheptatriene with SeO , and purified by bulb-to-bulb distillation ($20-110^\circ/0.05mm$) with Dry-Ice cooling, then fractionally redistilled at $0.25mm$ as a pale yellow oil. Its IR ($CHCl_3$) has ν_{max} at 3000, 1630, 1580, 1515, 1470, 1210 cm^{-1} , and 1H NMR ($CDCl_3$, 300MHz) with δ at 6.88-7.11 (several multiplets, 6H). [Dahnke & Paquette *Org Synth Coll Vol* **9** 396 1998, DOI: 10.15227/orgsyn.071.0181; *Beilstein* **7** IV 501.]

Tyramine (4-hydroxyphenethylamine) [51-67-2] $C_8H_{11}NO$, **M 137.2, m 160-162°, 164-165°, b 166°/2mm 175-181°/8mm, 205-207°/25mm $pK_1^{25} 9.74$ (OH), $pK_2^{25} 10.52$ (NH_2)**. Crystallise tyramine from $*benzene$ or EtOH; and/or distil it in a vacuum. Its solubility ($\sim 15^\circ$) in H_2O is 1.1w/v%, and in EtOH it is 10/v%. It is a strong base and should be stored under N_2 away from CO_2 , or better to keep it as the hydrochloride salt (see below). [*Beilstein* **13** IV 1788.] It can enter the catecholaminergic terminals and be released as a false transmitter [Bunzow et al. *Mol Pharmacol* **60** 1181 2001, DOI: 10.1124/mol.60.6.118] ***Tyramine hydrochloride*** [60-19-5] $C_8H_{11}NO.HCl$, **M 173.6, has m 269°, 271-274°, 274°, 274-276°**. Crystallise the hydrochloride from EtOH by addition of diethyl ether, or from concentrated HCl. An aqueous solution has a pH of ~ 7 . [*Beilstein* **3** II 355.] See previous entry for pharmacological activity.

Vanillin (4-hydroxy-3-methoxybenzaldehyde) [121-33-5] $C_8H_8O_3$, **M 152.2, m 81-83°, 81-84°, 81.5°, 82°, 83°, b 170°/15mm, 285°/atm, $pK^{25} 7.40$** . Crystallise vanillin from water or aqueous EtOH, or by distillation *in vacuo*. [*Beilstein* **8** IV 1763.] It is a food flavouring substance.

Veratraldehyde (vanillin methyl ether) [120-14-9] $C_9H_{10}O_3$, **M 166.2**, **m 40-43°, 42-43°, b 281°/atm**, Crystallise the ether from diethyl ether, petroleum ether, CCl_4 or toluene. [Beilstein 8 IV 1765.] It is a flavor and fragrance agent.

Variamine Blue RT [4-aminodiphenylamine diazonium HSO_4 , 4-(phenylamino)benzenediazonium sulfate (1:1)] [4477-28-5] $C_{12}H_{10}N_3.HSO_4$, **M 293.3**, **m (of the chloride) 120°(dec), 168-170° (dec), CI 37240, λ_{max} 377 nm**. Dissolve 10g of the dye in 100ml of hot water. Sodium dithionite (0.4g) is then added, followed by active carbon (1.5g) and is filtered hot. To the colourless or slightly yellow filtrate a solution of saturated NaCl is added, and the mixture is cooled. The needles are filtered off, washed with cold water, dried at room temperature, and stored in a dark bottle (light sensitive). [Anderson & Steedly *J Am Chem Soc* **76** 5144 1954, DOI: 10.1021/ja01649a043.] This diazonium salt is used in the spectrophotometric determination of uric acid in urine (or serum) based on an enzymatic method with Uricase which liberates H_2O_2 that reacts with the diazonium salt and changes its colour from yellow to pale yellow-green. The assay is performed in a pH 9.0 buffer, and the change in absorption at 269nm is measured, with a limit of sensitivity of 0.5–13mM. [Hamzah et al. *J Anal Bioanal Tech* **S7:011** 2155 2013, DOI: 10.4172/2155-9872.S7-011.] [For *Variamine blue as a gravimetric reagent: Determination of tungsten* see Erdey et al. *Talanta* **14** 515 1967, DOI: 10.1016/0039-9140(67)80078-X; Erdey *Chemist-Analyst* **48** 106 1959, *Beilstein* **16** H 602, **16** I 371, **16** II 307, **16** III 575.]

9-Vinylnanthracene [2444-68-0] C_9H_{12} , **M 204.3**, **m 61-64°, 62-65°, 64°, 65-67°, b 61-66°/10mm**. Purify it by vacuum sublimation. It has also been purified by chromatography on silica gel with cyclohexane as eluent, and recrystallised from EtOH [Werst et al. *J Am Chem Soc* **109** 32 1987, DOI: 10.1021/ja00235a005]. [*Beilstein* **5** IV 2415.]

4-Vinylbenzyl chloride (VBC, 4-chloromethylstyrene) [1592-20-7] C_9H_9Cl , **M 152.6**, **b 58-62°/0.4mm, 75.5-79°/2mm, 229°/760mm, d_{25}^{25} 1.083, n_D^{20} 1.572**. Purify 4-vinylbenzyl chloride by dissolving it in Et_2O , washing it with 0.5% of aqueous NaOH, separating, drying the organic layer (Na_2SO_4), evaporating and distilling the residual oil under N_2 *in vacuo*. Add 0.05% of 4-*tert*-butylcatechol as stabiliser. Store it at 4°. It is **lachrymatory**. [Nishikubo et al. *Tetrahedron Lett* **22** 3873 1981, DOI: 10.1016/S0040-4039(01)91332-2; Tanimoto et al. *Synth Commun* **4** 193 1974, DOI:10.1080/00397917408062071; *Beilstein* **6** IV 3818.]

1-Vinylnaphthalene [826-74-4, 92529-36-7] $C_{14}H_{10}$, **M 154.2**, **b 68-69°/0.4mm, 100-101°/3.7mm, 124-125°/15mm, 135-138°/atm, d^{25} 1.040g/ml, n_D^{20} 1.653**. Commercial material can contain up to 2% of poly-1-vinylnaphthalene. Fractionally distil it under reduced pressure using a spinning-band column, dry it with CaH_2 and again distil it under vacuum. Store it in sealed ampoules in a freezer. It has λ_{max} (cyclohexane) at 210nm and 320nm. The **picrate** has **m 105-106°**. [*Beilstein* **5** H 585, **5** III 1773, **5** IV 1833.] It reacts nicely with quinones [Corbett et al. *Aust J Chem* **18** 1775 1965, DOI: 10.1071/CH9651775]. It is an **irritant**.

2-Vinylnaphthalene [827-54-3] has **m 60-62°, 62-65°, 62-67°, 65-68°, 65°, 66°, b 76-81°/2.5mm, 128-131°/13mm, 135-137°/18mm**. Commercial material can contain up to 5% of MeOH as stabiliser. The flammable solid can be fractionally distilled under reduced pressure using a spinning-band column; dry it with CaH_2 and again distil it under vacuum. It crystallises from aqueous MeOH, EtOH (**m 66°**) or petroleum ether (**m 66-66.5°**). Store it in sealed ampoules in a freezer. The **picrate** has **m 91-92°** (from EtOH). [*Beilstein* **5** III 1775, **5** IV 1833.] It is an **irritant**.

Violanthrene (dibenzanthrene, 5,10-dihydroviolanthrene A) [81-31-2] $C_{34}H_{20}$, **M 428.5**. Purify violanthrene by vacuum sublimation over Cu in a muffle furnace at 450°/25mm in a CO_2 atmosphere [Scholl & Meyer *Chem Ber* **67** 1229 1934, DOI: 10.1002/cber.19340670722]. [*Beilstein* **5** I 392.] **Violanthrene A (anthro[9,1,2-cde]benzo[rsf]pentaphene)** [188-87-4] **M 426.5** has **m 506°**. [Clar *Chem Ber* **76** 458 1943, DOI: 10.1002/cber.19430760504; *Beilstein* **5** III 2778; for *Violanthrene B (violanthrene)* see *Beilstein* **7** I 466, **7** II 818, **7** III 4539.]

Xylene (dimethylbenzene) [1330-20-7] C_8H_{10} , **M 106.1 (mixed isomers)**. Usual impurities are ethylbenzene, paraffins, traces of sulfur compounds and water. With a very efficient still, *o*-xylene can be fractionally distilled from a mixture of isomers. Purify (and dry) by fractional distillation from $LiAlH_4$, P_2O_5 ,

CaH₂ or sodium. This treatment can be preceded by shaking successively with concentrated H₂SO₄, water, aqueous 10% NaOH, water and mercury, and drying with CaCl₂ for several days. [Beilstein 5 H 360.]

***o*-Xylene** [95-47-6] has **m -25.2°, -24°, b 84°/14mm, 144.4°/760mm, d₄²⁰ 0.88020, d₄²⁵ 0.87596, n_D²⁰ 1.50543, n_D²⁵ 1.50292**. The general purification methods listed under xylene are applicable [Clarke & Taylor *J Am Chem Soc* 45 830 1923, DOI: 10.1021/ja01656a043]. *o*-Xylene (4.4Kg) is sulfonated by stirring for 4 hours with 2.5L of conc H₂SO₄ at 95°. After cooling, and separating the unsulfonated material, the product is diluted with 3L of water and neutralised with 40% NaOH. On cooling, sodium *o*-xylene sulfonate separates and is recrystallised from half its weight of water. [A further crop of crystals is obtained by concentrating the mother liquor to one-third of its volume.] The salt is dissolved in the minimum amount of cold water, then mixed with the same amount of cold water, and with the same volume of concentrated H₂SO₄ and heated to 110°. *o*-Xylene is regenerated and steam distils. The distillate is saturated with NaCl, the organic layer is separated, dried and redistilled. [Beilstein 5 H 362, 5 I 179, 5 II 281, 5 III 807, 5 IV 917.]

***m*-Xylene** [108-38-3, 1330-20-7] has **m -48°, -47.9°, b 139.1°/760mm, d₄²⁰ 0.86417, d₄²⁵ 0.85990, n_D²⁰ 1.49721, n_D²⁵ 1.49464**. The general purification methods listed under *xylene* are applicable. The *o*- and *p*-isomers can be removed by their selective oxidation when a *m*-xylene sample containing them is boiled with dilute HNO₃ (one part concentrated acid to three parts water). After washing with water and alkali, the product can be steam distilled, collected as for *o*-xylene, then distilled and purified further by sulfonation. [Clarke & Taylor *J Am Chem Soc* 45 831 1923, DOI: 10.1021/ja01656a043.] *m*-Xylene is selectively sulfonated when a mixture of xylenes is refluxed with the theoretical amount of 50-70% H₂SO₄ at 85-95° under reduced pressure. By using a still resembling a Dean and Stark apparatus, water in the condensate can be progressively withdrawn while the xylene is returned to the reaction vessel. After cooling, then adding water, unreacted xylenes are distilled off under reduced pressure. The *m*-xylene sulfonic acid is subsequently hydrolysed by steam distillation up to 140°, the free *m*-xylene is washed, dried with silica gel and again distilled. It is stored over molecular sieves Linde type 4A. [Beilstein 5 H 370, 5 I 182, 5 II 287, 5 III 823, 5 IV 932.]

***p*-Xylene** [106-42-3] has **m 13.3, b 138.3°/760mm, d₄²⁰ 0.86105, d₄²⁵ 0.85669, n_D²⁰ 1.49581, n_D²⁵ 1.49325**. The general purification methods listed for *xylene* above are applicable. *p*-Xylene can readily be separated from its isomers by crystallisation from such solvents as MeOH, EtOH, isopropanol, acetone, butanone, toluene, pentane or pentene at low temperatures. It can be further purified by fractional crystallisation by partial freezing, and stored over sodium wire or molecular sieves Linde type 4A. [Stokes & French *JCS Faraday Trans 1* 76 537 1980, DOI: 10.1039/F19807600537; Beilstein 5 H 382, 5 I 185, 5 II 296, 5 III 845, 5 IV 951.]

HETEROCYCLIC COMPOUNDS

Acetaldehyde ammonia trimer (hexahydro-2,4,6-trimethyl-1,3,5-triazine trihydrate) [58052-80-5 ($3\text{H}_2\text{O}$), 76231-37-3, 638-14-2, 108-74-7 ($2\text{H}_2\text{O}$)] $(\text{C}_2\text{H}_7\text{NO})_3$, $\text{C}_6\text{H}_{15}\text{N}_3 \cdot 3\text{H}_2\text{O}$, **M 183.3**, **m 94-96°, 94-95°, 95-97°, 97°, b 110°/atm (partly dec)**. It crystallises from EtOH/Et₂O. When prepared (MeCHO + 15M aqueous NH_3 at $\sim 0^\circ$, then -25°), it separates as the **trihydrate** which can be dried in a vacuum over CaCl_2 at room temperature to give finally the **anhydrous** compound with the same melting point. Store away from light $<8^\circ$. The intermediate **dihydrate** melts at $25-28^\circ$, then resolidifies and melts again at $94-95^\circ$. On gentle warming (40° to 80°) it loses NH_3 to give $\text{MeCH}(\text{N}=\text{CHMe})_2$. It is used as a source of acetaldehyde in syntheses. The **monomer** [75-39-8] $\text{C}_2\text{H}_7\text{NO}$, **M 61.2**, is the intermediate in its preparation. Do not breathe in vapours — *IT IRRITATES THE EYES AND MUCOUS MEMBRANES*. [Prepn & NMR: Nielson et al. *J Org Chem* **38** 3288 1973, DOI: 10.1021/jo00959a010; *Beilstein* **26** III/IV 25.]

2-Acetamido-5-nitrothiazole (Acinitrazole) [140-40-9] $\text{C}_5\text{H}_5\text{N}_3\text{O}_3\text{S}$, **M 187.2**, **m 262-265°, 263°, 264-265°**. Recrystallise acinitrazole from hot absolute EtOH (charcoal) then cool to -10° , or AcOH. [Hurd & Wehrmeister *J Am Chem Soc* **71** 4007 1949, DOI: 10.1021/ja01180a038; *Beilstein* **27** III/IV 4676.] Used as a prophylactic in animal feeds. An analytical procedure was devised for estimating 0.03 to 0.07% in medicated feeds using λ_{max} at 340nm ($A_{1\text{cm}}^{1\%}$ 650) for Acinitrazole [Brown *Analyst* **91** 672 1966, DOI: 10.1039/AN9669100672].

4-Acetamido-2,2,6,6-tetramethylpiperidine-1-oxyl (acetamidoTEMPO) [14691-89-5] $\text{C}_{11}\text{H}_{21}\text{N}_2\text{O}_2$, **M 213.3**, **m 144-146°, 146-147°**. Dissolve the 1-oxyl in CH_2Cl_2 , wash it with saturated K_2CO_3 , then saturated aqueous NaCl , dry (Na_2SO_4), filter and evaporate. The red solid is recrystallised from aqueous MeOH, **m 147.5°**. [Ma & Bobbitt *J Org Chem* **56** 6110 1991, DOI: 10.1021/jo00021a027; Rozantsev & Kokhanov *Bull Acad Sci USSR, Div Chem Sci* **15** 1422 1966, DOI: 10.1007/BF01151392; *Beilstein* **22/8** V 174.]

5-Acetamido-1,3,4-thiadiazole-2-sulfonamide (Acetazolamide) [59-66-5] $\text{C}_4\text{H}_6\text{N}_4\text{O}_3\text{S}_2$, **M 222.3**, **m 256-259° (dec), 258-259° (dec)**. It is recrystallised from water. It is soluble in M NH_4OH (50mg/ml), Me_2SO , MeOH but slightly in EtOH. It is a carbonic anhydrase inhibitor. [Roblin & Clapp *J Am Chem Soc* **72** 4890 1950, DOI: 10.1021/ja01167a011; *Beilstein* **27** III/IV 8219.]

Acetoacetylpipecridide [1128-87-6] $\text{C}_9\text{H}_{15}\text{NO}_2$, **M 169.2**, **b 88.9°/0.1mm, 88-91°/0.1mm, n_D^{25} 1.4983**. Dissolve it in *benzene, extract with 0.5M HCl to remove basic impurities, wash with water, dry, and distil it at 0.1mm [Wilson *J Org Chem* **28** 314 1963, DOI: 10.1021/jo01037a009]. [*Beilstein* **20** IV 1121.]

4-Acetoxy-2-azetidinone [28562-53-0] $\text{C}_5\text{H}_7\text{NO}_3$, **M 129.1**, **m 38-40°, 38-41°**. Dissolve it in CHCl_3 , dry (MgSO_4), concentrate at $40^\circ/70\text{mm}$, or better at room temperature to avoid decomposition. Wash and stir the residual oil with hexane by decantation and discard the wash. Dry the oil at high vacuum when it should solidify, **m 34°**. It can be distilled at high vacuum, $80-82^\circ/10^{-3}\text{mm}$, but this results in extensive losses. The purity can be checked by TLC using Merck Silica Gel F_{254} and eluting with EtOAc. The azetidinone has R_F 0.38 (typical impurities have R_F 0.67). The spots can be detected by the **TDM spray**. This is prepared from (A) 2.5g of 4,4'-tetramethyldiaminodiphenylmethane (TDM) in 10ml AcOH and diluted with 50ml of H_2O , (B) 5g KI in 100ml of H_2O and (C) 0.3g ninhydrin in 10ml of AcOH and 90ml of H_2O . The spray is prepared by mixing (A) and (B) with 1.5ml of (C) and stored in a brown bottle. [Clauß et al. *Justus Liebigs Ann Chem* 539 1974, DOI: 10.1002/jlac.197419740403; Mickel et al. *Org Synth* **65** 135 1987, DOI: 10.15227/orgsyn.065.013; *Beilstein* **21/12** V 4.]

N-Acetylcaprolactam (1-acetylazepan-2-one, 1-acetylhexahydro-2H-azepin-2-one) [1888-91-1] $\text{C}_8\text{H}_{11}\text{NO}$, **M 155.2**, **b 85-86°/0.05mm, 119.5-222°/11mm, 134-135°/26mm, d_4^{25} 1.094, n_D^{20} 1.489**. Distil the lactam in a vacuum, but if the IR has broad OH bands (*ca* ν_{max} at $\sim 3300\text{cm}^{-1}$), then add *ca* one half volume or more of Ac_2O , reflux for 2 hours and distil or dissolve it in toluene add MeCOCl heat at $100^\circ/2$ hours and distil in a vacuum. Its IR has ν_{max} at 1730cm^{-1} (film). [Stoll & Griehl *Helv Chim Acta* **48** 1805 1965, DOI: 10.1002/hlca.19650480742; Bose et al. *Tetrahedron* **30** 3 1974, DOI: 10.1016/S0040-4020(01)97208-4; *Beilstein* **21** III/IV 3210, **21/6** V 460.]

3-Acetyl-2,5-dimethylthiophene [2530-10-1] $C_8H_{10}OS$, **M 154.2**, **b** 62°/0.25mm, 105-108°/15mm, 224°/atm, d_4^{25} 1.086, n_D^{20} 1.544. Distil the thiophene in a vacuum, or dissolve it in $*C_6H_6$, shake this with 6M HCl, then wash with H_2O , dry the organic layer, evaporate, and distil the residue preferably in a vacuum. [Glaze et al. *JCS Perkin Trans 1* 957 1985, DOI: 10.1039/P19850000957; *Beilstein* 17 H 298.]

N-Acetylimidazole [2466-76-4] $C_5H_6N_2O$, **M 110.1**, **m** 99-105°, 101.5-102.5°, **pK²⁵ 3.6**. It is recrystallised from isopropenyl acetate and dried in a vacuum over P_2O_5 . [Riordan & Vallee *Methods Enzymol* 25 500 1972, DOI: 10.1016/S0076-6879(72)25045-5, *Beilstein* 23/4 V 218.]

1-Acetylimidole [576-15-8] $C_{10}H_9NO$, **M 159.2**, **b** 123-125°/8mm, 148-150°/10mm, 152-153°/15mm, d_4^{25} 1.387, n_D^{20} 1.6100. It has been prepared by phase transfer catalysis of a mixture of $AcCl/NaOH$ and Indole [Illi *Synthesis* 387 1979, DOI: 10.1055/s-1979-28695]. Purify it by distillation in a vacuum, but if it is very discoloured, then distil it in steam, separate the layers, dry the organic layer (Na_2SO_4), filter it, and distil it in a vacuum. The *picrate* has **m 98-99°**. **1,3-Diacetylimidole** is formed when indole is treated with Ac_2O at 180-200°, and distils with 1-acetylimidole. The latter distils in steam, whereas the diacetylimidole remains behind. It is hydrolysed by boiling alkali to yield indole, and can be acetylated further to **1,3-diacetylimidole m 151°**. [For NMR see Elguero et al. *Org Magn Res* 7 445 1975, DOI: 10.1002/mrc.1270070910; Ciamician & Zatti *Chem Ber* 22 1976 1889, DOI: 10.1002/cber.1889022025; Zatti & Ferratini *Chem Ber* 23 1359 1890, DOI: 10.1002/cber.189002301214; Pleninger & Werst *Chem Ber* 89 2783, 2788 1956, DOI: 10.1002/cber.19560891220; *Beilstein* 20 H 309, 20 II 200, 20 III/IV 3182, 20/7 V 19.]

3-Acetylimidole [703-80-0] has **m 189°, 188-190°, 188-192°, 191-193°, 194°, pK²⁵ 12.99 (acidic NH)**. It is formed by reaction of indole- $MgCl$ and $AcCl$, and also by hydrolysis of 1,3-diacetylimidole (see above). It is purified by recrystallisation from $MeOH$, $*C_6H_6$ containing a little $EtOH$ or $*C_6H_6$; and by sublimation *in vacuo*. The *phenylureido* derivative has **m 154°**. The (*E*)-*oxime* has **m 144-145°** (also 149° reported) (from H_2O) [Kostyuchenko et al. *J Org Chem USSR (Engl Trans)* 8 2471 1972], and the (*Z*)-*oxime* has **m 95°** [Kostyuchenko et al. *J Org Chem USSR (Engl Trans)* 8 2469, 2473 1972]. The (*E*)-*O-methyloxime* crystallises from $EtOH$ with **127-129°**. [Oddo & Sessa *Gazz Chim Ital* 41 234 1911, Baker *J Chem Soc* 461 1946, DOI: 10.1039/JR9460000461; *Beilstein* 21 H 316, 21 I 300, 21 II 264, 21 III/IV 3775, 21/8 V 297.]

4-Acetylmorpholine [1696-20-4] $C_6H_{11}NO_2$, **M 129.2**, **m** 13.8-14°, 14°, 14.5°, **b** 96-97°/6mm, 113-128°/22mm, 242-247°/760mm, d_4^{20} 1.0963, n_D^{20} 1.4830. Distil it through an 8inch Fenske (glass helices packing) column with a manual take-off head. Purify it by fractional distillation. The *hydrobromide* has **m 172-175°**. [Brace *J Am Chem Soc* 75 357 1953, DOI: 10.1021/ja01098a030; deBenneville et al. *J Org Chem* 21 1072 1956, DOI: 10.1021/jo01116a006; *Beilstein* 27 III/IV 274.]

1-Acetylpiperazine [13889-98-0] $C_6H_{12}N_2O$, **M 128.2**, **m** 31-34°, 32-34°, 52°, **pK²⁵ 7.94**. Purify 1-acetylpiperazine by recrystallisation from 40% aqueous $EtOH$ or from $EtOH/Et_2O$. It is an **irritant**, and is **hygroscopic**. The *hydrochloride* has **m 191°** (from $EtOH$), and the *tosylate salt* has **m 148-149°** (from $EtOH/EtOAc$, 1:16). The free base, however, cannot be isolated by basifying the tosylate salt and extraction with CH_2Cl_2 . [Jacobi *Chem Ber* 66 113 1933, DOI: 10.1002/cber.19330660206; Mosher et al. *J Am Chem Soc* 75 4949 1953, DOI: 10.1021/ja01116a020; Hall *J Am Chem Soc* 78 2570 1956, DOI: 10.1021/ja01592a066; *Beilstein* 23 IV 201.]

1-Acetyl-4-piperidone [32161-06-1] $C_6H_{12}NO_2$, **M 141.2**, **b** 124-128°/0.2mm, 218°/760mm, d_4^{25} 1.1444, n_D^{20} 1.5023. Purify it by fractional distillation through a short Vigreux column (15mm). The **2,4-dinitrophenylhydrazone** has **m 212-213°** (from $EtOH$). It is freely soluble in H_2O but insoluble in Et_2O . [McElvain & McMahon *J Am Chem Soc* 71 901 1949, DOI: 10.1021/ja01171a038; *Beilstein* 21/6 V 426.]

3-Acetylpyridine [350-03-8] C_7H_7NO , **M 121.1**, **m** 13-14°, **b** 65-66°/1mm, 92-95°/8-9mm, 105°(113°)/16mm, 219-221°/760mm, d_4^{20} 1.1065, n_D^{20} 1.1065, **pK²⁵ 3.18**. It is purified by dissolving in HCl , extracting with Et_2O to remove the possible impurity of nicotinic acid, basified with $NaOH$ and extracted with Et_2O . The dried extract is filtered, evaporated and the residual oil is distilled. If the NMR spectrum indicates further impurities, then convert it to the *phenylhydrazone* (**m 137°**, yellow needles from $EtOH$). This is hydrolysed with HCl [Engler & Kiby *Chem Ber* 22 597 1889, DOI: 10.1002/cber.188902201132], the phenylhydrazine HCl is removed by filtration, $NaNO_2$ is added, the solution is basified with aqueous $NaOH$ and

extracted with Et₂O as before and distilled at atmospheric pressure to give 3-acetylpyridine as a colourless oil. Purification can also be achieved by shaking with 50% aqueous KOH, extracting with Et₂O, drying the extract and distilling it at atmospheric pressure or *in vacuo*. [Kloetzel & Chubb *J Am Chem Soc* **79** 4226 1957, DOI: 10.1021/ja01572a064.] The **hydrochloride** has **m 180-181°** (from MeOH/EtOH), the **picrate** has **m 133.8-134.8°** (from H₂O), and the **phenylhydrazone** has **m 137° (129-130°)** (needles, from EtOH) [Webb & Webb *J Am Chem Soc* **71** 2285 1949, DOI: 10.1021/ja01175a003]. The **ketoxime** has **m 112°** (from EtOH or *C₆H₆). [Strong & McElvain *J Am Chem Soc* **55** 816 1933, DOI: 10.1021/ja01329a062; Kolloff & Hunter *J Am Chem Soc* **63** 490 1941, DOI: 10.1021/ja01847a036; *Beilstein* **21/7** V 394.]

2-Acetylthiazole [24295-03-2] C₅H₅NOS, **M 127.2**, **b 89-91° (90-95°)/12mm**, **95-105°/15mm**, **d₄²⁰ 1.23**, **n_D²⁰ 1.55**. Check NMR spectrum; if it is not too bad, distil it through an efficient column in a vacuum. Otherwise purify it *via* the oxime. The **oxime** sublimes at 140-145°, **m 159°**, and when crystallised from H₂O has **m 163-165.5°**. [Erlenmeyer et al. *Helv Chim Acta* **31** 1142 1948, DOI: 10.1002/hlca.19480310421; Waisvisz et al. *J Am Chem Soc* **79** 4524 1957, DOI: 10.1021/ja01573a074; Menassé et al. *Helv Chim Acta* **40** 554 1957, DOI: 10.1002/hlca.19570400306; *Beilstein* **27** IV 2617.]

2-Acetylthiophene (methyl 2-thienyl ketone) [88-15-3] C₆H₆OS, **M 126.2**, **m 9.2-10.5°, 10.45°, 10-11°, b 77°/4mm**, **89-91°/9mm**, **94.5-96.5°/13mm**, **213-214°/atm**, **d₄²⁰ 1.17**, **n_D²⁰ 1.5666**. Fractionally distil the thiophene through a 12-plate column, and fraction **b 77°/4mm** is collected. Also wet the acetylthiophene in order to remove and free thiophene which forms an azeotrope with H₂O, **b 68°**. Store it in a brown bottle, and the clear colourless liquid remains thus for extended periods. [Kosak & Hartough *Org Synth Coll Vol* **3** 14 1955, DOI: 10.15227/orgsyn.028.0001; Hartough & Kosak *J Am Chem Soc* **69** 3093 1947, DOI: 10.1021/ja01204a049.] The red **4-nitrophenylhydrazone** crystallises from EtOH with **m 181-182°**. [*Beilstein* **17/9** V 387.]

3-Acetylthiophene (methyl 3-thienyl ketone, acetothienone) [1468-83-3] C₆H₆OS, **M 126.2**, **m 57°, 60-63°, b 106-107°/25mm**, **208-210°/748mm**. Recrystallise the thiophene from petroleum ether (b 30-60°) or EtOH. The **2,4-dinitrophenylhydrazone** crystallises from CHCl₃, **m 265°**, and the **semicarbazone** crystallises from EtOH, **m 174-175°**. [Campaigne & Le Suer *J Am Chem Soc* **70** 1555 1948, DOI: 10.1021/ja01184a078; *Beilstein* **17/9** V 399.]

Aconitine [302-27-2] C₃₄H₄₇NO₁₁, **M 645.8**, **m 198°, 204°, [α]_D²⁵ +20° (c 1, CHCl₃)**, **pK¹⁵ 8.35**. Crystallise it from EtOH, CHCl₃ or toluene. Its slightly soluble in H₂O (0.3mg/ml) and EtOH (35mg/ml). It is a very POISONOUS toxin from the *Aconitum* plant (devil's helmet or monkshood) and used against pain in Chinese herbal medicine. [*Beilstein* **21/6** V 310.]

Aconitine hydrobromide [6034-57-7] C₃₄H₄₇NO₁₁·HBr, **M 726.7**, **m 207°**. Crystallise the salt from water or EtOH/ether. [*Beilstein* **21/6** V 310.]

Acridine (2,3-benzoquinoline) [260-94-6] C₁₃H₉N, **M 179.2**, **m 107-110° (sublimes)**, **111° (sublimes)**, **b 346°/atm**, **pK²⁵ 5.58 (pK²⁵ of excited state 10.65)**. Acridine has been crystallised twice from *benzene/cyclohexane, or from aqueous EtOH (pale yellow crystals), then sublimed, removing and discarding the first 25% of the sublimate. The remainder is again crystallised and sublimed, discarding the first 10-15% [Wolf & Anderson *J Am Chem Soc* **77** 1608 1955, DOI: 10.1021/ja01611a060]. Acridine can also be purified by crystallisation from *n*-heptane and then from ethanol/water after pre-treatment with activated charcoal, or by chromatography on alumina with petroleum ether in a darkened room. Alternatively, acridine can be precipitated as the hydrochloride from *benzene solution by adding HCl, after which the base is regenerated, dried at 110°/50mm, and recrystallised to constant melting point from petroleum ether [Cumper et al. *J Chem Soc* 4518 1962, DOI: 10.1039/JR9620004518]. The **free base** may be chromatographed on basic alumina/C₆H₆, then vacuum-sublimed and zone-refined. Its UV has λ_{max}(log ε) 249(5.25), 338(3.83), 346(3.83), 354(4.02)nm at pH 8.0, and 255(5.10), 324(3.50), 338(3.99), 354(4.34), 386(3.50), 402(3.48) at pH 3.0. [Williams & Clarke, *JCS Faraday Trans 1* **73** 514 1977, DOI: 10.1039/F19777300514; Albert, *The Acridines* Arnold Press 1966.] It can exist in five crystalline forms and is steam volatile. It is a strong IRRITANT to skin and mucous membranes and can become a chronic irritant— handle it with CARE. [*Beilstein* **20/8** V 199.]

N-(9-Acridinyl)maleimide (NAM) [49759-20-8, 49761-68-4] $C_{17}H_{10}N_2O_2$, **M 274.3, m 248°, 255-258°**. Purify NAM by chromatography on silica gel using CH_2Cl_2 as eluant. Evaporation of pooled fractions that gave the correct NMR spectra gave a solid which was crystallised from Me_2CO as pale yellow prisms. Its IR(nujol) has ν_{max} at 1710 (imide); the UV (MeOH) has λ_{max} (nm) (ϵ $M^{-1}cm^{-1}$) at 251 (159 500), 343 shoulder (7 700), 360 (12 400) and 382 shoulder (47 000). It is used in fluorimetric assays and as a fluorescent tag. [Machida et al. *Chem Pharm Bull Jpn* **26** 596 1978, DOI: org/10.1248/cpb.26.596; Schuldiner et al. *Eur J Biochem* **25** 64 1972, DOI: 10.1111/j.1432-1033.1972.tb01667.x; Schäfer et al. *Analyt Biochem* **209** 53 1993, DOI: 10.1006/abio.1993.1081.]

Acridone (9,10-dihydro-9-oxoacridine, 9-hydroxyacridine) [578-95-0] $C_{13}H_9NO$, **M 195.2, m >300°, pK₁ - 0.32 (basic), pK₂ 12.80 (14.0, acidic, proton loss)**. Dissolve ~1g in ca 1% NaOH (100ml), add 3M HCl to pH 4 when acridone separates as a pale yellow solid with **m** just above **350°** (sharp). It can be recrystallised from large volumes of H_2O to give a few mg. It is soluble in 160 parts of boiling EtOH (540 parts at 22°) [Albert & Phillips *J Chem Soc* 1294 1956, DOI: 10.1039/JR9560001294]. A few decigrams are best crystallised as the **hydrochloride** from 400 parts of 10N HCl (90% recovery) from which the free base is obtained by washing the salt with H_2O . A small quantity can be recrystallised (as the neutral species) from boiling AcOH. Larger quantities are best recrystallised from a mixture of 5 parts of freshly distilled aniline and 12.5 parts of glacial acetic acid. Acridone distils unchanged at atmospheric pressure, but the boiling point was not recorded, and some sublimation occurs below 350°. Commercial acridone has been recrystallised three times from EtOH and dried at 70°/0.5mm. It has UV with λ_{max} (log ϵ) at 250(4.52), 294(3.36), 400(3.95)nm in EtOH. [see Albert, *The Acridines* Arnold Press pp 201, 372 1966; for pKa see Kalatzis *J Chem Soc (B)* 96 1969, DOI: 10.1039/J29690000096; *Beilstein* **23/9** V 7.]

Acriflavine [8048-52-0] $C_{13}H_{11}N_3$, **M 196.2, pK²⁵ >12**. Treat acriflavin twice with freshly precipitated AgOH to remove proflavine, then recrystallise it from absolute methanol [Wen & Hsu *J Phys Chem* **66** 1353 1962, DOI: 10.1021/j100813a501]. [*Beilstein* **22** III/IV 218.] (See following entry and the 3,6-diaminoacridine hydrochloride and sulfate below).

Acriflavin Mixture (Euflavin, Proflavin, 3,6-diamino-10-methylacridinium chloride + 3,6-diamino-acridine) [8048-52-0] $C_{14}H_{14}ClN_3$, $C_{13}H_{11}N_3$, **M 259.7, m 179-181°**. Purify it by dissolving in 50 parts of H_2O , shaking with a small excess of freshly precipitated and washed Ag_2O . The mixture is set aside overnight at 0° and filtered. The cake is not washed. The pH of the filtrate is adjusted to 7.0 with HCl and evaporated to dryness. The residue is then crystallised twice from MeOH, twice from H_2O and dried at 120°. Its UV has λ_{max} at 452nm and a loge value of 4.67. It is a red powder which readily absorbs H_2O . The solubility is increased in the presence of proflavin. The **dihydrochloride (Panflavin)** [8063-24-9] is a deep red crystalline powder. It is available as a mixture of 3,6-diaminoacridinium chloride (35%) and its 10-methochloride (65%). [See Albert, *The Acridines* Arnold Press p 346 1966, Benda *Chem Ber* **45** 1787 1912, DOI: 10.1002/cber.19120450251]. [*Beilstein* **22** III/IV 5488, **23** I 650.] These are antiseptics.

Adrenochrome (3-hydroxy-1-methyl-5,6-indoline-dione) [54-06-8] $C_9H_9NO_3$, **M 179.2, m 125-130° (dec)**. It was crystallised from MeOH/formic acid, as bright red crystals of the **hemihydrate (m ~115-120°)**, and stored in a vacuum desiccator. The much more stable **oxime sesquihydrate** [6055-73-8] $C_9H_{10}N_2O_4 \cdot 1.5 H_2O$, **m 278°**, crystallises in orange crystals from hot H_2O . The **mono-semicarbazone (Carbazochrome)** [69-81-8] **M 236.2**, crystallises as orange-red crystals from dilute EtOH with **m ~203°(dec)** and is haemostatic. [Heacock *Chem Rev* **59** 181 1959, DOI: 10.1021/cr50026a001; *Beilstein* **21** III/IV 6434.]

Aetioporphyryn I (Etioporphyryn, I 2,4,6,8-tetraethyl-1,3,5,7-tetramethylporphyrin) [448-71-5] **M 478.7, m 360-363°, pK²⁵ 18**. Purify it by chromatography on an Al_2O_3 column (300g/300mg of porphyrin) and elute with CH_2Cl_2 , evaporate the eluate and crystallise the residue from CH_2Cl_2 /MeOH, pyridine or $CHCl_3$ /petroleum ether (purple prisms, **m > 300°**) [Smith *JCS Perkin Trans 1* 1471 1972, DOI: 10.1039/P19720001471]. The copper salt crystallises as red needles from pyridine/AcOH. It complexes with metals. The **dihydrobromide** [69150-58-9] **M 640.5** separates from aetioporphyryn I in Et_2O on addition of 10% of aqueous HBr after 2 days [Fischer & Treibs *Justus Liebigs Ann Chem* **457** 209 1927, DOI: 10.1002/jlac.19274570108; Treibs & Dieter *Justus Liebigs Ann Chem* **513** 65 1934, DOI: 10.1002/jlac.19345130105]. [*Beilstein* **26** III/IV 1915.]

Agroclavin [548-42-5] $C_{16}H_{18}N_2$, M 238.3, m 198-203°(dec), 205-206°, 210-212°, $[\alpha]_D^{20}$ -155 (c 1, $CHCl_3$), pK_{Est} ~8.0. This ergot alkaloid crystallises from diethyl ether (colourless rods) or Me_2CO (needles). The **hydrochloride** crystallises from H_2O and has m 265-266°, $[\alpha]_D^{20}$ -110 ($CHCl_3$). [Plieninger et al. *Justus Liebigs Ann Chem* **743** 95 1971, DOI: 10.1002/jlac.19717430111; *Beilstein* **23** III/IV 1623.]

RS-Allantoin (5-ureidoimidazol-2,4-dione) [97-59-6] $C_4H_6N_4O_3$, M 158.1, m 230°(dec), 238°(dec). This purine metabolite in the uric acid pathway crystallises from water or EtOH [Hartman et al. *Org Synth Coll Vol II* 21 1943, DOI: 10.15227/orgsyn.013.0001]. It reacts with free radicals and is a useful biomarker for oxidative stress. [*Beilstein* **25** III/IV 4071.]

Alloxan [2,4,5,6(1H,3H)pyrimidine tetrone, 5,6-dihydroxyuracil] [50-71-5] $C_4H_2N_2O_4$, M 142.0, m ~170°(dec), pK^{25} 6.64. Crystallisation from water gives the **tetrahydrate**. **Anhydrous** crystals are obtained by crystallisation from acetone, glacial acetic acid or by sublimation *in vacuo*. [See below and *Beilstein* **24** H 500, **24** I 428, **24** II 301, **24** III/IV 2137.]

Alloxan monohydrate [2244-11-3] $C_4H_2N_2O_4 \cdot H_2O$, M 160.1, has m 255°(dec), pK^{25} 6.64. Recrystallisation from H_2O gave the **tetrahydrate** in large prisms or rhombs. On heating at 100°, or on exposure to air, this is converted to the **monohydrate**. Dissolve it in its own weight of boiling H_2O and cool it for several days below 0°; the **tetrahydrate** crystallises from solution much more slowly when free from HNO_3 . It is less soluble in bicarbonate solutions than in H_2O . Drying the solid over H_2SO_4 yields the **monohydrate**. The **anhydrous** crystals can be obtained by recrystallisation from dry Me_2CO or AcOH followed by washing with dry Et_2O , or by sublimation in a vacuum. On heating it turns pink at 230° and decomposes at ca 256°. It is acidic to litmus. [Hartman & Sheppard *Org Synth Coll Vol 3* 37 1955, DOI: 10.15227/orgsyn.023.0003.] It forms a compound with urea which crystallises from H_2O in yellow needles that become red at 170° and decompose at 185-186°. [*Beilstein* **24** H 500, **24** I 428, **24** II 301, **24** III/IV 2137.]

Alloxantin (5,5'-dihydroxy-5,5'-bibarbituric acid) [76-24-4] $C_8H_6N_4O_8$, M 286.2, m 253-255°(dec) (yellow at 225°). Alloxantin crystallises from water or EtOH and is kept under nitrogen. It turns red in air. It can be obtained from uric acid [Nightingale *Org Synth Coll Vol 3* 42 1955, DOI: 10.15227/orgsyn.023.0006], or from alloxan hydrate [Tipson et al. *Org Synth Coll Vol 4* 25 1963, DOI: 10.15227/orgsyn.033.0003]. [*Beilstein* **26** III/IV 2782.]

Alloxazine (6,7-benzopteridin-2,4(1H,3H)-dione) [490-50-5] $C_{10}H_6N_4O_2$, M 214.2, dec > 300°, $pK_{Est(1)}$ ~2.0 (proton gain), $pK_{Est(2)}$ ~7.92 (proto loss), $pK_{Est(3)}$ ~12.00 (proton loss). Prepared by cyclisation of 2-carbethoxyaminoquinoxaline-3-carboxamide with NaOMe/MeOH which deposits the bright yellow **sodium salt** that dissolves in hot 2% aqueous NaOH and deposits the Na salt as yellow needles in high yield. When the salt is dissolved in aqueous EtOH and acidified with AcOH, alloxazine separates as a pale yellow powder which decomposes without melting above 300°. It turns greyish-green on standing in light. It can be recrystallised from 50% aqueous AcOH. It is sparingly soluble in EtOH, insoluble in H_2O or Et_2O . The UV (EtOH) has λ_{max} 322nm (ϵ 5,900) and 246 (ϵ 19,000). It is characterised by conversion into **1,3-dimethylalloxazine** which crystallises from MeOH in bright yellow prisms with m 238-240° (234-236°), and has UV (EtOH) with λ_{max} 378nm (ϵ 7,200), 234.5nm (ϵ 6,900) and 245nm (ϵ 32,200). [Gowenlock et al. *J Chem Soc* 517 1948, DOI: 10.1039/JR9480000517; Tishler et al. *J Am Chem Soc* **67** 215 1945, DOI: 10.1021/ja01228a031; *Beilstein* **26** H 498.]

1-Allyl-6-amino-3-ethyluracil (Aminometradine) [642-44-4] $C_9H_{13}N_3O_2$, M 195.2, m 143-144° (anhydrous). It crystallises from water (as **monohydrate**, m 70-114°). It is a diuretic. [*Beilstein* **24** III/IV 4133.]

1-N-Allyl-3-hydroxymorphinan (Levallorphan) [152-02-3] $C_{19}H_{25}NO$, M 283.4, m 180-182°, $[\alpha]_D^{20}$ -89 (c 3, MeOH). It crystallises from aqueous EtOH. It is a narcotic antagonist. [Schnider & Grüssner *Helv Chim Acta* **34** 2211 1951, DOI: 10.1002/hlca.19510340715; Hellerbach et al. *Helv Chim Acta* **39** 429 1956, DOI: 10.1002/hlca.19560390212.]

5-Allyl-5-isobutylbarbituric acid (Butalbital) [77-26-9] $C_{11}H_{16}N_2O_3$, M 224.3, m 138-139°, 139°, 139-140°, 140-142°, pK^{18} 12.36. It can be recrystallised from H_2O or dilute EtOH, and sublimes at 100-120°/8-

12mm. It is soluble in C_6H_6 , cyclohexane, tetralin and petroleum ether at 20° . It is a controlled substance, a depressant, hypnotic and a sedative. [Butler et al. *J Am Chem Soc* **77** 1486 1955, DOI: 10.1021/ja01611a024; *Beilstein* **24** III/IV 2006.]

9-Aminoacridine (9-acridineamine, Aminacrine) [90-45-9] $\text{C}_{13}\text{H}_{10}\text{N}_2$, M 194.2, m 241° , pK^{20} 9.95. It crystallises from EtOH or acetone and sublimes at $170\text{--}180^\circ/0.04\text{mm}$ [Albert & Ritchie *Org Synth Coll Vol* **3** 53 1955, DOI: 10.15227/orgsyn.022.0005; for hydrochloride, see below]. [*Beilstein* **21** II 280, **21** III/IV 4174.]

9-Aminoacridine hydrochloride monohydrate (Acramine yellow, Monacrin) $\text{C}_{13}\text{H}_{10}\text{N}_2\cdot\text{HCl}\cdot\text{H}_2\text{O}$, [52417-22-8 hydrate, 134-50-9 anhydrous] M 248.7, has m $>355^\circ$, pK_1^{20} 4.7, pK_2^{20} 9.99. Recrystallise it from boiling H_2O (charcoal; 1g in 300 ml) to give pale yellow crystals with a neutral reaction. It is one of the *most fluorescent substances known*. At 1:1000 dilution in H_2O it is pale yellow with only a faint fluorescence, but at 1:100,000 dilution it is colourless with an intense blue fluorescence. [Albert & Ritchie *Org Synth Coll Vol* **3** 53 1955, DOI: 10.15227/orgsyn.022.0005; Falk & Thomas *Pharm J* **153** 158 1944, *Beilstein* **22** H 463, **21** II 280, **21** III/IV 4174.] See previous entry for the free base.

2-Amino-5-(2-aminoethyl)-4-methylthiazole dihydrobromide (Amthamine dihydrobromide) [142437-67-0 for free base] $\text{C}_6\text{H}_{11}\text{N}_3\text{S}\cdot 2\text{HBr}$, M 319.1, m $275^\circ(\text{dec})$, $\text{pK}_{\text{Est}(1)} \sim 5.0$, $\text{pK}_{\text{Est}(2)} \sim 9.7$. Prepared from 3-bromo-5-phthalimido-2-pentanone and thiourea in DMF at $120^\circ/3\text{hrs}$ which converted to 2-amino-4-methyl-5-(2-phthalimidoethyl)thiazole hydrobromide (55% yield, m $270.3\text{--}272.4^\circ$, from absEtOH/EtOAc), and the amino group was deprotected by refluxing in 30% HBr solution for 5 hours, evaporated (80/20mm), HBr was chased with toluene to give a 75.2% yield of 2-amino-5-(2-amino-methyl)-4-methylthiazole dihydrobromide after recrystallisation from hot EtOH/Et₂O. It has ^1H NMR (90MHz, DMSO-*d*₆) with δ at 2.16 (s, 3H), 2.84-3.12 (br m, 4H), 8.08 (br s, 3H), 9.32 (br s, 2H) ppm; MS *m/z* (rel intensity) 158(5), 157(7), 104(31), 30(2); exact mass $M^+ = 157.062$, calc for $\text{C}_6\text{H}_{11}\text{N}_3\text{S}$ 157.067. [Eriks et al. *J Med Chem* **35** 3239 1992, DOI: 10.1021/jm00095a021]. *Amthamine* is a highly selective H_2 agonist that is slightly more potent than histamine, but only a weak antagonist at H_3 but no activity at H_1 receptors. [Coruzzi et al. *Arch Pharmacol* **348** 77 1993, DOI: 10.1007/BF00168540; for allergy and Histamine see Poli et al. *Agents Actions* **40** 44 1993, DOI: 10.1007/BF01976750.]

2-Amino-4-anilino-s-triazine (Amanozine) [537-17-7] $\text{C}_9\text{H}_9\text{N}_5$, M 187.2, m $235\text{--}236^\circ$, $\text{pK}_{\text{Est}} \sim 5.5$. It crystallises from dioxane or 50% aqueous EtOH. [*Beilstein* **26** III/IV 1195.]

4-Aminoantipyrine (Ampyrone, 4-amino-2,3-dimethyl-1-phenylpyrazol-5-one) [83-07-8] $\text{C}_{11}\text{H}_{13}\text{N}_3\text{O}$, M 203.3, m $105\text{--}110^\circ$, 109° . It crystallises from EtOH or EtOH/ether. It is antipyretic and an analgesic. [*Beilstein* **25** III/IV 3554.]

2-Aminobenzothiazole [136-95-8] $\text{C}_7\text{H}_6\text{N}_2\text{S}$, M 150.2, m $126\text{--}129^\circ$, 132° , pK^{20} 4.48. The thiazole crystallises from H_2O , aqueous EtOH, C_6H_6 or petroleum ether. The *hydrochloride* crystallises from dilute HCl and has m 240.5° . [*Beilstein* **27** H 182, **27** III/IV 4824.]

6-Aminobenzothiazole [533-30-2] $\text{C}_7\text{H}_6\text{N}_2\text{S}$, M 150.2, m 87° , $87\text{--}91^\circ$, $\text{pK}_{\text{Est}} \sim 3$. It crystallises from aqueous EtOH, petroleum ether or C_6H_6 /petroleum ether. The *hydrochloride* has m $305^\circ(\text{dec})$ from dilute HCl, and the *picrate* has m $185^\circ(\text{dec})$ from Me_2CO . [Boggust & Cocker *J Chem Soc* 355 1949, DOI: 10.1039/JR9490000355; *Beilstein* **27** III/IV 4884.]

3-o-Aminobenzyl-4-methylthiazolium chloride hydrochloride [534-94-1] $\text{C}_{11}\text{H}_{14}\text{N}_2\text{SCl}_2$, M 277.2, m $213^\circ(\text{dec})$. The *hydrochloride* crystallises from aqueous EtOH, and the *iodide hydroiodide* has m 273° (from aqueous HI). [*Beilstein* **27** III/IV 973.]

4-Amino-1-benzylpiperidine [50541-93-0] $\text{C}_{12}\text{H}_{18}\text{N}_2$, M 190.3, b $\sim 180^\circ/20\text{mm}$, d_4^{20} 0.933, n_D^{20} 1.543, $\text{pK}_{\text{Est}(1)} \sim 8.3$, $\text{pK}_{\text{Est}(2)} \sim 10.4$. Purify it by distillation *in vacuo* and store it under N_2 because it absorbs CO_2 . The *dihydrochloride salt* [1205-72-7] has m $270\text{--}273^\circ$ (255°) after recrystallisation from MeOH/EtOAc or EtOH. [Brookes et al. *J Chem Soc* 3165, 3172 1957, DOI: 10.1039/JR9570003165.]

The 4-methylamino-1-benzylpiperidine derivative has b $168\text{--}172^\circ/17\text{mm}$, n_D^{20} 1.5367 [Reitsema & Hunter *J Am Chem Soc* **70** 4009 1948, DOI: 10.1021/ja01192a011].

The *1-(1-benzyl-4-piperidinyl)-3-cyano-2-methylisothiourea* derivative has **m 160°** from $\text{CHCl}_3/\text{Et}_2\text{O}$ [Preparation, IR, NMR: Ried et al. *Chem Ber* **116** 1547 1983, DOI: 10.1002/cber.19831160431; *Beilstein* **22** III/IV 3752].

2-Amino-4-chloro-6-methylpyrimidine [5600-21-5] $\text{C}_5\text{H}_6\text{ClN}_3$, **M 143.6**, **m 183-186°, 184-186°, $\text{pK}_{\text{Est}} \sim 1.0$** . Recrystallise it from EtOH. [*Beilstein* **24** H 84, **25** IV 2171.]

2-Amino-5-chloropyridine [1072-98-6] $\text{C}_5\text{H}_5\text{ClN}_2$, **M 128.6**, **m 135-136°, 135-138°, 127-128°/11mm, $\text{pK}^{25}_{\text{Est}} 4.38$** . Recrystallise this base from petroleum ether. It sublimes at 50°/0.5mm. [*Beilstein* **22** II 332, **22/8** V 541.]

2-Amino-4-chloropyrimidine [3993-78-0] $\text{C}_4\text{H}_4\text{ClN}_3$, **M 129.55**, **m 168-169°, m 170°, $\text{pK}_{\text{Est}} \sim 1.2$** . The pyrimidine crystallises in glistening plates from EtOH (**m 170°**, sintering at 167°). It has also been purified by sublimation in a vacuum and recrystallisation from H_2O . [Hilbert & Johnson *J Am Chem Soc* **52** 1152 1930, DOI: 10.1021/ja01366a051]. Cf. Littke et al. *Org Lett* **9** 1711 2007, DOI: 10.1021/ol070372d for its use as a substrate in a palladium-catalysed cyanation with $\text{Z}(\text{CN})_2$. [*Beilstein* **24** H 80, **25** III/IV 2117.]

2-Amino-3,5-dibromopyridine [35486-42-1] $\text{C}_5\text{H}_4\text{Br}_2\text{N}_2$, **M 251.9**, **m 103-104°, 104-105°, $\text{pK}_{\text{Est}} \sim 2.4$** . Steam distil it and recrystallise it from aqueous EtOH or petroleum ether. [*Beilstein* **22** II 333, **22** III/IV 4041.]

3-Amino-2,6-dichloropyridine [62476-56-6] **M 164.0**, **m 119°, b 110°/0.3mm, $\text{pK}_{\text{Est}} \sim 2.0$** . Recrystallise it from water. [*Beilstein* **22** III/IV 4093.]

2-Amino-4,6-dimethylpyridine (6-amino-2,4-lutidine) [5407-87-4] $\text{C}_7\text{H}_{10}\text{N}_2$, **M 122.2**, **m 63-64°, 69-70.5°, 235°/atm, $\text{pK}^{25}_{\text{Est}} 7.84$** . Recrystallise this base from hexane, ether/petroleum ether or *benzene. Residual *benzene is removed over paraffin-wax chips in an evacuated desiccator. The *dipicrate* crystallises from EtOH and has **m 205-207°(dec)**. [*Beilstein* **22** III/IV 4210.]

2-Amino-4,6-dimethylpyrimidine [767-15-7] $\text{C}_6\text{H}_9\text{N}_3$, **M 123.2**, **m 151-153°, 152-153°, $\text{pK}^{25}_{\text{Est}} 4.95$** . Recrystallisation from water gives the pyrimidine with **m 197°**, but recrystallisation from acetone gives **m 153°**. [*Beilstein* **25** III/IV 2205.]

2-(Aminomethyl)piperidine [22990-77-8] $\text{C}_6\text{H}_{14}\text{N}_2$, **M 114.2**, **b 66-67°/12mm, 80-81°/18mm, $d_4^{20} 0.9406$, $n_D^{20} 1.4854$, $\text{pK}_1^{20} 6.33$, $\text{pK}_2^{20} 9.70$** . Dry (over Na_2SO_4) and distil the piperidine under vacuum from KOH. It has been purified via the *Reineke salt* (**m 173-174°**) and its *dipicrate salt* (**m 201°**, from H_2O). [Norton et al. *J Am Chem Soc* **68** 1330 1946, DOI: 10.1021/ja01211a071; Mortimer *Aust J Chem* **11** 82 1958, DOI: 10.1071/CH9580082; Augustine *J Am Chem Soc* **81** 4664 1959, DOI: 10.1021/ja01526a054; *Beilstein* **22** III/IV 3765.]

4-Amino-3-hydrazino-5-mercapto-1,2,4-triazole (Purpald) [1750-12-5] $\text{C}_2\text{H}_6\text{N}_6\text{S}$, **M 146.2**, **m 228-230°(dec), 234-235°(dec), $\text{pK}_{\text{Est}(1)} \sim 2$, $\text{pK}_{\text{Est}(2)} \sim 3$ (NH_2), $\text{pK}_{\text{Est}(3)} \sim 8$ (SH)**. Recrystallise Purpald from H_2O (0.6g in 300-400ml). The *benzylidene* derivative has **m 245-246°(dec)** from *i*-PrOH [Hoggarth *J Chem Soc* 4817 1952, DOI: 10.1039/JR9520004817; Dickinson & Jacobson *JCS Perkin Trans I* 975 1975, DOI: 10.1039/P19750000975; [*Beilstein* **26** III/IV 547.]

5-Amino-8-hydroxyquinoline hydrochloride [3881-33-2] $\text{C}_9\text{H}_8\text{N}_2\text{O} \cdot \text{HCl}$, **M 196.7**, **$\text{pK}_1^{20} 5.67$, $\text{pK}_2^{20} 11.24$** . Dissolve the hydrochloride in the minimum volume of MeOH, then add Et_2O to initiate crystallisation. The crystals are filtered off and dried [Lovell et al. *J Phys Chem* **88** 1885 1984, DOI: 10.1021/j150653a042]. The *dihydrochloride* [21302-43-2] has **M 233.1, m 279°(dec)**. [*Beilstein* **22** III/IV 5866.]

4-Aminoimidazole-5-carboxamide hydrochloride (AICAR HCl) [72-40-2] $\text{C}_4\text{H}_6\text{N}_4\text{O} \cdot \text{HCl}$, **M 162.6**, **m 250-252°(dec), 255-256°(dec), $\text{pK}_{\text{Est}(1)} \sim 3.5$, $\text{pK}_{\text{Est}(2)} \sim 9.4$** . Recrystallise the hydrochloride from EtOH. [Kuroda & Suzuki *J Heterocycl Chem* **30** 593 1993, DOI: 10.1002/jhet.5570300302; Alhede et al. *J Org Chem* **56** 2139 1991, DOI: 10.1021/jo00006a033; Chern et al. *Heterocycles* **34** 1133 1992, DOI: 10.3987/COM-92-6000; *Beilstein* **25** II 221, **25** III/IV 4329.]

6-Aminoindazole [6967-12-0] $C_7H_7N_3$, M 133.2, m 204-206°(dec), 210°, pK^{25} 3.99. It is recrystallised from H_2O or EtOH and sublimes in a vacuum. [Beilstein 25 H 317.]

2-Amino-3-iodopyridine [104830-06-0] $C_5H_5IN_2$, M 220.1, m 78-91°, 90-91.5°, pK_{Est} ~4.9. Purify this pyridine by recrystallisation from hexane. The *N-Me* derivative [113975-23-8] has m 50°, and distils at b 129°/14mm. [Sakamoto et al. *Chem Pharm Bull, Japan* 33 4764 1985, DOI: org/10.1248/cpb.33.4764; Estel et al. *J Org Chem* 53 2740 1988, DOI: 10.1021/jo00247a016.]

2-Amino-4-iodopyridine [552331-00-7] has m 163-164°, pK_{Est} ~5.1. Purify this pyridine by recrystallisation from H_2O . The *picrate* has m 253-254° (from H_2O), the *N-acetyl* derivative has m 150° (from H_2O), and the *N-benzoyl* derivative has m 167-168° (from aqueous EtOH). [Graf *Chem Ber* 64 21 (25) 1931, DOI: 10.1002/cber.19310640103.]

2-Amino-5-iodopyridine [20511-12-0] has m 128-131°, 129-130°, 130°, pK_{Est} ~4.5. The pyridine can be purified by steam distillation. Separate the solid from the cooled distillate by filtration, acidify the filtrate, decolorise it with charcoal, concentrate it to ~200ml, make alkaline with KOH and cool. Filter the solid, add it to the original solid that was collected, and dry it *in vacuo*. It crystallises from $*C_6H_6$ in white needles. The *picrate* separates as yellow needles from hot EtOH or Me_2CO , m 240°. [Caldwell et al. *J Am Chem Soc* 66 1479 1944, DOI: 10.1021/ja01237a018.] [Beilstein 22 II 334.]

5-Amino-2-iodopyridine [29958-12-1] has m 63-65°, pK_{Est} ~2.6. Purify it by recrystallisation from EtOH (white needles). [Caldwell et al. *J Am Chem Soc* 66 1479 1944, DOI: 10.1021/ja01237a018.]

3-Amino-5-mercapto-1,2,4-triazole [16691-43-3] $C_2H_4N_4S$, M 116.1, m 298°, ~300°, $pK_{Est(1)}$ ~3.0, $pK_{Est(2)}$ ~9. Recrystallise the triazole from H_2O and dry it *in vacuo*. The *acetyl* derivative has m 325°(dec) after recrystallisation from H_2O . [Beilstein 26 III/IV 1351.] It has also been recrystallised from EtOH/ H_2O (3:1, 1g in 50 ml, 50% recovery), m 300-302° (dec subject to heating rate), (λ_{max} 263nm, log ϵ 4.12). The *S-benzyl* derivative, when crystallised from $*C_6H_6$ /EtOH (20:1), or $CHCl_3$ /Et $_2O$ has m 109-111° [Godfrey & Kurzer *J Chem Soc* 3437 1960, DOI: 10.1039/JR9600003437; Beilstein 26 III/IV 1351.]

2-Amino-4-methoxy-6-methylpyrimidine [7749-47-5] $C_6H_9N_3O$, M 139.2, m 156-158°, 157-159°, 158-158.5°, 158-160°, pK_{Est} ~6.0. Recrystallise it from H_2O . The *picrate* has m 220-221°(dec). [Braker et al. *J Am Chem Soc* 69 3072, 3075 1947, DOI: 10.1021/ja01204a044; Sirakawa et al. *Yakugaku Zasshi* 73 598 1953, Backer & Grevenstuk *Recl Trav Chim Pays-Bas* 61 291 1942, DOI: 10.1002/recl.19420610408; Beilstein 25 III/IV 3385.]

8-Amino-6-methoxyquinoline [90-52-8] $C_{10}H_{10}N_2O$, M 174.2, m 41-42°, 51°, b 137-138°/1mm, $pK^{70.1}$ 3.38. Distil it under N_2 and at high vacuum, then recrystallise it several times from MeOH (0.4ml/g). It remains colourless for several months when purified in this way [Elderfield & Rubin *J Am Chem Soc* 75 2963 1953, DOI: 10.1021/ja01108a052]. The *hydrobromide* [312693-53-1] M 255.1 has m 238°(dec). [Beilstein 22 III/IV 5934.]

7-Amino-4-methylcoumarin (Coumarin 120) [26093-31-2] $C_{10}H_9NO_2$, M 175.2, m 223-226°, 224-229°(dec), pK_{Est} ~3.2. Dissolve it in 5% HCl, filter and basify with 2M ammonia. The precipitate is dried in a vacuum and recrystallised from dilute EtOH. It yields a blue solution and is light sensitive. [Khammungskhune & Sigler *Synthesis* 614 1980, DOI: 10.1055/s-1980-29136; Kanaoka et al. *Chem Pharm Bull Jpn* 30 1485 1982, DOI: org/10.1248/cpb.30.1485]. Useful laser dye with λ_{max} at 354nm, and in fluorescent labeling of trace enzymes [Aldrichimica Acta 15 42 1982, for 7-amino-4-methyl-6-sulfocoumarin-3-acetic acid (AMCA-S), see Leung et al. *Biorg Med chem Lett* 9 2229 1999, DOI: 10.1016/S0960-894X(99)00364-9]. [Beilstein 18/11 V 445.]

2-Amino-3-methylpyridine (2-amino-3-picoline) [1603-40-3] $C_6H_8N_2$, M 108.1, m 29-31°, 33.2°, b 221-222°/atm, d_4^{25} 1.073, n_D^{20} 1.5823. pK^{25} 7.24. Recrystallise the picoline three times from $*benzene$, most of the residual $*benzene$ being removed from the crystals by standing over paraffin wax chips in an evacuated desiccator. The amine is also transferred to a separating funnel under N_2 , and left in contact with NaOH pellets for 3 hours with occasional shaking. It is then placed in a vacuum distilling flask where it is refluxed gently in a

stream of dry N₂ before fractionally distilling it. [Mod et al. *J Phys Chem* **60** 1651 1956, DOI: 10.1021/j150546a013; *Beilstein* **22/9** V 212].

2-Amino-4-methylpyridine (2-amino-4-picoline) [695-34-1] C₆H₈N₂, M 108.1, m 96-99°, 99.2°, 100-100.5°, b 115-117°/11mm, 230°/atm, pK²⁵ 7.48. Crystallise it from EtOH, a 2:1 *benzene/acetone mixture, or petroleum ether, and dry it under vacuum as in the previous entry. Sublimes in a vacuum. It is cardiotoxic. [*Beilstein* **22/9** V 325.]

2-Amino-5-methylpyridine (6-amino-3-picoline) [1603-41-4] has m 76-77°, 76.5°, b 227°/atm, pK²⁵ 7.22. Crystallise it from acetone, and/or distil it. [*Beilstein* **22/9** V 289.]

2-Amino-6-methylpyridine (6-amino-2-picoline) [1824-81-3] has m 40-44°, 44.2°, b 208-209°/atm, pK²⁵ 7.41. Crystallise it three times from acetone and dry it under vacuum at ca 45°. Alternatively, keep it in contact with NaOH pellets for 3 hours, with occasional shaking, decant and fractionally distil it [Mod et al. *J Phys Chem* **60** 1651 1956, DOI: 10.1021/j150546a013]. It also recrystallises from CH₂Cl₂ on addition of petroleum ether. [Marzilli et al. *J Am Chem Soc* **108** 4830 1986, DOI: 10.1021/ja00276a021; *Beilstein* **22/9** V 210.]

2-Amino-4-methylpyrimidine [108-52-1] C₅H₇N₃, M 109.1, m 158-160°, 159-160°, 161°, pK²⁰ 4.15. Crystallise the pyrimidine from H₂O or EtOH and sublime it in a vacuum. The *picrate* crystallises from EtOH and has m 235-236°(dec). [*Beilstein* **24** H 84, **25** III/IV 2152.]

2-Amino-5-methylpyrimidine [50840-23-8] has m 193.5°, pK_{Est} ~4.0. Crystallise it from water or *benzene/petroleum ether and sublime it at 50°/0.5mm. [*Beilstein* **24** H 87.]

4-Amino-2-methylquinoline (4-aminoquinaldine) [6628-04-2] C₁₀H₁₀N₂, M 158.2, m 162-166°, 168°, b 333°/760mm, pK²⁰ 9.42. Recrystallise it from *benzene/petroleum ether. [*Beilstein* **22/10** V 347.]

6-Aminonicotinic acid [3167-49-5] C₆H₆N₂O₂, M 138.1, m 312°(dec), pK_{Est(1)} ~2.2 (CO₂H), pK_{Est(2)} ~6.5. Crystallise the acid from aqueous acetic acid. Dry it *in vacuo* at 70°. It decarboxylates cleanly above its melting point to give 2-aminopyridine. [*Beilstein* **22** III/IV 6726.]

(+)-6-Aminopenicillanic acid [551-16-6] C₈H₁₂N₂O₃S, M 216.3, m 198-200°(dec), 208-209°, [α]_D²² +267.3 (c 1.2, 0.1M HCl), [α]₅₄₆ +327 (in 0.1M HCl), pK₁²⁵ 2.30, pK₂²⁵ 4.90. This acid crystallises from water or aqueous HCl. [Kleppe & Strominger *J Biol Chem* **254** 4856 1979, <http://www.jbc.org/content/254/11/4856>; *Beilstein* **27** III/IV 2858.]

2-Aminoperimidine [28832-64-6] C₁₁H₉N₃, M 183.1, m 239°, b 170-175°/1.5mm, pK_{Est} ~7.9 (free base). It crystallises from EtOH/H₂O (1:1). It precipitates as the *hydrochloride* with dilute HCl which has m 282°. [Dasgupta et al. *Anal Chim Acta* **94** 205 1977, DOI: 10.1016/S0003-2670(01)83651-5; *Beilstein* **24** H 193, **25** III/IV 2677.] **2-Aminoperimidine hydrobromide** [40835-96-9, 313223-13-1 hydrate] C₁₁H₉N₃·HBr·xH₂O, M 264.1(anhydr), has m 299°, pK_{Est} ~7.9 (free base). Purify the hydrobromide by boiling a saturated aqueous solution with charcoal, filtering and leaving the salt to crystallise. Store this *dihydrate salt* in a cool, stoppered flask in the dark place. The *anhydrous salt* is obtained by heating at 80°/4 hours, and it is hygroscopic. The solubilities of the hydrobromide at 26° are 2.4% in EtOH, 0.6% in H₂O, 0.3% in Et₂O, 0.1% in Me₂CO and 0.003% in *C₆H₆. [Dasgupta et al. *Anal Chim Acta* **94** 205 1977, DOI: 10.1016/S0003-2670(01)83651-5; Dasgupta & West *Microchim Acta* **70** 505 1978, DOI: <http://www.jbc.org/content/177/1/357>; Dasgupta et al. *Anal Chem* **50** 1793 1978, DOI: 10.1021/ac50035a020; *Beilstein* **24** H 193, **25** III/IV 2677.]

2-Aminopyridine [504-29-0] C₅H₆N₂, M 94.1, m 54-58°, 58°, 58.1°, b 204-210°/atm, 210.6°/atm, pK₁²⁵ -7.6, pK₂²⁵ 6.71. It crystallises from *benzene/petroleum ether (b 40-60°) or CHCl₃/petroleum ether. [*Beilstein* **22/8** V 280.] IRRITANT.

3-Aminopyridine [462-08-8] has m 64°, b 248°/atm, 250-252°/atm, pK₁²⁵ -1.5, pK₂²⁵ 6.03. It crystallises from *benzene, CHCl₃/petroleum ether (b 60-70°), or *benzene/petroleum ether (4:1). [Allen & Wolf *Org Synth Coll Vol* **4** 45 1963, DOI: 10.1522/orgsyn.030.0003; *Beilstein* **22/9** V 3.] IRRITANT.

4-Aminopyridine [504-24-5] has m 155-158°, 158-159°, 160°, b 180°/12-13mm, 273°/atm, pK₁²⁵ -6.55, pK₂²⁵ 9.11 (9.18). Crystallise the aminopyridine from *benzene/EtOH, then recrystallise it twice from water, then crush and dry it for 4 hours at 105° [Bates & Hetzer *J Res Nat Bur Stand* **64A** 427 1960, DOI:org/10.6028/jres.064A.044]. It has also been crystallised from EtOH, *benzene, *benzene/petroleum

ether, toluene and sublimes in a vacuum. It is a relatively strong organic base. [*Beilstein* 22/9 V 106.] It is of some effect in demyelinating nerve fibres and in multiple sclerosis.

2-Aminopyrimidine [109-12-6] $C_4H_5N_3$, M 95.1, m 122-126°, 126-127.5°, pK²⁰ 3.45. Crystallise 2-aminopyrimidine from *C₆H₆, EtOH or H₂O. [*Beilstein* 25 III/IV 2071.]

4-Aminopyrimidine [591-54-8] has m 149-151°, 151-152°, 154-156°, pK²⁵ 5.69. Recrystallise 10.6g of aminopyrimidine from hot EtOAc (200ml) to give 7.4g colourless needles as first crop; evaporation to 25ml gives a second crop of 1.7g. It also crystallises from ~200 parts of light petroleum (b 100-120°) or 50 parts of isobutyl methyl ketone (charcoal). The *hydroiodide* has m 180°. The *picrate* has m 225°, and the *1-methyl-iodide* has m 204-205° (crystallise from 20 parts of EtOH) [Brown et al. *J Chem Soc* 4035 1955, DOI: 10.1039/JR9550004035]. [Synth: Brown *J Soc Chem Ind* (London) 69 353 1950; also prepared by decarboxylation of 4-amino-5-carboxypyrimidine Brown & Short *J Chem Soc* 331 1953, DOI: 10.1039/JR9530000331; *Beilstein* 24 H 81, 24 III/IV 2130.]

5-Aminopyrimidine [591-55-9] has m 171-172° (with sublimation), pK²⁵ 2.52. It is purified by conversion to the MgCl₂ complex in a small volume of H₂O. The complex (~5g) is dissolved in the minimum volume of hot H₂O, passed through a column of activated Al₂O₃ (200g), and the column is washed with EtOH. Evaporation of the EtOH gives a colourless residue of the aminopyrimidine which is recrystallised from *C₆H₆ (toluene could also be used) which forms needles at first, then prisms. It melts with sublimation. Acetylation yields *5-acetamidopyrimidine* which crystallises from *C₆H₆, m 148-149°. The NH₂ group behaved more like the one in aniline. [Whittaker & Jones *J Chem Soc* 1565 1951, DOI: 10.1039/JR9510001565.]

Aminopyrine (4-dimethylaminoantipyrine) [58-15-1] M 231.3, m 107-109°, 108°, pK₁²⁵ -2.22, pK₂²⁵ 4.94. It crystallises from petroleum ether, sublimes between 80° and 90°, and forms metal complexes. [*Beilstein* 25 H 452, 25 III/IV 3555.]

3-Aminoquinoline [580-17-6] $C_9H_8N_2$, M 144.2, m 91-92°, 93.5°, pK₁²⁰ -0.58, pK₂²⁰ 4.94. It crystallises from *C₆H₆, toluene, hexane and aqueous EtOH. [*Beilstein* 22 III/IV 4605, 22/10 V 233.]

4-Aminoquinoline [578-68-7] has m 155-155.5°, 158°, pK₁²⁵ -7.11(5.99), pK₂²⁰ 9.13. It has been purified by zone refining and recrystallisation from *C₆H₆, EtOH or H₂O. The *hydrochloride* has m 308° (from MeOH), and the *picrate* has m 277° (from EtOH). [Albert et al. *J Chem Soc* 2240 1948, DOI: 10.1039/JR9480002240; *Beilstein* 22 III/IV 4611, 22/10 V 341.]

5-Aminoquinoline [611-34-7] has m 106-109°, 110°, b 184°/10mm, 310°/760mm, pK₁²⁰ 0.97(0.49), pK₂²⁰ 5.42. It crystallises from pentane and from *benzene or EtOH. The *picrate* has m 209-210°(dec) (202° dec) (from aqueous EtOH). [*Beilstein* 22 III/IV 4669, 22/10 V 297.]

6-Aminoquinoline [580-15-4] has m 115-119°, 117-119°, 120°, b 146°/0.3mm, 192-195°/14mm, pK₁²⁰ 1.63, pK₂²⁰ 5.59. It is purified by column chromatography on a SiO₂ column using CHCl₃/MeOH (4:1) as eluent. It crystallises from *C₆H₆ or *C₆H₆/petroleum ether and is an irritant. The *styphnate* has m 239-240° (from EtOH) and m 242-243° (from aqueous Me₂CO). [Barrett et al. *J Chem Soc* 50, 57 1953, DOI: 10.1039/JR9530000050; *Beilstein* 22 III/IV 4681, 22/10 V 303.]

8-Aminoquinoline [578-66-5] $C_9H_8N_2$, M 144.2, m 60-65°, 70°, b 140.5-141°/7mm, 123°/5mm, 174°/26mm, pK₁²⁰ -0.52, pK₂²⁰ 3.95. 8-Aminoquinoline crystallises from EtOH, ligroin, octane or H₂O, and complexes with metals. [*Beilstein* 22 III/IV 4708, 22/10 V 316.]

2-Amino-5-sulfanilylthiazole (thiazolsulfone, Promizole) [473-30-3] $C_9H_9N_3O_2S_2$, M 255.3, m 219-221°(dec), pK_{Est} ~4.5 (OH). If too impure, it may contain some 'iron-mud', then extract it with Me₂CO, filter and evaporate the filtrate *in vacuo*. Dissolve the residue in boiling absolute EtOH, treat it with charcoal (ca 10% w/w) filter and allow to cool. Recrystallise the thiazole again from EtOH (needles). It possesses tuberculotherapeutic activity. [Bambas *J Am Chem Soc* 67 671 1945, DOI: 10.1021/ja01220a050.]

2-Aminothiazole [96-50-4] $C_3H_4N_2S$, M 108.1, m 91-93°, 93°, b 140°/11mm, pK²⁰ 5.36. It crystallises from petroleum ether (b 100-120°), or EtOH. It undergoes Ulmann coupling with 2-chlorobenzoic acid in the presence of ultrasonic irradiation [Pellón et al. *Synth Commun* 37 1853 2007, DOI:10.1080/00397910701319056]. [*Beilstein* 27 III/IV 4574.] It inhibits thyroid growth.

2-Amino-4-thiazoleacetic acid (2-amino-4-carboxymethylthiazoline) [29676-71-9] $C_5H_6N_2O_2S$, **M 158.2**, **m 130°(dec)**, $pK_{Est(1)} \sim 4.5$ (COOH), $pK_{Est(2)} \sim 5.3$ (2-NH₂). Purified by recrystallisation from hot H₂O (solubility is ~ 6.5 g/L at $\sim 20^\circ$) or EtOH. It has been prepared by mixing ethyl chloroacetoacetate and ammonium dithiocarbamate in H₂O for 5 hours at room temperature then refluxed for 2 hours which gives an 81% yield of the **ethyl ester** [53266-94-7] $C_7H_{10}N_2O_2S$, **M 186.2**, **m 92-94°**, recrystallised from di-*iso*-propyl ether, and hydrolysed by 2.5N NaOH at 55° for 1 hour followed by acidification. [Bolchi et al. *Bioorg Med Chem Lett* **21** 5408 2011, DOI: 10.1016/j.bmcl.2011.07.003; Steude *Justus Liebigs Ann Chem* **261** 22 1891, DOI: 10.1002/jlac.18912610103]. The acid and ester are **IRRITANTS**. They are used in the pharmaceutical industry and as pesticides. The acid forms Ni II [He et al. *Acta Crystallogr Sect E* **65** m666 2009, DOI: 10.1107/S1600536809017978, PMID: 21583027], CdCl₂ [with the ethyl ester Zhang et al. *Acta Crystallogr Sect E* **68** m788 2012, DOI: 10.1107/S1600536812021976, PMID: 22719339] and Zn II [Zhang et al. *Acta Crystallogr Sect E* **65** m1517 2009, DOI: 10.1107/S1600536809045589, PMID: 21578564] crystalline complexes. [Beilstein **27** H 336.]

The **2-amino-4-thiazoleacetic acid fluoroborate salt** [110295-78-8 *zwitterion*] $C_5H_7N_2O_2SBF_4$, prepared by hydrolysis of the ethyl ester with 34% aqueous fluoroboric acid overnight at $\sim 25^\circ$, and the salt which crystallise upon addition of Et₂O to the mixture, had **m 180-185°** and ¹H NMR (Me₂SO-*d*₆, TMS) with δ at 3.55 (2H, s), 6.55 (1H, s), 7.8-8.2 NH₂, br [Bouchet et al. *J Med Chem* **30** 2222 1987, DOI: 10.1021/jm00395a008].

2-Amino-4-thiazoleacetic acid hydrazide is prepared by boiling the **ethyl ester** (0.4mol, above) and hydrazine hydrate (0.4mol) in absolute EtOH (300ml) for 6 hours, cooled and allowed to crystallise over 3 days. The hydrazide is then recrystallised from EtOH and dried in a desiccator over silica gel. It has antifungal and antitubercular activities and it forms light brown crystals of the **CuII complex**. [Enedoh *Int J Sci Eng Res* **6** (4) 110 2015, ISSN 2229-5518.]

2-Amino-5-thiazoleacetic acid ethyl ester (2-amino-5-ethoxycarbonylmethyl-1,4-thazole) [62557-32-8] $C_7H_{10}N_2O_2S$, **M 186.2**, has **m 100-101°**. The ester was prepared by condensing thiourea (1.4g) with β -bromo- β -aldehydo-propionic acid ethyl ester (5.2g, = bromosuccinic acid semialdehyde ethyl ester) in EtOH (25ml) during 3 hours (boiling water bath), evaporating *in vacuo*; the crystalline residue was dissolved in dilute HCl (2ml concentrated HCl and 25ml H₂O), extracted into Et₂O (2 x 5ml), washed with aqueous NaHCO₃, filtered and evaporated. The residue was recrystallised from *C₆H₆/petroleum ether or CHCl₃/petroleum ether to give pure **ester** as white plates (3.1g, 90%) which were fairly soluble in H₂O, Et₂O, EtOH, *C₆H₆ and CHCl₃ but insoluble in petroleum ether. The **ester-picrate** forms yellow needles from EtOH with **m 204-205°(dec)**, and the **ester-picolonate** has **m 216-218°(dec)** (golden yellow plates from absolute EtOH). The **amide** was formed by shaking the ester with concentrated aqueous NH₃, evaporating to dryness (hot water bath), and the residual solid was purified by subliming at 140-150°/0.05mm or distilling at 180°/0.5mm to give white crystals **m 174-175°(dec)** of the amide which are slightly soluble in H₂O but insoluble in most organic solvents. [Mory & Schenkel *Helv Chim Acta* **33** 405 1950, DOI: 10.1002/hlca.19500330225.]

2-Amino-5-thiazoleacetic acid fluoroborate salt [110295-83-5 *zwitterion*] $C_5H_7N_2O_2SBF_4$, was prepared by hydrolysis of the ethyl ester with 34% aqueous fluoroboric acid overnight at $\sim 25^\circ$, and the salt was crystallised by addition of Et₂O to the mixture. It had **m 215°** and ¹H NMR (Me₂SO-*d*₆) with δ at 3.72 (2H, s), 6.2-6.8 NH₂, br) and 7.0 (1H, s) [Bouchet et al. *J Med Chem* **30** 2222 1987, DOI: 10.1021/jm00395a008].

1-Amino-1,2,4-triazole [24994-60-3] $C_2H_4N_4$, **M 84.1**, **m 91-93°**, $pK_{Est} \sim 2$. The triazole crystallises from water. [Barszcz et al. *JCS Dalton Trans* 2025 1986, DOI: 10.1039/DT9860002025; Temple & Montgomery *1,2,4-Triazoles – The Chemistry of Heterocyclic Compounds* Vol 37 (Weissberger & Taylor eds.). Wiley & Sons NY 1981, ISBN 0-471-0656-6.]

3-Amino-1,2,4-triazole (3-AT, Amitrol) [61-82-5] has **m 159°**, pK_1^{20} **4.04**, pK_2^{20} **11.08**. It crystallises from EtOH (charcoal), then three times from dioxane [Williams et al. *J Phys Chem* **61** 261 1957, DOI: 10.1021/j150549a002]. [Beilstein **26** H 137.] **Possible carcinogen**. [Sjostedt & Gringas *Org Synth Coll Vol* **3** 95 1955, DOI: 10.15227/orgsyn.026.0011; Beilstein **26** H 137, Temple & Montgomery *1,2,4-Triazoles – The Chemistry of Heterocyclic Compounds* Vol 37 (Weissberger & Taylor eds.). Wiley & Sons NY 1981, ISBN 0-471-0656-6.] The **hydrochloride**, **m 153°**, crystallises from EtOH, and has herbicidal activity. **IRRITANT**.

4-Amino-1,2,4(4H)-triazole [584-13-4] has **m 80-81°**, **84-86°**, pK^{25} **3.23**. It crystallises from EtOH/ Et₂O or H₂O. The **hydrochloride** has **m 151-152°** (from EtOH, 1g/10ml). [Allen & Bell *Org Synth Coll Vol* **3** 96 1955, DOI: 10.15227/orgsyn.024.0012; Barszcz et al. *JCS Dalton Trans* 2025 1986, DOI: 10.1039/DT9860002025; Beilstein **26** H 16, **26** II 7, **26** III/IV 40, Temple & Montgomery *1,2,4-Triazoles – The Chemistry of Heterocyclic Compounds* Vol 37 (Weissberger & Taylor eds.). Wiley & Sons NY 1981, ISBN 0-471-0656-6.]

7-Amino-4-(trifluoromethyl)coumarin (Coumarin 151) [53518-15-3] $C_{10}H_6F_3NO_2$, M 229.1, m 221-222°, 222°, $pK_{Est} \sim 3.1$. Purify the coumarin by column chromatography on a C18 column, elute with acetonitrile/0.01M H_2O/HCl (1:1), and recrystallise it from isopropanol. Alternatively, it is eluted from a silica gel column with CH_2Cl_2 , or by extracting a CH_2Cl_2 solution (4g/L) with 1M aqueous NaOH (3 x 0.1L), followed by drying ($MgSO_4$), filtration and evaporation. It is a suitable laser dye with λ_{max} at 207nm. [Bissell *J Org Chem* **45** 2283 1980, DOI: 10.1021/jo01300a003; Zimmermann et al. *Anal Biochem* **70** 258 1976 DOI: 10.1016/S0003-2697(76)80066-8.]

4(6)-Aminouracil (4-amino-2,6-dihydroxypyrimidine) [873-83-6] $C_4H_5N_3O_2$, M 127.1, m >350°, ~360°, pK_1^{20} 0.00 (basic), pK_2^{20} 8.69 (acidic), pK_3^{20} 15.32 (acidic). Purify the aminouracil by dissolving it in 3M aqueous NH_3 , filtering hot, and adding 3M formic acid until precipitation is complete. Cool, filter off (or centrifuge), wash well with cold H_2O , then EtOH and dry it in air. Dry it further in a vacuum at ~80°. [Barlin & Pfeleiderer *J Chem Soc (B)* 1425 1971, DOI: 10.1039/J29710001425; Beilstein **25** III/IV 4107.]

Amodiaquin [SN 10,751; **4-(3-dimethylaminomethyl-4-hydroxyanilino)-7-chloroquinoline, Camoquin**] [86-42-0] $C_{20}H_{22}ClN_3O$, M 355.9, m 208°(dec). Amodiaquin crystallises from 2-ethoxyethanol, *iso*PrOH or EtOH. It was prepared in various ways, e.g. by reacting 4,7-dichloroquinoline with 4-aminophenol in AcOH, and without isolating the intermediate, the mixture was treated with formaldehyde and Et_2NH in aqueous HCl. Alternatively, 4-acetaminophenol was converted, with formaldehyde and Et_2NH in aqueous HCl, into 4-amino-2-dimethylaminomethylphenol which was condensed with 4,7-dichloroquinoline in DMF/ Na_2CO_3 then aqueous HCl, all in a 'one pot synthesis' in a *Green Chemistry* US Patent Synthesis [Burckhalter et al. US Patent 2474819, 2474821 1949 to Parke Davis; WO 2013138200 A1, Kulkarni et al. Sep 19 2013 Howard University]. It also crystallises in pale yellow crystals from H_2O with **2HCl. 2H₂O** [6398-98-7] m 150-160°(dec) (under the names Camoquin, CAM-AQ1 or Flavoquin); forms yellow crystals from Me_2CO/H_2O with **2HCl. 1H₂O**, m 183°(dec), and when recrystallised from MeOH it also forms yellow crystals with **2HCl. 0.5H₂O**, m 243°(dec). The *free base* is *anhydrous* as it crystallises from absolute EtOH with m 208°(dec). They have UV with λ_{max} mn ($E_{1\%}^{1cm}$) at 342(349) in MeOH; 341.5(389) in H_2O and 342(396) in 0.1N HCl. They have some solubility in H_2O , more so in aqueous acid due to the dialkylamino group and 4-arylaminoquinoline moiety, also in alkaline medium due to the phenolic group; low solubility in alcohols, but quite insoluble in C_6H_6 , $CHCl_3$ or Et_2O . [Burckhalter et al. *J Am Chem Soc* **70** 1363 1948, DOI: 10.1021/ja01184a023; for UV spectra see Krácmár et al. *Pharmazie* **29** 773 1974, PMID: 4460004; Beilstein **22** III/IV 4647.]. They have antimalarial activity, and their activities, e.g. against *Gallinaceum* malaria in chicks are ~25 times those of quinine; and if the diethylamino groups are replaced by di-*iso*-propyl groups their activities can be up to ~75 times those of quinine.

2-n-Amylpyridine (2-n-pentylpyridine) [2294-76-0] $C_{10}H_{15}N$, M 149.2, b 63.0°/2mm, 206.5-207°/~760mm, n_D^{26} 1.4861, pK^{25} 6.00. Dry it with NaOH for several days, then distil it from CaO under reduced pressure, taking the middle fraction and redistilling it. The *picrate* has m 72-72.8° (from EtOH). [Beilstein **20** III/IV 2835.]

3-n-Amylpyridine [1802-20-6] $C_{10}H_{15}N$, M 149.2, b 110-112°/20mm, 224-226°/748mm, $pK_{Est} \sim 5.8$. Purify as the 2-isomer, and has the *picrate* has m 79.5-80°(from EtOH). [Beilstein **20** III/IV 2835.] Flavouring agent.

4-n-Amylpyridine [2961-50-4] $C_{10}H_{15}N$, M 149.2, b 78.0°/2.5mm, 229-230°/~760mm, n_D^{20} 1.4908, $pK_{Est} \sim 6.1$. It is dried with NaOH for several days, then distilled from CaO under reduced pressure, taking the middle fraction and redistilling it. The *picrate* has m 104° (from EtOH). [Beilstein **20** III/IV 2836.]

Antipyrine (2,3-dihydro-1,5-dimethyl-3-oxo-2-phenylpyrazole) [60-80-0] $C_{11}H_{12}N_2O$, M 188.2, m 111-113°, 114°, b 319°, pK^{25} 1.45. Antipyrine crystallises from EtOH/water mixture, *benzene, *benzene/petroleum ether or hot water (charcoal), and the crystals are dried under a vacuum. [Beilstein **24** H 27, **24** III/IV 75.] It is an analgesic.

Aspergillilic acid [2-hydroxy-3-isobutyl-6-(1-methylpropyl)pyrazine 1-oxide] [490-02-8] $C_{12}H_{20}N_2O_2$, M 224.3, m 97-99°, pK^{25} 5.5, $[\alpha]_D^{20}$ +13.3 (c 4, EtOH). It is recrystallised from MeOH (yellow needles), and is sublimed at 80°/10⁻³mm. [Dutcher *J Biol Chem* **171** 321 1947, <http://www.jbc.org/content/254/11/4856>; Beilstein **24** III/IV 235.]

8-Azaadenine (7-amino-1-*v*-triazolo[*d*]pyrimidine) [1123-54-2] $C_4H_4N_6$, **M 136.1**, **m 345°(dec)**, **pK₁²⁰ 2.65**, **pK₂²⁰ 6.29**. 8-Azaadenine crystallises from H₂O. Antimetabolite of purines. [See Cavalieri et al. *J Am Chem Soc* **70** 3875 1948, DOI: 10.1021/ja01191a102; for spectra, *Beilstein* **25** III/IV 4157.]

2-Azacyclotridecanone (lauro lactam) [947-04-6] $C_{12}H_{23}NO$, **M 197.3**, **m 150-153°, 152°**. 2-Azacyclotridecanone crystallises from CHCl₃ and is stored over P₂O₅ in a vacuum desiccator. [*Beilstein* **26**/I V 566.]

8-Azaguanine (5-amino-7-hydroxy-1-*v*-triazolo[*d*]pyrimidine) [134-58-7] $C_4H_4N_6O$, **M 152.1**, **m >300°**, **pK₁²⁰ 1.04**, **pK₂²⁰ 6.29**. Dissolve it in hot M NH₄OH, filter, and cool whereby colourless crystals separate out. Recrystallise it, and wash it with water, then dry it in a vacuum. Antimetabolite of purines. [Roblin et al. *J Am Chem Soc* **67** 290 1945, DOI: 10.1021/ja01218a043; for spectra see Cavalieri et al. *J Am Chem Soc* **70** 3857 1948, DOI: 10.1021/ja01191a102; *Beilstein* **26** III/IV 4171.]

7-Azaindole (1*H*-pyrrolo[2,3*b*]pyridine) [271-63-6] $C_7H_6N_2$, **M 118.1**, **m 105-106°**, **pK²⁰ 4.57**. Recrystallise it repeatedly from EtOH, then sublime it in a vacuum [Tokumura et al. *J Am Chem Soc* **109** 1346 1987, DOI: 10.1021/ja00239a010]. It is used as a pharmaceutical synthetic building block [Wang et al. *J Org Chem* **71** 4021 2006, DOI: 10.1021/jo0602571]. The *N*-acetate has **m 65-66°** (from *C₆H₆), and the *picrate* has **m 232-233°** (from Me₂CO) [Clemo & Swan *J Chem Soc* 603 1945, DOI: 10.1039/JR9450000603; *Beilstein* **23** III/IV 1105.]

1-Azaindolizine (1,7a-diazaindene, imidazo[1,2-*a*]pyridine) [274-76-0] $C_7H_6N_2$, **M 118.1**, **b 72-73°/1mm**, **103°/1mm**, **d₄²⁵ 1.165**, **n_D²⁰ 1.626**, **pK²⁰ 1.43**. (in aqueous HCl). 1-Azaindolizine is purified by distillation or gas chromatography. [Bower & Ramage *J Chem Soc* 4506 1957, DOI: 10.1039/JR9570004506; Armarego *J Chem Soc* 4226 1964, DOI: 10.1039/JR9640004226; *Beilstein* **23** II 1554, **23** III/IV 1104.]

8-Azapurine (1*H*-1,2,3-triazolo[4,5-*d*]pyrimidine, 1,2,3,4,6[3*H*]penta-azaindene) [273-40-5] $C_4H_3N_5$, **M 121.1**, **m 174-175°** (effervescence, **m** depends on heating rate), **pK₁²⁰ 2.05** (equilib with covalent hydrate), **pK₂²⁰ 4.84**. Sublime 8-azapurine at 120-130°/0.01mm and recrystallise it from 3 parts of EtOH. [Albert *J Chem Soc(B)* 427 1966, DOI: 10.1039/J29660000427; *Beilstein* **26** III/IV 4108.]

Azetidine (trimethyleneimine) [503-29-7] C_3H_7N , **M 57.1**, **b 19°/132.5mm**, **61.3-61.5°/760mm**, **d₄²⁰ 0.846**, **n 1.432**, **pK²⁵ 11.29**. Azetidine is a flammable, hygroscopic liquid smelling of ammonia, which absorbs CO₂ from air and should be kept under Argon. Purify it by drying it over solid KOH and distilling it through a short Vigreux column (p 11) at atmospheric pressure (under Argon) and keeping the pot temperature below 210°. It is moisture sensitive. The *hydrochloride* [36520-39-5] **M 93.6** has **m > 300°** and the *hydroiodide* has **m 146.5°** (from EtOH). The *N*-Me derivative has **m 112°** (from *C₆H₆/petroleum ether), and the *N*-phenylcarbonyl derivative has **m 189-190°** (from EtOH). [Searles et al. *J Am Chem Soc* **78** 4917 1956, DOI: 10.1021/ja01600a029; *Beilstein* **20** H 2, **20** I 3, **20** II 3, **20** III/IV 53, **20**/I V 136.]

Aziridine (ethyleneimine) [151-56-4] C_2H_5N , **M 43.1**, **b 55-56°/756mm**, **56°/760mm**, **d₄²⁴ 0.8321**, **pK²⁵ 8.00**. Redistil it in an argon or N₂ atmosphere in a fume hood, and store it over KOH in sealed bottles in a refrigerator. Commercial aziridine has been dried over sodium and distilled from the metal through an efficient column before use [Jackson & Edwards *J Am Chem Soc* **83** 355 1961, DOI: 10.1021/ja01463a023; Wenker *J Am Chem Soc* **57** 2328 1935, DOI: 10.1021/ja01314a504]. It is a weaker base than Me₂NH (pK²⁵ 10.87) but is caustic to the skin. It should not be inhaled, causes inflammation of the eyes, nose and throat, and one may become sensitised to it. It is soluble in H₂O, has an ammoniacal smell and reacts with CO₂. Pure aziridine is comparatively stable but polymerises in the presence of traces of H₂O and is occasionally explosive in the presence of acids. CO₂ is sufficiently acidic to cause polymerisation (forms linear polymers) which is not free radical promoted. It is stable in the presence of bases. The violet 2:1 **Cu complex** crystallises from EtOH containing a few drops of aziridine and adding Et₂O, and has **m 142°(dec)**. The *picrate* has **m 142°**. [O'Rourke et al. *J Am Chem Soc* **78** 2159 1956, DOI: 10.1021/ja01591a035.] It has also been dried over BaO and has been distilled from sodium under nitrogen. [Allen et al. *Org Synth Coll Vol* **4** 433 1963, DOI: 10.15227/orgsyn.030.0038; *Beilstein* **20** III/IV 1.] **TOXIC**.

Azuleno(1,2-b)thiophene [25043-00-9] $C_{12}H_8S$, **M 184.2**. It is crystallised from cyclohexane, then sublimed *in vacuo*. **Azuleno(2,1-b)thiophene** [248-13-5] is recrystallised from cyclohexane, then sublimed *in vacuo*.

Azure A (3-amino-7-dimethylaminophenazin-5-ium chloride) [531-53-3] $C_{14}H_{14}ClN_3S$, **M 291.8**, **CI 52005**, **m > 290°(dec)**, λ_{max} 633nm, **pK²⁵ 7.2**. This biological stain, Azure A, has been twice recrystallised from H₂O and dried at 100°/1 hour in an oven. The green crystals give a blue aqueous solution. [Beilstein 27 III/IV 5151.]

Azure B (3-methylamino-7-dimethylaminophenazin-5-ium chloride) [531-55-5] $C_{15}H_{16}ClN_3S$, **M 305.8**, **CI 52010**, **has m > 201°(dec), 205-210°(dec)**, λ_{max} 648nm, **pK²⁵ 7.4**. This biological stain, Azure B, has been twice recrystallised from H₂O and dried at 100°/1 hour in an oven. The green crystals give a blue aqueous solution. [Beilstein 27 III/IV 5151.]

Azure C (3-amino-7-methylaminophenazin-5-ium chloride) [531-57-7] $C_{13}H_{12}ClN_3S$, **M 277.8**, **CI 52002**, **has λ_{max} 616nm, pK²⁵ 7.0**. This biological stain, Azure C, has been twice recrystallised from H₂O, and dried at 100°/1 hour in an oven. The green crystals give a blue aqueous solution. [Beilstein 27 III/IV 5151.]

Barbituric acid (6-hydroxypyrimidin-2,4-dione) [67-52-7] $C_4H_4N_2O_3$, **M 128.1**, **m 248-250°(dec), 250°(dec)**, **pK₁²⁵ 3.99, pK₂²⁵ 12.5**. Recrystallise it twice from H₂O, then dry it for 2 days at 100°. [Beilstein 24 III/IV 1873.] It is used as a buffer reagent.

Benzimidazole [51-17-2] $C_7H_6N_2$, **M 118.1**, **m 169-171°, 170-172°, 172-173°**, **pK₁²⁵ 5.53, pK₂²⁵ 11.70**. It crystallises from boiling water (1g/15ml) or aqueous EtOH (charcoal) and is dried at 100° for 12 hours. [Wagner & Millett *Org Synth Coll Vol* 2 65 1943, DOI: 10.15227/orgsyn.019.0012; Beilstein 23 H 131, 23/6 V 196.]

2-Benzimidazolylacetoneitrile [4414-88-4] $C_9H_7N_3$, **M 157.2**, **m 200-205°(dec), 209.7-210.7°(corrected), 210°**. It is recrystallised from aqueous EtOH. It has also been recrystallised from hot H₂O using charcoal, and finally from aqueous EtOH. [Copeland & Day *J Am Chem Soc* 65 1072 1943, DOI: 10.1021/ja01246a019; Beilstein 25 III/IV 820.]

Benzo-15-crown-5 [14098-44-3] $C_{14}H_{20}O_5$, **M 268.3**, **m 78-80°**. It is recrystallised from *n*-heptane. **IRRITANT**. [Vögtle ed, Host Guest Complex Chemistry in *Topics in Current Chemistry* 98 1981, DOI: 10.1007/BFb0111244; Beilstein 19/10 V 618.]

Benzo-18-crown-6 [14098-24-9] $C_{16}H_{24}O_6$, **M 312.2**, **has m 42-45°, 43-43.5°**. Purify it by passage through a DEAE cellulose column in cyclohexane. It recrystallises from *n*-hexane. Its **thiourea complex** has **m 127°** [5-6 mol of urea to ether, Pedersen *J Org Chem* 36 1690 1971, DOI: 10.1021/jo00811a027]. The stability constants of the Na⁺, K⁺, Rb⁺, Cs⁺, Tl⁺ and Ba²⁺ complexes are described in Hofmanova et al. *Inorg Chim Acta* 28 73 1978, DOI:10.1016/S0020-1693(00)87416-2. [NMR: Live & Chan *J Am Chem Soc* 98 3769 1976, DOI: 10.1021/ja00429a006]. [Beilstein 19/12 V 618.] **IRRITANT**.

Benzo[3,4]cyclobuta[1,2-b]quinoxaline [259-57-4] $C_{14}H_8N_2$, **M 204.2**, **m dec >250°**. It is purified by sublimation under reduced pressure. For the He(I) photoelectron spectra see Yamaguchi & Baumann *Spectrochim Acta Part A: Molecular Spectroscopy* 43 683 1987, DOI: 10.1016/0584-8539(87)80151-4.

Benzofuran (coumarone) [271-89-6] C_8H_6O , **M 118.1**, **b 62-63°/15mm, 97.5-99.0°/80mm, 169°/760mm, 170-173°/atm, 173-175°/760mm, d₄²⁰ 1.0945, n_D²⁰ 1.565**. Benzofuran is steam distilled, dissolved in Et₂O, washed with 5% aqueous NaOH, saturated NaCl, dried (Na₂SO₄), evaporated and redistilled. The UV has λ_{max} at 245, 275, 282nm (log ϵ 4.08, 3.45, 3.48). The **picrate** has **m 102-103°**. [Burgstahler & Worden *Org Synth Coll Vol* 5 251 1973, DOI: 10.15227/orgsyn.046.0028; NMR: Black & Heffernan *Aust J Chem* 18 353 1965, DOI: 10.1071/CH9650353; Beilstein 17/2 V 3.]

2-Benzofurancarboxylic acid (Coumanilic acid) [496-41-3] $C_9H_6O_3$, **M 162.1**, **m 192-193°, 193-196°, b 310-315°/atm with some dec, pK_{Est} ~3.2**. Purified by reprecipitation from alkaline solution with acid, and crystallised from hot water. [Fuson et al. *Org Synth Coll Vol* 3 209 1955, DOI: 10.15227/orgsyn.024.0033; Beilstein 18/6 V 419.]

Benzofurazan (benz[1,2,5]oxadiazole) [273-09-6] $C_6H_4N_2O$, M 120.1, m 47-51°, 55°, 75-78°/20mm. Purify benzofurazan by steam distillation from a dilute alkaline solution, crystallisation from EtOH (long white needles) and by sublimation. [Green & Rowe *J Chem Soc* **101** 2452 1912, DOI: 10.1039/CT9120102452; Green & Rowe *J Chem Soc* **111** 612 1917, DOI: 10.1039/CT9171100612; Ghosh et al. *J Med Chem* **15** 255 1972, DOI: 10.1021/jm00273a012; *Beilstein* **27** H 568, **27** I 573, **27** III/IV 7115.]

Benzofuroxan (benzofurazan-1-oxide, benz[1,2,5]oxadiazol-1-oxide) [480-96-6] $C_6H_4N_2O_2$, M 136.1, m 70-71°, 72-73°. Purify the 1-oxide by dissolving 3.6g in 45ml of 95% EtOH + 15ml H_2O , boil, filter hot and cool to 25°. The yellow crystalline solid is steam volatile (but less so than benzofurazan, above) and has a peculiar pungent odour. Its UV has λ_{max} at 355nm (EtOH). It was used as a dehydrogenation oxidant (Pätzold et al. *Synth Commun* **22** 281 1992, DOI:10.1080/00397919208021304). [Mallory *Org Synth Coll Vol* **4** 74 1963, DOI: 10.15227/orgsyn.037.0001; Boulton & Ghosh *Adv Heterocyclic Chem* **10** 1 1969, DOI:10.1016/S0065-2725(08)60494-8; Boyer et al. *J Am Chem Soc* **79** 1748 1977, DOI: 10.1021/ja01564a059; *Beilstein* **27** I 622, **27** II 629, **27** III/IV 7115.]

5,6-Benzoquinoline (benzo[f]quinoline) [85-02-9] $C_{13}H_9N$, M 179.2, m 93°, 94°, b 350°/atm, pK^{20} 5.11. Purify as for phenanthridine (benzo[c]quinoline) below. It forms an insoluble complex $[(C_{13}H_9N)_2 \cdot H_2CdI_4]$ with Cd in the presence of KI and mineral acids. The *picrate* has m 258.1-259° (from EtOH or H_2O). [Albert et al. *J Chem Soc* 2240 1948, DOI: 10.1039/JR9480002240; *Beilstein* **20** III/IV 4009, **20**/8 V 220.]

7,8-Benzoquinoline (benzo[h]quinoline) [230-27-3] has m 48-50°, 52.0-52.5°, b 238°/719mm, pK^{20} 4.21. Purify it as for phenanthridine (benzo[c]quinoline) below. The *picrate* has m 196° (from Me_2CO). [*Beilstein* **20** H 463, **20** III/IV 4003, **20**/8 V 215.]

1,2,3-Benzothiadiazole [273-77-8] $C_6H_4N_2S$, M 136.2, m 35°, 36-37°, b 63°/0.5mm, $pK_{Est} \sim <0$. 1,2,3-Benzothiadiazole crystallises from petroleum ether, and has λ_{max} at 264 and 306nm in hexane. [Overberger et al. *J Org Chem* **24** 1407 1959, DOI: 10.1021/jo01092a004; *Beilstein* **27** III/IV 7113.]

2,1,3-(1,2,5)-Benzothiadiazole [272-13-2] has m 42-44°, 44°, b 206°/760mm, $pK_{Est} <0$. 2,1,3-Benzothiadiazole crystallises from petroleum ether and has UV with λ_{max} at 221-222, 304 and 330nm in EtOH. [Sawicki & Carr *J Org Chem* **22** 503 1959, DOI: 10.1021/jo01356a007; *Beilstein* **27** III/IV 7118.]

1-Benzothiophene (benzo[b]thiophene, thianaphthene) [95-15-8] C_8H_6S , M 134.2, m 29-32°, 30°, 31-33°, 31-32°, 32°, b 100°/16mm, 103-105°/20mm, 221-222°/760mm, $d_4^{32.2}$ 1.1484, n_D^{39} 1.6306. 1-Benzothiophene has the odour of naphthalene. If the IR spectrum is not very good, then suspend it in a faintly alkaline aqueous solution and steam distil it. Extract the distillate with Et_2O , dry the extract ($CaCl_2$), filter, evaporate the solvent and fractionate the residue. The distillate sets solid. The *sulfoxide* has m 142°, the *picrate* has m 148-149° (yellow crystals from EtOH) and the *styphnate* has m 136-137°. [Hansch & Lindwall *J Org Chem* **10**, 381 1945, DOI: 10.1021/jo01181a001; Meyer & Meyer *Chem Ber* **52B** 1249 1919, DOI: 10.1002/cber.19190520702; Weisgerber & Kruber *Chem Ber* **53** 1551 1920, DOI: 10.1002/cber.19200530836; Iddon & Scrowston *Adv Heterocycl Chem* **11** 177 1970, DOI: 10.1016/S0065-2725(08)60776-X; *Beilstein* **17**/2 V 6.]

1,2,3-Benzotriazole [95-14-7] $C_6H_5N_3$, M 119.1, m 96-97°, 97-99°, 98.5°, 100°, b 159°/0.2mm, 204°/15mm, pK_1^{20} 1.6, pK_2^{20} 8.64. 1,2,3-Benzotriazole crystallises from toluene, $CHCl_3$, Me_2NCHO or a saturated aqueous solution, and is dried at room temperature or in a vacuum oven at 65°. Losses are less if the material is distilled in a vacuum. **CAUTION: may EXPLODE during distillation; necessary precautions must be taken.** [Damschroder & Peterson *Org Synth Coll Vol* **3** 106 1955, DOI: 10.15227/orgsyn.020.0016; *Beilstein* **26** III/IV 93.]

O-Benzotriazol-1-yl-N,N,N',N'-tetramethyluronium hexafluorophosphate (HBTU) [94790-37-1] $C_{11}H_{16}F_6N_5OP$, M 379.2, m 200° (dec), 250°, 254°(dec). Wash the salt with H_2O (3x), CH_2Cl_2 (3x), dry and recrystallise it from MeCN. Dry it in a vacuum and store it cold in the dark [Dourtoglou et al. *Tetrahedron Lett* 1269 1978, DOI: 10.1016/0040-4039(78)80103-8; NMR: Dourtoglou et al. *Synthesis* 572 1984, DOI: 10.1055/s-1984-30895]. Useful coupling reagent for peptide synthesis [Carpino et al. *Angew Chem Int Ed* **41** 441 2002, DOI: 10.1002/1521-3773(20020201)41:3<441::AID-ANIE441>3.0.CO;2-N; Knorr et al. *Tetrahedron Lett* **30** 1927 1989, DOI: 10.1016/S0040-4039(00)99616-3.]

Benzoxazolinone (2-hydroxybenzoxazole) [59-49-4] $C_7H_5NO_2$, M 135.1, m 137-139°, 142-143°(corrected), b 121-213°/17mm, 335-337°/760mm. Benzoxazolinone is purified by recrystallisation from aqueous Me_2CO followed by distillation at atmospheric pressure, then in a vacuum. The *methyl mercury salt* recrystallises from aqueous EtOH and has m 156-158°. [Bywater et al. *J Am Chem Soc* **67** 905 1945, DOI: 10.1021/ja01222a008; *Beilstein* **27** III/IV 2677.]

S-(+) and R-(-) 1-Benzoyl-2-tert-butyl-3-methyl-4-imidazolinone [R- 101055-57-6] [S-101055-56-5] M 260.3, m 142-143°, 145.6-146.6°, 145-147°, $[\alpha]_{546}^{20}$ (+) and (-) 155, $[\alpha]_D^{20}$ (+) and (-) 133 (c 1, $CHCl_3$). Recrystallise these chiral imidazolinones from boiling EtOH (solubility is 1.43g/ml) or better by dissolving in CH_2Cl_2 and adding pentane, filtering and drying for at least 12 hours at 60°/0.1mm and sublime them at 135°/0.01mm. They have also been purified by flash column chromatography through Merck silica gel at 0.04-0.063mm and using Et_2O /petroleum ether/MeOH (60:35:5) as eluent. They are then recrystallised from EtOH/petroleum ether. [IR, NMR: Seebach et al. *Helv Chim Acta* **70** 237 1987, DOI: 10.1002/hlca.19870700129; Fitzi & Seebach *Angew Chem Int Ed* **25** 345 1986, DOI: 10.1002/anie.198603451.] The *racemate* is purified in a similar manner and has m 104-105° [NMR: Seebach et al. *Helv Chim Acta* **68** 949 1985, DOI: 10.1002/hlca.19850680418].

2-Benzoylpyridine [91-02-1] $C_{12}H_9NO$, M 183.2, m 41-43°, 48-50°, 72°/0.02mm, 104-105°/0.01mm, 317°/atm, n_D^{24} 1.6032, $pK_{Est} \sim 2.4$. Dissolve 2-benzoylpyridine in Et_2O , shake it with aqueous $NaHCO_3$, H_2O , dry it over $MgSO_4$ and evaporate. The residue solidifies on cooling. The solid can be recrystallised from petroleum ether. Its *hydrochloride* crystallises from Me_2CO , m 126-127°, and the **2,4-dinitrophenylhydrazone** has m 193-195°. It distils at high vacuum. [Pinkerton & Thames *J Organomet Chem* **24** 623 1970, DOI:10.1016/S0022-328X(00)84491-5; *Beilstein* **21/8** V 566.]

N⁶-Benzyladenine [1214-39-7] $C_{12}H_{11}N_5$, M 225.3, m 230-233°, 231-232°, 232.5°(dec), $pK_{Est(1)} \sim 4.2$, $pK_{Est(2)} \sim 10.1$. It is purified by recrystallisation from aqueous EtOH. It has λ_{max} at 207 and 270nm (H_2O), 268 nm (pH 6), 274nm (0.1 N HCl) and 275nm (0.1 N NaOH). [Daly & Christensen *J Org Chem* **21** 1553 1956, DOI: 10.1021/jo01118a630; Bullock et al. *J Am Chem Soc* **78** 3693 1956, DOI: 10.1021/ja01596a037; *Beilstein* **26** III/IV 3575.]

1-Benzyl-1-aza-12-crown-4 (10-benzyl-1,4,7-trioxa-10-azacyclododecane) [84227-47-4] $C_{15}H_{23}NO_3$, M 265.4, 122-125°/0.03mm, 140-143°/0.05mm, d_4^{20} 1.09, n_D^{20} 1.52, $pK_{Est} \sim 7.7$. Dissolve it in CH_2Cl_2 or CCl_4 (1g in 30ml), wash it with H_2O (30ml), brine (30ml), H_2O (30 ml) again, dry ($MgSO_4$ or Na_2SO_4), and evaporate. The residue in CH_2Cl_2 is chromatographed through Al_2O_3 (eluting with 10% EtOAc in hexane); evaporate, collect the correct fractions and distil (Kügelrohr) them. Log K_{Na} in dry MeOH at 25° for Na^+ complex is 2.08. Also a good complexing agent for Lithium ions. [White et al. *Tetrahedron Lett* **26** 151 1985, DOI:10.1016/S0040-4039(00)61866-X; Arnold et al. *J Org Chem* **53** 5652 1988, DOI: 10.1021/jo00259a008.]

2-Benzyl-1,3-dioxolane [101-49-5] M 164.2, b 98-99°/1mm, 110°/5mm, 137-138°/34mm, 240-242°/atm, d_4^{20} 1.087, n_D^{20} 1.532. Dissolve 2-benzyl-1,3-dioxolane in CH_2Cl_2 , wash well with 1M NaOH, dry over K_2CO_3 , filter, evaporate and distil it through a short path still (Kügelrohr). It has also been purified by preparative gas chromatography. [Raber & Guida *Synthesis* 808 1974, DOI: 10.1055/s-1974-23438; Lloyd & Luberoft *J Org Chem* **34** 3949 1969, DOI: 10.1021/jo01264a043; *Beilstein* **19** III/IV 220.]

S-(+)- and R-(-)- Benzyl glycidyl ether (1-benzyloxyoxirane) [S:14618-80-5] [R:16495-13-9] $C_{10}H_{12}O_2$, M 164.2, b 68°/10⁻⁴ mm, 105°/0.4mm, d_4^{20} 1.072, n_D^{20} 1.517, $[\alpha]_{546}^{20}$ (+) and (-) 5.5, $[\alpha]_D^{20}$ (+) and (-) 5.1 (c 5, toluene), $[\alpha]_D^{20}$ (+) and (-) 1.79 (c 5.02, $CHCl_3$), $[\alpha]_D^{21}$ (+) and (-) 15.3 (neat). This ether in EtOAc is dried (Na_2SO_4), then purified by flash chromatography using petroleum ether/EtOAc (5:1) as eluent. The ether distils through a short path distillation apparatus (Kügelrohr) as a colourless liquid. Alternatively, dissolve it in $CHCl_3$, wash it with H_2O , dry (Na_2SO_4), evaporate and purify by silica gel chromatography. [Anisuzzamen & Owen *J Chem Soc (C)* 1021 1967, DOI: 10.1039/J39670001021; Takano et al. *Heterocycles* **16** 381 1981, DOI: 10.3987/R-1981-03-0381; Lipshutz et al. *Org Synth* **69** 80 1990, DOI: 10.15227/orgsyn.069.0080; Takano et al. *Synthesis* 539 1989, DOI: 10.1055/s-1989-27310; Honda et al. *Chem Pharm Bull Jpn* **39** 1385 1991, DOI: org/10.1248/cpb.39.1385; *Beilstein* **12** IV 2277.] The *racemate* [RS 2930-05-4] has b 70-73°/11mm, d_4^{25} 1.077, n_D^{20} 1.5170, and is purified in the same manner.

3-Benzyl-5-(2-hydroxyethyl)-4-methylthiazolinium chloride [4568-71-2] $C_{13}H_{16}ClNOS$, M 269.8, m 142-144°, 145-147°. Purify the chloride by recrystallisation from EtOH or H_2O . If placed in a bath at 125° and heated at 2°/minute, the melting point is 140.5-141.4°. [Livermore & Sealock *J Biol Chem* **167** 699 1947, <http://www.jbc.org/content/254/11/4856>; Maier & Metzler *J Am Chem Soc* **79** 4386 1957, DOI: 10.1021/ja01573a040; *Beilstein* **27** III/IV 1758.]

5-Benzyloxyindole [1215-59-4] $C_{15}H_{13}NO$, M 223.3, m 96-97°, 100-103°, 100-104°, 104-106°, $pK^{25} < 0$. It is recrystallised from $*C_6H_6$ /petroleum ether or petroleum ether. The *picrate* forms red crystals from $*C_6H_6$ and has m 142-143°. [Burton & Leong *Chem Ind (London)* 1035 1953, Ek & Witkop *J Am Chem Soc* **76** 5579 1954, DOI: 10.1021/ja01651a001; fluorescence: Bridges & Williams *Biochem J* **107** 225 1968, DOI: 10.1042/bj1070225; *Beilstein* **27** III/IV 1758, **21/3** V 19]

1-Benzyl-4-piperidone [3612-20-2] $C_{12}H_{15}NO$, M 189.3, b 107-108°/0.2mm, 114-116°/0.3mm, 143-146°/5mm, 157-158°/11mm, d_4^{20} 1.059, n_D^{20} 1.538. If the physical properties show contamination, then dissolve it in the minimum volume of H_2O , made strongly alkaline with aqueous KOH, extract it with toluene several times, dry the extract with K_2CO_3 , filter, evaporate and distil the residue at high vacuum using a bath temperature of 160-190°, and redistil it. [Brookes & Walker *J Chem Soc* 3173 1957, DOI: 10.1039/JR9570003173; Bolyard *J Am Chem Soc* **52** 1030 1930, DOI: 10.1021/ja01366a029.] The *hydrochloride* has m 159-161° (from Me_2CO/Et_2O), and the *picrate* has m 174-182° (from Me_2CO/Et_2O). [Grob & Brenneisen *Helv Chim Acta* **41** 1184 1958, DOI: 10.1002/hlca.19580410503; *Beilstein* **21/6** V 424.]

2-Benzylpyridine [101-82-6] $C_{12}H_{11}N$, M 169.2, m 8-10°, b 98.5°/4mm, 276°/atm, d_4^{20} 1.054, n_D^{26} 1.5771, pK^{25} 5.13. Dry it with NaOH for several days, then distil it from CaO under reduced pressure, and redistil the middle fraction. [*Beilstein* **20/7** V 556.]

4-Benzylpyridine [2116-65-6] $C_{12}H_{11}N$, M 169.2, b 110.0°/6mm, 287°/atm, d_4^{20} 1.065, n_D^{26} 1.5814, pK^{25} 5.59. Dry it with NaOH for several days, then distil it from CaO under reduced pressure, and redistil the middle fraction. [*Beilstein* **20/7** V 561.]

Berberamine [478-61-5] $C_{37}H_{40}N_2O_6$, M 608.7, m 197-210°, $[\alpha]_D^{20} +115$ ($CHCl_3$), pK^{20} 7.33. (70% aqueous EtOH) Crystallise berberamine from petroleum ether or $*C_6H_6$ (with 1 $*C_6H_6$ m 129-134°). It also crystallises as the *monohydrate* from aqueous EtOH which sinters at ~147°. [Bick et al. *Aust J Chem* **9** 111, 118 1956, DOI: 10.1071/CH9560111; *Beilstein* **27** II 891, **27** III/IV 8732.]

Berberine [2086-83-1] $C_{20}H_{18}NO_4$, M 336.4, m 145°, 147-148°, pK_1^{20} 2.47, pK_2^{20} 11.73 (pseudobase?). Berberine crystallises from petroleum ether or ether as yellow needles or from H_2O . It is an antimalarial and is antibacterial as well. [*Beilstein* **27** II 567, **27** III/IV 6539.]

Berberine chloride (2H₂O) (Neutral Yellow 18) [633-65-8 (anhydrous), 5956-60-5 (2H₂O)] $C_{20}H_{18}NO_4.Cl$, M 407.9, has m 204-206°(dec), CI 75160, pK^{20} 2.47. Berberine chloride crystallises from water to give the *dihydrate*. The anhydrous salt may be obtained by recrystallisation from EtOH/ Et_2O , wash the crystals with Et_2O and dry them in a vacuum. The *iodide* has m 250°(dec) (from EtOH). [Perkin *J Chem Soc* **113** 492 1918, DOI: 10.1039/CT9181300492; Kametani et al. *J Chem Soc(C)* 2036 1969, DOI: 10.1039/J39690002036; *Beilstein* **27** I 515, **27** II 567.]

Bilirubin [635-65-4] $C_{33}H_{36}N_4O_6$, M 584.7, m >360°, ϵ_{450nm} 55,600 in $CHCl_3$, pK_{Est} ~3.0. An acyclic tetrapyrrole bile pigment with impurities which can be eliminated by successive Soxhlet extraction with diethyl ether and MeOH. It crystallises from $CHCl_3$ as deep red-brown rhombs, plates or orange-red prisms from chlorobenzene (m 330° dec) and is dried to constant weight at 80° under vacuum. [Gray et al. *J Chem Soc* 2264, 2276 1961, DOI: 10.1039/JR9610002264; *Beilstein* **26** III/IV 3268.]

Biliverdine [114-25-0] $C_{33}H_{34}N_4O_6$, M 582.6, m >300°, pK^{25} 3.0. This is the precursor of bilirubin (above) and forms dark green plates or prisms, with a violet reflection, from MeOH. The *dimethyl ester*, when crystallised from MeOH has m 215°(209°), and when crystallised from $CHCl_3$ /petroleum ether gives blue-green crystals with m 202°. [Gray et al. *J Chem Soc* 2264 1961, DOI: 10.1039/JR9610002264; Sheldrick *JCS Perkin Trans 2* 1457 1976, DOI: 10.1039/P29760001457; *Beilstein* **26** III/V 3272.]

2-(4-Biphenyl)-5-phenyl-1,3,4-oxadiazole (BPD) [852-38-0] $C_{20}H_{14}N_2O$, M 298.4, m 166-167°, 167-169°, 167-170°. BPD is recrystallised from toluene. It is a good scintillation material and is suitable as a laser dye. [Brown et al. *Discussion Faraday Soc* 27 43 1959, DOI: 10.1039/DF9592700043]. [Beilstein 27 III/IV 7283.]

2,2'-Bipyridyl [366-18-7] $C_{10}H_8N_2$, M 156.2, m 70-73°, 70.5°, b 273°/atm, pK_1^{25} -0.52, pK_2^{25} 4.44. 2,2'-Bipyridyl crystallises from hexane, or EtOH, or (after charcoal treatment of a $CHCl_3$ solution) from petroleum ether. Also, it precipitates from a concentrated solution in EtOH by addition of H_2O . Dry it in a vacuum over P_2O_5 . It can be further purified by chromatography on Al_2O_3 or by sublimation. Its UV (EtOH) has λ_{max} at 280nm (log ϵ 4.13). It is a metalloprotease inhibitor and high affinity chelator of iron and other transition metal ions. [Airoldi et al. *JCS Dalton Trans* 1913 1986, DOI: 10.1039/DT9860001913; Beilstein 23/8 V 16.]

4,4'-Bipyridyl [553-26-4] has m 73°(hydrate) [553-26-4, 123333-55-1], 109-112°, 114° (anhydrous), b 305°/760mm, 293°/743mm, pK_1^{20} 3.17, pK_2^{20} 4.82. It crystallises from water, *benzene/petroleum ether, ethyl acetate and sublimes *in vacuo* at 70°. Also purify it by dissolving in 0.1M H_2SO_4 and twice precipitating by addition of 1M NaOH to pH 8. Then recrystallise it from EtOH. For the *dihydrochloride* see Viologen below. [Collman et al. *J Am Chem Soc* 109 4606 1987, DOI: 10.1021/ja00249a025; Beilstein 23 H 800, 23 III/IV 1371, 23/8 V 28.]

2,2'-Biquinolin-4,4'-dicarboxylic (2,2'-bicinchoninic) acid [1245-13-2] $C_{20}H_{12}N_2O_4$, M 344.3, m 367°, $pK_{Est(1)}$ ~1.5, $pK_{Est(2)}$ ~4.0. Dissolve the acid in dilute NaOH and precipitate it with acetic acid, filter, wash swell with H_2O and dry it at 100° in a vacuum oven. Attempts to form a picrate failed. The *methyl ester* ($SOCl_2$ -MeOH) has m 165.6-166°. [Lesesne & Henze *J Am Chem Soc* 64 1897 1942, DOI: 10.1021/ja01260a041; Brown et al. *J Am Chem Soc* 68 2705 1946, DOI: 10.1021/ja01216a087; Beilstein 25 III/IV 1148.] For the di-K salt see entry in 'Metal-organic Compounds' in Chapter 4.

2,2'-Biquinolyl (Cuproin, α,α' -diquinolyl) [119-91-5] $C_{18}H_{12}N_2$, M 256.3, m 192-195°, 196°, pK_{Est} ~4.2. Decolourise 2,2'-biquinolyl in $CHCl_3$ solution (charcoal), then crystallise it to constant melting point from EtOH or petroleum ether [Cumper et al. *J Chem Soc* 1188 1962, DOI: 10.1039/JR9620001188]. [Beilstein 23/10 V 8.]

2,5-Bis(4-aminophenyl)-1,3,4-oxadiazole (BAO) [2425-95-8] $C_{14}H_{12}N_4O$, M 252.3, m 252-255°, 254-255°. BAO is recrystallised from EtOH using charcoal and under N_2 to avoid oxidation. It is a fluorescent stain for DNA [Yataghanas et al. *Exptl Cell Res* 56 59 1969, DOI: 10.1016/0014-4827(69)90394-2]. [Beilstein 27 III/IV 8158.]

2,5-Bis(2-benzothiazolyl)hydroquinone [33450-09-8] $C_{20}H_{12}N_2O_2S_2$, M 376.2, m dec >200°. Purify the hydroquinone by repeated crystallisation from dimethylformamide followed by sublimation in vacuum [Ernsting et al. *J Phys Chem* 91 1404 1987, DOI: 10.1021/j100290a026].

2,5-Bis(4-biphenyl)-1,3,4-oxadiazole (BBOD) [2043-06-3] $C_{26}H_{18}N_2O$, M 374.5, m 229-230°, 235-238°. BBOD is recrystallised from heptane or toluene. It is a good scintillant. [Hayes et al. *J Am Chem Soc* 77 1850 1955, DOI: 10.1021/ja01612a041.]

3,3-Bis(chloromethyl)oxacyclobutane (3,3-bis-[chloromethyl]oxetane) [78-71-7] $C_5H_8Cl_2O$, M 155.0, m 18.9°, b 65°/5mm, 80°/10mm, 103°/30mm, 198°/atm, d_4^{20} 1.290, n_D^{20} 1.486. Shake it with aqueous $NaHCO_3$ or $FeSO_4$ to remove peroxides, separate, dry with anhydrous Na_2SO_4 , then distil it under reduced pressure from a little CaH_2 [Dainton et al. *Trans Faraday Soc* 56 1784 1960, DOI: 10.1039/TF9605601784; Farthing *J Chem Soc* 3648 1955, DOI: 10.1039/JR9550003648]. The 3,3-bis-(phenoxymethyl) derivative is described below. [Beilstein 17 III/IV 68.] **Lachrymatory.**

***N,N'*-Bis(nicotinic acid) hydrazide** [840-78-8] $C_{12}H_{10}N_4O_2$, M 242.2, m dec 200°, pK_{Est} ~3.3. The hydrazide crystallises from water; it is also soluble in hot EtOH but insoluble in * C_6H_6 , petroleum ether or $CHCl_3$. [Graf *J Prakt Chem* [2] 138 289 1933, DOI: 10.1002/prac.19331381101; Beilstein 22 III/IV 455.]

3,3-Bis(phenoxymethyl)oxacyclobutane (3,3-bis-[phenoxymethyl]oxetane) [1224-69-7] $C_{17}H_{18}O_3$, M 270.3, m 67.5-68°, b 143°/0.05mm. Distil it under high vacuum, then crystallise the solidified distillate from MeOH. [Farthing *J Chem Soc* 3648 1955, DOI: 10.1039/JR9550003648; Beilstein 17 III/IV 2010.]

Blue Tetrazolium (BTC, tetrazolium blue chloride) [1871-22-3] $C_{40}H_{32}Cl_2N_8O_2$, **M 727.7, m 254-255°(dec), 255°(dec)**. Crystallise the chloride from 95% EtOH/anhydrous diethyl ether to constant absorbance at 254nm. [Beilstein **26** III/IV 1789.]

1-N-Boc-piperidine (*N-tert-butoxycarbonylpiperidine, 1-piperidincarboxylic acid 1,1-dimethylethyl ester*) [75844-69-8] $C_{10}H_{19}NO_2$, **M 185.3, b 79°/0.05mm, d_4^{25} 0.964, n_D^{20} 1.454**. It is purified by bulb-to-bulb distillation at 65°/1mm. If it is discoloured then dissolve it in Et₂O, wash it with brine, separate, dry the Et₂O layer over K₂CO₃, filter, evaporate, and distil in a vacuum. Its IR has ν_{max} at 1695 (CO) cm⁻¹ (film). [IR, NMR, MS: Dieter & Li *J Org Chem* **62** 7726 1997, DOI: 10.1021/jo970985b; Beak & Lee *J Org Chem* **58** 1109 1993, DOI: 10.1021/jo00057a024].

1-N-Boc-2-piperidone (1-N-tert-butoxycarbonyl-2-piperidone) [85908-96-9] $C_{10}H_{17}NO_3$, **M 199.3, m 29-36°, 30-33°, 32-34°, 33-35°, 36°, b 110°/0.1mm**. This useful synthetic starting material is prepared by using the same principles as for the 4-oxo isomer below. Thus to a solution of γ -valerolactam (2-piperidone, 6.0g, 60.5mmol, [675-20-7]), DMAP (1.85g, 15.1mmol, 0.2 equiv, [1122-58-3] and Boc₂O (26.6g, 121.0mmol, 2.0 equiv, [24074-26-8]) in CH₂Cl₂ (70ml) at ~25° is added Et₃N (17.8g, 127.7mmol, 2.1 equiv) and is stirred for 32 hours, then quenched with 1.2N HCl (10ml). The aqueous layer is extracted with CH₂Cl₂, the combined CH₂Cl₂ solutions are washed with saturated aqueous NaHCO₃ (20ml), brine (30ml), dried (MgSO₄), filtered and evaporated *in vacuo*. The crude product is purified by flash chromatography through Kieselgel 60 (230-240 mesh with cyclohexane/EtOAc 8:2 elution) to give **1-N-Boc-2-piperidone** (9.71g, 48.7mmol, 81%) as a colourless low melting solid **m 36°**. It has R_F 0.40 (TLC on Kieselgel 60F₂₅₄, cyclohexane/EtOAc 7:3), and its IR (KBr) has ν_{max} at 1713, 1302, 1249, 1159 and 1138 cm⁻¹; the ¹H NMR (300MHz, CDCl₃, with residual CHCl₃ as internal standard at δ 7.27) has δ at 3.66 (m, 2H, H-6,6), 2.50 (m, 2H, H-3,3), 1.90-1.74 (4H, H-3,3,4,4) and 1.53 (s, 9H, *t*-Bu); the ¹³C NMR (75MHz, CDCl₃, with residual CDCl₃ as internal standard at δ 77.1) has δ at 171.1 (s), 152.4 (s), 82.5 (s), 46.0 (t), 34.6 (t), 27.7 (q), 22.5 (t) and 20.2 (t); and the EI MS has m/z (relative intensity) at 199 (M+ 0.3), 144 (52), 126 (21), 100 (38), 99 (31), 98 (31), 82 (24), 57 (100), 56 (38) and 55 (30). [Cossy et al. *New J Chem* **27** 475 2003.] This synthesis has also been carried out in MeCN/DMAP at ~25°/24 hours (74% yield) without further base [Moody & Taylor *JCS Perkin Trans 1* 721 1989, DOI: 10.1039/P19890000721].

1-N-Boc-2-piperidone is hydrolysed at ambient temperature by LiOH (30 minutes, 90%), or undergoes methanolysis by MeONa/MeOH (15 minutes, 94%) to provide γ -BocNH(CH₂)₄CO₂H (after acidification) or γ -BocNH(CH₂)₄CO₂Me without losing the Boc protecting group, respectively [Flynn et al. *J Org Chem* **48** 2424 1983, DOI: 10.1021/jo00162a028].

1-N-Boc-3-piperidone (1-N-tert-butoxycarbonyl-3-piperidone) [98977-36-7] has **m 35-40°, b 104-105°/0.4mm**. 3-Piperidone used to prepare the Boc derivative was obtained by hydrogenolysis of *N*-benzyl-3-piperidone hydrochloride hydrate (4.2g, 18.6mmol, [50606-58-1]) catalyzed by 10% Pd/C (0.8g) and H₂ at 55 psi while stirring for 16 hours in degassed MeOH (200ml). This was filtered (Celite) and evaporated *in vacuo*. The crude oily 3-piperidone was dissolved in THF (200ml), treated with Boc₂O (5.27g, 24.1mmol) and saturated aqueous Na₂CO₃ (50ml), stirred for 4 hours and evaporated *in vacuo*. The white solid was partitioned between EtOAc and 1N HCl, the organic layer was collected, washed with 1N NaOH and brine, dried (MgSO₄), filtered, and evaporated *in vacuo* to give an oil which was purified by flash chromatography (silica gel, hexane/EtOAc 3:1) to give **1-N-Boc-3-piperidone** (2.93g, ~100%) as a colourless oil. [Lucca et al. *J Med Chem* **48** 2194 2005, DOI: 10.1021/jm049530m.] Its ¹H NMR (300MHz, CDCl₃, TMS) has δ at 3.99 (s, 2H, 2,2-H), 3.58 (t, *J* = 6.3 Hz, 2H, 6,6-H), 2.46 (t, *J* = 6.3 Hz, 2H 4,4-H), 1.97 (t, *J* = 6.3 Hz, 2H, 5,5-H), and 1.45 (s, 9H, *t*-Bu); and the HRMS has m/z 200.1285 and calculated for C₁₀H₁₈NO₃ (M + H) was 200.1287. Alternatively, and almost similar preparation, was carried out except that saturated aqueous Na₂CO₃ was replaced by Et₃N (3 equiv) to give an 81% yield of **1-N-Boc-3-piperidone** which was distilled at high vacuum undecomposed. [Brehm et al. *J Med Chem* **29** 224 1986, DOI: 10.1021/jm00152a010.]

In a completely different synthesis, involving catalysed intramolecular cyclisation, **(5-oxo-6-dimethylsulfoxonium-hexyl)-carbamic acid tert-butyl ester** (250mg, 0.901mmol) in CH₂Cl₂ (8.0ml, purged with N₂ for 30 minutes) was added *via* a syringe pump over 18 hours to a solution of [Ir(COD)Cl]₂ (6.05mg, 9.01 μ mol, 0.01 equiv, [12112-67-3]) in degassed CH₂Cl₂ (4.0ml) at 70°. The mixture was then evaporated *in vacuo*, and the residue was purified through a silica gel column and eluting with hexanes/EtOAc (3:2) to afford **1-N-Boc-3-piperidone** (147mg, 82%) identical with the above. [Mangion et al. *Org Lett* **11** 3566 2009, DOI: 10.1021/ol901298p.]

1-*N*-Boc-4-piperidone (1-*N*-*tert*-butoxycarbonyl-4-piperidone) [79099-07-3] has **m 68-69°, 70-72°, 72°, 73-75°, 74-75°, 74.4-75.2°**. It is prepared by dissolving 4-piperidone (1g, 6.5mmol [41661-47-6]) and EtN(*iso*-Pr)₂ (2.8ml, 16mmol, DIPEA [7087-68-5]) in dioxane/H₂O (4:1, 10ml), adding Boc₂O (2.2g, 9.8mmol, [24074-26-8]) slowly with stirring at ~25° for 24 hours. The solvent is removed *in vacuo*, the residue is dissolved in CH₂Cl₂, washed with H₂O (3 x), dried (MgSO₄), filtered and evaporated *in vacuo*. The residue is triturated with Et₂O, the white residue is collected by filtration and recrystallised from hexane to give the **Boc-piperidone** (75%), **m 73.5°**. It also crystallises from petroleum ether, toluene or EtOAc and can be dried in a vacuum desiccator over shredded paraffin wax. It has R_F 0.74 (TLC on Kieselgel 60F₂₅₄, hexane/EtOAc 4:6), and its IR (KBr) has ν_{max} at 1715 and 1680 cm⁻¹. Its ¹H NMR (300MHz, CDCl₃) has δ at 3.72 (t, *J* = 6.3 Hz, 4H, 3,5-H), 2.44 (t, *J* = 6.3 Hz, 4H 2,6-H) and 1.50 (s, 9H, *t*-Bu); and the ¹H NMR (300MHz, DMSO-d₆) has δ at 3.6 (t, 4H, 3,5-H), 2.4 (t, 4H, 2,6-H) and 1.4 (s, 9H, *t*-Bu). It has been used extensively to introduce a piperidine ring for making drugs because the 4-oxo group can be functionalised in various ways and deprotection of the nitrogen allows bonding to this atom. [Houssin et al. *J Med Chem* **45** 533 2002, DOI: 10.1021/jm010297r; Ellis et al. *J Med Chem* **51** 2170 2008, DOI: 10.1021/jm701435h.]

Brazilin (6a*S*-cis-7,11b-dihydrobenzo[*b*]indeno[1,2-*d*]pyran-3,6a,9,10(6*H*)-tetraol) [474-07-7] C₁₆H₁₄O₅, **M 286.3, m 130°(dec), 250°, pK_{Est(1)} ~9.3, pK_{Est(2)} ~10.0, pK_{Est(3)} ~12.5 (all phenolic), CI 75280**. Brazilin crystallises from EtOH as yellow crystals which become orange when exposed to light and air, and is yellow in dilute acid but crimson in dilute alkali. When crystallised from H₂O, it has **m 247-248°**. It forms coloured metal salts and is oxidised in air to *Brazilein*, the quinonoid form. The (±)-form has been resolved, and the (+)-enantiomer has [α]_D²⁰ +121° (c 1, MeOH). [Craig et al. *J Org Chem* **30** 1573 1965, DOI: 10.1021/jo01016a058; Morsingh & Robinson *Tetrahedron* **26** 281 1970, DOI: 10.1016/0040-4020(70)85029-3; Beilstein **17** H 194, **17** II 244, **17** III/IV 2711.]

Brilliant Cresyl Blue (NN-diethyl-3-imino-8-methyl-3*H*-phenoxazin-7-amine hydrochloride) [4712-70-3] **M 332.8, pK₂₅ 3.2**. Crystallise the dye from petroleum ether. It has λ_{max} at 625nm (H₂O) and 622nm (95% aqueous EtOH). [Beilstein **27** H 400, **27** II 454, **27** III/IV 5160.]

5-Bromocytosine [2240-25-7] C₄H₄BrN₃O, **M 190.0, m 240-243°(dec), 245-255°(dec), 250°(dec), pK_I²⁵ 3.04, pK₂²⁵ 10.33**. 5-Bromocytosine is recrystallised from H₂O or 50% aqueous EtOH. Alternatively, dissolve ca 3g in conc HCl (10ml) and evaporate to dryness. Dissolve the residual hydrochloride in the minimum volume of warm H₂O and make faintly alkaline with aqueous NH₃. Collect the crystals and dry them in a vacuum at 100°. [Hilbert & Jansen *J Am Chem Soc* **56** 134 1934, DOI: 10.1021/ja01316a041; Beilstein **25** III/IV 3689.]

2-(2-Bromoethyl)-1,3-dioxane [33884-43-4] C₆H₁₁BrO₂, **M 195.1, b 67-70°/2.8mm, 71-72°/4mm, 95°/15mm, d₄²⁰ 1.440, n_D²⁰ 1.4219**. Purify it by vacuum fractionation. Also dissolve it in Et₂O, wash with aqueous NaHCO₃, dry the extract (Na₂SO₄), filter and fractionate at high vacuum. Its ¹H NMR in CCl₄ has δ at 1.3 (m, 1H), 2.1 (m, 3H), 3.36 (t, 2H), 3.90 (m, 4H) and 4.57 (t, H). [Stowell *J Org Chem* **41** 560 1976, DOI: 10.1021/jo00865a034; NMR, MS: Schwarz et al. *Tetrahedron* **35** 1969 1979, DOI: 10.1016/0040-4020(70)85029-3; Kriesel & Gisvold *J Pharm Sci* **60** 1250 1971, DOI: 10.1002/jps.2600600833; Beilstein **19/1** V 69; Fieser **7** 37, **11** 78.]

2-(2-Bromoethyl)-1,3-dioxolane [18742-02-4] C₅H₉BrO₂, **M 181.1, b 68-70°/8mm, 68-73°/10mm, 78-80°/20mm, d₄²⁰ 1.510, n_D²⁰ 1.479**. Dissolve it in pentane, wash with 5% aqueous NaHCO₃, dry (Na₂SO₄), and evaporate. Distil the residue. [NMR: Buechi & Wuest *J Org Chem* **34** 1122 1969, DOI: 10.1021/jo01256a076; Kriesel & Gisvold *J Pharm Sci* **60** 1250 1971, DOI: 10.1002/jps.2600600833; Beilstein **19/1** V 69; Fieser **7** 37.]

6-Bromo-5-fluoro-1*H*-indole [259860-08-7] C₈H₅BrFN, **M 214.0, m 82-84°, 83-84°, pK_{Est(1)} ~ -4.05 (basic), pK_{Est(2)} ~ 15.9 (acidic)**. There are several preparations of this indole. In a patent (to Vernallis Research Ltd, US6380238, 2002) *N,N*-dimethylformamide dimethylacetal (8.5ml, 60mmol) was added to a stirred solution of 4-bromo-5-fluoro-2-nitrotoluene (11.8g, 50mmol) in DMF (30ml) in one portion under argon at ~25°, then heated at 120° for 16 hours and evaporated *in vacuo*. The residual oil crystallised from MeOH/CH₂Cl₂ (4:1) to give a purple solid (4.5g) which was dissolved in MeOH/THF (1:1, 30ml), mixed with Raney Ni (1g), cooled to 0° and N₂H₄·H₂O (0.8ml, 16mmol) was added all at once. After stirring for 90min,

more $\text{N}_2\text{H}_4\cdot\text{H}_2\text{O}$ (0.8ml) was added and stirred at 0° for 30 minutes. The mixture was filtered through Celite and the filter cakes was washed with THF. The combined filtrates were evaporated *in vacuo*, and the residue was purified by chromatography [SiO_2 with heptane/ CH_2Cl_2 (4:1) as eluent] to give the **indole** (1.7g, 16%) as an off-white solid. In a second patent [to M.P. Dillon et al. US2004/224973, 2004, see also Batcho & Leimgruber *Org Synth* **63** 214 1985, DOI: 10.15227/orgsyn.063.0214] crude [2-(4-bromo-5-fluoro-2-nitrophenyl)vinyl]-dimethylamine and Raney Ni in THF gave after flash chromatography through a silica gel column (eluting with 10% EtOAc in hexane) which gave the indole as a light green solid (61%). However, when the same enamine (1.89g, 4.38mmol) in EtOH (35ml) was treated with a mixture of Fe (4.42g, 79mmol) in AcOH (35ml) and stirred at 90° overnight, then filtered, evaporated and the residue purified by chromatography as above, pure **6-bromo-5-fluoro-1H-indole** (0.83g, 89%) was obtained as a yellow solid [Patent to the Board of Trustees of the University of Illinois; WO2008/77138 2008]. It had IR (nujol) with ν_{max} at 3395, 2925, 2855, 1570, 1469, 1451, 1408, 1314, 1145, 1105, 865, 763 and 505 cm^{-1} ; and the ^1H NMR (400MHz, CDCl_3) had δ at 7.85 (br s, 1H), 7.55 (dd, 1H, $J = 5.6$ and 0.9 Hz), 7.34 (d, 1H, $J = 9.0\text{ Hz}$), 7.23 (t, 1H, $J = 2.8\text{ Hz}$) and 6.45-6.51 (m, 1H).

6-Bromo-5-fluoro-1-methylindole was obtained as a colourless solid in 99% yield by converting the indole above to its *N-sodio- derivative* with NaH (1.5mol) in DMF at $0^\circ/30$ minutes, followed by MeI (1.2mol) for 1 hour at $\sim 25^\circ$, then purified by column chromatography as above. Its ^1H NMR (400MHz, CDCl_3) had δ at 3.66 (s, 3H, *N*-Me), 6.43 (d, 1H, $J = 2.7\text{ Hz}$), 7.05 (d, 1H, $J = 2.7\text{ Hz}$), 7.34 (d, 1H, $J = 9.3\text{ Hz}$) and 7.44 (d, 1H, $J = 5.4\text{ Hz}$) [Patent to the Board of Trustees of the University of Illinois; WO2008/77138 2008].

3-Bromofuran [22037-28-1] $\text{C}_4\text{H}_3\text{BrO}$, M 147.0, b $38.5^\circ/40\text{mm}$, $50^\circ/110\text{mm}$, $102.5\text{--}103^\circ/\text{atm}$, d_4^{20} 1.661, n_D^{20} 1.4970. Purify 3-bromofuran by two steam distillations and dry it over fresh CaO. It can be dried over Na metal (no obvious reaction) and fractionated. It is not very soluble in H_2O but is soluble in organic solvents. When freshly distilled, it is a clear oil, but darkens on standing and eventually resinifies. It can be stored for long periods by covering the oil with an alkaline solution of hydroquinone and is redistilled when required. It forms a characteristic **maleic anhydride adduct**, m $131.5\text{--}132^\circ$. [Shepard et al. *J Am Chem Soc* **52** 2083 1930, DOI: 10.1021/ja01368a057; Hughes & Johnson *J Am Chem Soc* **53** 737 1931, DOI: 10.1021/ja01353a044; adduct: van Campen & Johnson *J Am Chem Soc* **55** 430 1933, DOI: 10.1021/ja01328a512; *Beilstein* **17/1** V 295.]

5-Bromoindole [10075-50-0] $\text{C}_8\text{H}_6\text{BrN}$, M 196.1, m $90.5\text{--}91^\circ$, $90\text{--}92^\circ$, pK^{25} 16.13 (NH). Purify it by steam distillation from a faintly alkaline solution. Cool the aqueous distillate, collect the solid, dry it in a vacuum desiccator over P_2O_5 and recrystallise it from aqueous EtOH (35% EtOH) or petroleum ether/ Et_2O . Its UV in MeOH has λ_{max} at 279, 287 and 296nm ($\log \epsilon$ 3.70, 3.69 and 3.53). The **picrate** has m $137\text{--}138^\circ(\text{dec})$ (from Et_2O /petroleum ether). [UV: Thesing et al. *Chem Ber* **95** 2205 1962, DOI: 10.1002/cber.19620950917; UV and NMR: Lallemand & Bernath *Bull Soc Chim Fr* 4091 1970, *Beilstein* **20/7** V 36.]

5-Bromoisatin [87-48-9] $\text{C}_8\text{H}_4\text{BrNO}_2$, M 226.0, m 245° , $247\text{--}152^\circ$, $251\text{--}153^\circ$, $255\text{--}256^\circ$. 5-Bromoisatin forms red prisms or needles from EtOH. The *N-acetate* crystallises as yellow prisms from $^*\text{C}_6\text{H}_6$, m $170\text{--}172^\circ$, and the *N-methyl derivative* forms orange-red needles from MeOH, m $172\text{--}173^\circ$. [Heller *Chem Ber* **53** 1545 1920, DOI: 10.1002/cber.19200530835; Buu-Hoi *Recl Trav Chim Pays-Bas* **73** 197 1954, DOI: 10.1002/recl.19540730305; Baker et al. *Tetrahedron Lett* 219 1978, DOI: 10.1016/S0040-4039(01)85088-7; *Beilstein* **21** H 453, **21** III/IV 5009.]

6-Bromoisatin [6326-79-0] has m 270° , pK^{25} 10.35. 6-Bromoisatin recrystallises from AcOH (yellow needles). It is a plant growth substance. Its IR (CHCl_3) has ν_{max} at 1320 and 3440cm^{-1} . [Sadler *J Org Chem* **21** 169 1956, DOI: 10.1021/jo01108a004; *Beilstein* **21** III/IV 5012.]

2-Bromo-3-methylindole (2-bromoskatole) [1484-28-2] $\text{C}_9\text{H}_8\text{BrN}$, M 210.1, m $102\text{--}104^\circ$, $\text{pK}_{\text{Est}} < 0$. Purify 2-bromoskatole by chromatography on silica gel in CHCl_3 /petroleum ether (1:2) followed by crystallisation from aqueous EtOH. [Phillips & Cohen *J Am Chem Soc* **108** 2023 1986, DOI: 10.1021/ja00268a049; cf. *Beilstein* **20** III/IV 3205.]

4-(Bromomethyl)-7-methoxycoumarin [35231-44-8] $\text{C}_{11}\text{H}_9\text{BrO}_3$, M 269.1, m $208\text{--}209^\circ$, $213\text{--}215^\circ$, $216\text{--}218^\circ$. The coumarin is crystallised from boiling AcOH, the crystals are washed with AcOH, EtOH and dried in a vacuum. The ^1H NMR (TFA) has δ at 3.97s, 4.57s, 6.62s, 6.92-7.19m and 7.80d. A useful fluorescent label,

e.g. for a wide range of acids. [Secrist et al. *Biochem Biophys Res Commun* **45** 1262 1971, DOI: 10.1016/0006-291X(71)90154-9; Duenges *Anal Chem* **49** 442 1977, DOI: 10.1021/ac50011a028; *Beilstein* **18** III/IV 348.]

5-Bromonicotinic acid [20826-04-4] $C_8H_4BrNO_2$, **M 202.0**, **m 178-180°, 178-182°, 189-190°, $pK_{Est} \sim 4.4$, $pK^{25} 4.02$ (50% aqueous EtOH).** The acid is recrystallised from H_2O and then from EtOH using charcoal. The **amide** has **m 219-219.5°** (from aqueous EtOH), and the **methyl ester**, prepared by addition of ethereal diazomethane, can be purified by sublimation in a vacuum and has **m 98-99°**. The **acid chloride** also can be sublimed in *vacuo* and has **m 74-75°** and gives the **methyl ester** in MeOH. [Graf *J Prakt Chem* **138** 244 1933, DOI: 10.1002/prac.19331380904; Bachman & Micucci *J Am Chem Soc* **70** 2381 1948, DOI: 10.1021/ja01187a020; Garcia et al. *J Am Chem Soc* **82** 4430 1960, DOI: 10.1021/ja01501a079; Misić-Voković et al. *JCS Perkin Trans 2* **34** 1978, DOI: 10.1039/P29780000034; *Beilstein* **22/2** V 181.]

2-Bromopyridine [109-04-6] C_5H_4BrN , **M 158.0**, **b 49.0°/2.7mm, 192-194°/atm, $d_4^{20} 1.660$, $n_D^{20} 1.5713$, $pK^{25} 0.90$.** Dry 2-bromopyridine over KOH for several days, then distil it from CaO under reduced pressure, and taking the middle fraction. [*Beilstein* **20/5** V 422.]

8-Bromotheophylline (bromo-1,3-dimethyl-2,6(1*H*,3*H*)-purinedione) [10381-75-6] **M 259.1, m 309°, 315-320° (with browning and dec), $pK_{Est(1)} \sim 5.5$, $pK_{Est(2)} \sim 9.2$.** It is purified by dissolving in the minimum volume of dilute NaOH (charcoal), filtering and acidifying to pH ca 3.5-4. The solid that separates is collected, dried in *vacuo* at 100° and stored in a dark container. It has also been recrystallised from EtOH or AcOH. [Blitz & Beck *J Prakt Chem* [2] **118** 149 1928, DOI: 10.1002/prac.19281180116; Fischer & Ach *Chem Ber* **28** 3135 1895, DOI: 10.1002/cber.189502803156; *Beilstein* **26** H 476, **26** II 227, **26** III/IV 2447.]

5-Bromothiazole [3034-55-7] C_3H_2BrNS , **M 164.0**, **b 62-63°/15mm, $d_4^{20} 1.835$, $n_D^{20} 1.5955$, $n_D^{25} 1.5976$, $pK_{Est} \sim 2.0$.** If bromothiazole is too coloured, then suspend it in dilute NaOH and steam distil it. Add NaCl to the aqueous distillate, extract it with Et_2O , dry it (Na_2SO_4), evaporate and fractionate the residue in a vacuum. The **$HgCl_2$ salt** crystallises from EtOH with **m 148° (dec)**. [Beyerman et al. *Recl Trav Chim Pays-Bas* **73** 325 1954, DOI: 10.1002/recl.19540730405; *Beilstein* **27** III/IV 962.]

2,3-(di)-Bromothiophene [3140-93-0] $C_4H_2Br_2S$, **M 241.9**, **m -17.5°, b 89-91°/13mm, 212-213°/atm, 218.6-219.6°/atm, $d_4^{25} 2.137$, $n_D^{20} 1.632$.** Purify the dibromothiophene by fractional distillation, preferably in a vacuum. Low temperature crystallisation from a small volume of petroleum ether (use Et_2O/CO_2 bath) removes the more soluble 2,4-dibromothiophene isomer. Nitration with Ac_2O/HNO_3 at 50-55° yields **2,3-dibromo-5-nitrothiophene m 75°** (from EtOH). [Steinkopf et al. *Justus Liebigs Ann Chem* **512** 136 1934, DOI: 10.1002/jlac.19345120113; Gronowitz et al. *Acta Chem Scand* **46** 654 1992, DOI: 10.3891/acta.chem.scand.46-0654; NMR: Fujiwara et al. *Bull Chem Soc Jpn* **32** 201 1959, DOI: org/10.1246/bcsj.32.201; *Beilstein* **17/1** III/IV 247, **17/1** V 308.]

4-tert-Butylcalix[4]arene [60705-62-6] $C_{44}H_{56}O_4$, **M 648.9**, **m >300°(dec), 380°(dec), 344-346°.** The calixarene recrystallises from $CHCl_3$ in large solvated prisms (**m 380° dec**); it effloresces on drying in air. Its **tetra-acetate** crystallises from Ac_2O in colourless prisms **m 332-333°(dec)**. It crystallises from CCl_4 or chlorobenzene /EtOH (**m >300°**) and the **tetra-acetate** crystallises from $CHCl_3$ /EtOH **m >290°(dec)**. It also crystallises from toluene in white plates with toluene of crystallisation **m 344-346° (330-332°)**; the **tetra-acetate** crystallises with 1AcOH of crystallisation **m 383-386°** (softening at **330-340°**, also **m 283-286°**), but acetylation with $Ac_2O/NaOAc$ gives the **triacetate** which recrystallises from AcOH with 1AcOH of crystallisation **m 278-281°**. 4-tert-Butylcalix[4]arene (100mg) is unchanged after boiling for 4 hours with 10N KOH (0.04ml) in xylene (4ml). [Cornforth et al. *Br J Pharmacol Chemother* **10** 73 1955, PMID: PMC1509476; Kämmerer et al. *Monatsh Chem* **109** 767 1978, DOI: 10.1007/BF00907297; Gutsche et al. *J Am Chem Soc* **103** 3782 1981, DOI: 10.1021/ja00403a028; see also Kluwaver in *Calixarenes*, Vicens & Böhner eds Academic Press 1991, *Beilstein* **6** IV 7858.]

4-tert-Butylcalix[6]arene [78092-53-2] $C_{66}H_{84}O_6$, **M 972.3**, has **m >300°, 380-381°**. It is recrystallised from $CHCl_3$ or $CHCl_3/MeOH$ to give a white solid from the mother liquors of the calix[8]arene preparation. The **hexa-acetate** (Ac_2O/H_2SO_4) crystallises from $CHCl_3/MeOH$ with **m 360-362°(dec)**, and the **(SiMe₃)₆ derivative** crystallises from $CHCl_3/MeOH$ with **m 410-412°**. Its stability in KOH-xylene is the same as for the 4-tert-

butylcalix[4]arene. [Gutsche et al. *J Am Chem Soc* **103** 3782 1981, DOI: 10.1021/ja00403a028. See also Kluawer in *Calixarenes*, Vicens & Böhner eds Academic Press 1991, *Beilstein* **6** IV 7858.]

4-tert-Butylcalix[8]arene [68971-82-4] $C_{88}H_{112}O_8$, **M 1297.8**, has **m 411-412°**. The calixarene recrystallises from $CHCl_3$ in fine colourless, glistening needles. It melts sharply between 400-401° and 411-412° depending on the sample and is sensitive to traces of metal ions. On TLC with silica gel (250 μ m thick) and elution with $CHCl_3$ /hexane (3:4) it has R_F 0.75. The **octa-acetate** is prepared from 8g in Ac_2O (50ml) and 2 drops of concentrated H_2SO_4 and refluxed for 2 hours. On cooling, a colourless precipitate separates and is recrystallised from Ac_2O (1.2g 48%) with **m 353-354°**. The **(SiMe₃)₈ derivative** is prepared from 4-tert-butylcalix[8]arene (0.65g) in pyridine (4ml) with excess of hexamethyldisilazane (1ml) and trimethylchlorosilane (0.5ml) and refluxed under N_2 for 2 hours. Cool, evaporate the pyridine, triturate the gummy residue with MeOH. Chromatograph on silica gel using hexane/ CH_2Cl_2 gave 0.5g (61%) with one spot on TLC. Recrystallise it from hexane/ Me_2CO to give colourless needles **m 358-360°**. [Gutsche et al. *J Am Chem Soc* **103** 3782 1981, DOI: 10.1021/ja00403a028; Gutsche & Muthukrishnan *J Org Chem* **43** 4905 1978, DOI: 10.1021/jo00419a052; Muthukrishnan & Gutsche *J Org Chem* **44** 3962 1979, DOI: 10.1021/jo01336a045; Andretti et al. *JCS Chem Commun* 533 1981, DOI: 10.1039/C39810000533; see Kluawer in *Calixarenes*, Vicens & Böhner eds Academic Press 1991.]

8-sec-Butylmetrazole [25717-83-3] **M 194.3**, **m 70°**. Crystallise it from petroleum ether and dry it for 2 days under vacuum over P_2O_5 . [*Beilstein* **26** II 213 for Metrazole.]

N-(n-Butyl)-5-nitro-2-furamide [14121-89-2] **M 212.2**, **m 89-90°**, **b 190°/10mm**. Distil the amide in a vacuum and recrystallise it twice from EtOH/water mixture or petroleum ether. [Gilman & Yale *J Am Chem Soc* **72** 3593 1950, DOI: 10.1021/ja01164a075; *Beilstein* **18** III/IV 3995.]

Butyloxirane (1-hexene oxide) [1436-34-6] $C_6H_{12}O$, **M 100.2**, **b 116-117°/atm**, **116-119°/atm**, **118-120°/atm**, **d₄²⁰ 0.833**, **n_D²⁰ 1.44051**. Purify it by fractional distillation through a 2ft helices-packed column at atmospheric pressure in a N_2 atmosphere. [Pasto & Cumbo *J Org Chem* **30** 1271 1965, DOI: 10.1021/jo01015a523; Emmons & Pagaro *J Am Chem Soc* **77** 89 1955, DOI: 10.1021/ja01606a029; ¹³C NMR Davies & Whitham *JCS Perkin Trans 2* 861 1975, DOI: 10.1039/P29750000861; *Beilstein* **17/1** V 103.]

4-tert-Butylpyridine [3978-81-2] $C_9H_{13}N$, **M 135.2**, **m -44.4°**, **b 194-197°/atm**, **196-197°/atm**, **197°/765mm**, **d₄²⁰ 0.923**, **n_D²⁰ 1.495**, **pK²⁵ 5.82**. Dry 4-tert-butylpyridine over solid KOH and purify it by fractional distillation through an efficient column under dry N_2 . Its **picrate** has **m 153.9-154°**, and the **hydrochloride** has **m 151.7-154.8°** (from Me_2CO). [Brown & Murphey *J Am Chem Soc* **73** 3308 1951, DOI: 10.1021/ja01151a093; Arnett & Chawla *J Am Chem Soc* **100** 214 1978, DOI: 10.1021/ja00469a037; Kyle et al. *J Chem Soc* 4454 1960, DOI: 10.1039/JR9600004454; *Beilstein* **20/6** V 123.]

Cacotheline (2,3-dihydro-4-nitro-2,3-dioxo-9,10-secostrychnidin-10-oic acid) [561-20-6] $C_{21}H_{21}N_3O_7$, **M 427.4**, **pK_{Est(1)} ~4.4 (CO₂H)**, **pK_{Est(2)} ~10.2 (proton gain)**. Cacotheline gives yellow crystals from H_2O . It is then dried over H_2SO_4 which gives the **dihydrate**, and in a vacuum over H_2SO_4 at 105° it forms the **anhydrous** compound. The **hydrochloride** separates as the **hydrate** (on heating in vacuum at 80°) in orange-yellow prisms or plates, **m 250°(dec)** and forms a **resorcinol complex** which gives brown crystals from EtOH, **m 325°**; and a **hydroquinone complex** as dark red crystals from EtOH, **m 319°**. [Leuchs & Leuchs *Chem Ber* **43** 1042 1910, DOI: 10.1002/cber.191004301179; Teuber *Chem Ber* **86** 232, (UV: 242) 1953, DOI: 10.1002/cber.19530860218; complexes: Gallo *Gazz Chim Ital* **85** 1441 1955.] It is used in the titrimetric estimation of Sn^{2+} ions [Szarvas & Lantos *Talanta* **10** 477 1963, DOI: 10.1016/0039-9140(63)80055-7]. [*Beilstein* **27** III/IV 8014.]

1S(-)-Camphorsulfonylimine {(7S)-10,10-dimethyl-5-thia-4-azatricyclo[5.2.1.0^{3,7}]dec-3-ene-5,5-dioxide} [60886-80-8] $C_{10}H_{15}NO_2S$, **M 213.3**, **m 228-230°**, **[α]_D²⁰ -34 (c 1, $CHCl_3$)**. The method described here is that of Davis and coworkers [*Org Synth Coll Vol* **8** 110 1993, *Org Synth* **69** 158 1990, DOI: 10.15227/orgsyn.069.0158]: Under N_2 , and with stirring, 1S(+)-camphorsulfonamide (41.5g, see [60933-63-3] above) in toluene (500ml) and Amberlyst 15 ion-exchange resin (5g) are boiled under a Dean-Stark water separator and reflux for 4 hours, after which time H_2O separation was complete. While still warm (at 40-50°),

CH₂Cl₂ (200ml) is added slowly to dissolve any sulfonylimine that separated and the solution is filtered through a 150ml coarse porosity sintered glass funnel, and the funnel is washed with CH₂Cl₂ (75ml). The combined CH₂Cl₂ solutions (containing toluene) are evaporated and the solid residue being **1S(-)-camphorsulfonylimine** is recrystallised from absolute EtOH (750ml) to provide white crystals (34.5-36.4g, 90-95% yield) with **m 225-228°**, $[\alpha]_D^{20}$ -32.7 (c 1.9, CHCl₃). Its IR (CHCl₃) has ν_{\max} at 3030, 2967, 1366 cm⁻¹; ¹H NMR (CDCl₃) has δ at 1.03 (s, 3H, CH₃), 1.18 (s, 3H, CH₃), 1.45-2.18 (m, 6H), 2.65 (m, 1H), 3.10 and 3.28 (AB quartet, 2H CH₂-SO₂, J = 14Hz); ¹³C NMR (CDCl₃) has δ at 19.01 (q, CH₃), 19.45 (q, CH₃), 26.64 (t), 28.44 (t), 35.92 (t), 44.64 (d), 48.00 (s), 49.46 (t), 64.52 (s), 195.52 (s). The **1R(+)- enantiomer** was similarly prepared. [Beilstein **27** III/IV 1007.]

2R,8aS-(+)-10-Camphorsulfonyloxaziridine {**4H-4A,7-methanooxazirino[3,2-*i*][2,1]benzisothiazole, tetrahydro-9,9-dimethyl, 3,3-dioxide [4aS-(4a α ,7 α ,8aR*)]**} [104322-63-6] C₁₀H₁₅NO₃S, **M 229.3, m 165-167°**, $[\alpha]_D^{20}$ +45 (c 2, CHCl₃). The method described here is that of Davis et al. [Org Synth Coll Vol **8** 110 1993, Org Synth **69** 158 1990, DOI: 10.15227/orgsyn.069.0158]: To a stirred solution of the preceding **1S(-)-camphorsulfonylimine** (39.9g, 0.187mol) in toluene (500ml) in a 5L three necked flask, a solution of anhydrous K₂CO₃ (543g, 7 equivalents based on oxone) in H₂O (750ml) is added with vigorous stirring, followed by a solution of **oxone** [345g, 0.56mol, 6 equivalents of KHSO₅, see [37222-66-5, 70693-62-8 (triple salt)] in 'Inorganic Compounds', Chapter 4] in H₂O (1250ml) dropwise in three volumes over 45 minutes. [Note that the solution turns milky, and the reaction time depends on the activity of oxone; if it is below parr (e.g. if previously exposed to moisture, but if stored <0° under N₂ it should be stable for at least 6 months), then more oxone should be added making sure that the pH is maintained at ~9.] Completion of reaction should occur after all the oxone is added and should be checked by TLC. Thus an aliquot of the reaction mixture (~0.5ml) is spotted on a TLC silica gel plate (250 μ m thick), eluted with CH₂Cl₂ and developed with 10% molybdophosphoric acid in EtOH and heated for a few minutes in 100° oven, whereby the absence of a spot with R_F = 0.28 (for the *sulfonylimine*) indicates completion of reaction. {Note also, that if the reaction becomes brownish in colour and the addition of oxone is not complete after 30 minutes, the mixture should be filtered through a 150ml coarse porosity sintered glass funnel, the solids are washed with CH₂Cl₂ (50ml). The filtrate is returned to the original flask and stirred vigorously with more oxone (52g, 0.08mol, 1 equivalent of KHSO₅) added within 5 minutes and stirring is continued until the reaction is complete (~10-15 minutes).} At completion, the mixture is filtered through a 150ml coarse porosity sintered glass funnel to remove solids, the filtrate is transferred to a separating funnel (3L), the toluene phase is collected, the aqueous phase is washed with CH₂Cl₂ (100ml x 3). The solids together with the solids remaining in the original reaction flask are combined and washed with more CH₂Cl₂ (200ml). The combined organic liquids are washed with saturated sodium sulfite, dried (MgSO₄) for 15-20min, filtered and evaporated to dryness. The residual solid is recrystallised from hot *iso*-propanol (~500ml), which after drying *in vacuo* gave the **(+)-oxaziridine** (35.9g, 84% yield) as white needles with **m 165-167°**, $[\alpha]_D^{20}$ +44.7 (c 2.2, CHCl₃). Its ¹H NMR (CDCl₃) has δ at 1.03 (s, 3H, CH₃), 1.18 (s, 3H, CH₃), 1.45-2.18 (m, 6H), 2.65 (m, 1H), 3.10 and 3.28 (AB quartet, 2H, CH₂-SO₂, J = 14Hz); ¹³C NMR (CDCl₃) has δ at 19.45 (q, CH₃), 20.42 (q, CH₃), 26.55 (t), 28.39 (t), 33.64(t), 45.78 (d), 48.16 (s), 48.32 (t), 45.07 (s), 98.76 (s). The **(-)-oxaziridine enantiomer** [104372-31-8] was prepared similarly and had **m 166-167°**, $[\alpha]_D^{20}$ -43.6 (c 2.2, CHCl₃). Note the inversion of rotation on oxidation of the *sulfonylimine*.

These oxaziridines are selective aprotic oxidising agents [see Davis & Jenkins on the chemistry of *N*-sulfonyloxaziridines in *Asymmetric Synthesis* Morrison Ed. Academic Press Vol 4 Chapter 4 1984, ISBN 10: 012431452X, 13: 978-0124314528; and Davis & Sheppard *Tetrahedron* **45** 5703 1989, DOI: 10.1016/S0040-4020(01)89102-X], and stereoselectively oxidise sulfides to sulfoxides [in 30-91% ee, Davis et al. *J Am Chem Soc* **109** 3370 1987, DOI: 10.1021/ja00245a030], selenides to selenoxides [8-9% ee, Davis et al. *Tetrahedron* **41** 4747 1985, DOI: 10.1016/S0040-4020(01)96713-4], sulfides to thiolsulfonates [2-13% ee, Davis et al. *J Am Chem Soc* **104** 5412 1982, DOI: 10.1021/ja00384a028], used in asymmetric epoxidation of alkenes [19-65% ee, Davis & Chattopadhyay *Tetrahedron Lett* **27** 5079 1986, DOI: 10.1016/S0040-4039(00)85137-0; Davis et al. *J Am Chem Soc* **105** 3123 1983, DOI: 10.1021/ja00348a029] and in the asymmetric oxidation of enolates [35-82% ee, Davis et al. *J Org Chem* **51** 2402 1986, DOI: 10.1021/jo00362a053; 51-84% ee, Davis & Haque *J Org Chem* **51** 4083 1986, DOI: 10.1021/jo00371a038, Davis et al. *J Org Chem* **54** 2021 1989, DOI: 10.1021/jo00269a054; 55-80% ee, Davis et al. *J Am Chem Soc* **112** 6679 1990, DOI: 10.1021/ja00174a035]. [Fieser **13** 64, **14** 72, **16** 61, **17** 320.]

2,10-Camphorsultam [(3*aS*)-4,5,6,7-tetrahydro-8,8-dimethyl-1,2-dioxide-3*H*-3*a*,6-methano-2,1-benzisothiazole] [*IR*-(+)- 108448-77-7, *1S*-(-)- 94594-90-8] $C_{10}H_{17}NO_2S$, **M 215.3**, **m 181-183°**, **183-184 185-187°**, $[\alpha]_D^{20}$ (+) and (-) **32** (c 5, $CHCl_3$). The (-)-enantiomer has been prepared by reduction of (-)-*camphorsulfonylimine* (see above) with $LiAlH_4$. The (-)-enantiomer is recrystallised from 95% EtOH and dried in a vacuum desiccator. Alternatively dissolve the sultam (33.5g) in hot absolute EtOH (~60 mL), cool, collect the solid on a coarse porosity sintered glass funnel and dry it in a vacuum desiccator to give the pure sultam (31.1g, + 1.4g from the mother liquors) as white crystals. It dissolves in dilute aqueous NaOH and can be precipitated without hydrolysis by acidifying. It forms the *N*-**Na salt** in EtOH (by addition of Na to the EtOH solution), and the salt can be methylated with MeI to give the (-)-*N*-**Me lactam** with **m 80°** after recrystallisation from hot H_2O , and has $[\alpha]_D^{25}$ -59.6 (c 5, $CHCl_3$) [Shriner et al. *J Am Chem Soc* **60** 2794 1938, DOI: 10.1021/ja01278a072]. [Oppolzer et al. *Helv Chem Acta* **69** 1542 1986, DOI: 10.1002/hlca.19860690703; for prep and 1H NMR see: Weismiller et al. *Org Synth* **69** 154 1955, Coll Vol **8** 110 1993, DOI: 10.15227/orgsyn.069.0154; *Beilstein* **27** III/IV 1007.]

ϵ -Caprolactam (azepan-2-one, aza-2-cycloheptanone, 2-oxohexamethyleneimine) [105-60-2] $C_6H_{11}NO$, **M 113.2**, **m 68-70°**, **70°, 70.5-71.5°, 70-71°, 136-138°/10mm, 262.5°/760mm**. The lactam is distilled under reduced pressure, recrystallised from acetone or petroleum ether and redistilled. It can be purified by zone melting. It is very *hygroscopic* and discolours in contact with air unless small amounts (0.2g/L) of NaOH, Na_2CO_3 or $NaBO_2$ are present. It has been crystallised from a mixture of petroleum ether (185ml of **b 70°**) and 2-methyl-2-propanol (30ml), from acetone, or petroleum ether. It is then distilled under reduced pressure and stored under nitrogen. [Pellegata et al. *Synthesis* 614 1978, DOI: 10.1055/s-1978-24834; *Beilstein* **21/6** V 444.]

Caprylolactam (azanon-2-one, azacyclononan-2-one, 8-aminooctanoic acid lactam, cyclooctanone isoxime) [935-30-8] $C_8H_{15}NO$, **M 141.2**, **m 72°, 73°, 74-76°, 75°, 76-77°, b 119-122°/0.7mm, 150-151°/7-8mm, 164°/14mm, d_4^{73} 1.009, n_D^{73} 1.489, pK^{25} 0.55 (AcOH)**. Dissolve it in $CHCl_3$, decolorise it with charcoal, evaporate to dryness and recrystallise it from $CHCl_3$ /hexane. Sublime it at high vacuum. The *oxime* has **m 117°** (from $*C_6H_6$ or petroleum ether). [Guggisberg et al. *Helv Chim Acta* **61** 1050 1978, DOI: 10.1002/hlca.19780610314; Olah et al. *Synthesis* 537 1979, DOI: 10.1055/s-1979-28752; Behringer & Meier *Justus Liebigs Ann Chem* **607** 67 1957, DOI: 10.1002/jlac.19576070109; *Beilstein* **21** III/V 3260.]

Carbazole [86-74-8] C_8H_7NO , **M 167.2**, **m 240-243°, 243-246°, b 355°/atm, pK^{25} <0**. Dissolve carbazole (60g) in conc H_2SO_4 (300ml), extract with three 200ml portions of *benzene*, then stir this into 1600ml of an ice-water mixture. The precipitate is filtered off, washed with a little water, dried, recrystallised from *benzene* and then from pyridine/*benzene* [Feldman et al. *J Am Chem Soc* **73** 4341 1951, DOI: 10.1021/ja01153a091]. It has also been recrystallised from EtOH or toluene, sublimed in vacuum, zone-refined, and purified by TLC. [UV: Armarego in *Physical Methods in Heterocyclic Chemistry* (Ed Katritzky, Academic Press, LCCC No 62-12037, Internet Public Domain) **Vol III** 158 1971; *Beilstein* **20/8** V 9.]

9-Carbazoleacetic acid [524-80-1] $C_{14}H_{11}NO_2$, **M 225.2**, **m 215°, pK_{Est} ~3.5**. Crystallise the acid from ethyl acetate. It yields **9-methylcarbazole** [1484-12-4] $C_{13}H_{11}N$, **M 181.2**, **m 90-92°**, on decarboxylation at **240-260°**. [Fan et al. *Anal Chim Acta* **367** 81 1998, DOI: 10.1016/S0003-2670(98)00125-1; *Beilstein* **20** III/IV 3841, and *Beilstein* **20** H 436 for 9-methylcarbazole] The *ethyl ester*, $C_{16}H_{15}NO_2$, **m 97°**, crystallises from EtOH.

Carbazole-9-carbonyl chloride [73500-82-0] **M 300.0**, **m 100-103°, 103.5-104.5°**. Recrystallise the acid chloride from $*C_6H_6$. If it is not very pure (presence of OH or NH bands in the IR), dissolve it in pyridine, shake it with phosgene in toluene, evaporate and recrystallise the residue. Carry out this experiment in a good fume cupboard as $COCl_2$ is very **TOXIC**, and store the product in the dark. It is moisture sensitive. The *amide* has **m 246.5-247°**, and the *dimethylaminoethylamide hydrochloride* has **m 197-198°**. [Weston et al. *J Am Chem Soc* **75** 4006 1953, DOI: 10.1021/ja01112a038; *Beilstein* **20** III/IV 3841.]

1-Carbethoxy-4-methylpiperazine hydrochloride [532-78-5] **M 204.7**, **m 168.5-169°, pK^{25} 7.31**. Crystallise the hydrochloride from absolute EtOH. Its solubility is 280g/100g of H_2O at 25°. The *free base ester* has **b 97-98°/8mm**. [*Beilstein* **23** III/IV 223.]

γ -Carboline (9*H*-pyrido[3,4-*b*]indole) [244-69-9] $C_{11}H_8N_2$, **M 168.2**, **m 225°**, **230-233°**, **pK²⁵ ~0**. Crystallise it from water or EtOAc. [Robinson & Thornley *J Chem Soc* **125** 2169 1924, DOI: 10.1039/CT9242502169; Dalton et al. *Aust J Chem* **22** 185 1969, DOI: 10.1071/CH9690185; Staab & Wendel *Org Synth* **48** 44 1968, DOI: 10.15227/orgsyn.048.0044; *Beilstein* **23** II 223, **23** III/IV 1572.]

***N,N'*-Carbonyldiimidazole (CDI)** [530-62-1] $C_7H_6N_4O$, **M 162.2**, **m 115-122°**, **115.5-116°**, **117-122°**. Crystallise it from *benzene or tetrahydrofuran in a dry-box and store it dry. [*Beilstein* **23/4** V 245.] Very useful for activating acids (as imidazolides) for the synthesis of esters, amides, and ketones, and for immobilising enzymes on affinity ligands [Staab *Angew Chem* **74** 407 1962, DOI: 10.1002/ange.19620741203; Hearn *Methods Enzymol* **135** 102 1987, DOI: 10.1016/0076-6879(87)35068-2]. It has also been used as a carbonyl-transfer reagent for synthesising various heterocyclic compounds [Barluenga et al. *J Org Chem* **56** 6751 1991, DOI: 10.1021/jo00024a011; Kang et al. *Synth Commun* **23** 2219 1993, DOI: 10.1080/00397919308013776]. [*Beilstein* **23/4** V 245.]

1,1'-Carbonyldi(1,2,4-triazole) (CDT) [41864-22-6] $C_5H_4N_6O$, **M 164.1**, **m 134-136°**, **145-150°**. Dissolve CDT in tetrahydrofuran and evaporate at 10mm until it crystallises. Wash the crystals with cold tetrahydrofuran and dry them in a vacuum desiccator over P_2O_5 in which it can be stored for months. Its applications are similar to those of CDI (see preceding entry). [Bergmann & van Den Brink *Recl Trav Chim Pays-Bas* **80** 1372 1961, DOI: 10.1002/recl.19610801209; Potts & Crawford *J Org Chem* **27** 2631 1962, DOI: 10.1021/jo01054a502; Staab *Justus Liebigs Ann Chem* **609** 75 1957, DOI: 10.1002/jlac.19576090108.] [*Beilstein* **24** IV 38.]

(±)-Catechin (3,3',4',5,7-pentahydroxyflavan) [7295-85-4] $C_{15}H_{14}O_6$, **M 290.2**, **m 177° (anhydrous)**. Crystallise it from hot water or hot H_2O + AcOH, and dry it at 100° in a vacuum. **(-)-Catechin** [18829-70-4] has **m 93-96°** (hydrate from H_2O + AcOH) and **m 175-177° (anhydrous)** and $[\alpha]_D -16.8$ (H_2O); and the enantiomer **(+)-Catechin** [154-23-4] (which occurs as the hydrate) has **m 93-96°** (hydrate from H_2O + AcOH) and **m 175-177° (anhydrous)** and $[\alpha]_D^{18} +16.8$ to +18 (H_2O). [*Beilstein* **17/8** V 448, for stereochemistry see Clark-Lewis *J Chem Soc* 2433 1960, DOI: 10.1039/JR9600002433.]

Cetylpyridinium chloride (H_2O) (hexadecylpyridinium chloride) [6004-24-6] $C_{21}H_{38}ClN$, **M 358.0**, **m 80-83°**. Crystallise the chloride from MeOH or EtOH/diethyl ether and dry it *in vacuo*. [Moss et al. *J Am Chem Soc* **108** 788 1986, DOI: 10.1021/ja00264a035; Lennox & McClelland *J Am Chem Soc* **108** 3771 1986, DOI: 10.1021/ja00273a035; *Beilstein* **20** V 233.]

Chelerythrine (1,2-dimethoxy-12-methyl[1,3]benzodioxolo[5,6-*c*]phenanthridinium) [34316-15-9] $C_{21}H_{18}NO_4^+$, **M 389.4**, **m 207°**. Chelerythrine crystallises from $CHCl_3$ on addition of MeOH [Manske *Can J Res* **21B** 140 1943, DOI: 10.1139/cjr43b-018; UV: Hruban et al. *Coll Czech Chem Commun* **35** 3420 1970, DOI: 10.1135/cccc19703420]. The *pseudo base* is colourless while the salts are yellow in aqueous solution and are fluorescent.

Chelidonic acid (4-oxopyran-2,6-dicarboxylic acid) [99-32-1] $C_7H_4O_6$, **M 184.1**, **m 262°**, **265°(dec)**, **pK₂²⁵ 2.36**. The acid crystallises from aqueous EtOH. Dry it first at 100°/2 hours, then at 160° to constant weight to remove water of crystallisation. It decarboxylates at 220-230° in a vacuum. [Riegel & Zwillgmeyer *Org Synth Coll Vol* **2** 126 1943, DOI: 10.15227/orgsyn.017.0040; *Beilstein* **18** H 490, **18/8** V 646.]

2-Chlorobenzothiazole [615-20-3] C_7H_4ClS , **M 169.6**, **m 21°**, **21-23°**, **90-91.4°/4mm**, **135-136°/28mm**, **141°/30mm**, **d₄²⁰ 1.303**, **n_D²⁰ 1.6398**. It is purified by fractional distillation *in vacuo*. The **2-chloro-3-methylbenzothiazolinium 2,4-dinitro-benzenesulfonate** crystallises from Ac_2O , **m 162-163°(dec)**. [Young & Amstutz *J Am Chem Soc* **73** 4773 1951, DOI: 10.1021/ja01154a088; Brower et al. *J Org Chem* **19** 1830 1954, DOI: 10.1021/jo01376a018; Hunter & Jones *J Chem Soc* 2190 1930, DOI: 10.1039/JR930000219; *Beilstein* **27** H 44, **27** II 18, **27** III/IV 1072.]

***o*-Chlorobenzotrifluoride** [88-16-4] $C_7H_4ClF_3$, **M 180.6**, **b 152.3°/atm**, **d₄²⁰ 1.371**, **n_D²⁰ 1.456**. Dry it over $CaSO_4$, and distil it at high reflux ratio. [*Beilstein* **5** III 692, **5** IV 814.]

2-Chlorobenzoxazole [615-18-9] C_7H_4ClNO , M 153.6, m 7°, b 95-96°/20mm, 198-202°/atm, d_4^{20} 1.331, n_D^{20} 1.570. Purify it by fractional distillation, preferably in a vacuum. [Seidel *J Prakt Chem* 42 445 1890, DOI: 10.1002/prac.18900420140; Katz *J Am Chem Soc* 75 712 1953, DOI: 10.1021/ja01099a059; Meyer & Sigel *J Org Chem* 42 2769 1977, DOI: 10.1021/jo00436a024; *Beilstein* 27 H 43, 27 II 17.]

2-(4-Chlorobutyl)-1,3-dioxolane [118336-86-0] $C_7H_{13}ClO_2$, M 164.6, b 56-58°/0.1mm, d_4^{20} 1.106, n_D^{20} 1.457. If the IR has no CHO band, then just distil it in a vacuum. If it is present, then dissolve it in Et_2O , wash it with H_2O , then saturated $NaHCO_3$, dry over $MgSO_4$, evaporate and distil it. [cf. Kriesel & Gisvold *J Pharm Sci* 60 1250 1971, DOI: 10.1002/jps.2600600832; Loftfield *J Am Chem Soc* 73 1365 1951, DOI: 10.1021/ja01147a519.]

4-Chloro-2,6-diaminopyrimidine (2,4-diamino-6-chloropyrimidine) [156-83-2] $C_4H_5ClN_4$, M 144.6, m 198°, 199-202°, pK^{25} 3.57. It recrystallises from boiling H_2O (charcoal) as needles; it also crystallises from Me_2CO . [Büttner *Chem Ber* 36 2227 1903, DOI: 10.1002/cber.190303602145; Roth et al. *J Am Chem Soc* 72 1914 1950, DOI: 10.1021/ja01161a017; UV: Brown & Jacobsen *J Chem Soc* 3172 1962, DOI: 10.1039/JR9620003172; *Beilstein* 24 H 318, 25 III/IV 2788.]

2-Chloro-4,6-dimethylpyrimidine [4472-44-0] $C_6H_7ClN_2$, M 142.6, m 34-38°, 38°, b 223°/atm, pK^{20} -0.68. The chloro-pyrimidine has been distilled at atmospheric pressure, and solidifies on cooling. Purify it by recrystallisation from petroleum ether (b 40-60°). The 2-chloro-substituent is readily subject to nucleophilic substitution, and the kinetics of reaction with morpholine and piperidine are reported in detail (cf. Chapman & Rees). It is a very weak base and requires 2N HCl to form the cation species. The UV has λ_{max} ($\log \epsilon$) at 255 (3.65) and 263 (infl 3.49) nm at pH 2.0 (neutral species); and 261nm (3.87) in ~2N HCl (H_o -3.0, cation). [Boarland & McOmie *J Chem Soc* 1218 1951, DOI: 10.1039/JR9510001218; Boarland & McOmie 3722 1952, DOI: 10.1039/JR9520003722; Chapman & Rees *J Chem Soc* 1190 1954, DOI: 10.1039/JR9540001190; Brown & England *J Chem Soc, C* 1922 1967, DOI: 10.1039/J39670001922; Matsukawa & Ohta *J Pharm Soc, Jpn* 69 491 1949, *Chem Abstr* 44 3456 1950, Andrisano & Modena *Gazz. Chem Ital* 81 405 1951, Angerstein *Chem Ber* 34 3956 1901, DOI: 10.1002/cber.190103403119; *Beilstein* 23 H 95, 23 IV 918.]

2-Chloro-3,5-dinitropyridine [2578-45-2] $C_5H_2ClN_3O_3$, M 203.5, m 62-65°, 63-65°, 64°, pK_{Est} <-5. Dissolve it in $CHCl_3$, shake it with saturated $NaHCO_3$, dry ($MgSO_4$), evaporate and apply to an Al_2O_3 column, elute with petroleum ether (b 60-80°), evaporate and recrystallise it from $*C_6H_6$ or petroleum ether. [Ochiai & Kaneko *Chem Pharm Bull Jpn* 8 28 1960, DOI: org/10.1248/cpb.8.28; Plazek *Recl Trav Chim Pays-Bas* 72 569 1953, DOI: 10.1002/recl.19530720706; *Beilstein* 20/5 V 458.]

1-(2-Chloroethyl)pyrrolidine hydrochloride [7250-67-1] $C_6H_{12}Cl.HCl$, M 170.1, m 167-170°, 173.5-174°, pK_{Est} ~8.5 (free base). Purify the hydrochloride by recrystallisation from isopropanol/di-isopropyl ether (charcoal) and recrystallise it twice more. The *free base*, b 55-56°/11mm, 60-63°/23mm and 90°/56mm, is relatively unstable and should be converted to the hydrochloride immediately, by dissolving in isopropanol and bubbling dry HCl through the solution at 0°, filtering off the hydrochloride and recrystallising it. The *picrate* has m 107.3-107.8° (from EtOH) [Cason et al. *J Org Chem* 24 247 1959, DOI: 10.1021/jo01084a624; Wright et al. *J Am Chem Soc* 70 3098 1948, DOI: 10.1021/ja01189a078]. [*Beilstein* 20 III/IV 66.]

5-Chloroindole [17422-32-1] C_8H_6ClN , M 151.6, m 69-71°, 72-73°, b 120-130°/0.4mm, pK_{Est} <0 It is distilled at high vacuum and recrystallises from petroleum ether (b 40-60°) or (b 80-100°) as glistening plates. The *picrate* has m 147° (146.5-147.5°) (from $*C_6H_6$). [Rydon & Tweddle *J Chem Soc* 3499 1955, DOI: 10.1039/JR9550003499; Sugawara et al. *J Org Chem* 44 578 1979, DOI: 10.1021/jo01318a021; *Beilstein* 20/4 V 34.]

2-Chloro-6-(methylamino)purine [82499-02-3] $C_6H_6ClN_5$, M 183.6, m >220°, > m 300°, $pK_{Est(1)}$ ~3.0, $pK_{Est(2)}$ ~10.0. Purify by recrystallisation from glacial acetic acid or $*C_6H_6/EtOCH_2CH_2OH$, and dry *in vacuo*. Its UV has $\lambda_{max}(\epsilon)$ at 267(14400) (0.1N HCl), 271(15000) (aqueous pH 7.0); and 226(15300) and 272(5340) (0.1N NaOH in EtOH) [Montgomery & Holum *J Am Chem Soc* 80 404 1958, DOI: 10.1021/ja01535a040]. [Kim et al. *J Med Chem* 43 746 2000, DOI: 10.1021/jm9905211; *Beilstein* 26 III/IV 3724.]

2-Chloro-3-methylindole (2-chloroskatole) [51206-73-6] C_9H_8ClN , M 165.6, m 114.5-115.5°, $pK_{Est} <0$. Purify 2-chloroskatole by chromatography on silica gel in CH_2Cl_2 /petroleum ether (1:2), followed by recrystallisation from aqueous EtOH, aqueous AcOH or petroleum ether (m 113.5°). The *picrate* has m 121-122°(dec) (orange-red crystals from petroleum ether). [Phillips & Cohen *J Am Chem Soc* **108** 2023 1986, DOI: 10.1021/ja00268a049; *Beilstein* **20** H 317, **20** III/IV 3212.]

4-(Chloromethyl)pyridine hydrochloride [1822-51-1] $C_6H_6ClN.HCl$, M 164.0, m 166-173°, 170-175°, 172-173°, $pK_{Est} \sim 5.6$. Purify it by recrystallisation from EtOH or EtOH/dry Et_2O . It melts between 171° and 175°, and the clear melt resolidifies on further heating at 190° and turns red to black at 280° but does not melt again. The *picrate-hydrochloride* (prepared in EtOH) has m 146-147°. The *free base* is an oil. [Mosher & Tessieri *J Am Chem Soc* **73** 4925 1951, DOI: 10.1021/ja01154a135; *Beilstein* **20** III/IV 2752.]

2-Chloro-1-methylpyridinium iodide [14338-32-0] C_6H_7ClIN , M 255.5, m 200°(dec), 203-205°, 205-206°(dec), 207°. Purify it by dissolving in EtOH and adding dry Et_2O . The solid is washed with Me_2CO and dried at 20°/0.35mm. Store it in the dark. Attempted recrystallisation from Me_2CO /EtOH/petroleum ether (b 40-60°) causes some exchange of the Cl substituent by I. The *picrate* has m 106-107°, and the *perchlorate* has m 212-213°. [Jones et al. *J Am Chem Soc* **111** 1157 1989, DOI: 10.1021/ja00185a071; UV and solvolysis: Barlin & Benbow *JCS Perkin Trans 2* 790 1974, DOI: 10.1039/P29740000790; *Beilstein* **20/5** V 405.]

6-Chloronicotinic acid [5326-23-8] $C_6H_4ClNO_2$, M 157.6, m 190°(dec), 190-193°, 198-199°(dec), pK^{25} 4.22 (50% aqueous EtOH). Purify it by recrystallisation from hot H_2O and sublime it in a vacuum. [Pechmann & Welsh *Chem Ber* **17** 2384 1884, DOI: 10.1002/cber.188401702153; Herz & Murty *J Org Chem* **26** 122 1961, DOI: 10.1021/jo01060a029; *Beilstein* **22/2** V 177.]

4-Chloro-7-nitrobenzofurazane (7-chloro-4-nitrobenzoxadiazole, NBD-chloride) [10199-89-0] $C_6H_2ClN_3O_3$, M 199.6, m 96.5-97°, 97°, 97-99°, 99-100°. Wash the solid with H_2O , and it recrystallises from aqueous EtOH (1:1) as pale yellow needles. It sublimes in a vacuum [Ghosh & Whitehouse *Biochem J* **108** 155 1968, DOI: 10.1042/bj1080155; UV, NMR: Boulton et al. *J Chem Soc (B)* 1004 1966, DOI: 10.1039/J29660001004].

2-Chloro-3-nitropyridine [5470-18-8] $C_5H_3ClN_2O_2$, M 158.5, m 100-103°, 101-102°, 103-104° (sublimes), pK^{20} -2.6. It forms needles from H_2O . Purify it by continuous sublimation over a period of 2 weeks at 50-60°/0.1mm [Barlin *J Chem Soc* 2150 1964, DOI: 10.1039/JR9640002150]. The *N-oxide* has m 100°(from CH_2Cl_2/Et_2O). [Taylor & Driscoll *J Org Chem* **25** 1716 1960, DOI: 10.1021/jo01080a008; Ochiai & Kaneko *Chem Pharm Bull Jpn* **8** 28 1960, DOI: org/10.1248/cpb.8.28; *Beilstein* **20/5** V 451.]

2-Chloro-5-nitropyridine [4548-45-2] has m 105-108°, 108°, $pK_{Est} \sim 2.6$. It crystallises from *benzene or *benzene/petroleum ether. [*Beilstein* **20/5** V 452.]

N-(3-Chlorophenyl)-6,7-dimethoxyquinazolinamine (Tyrphostin AG 1478) [175178-82-2 *free-base*; 153436-53-4 *hydrochloride*] $C_{16}H_{14}ClN_3O_2$, M 315.7 (free base), decoposes $\geq 275-280$, pK_a (very weak base). This and related bio-active 4-anilinoquinazolines [see Barker & Davies EP 0520722 A1 30 Dec 1992 to Zeneca; Hess US 3511836 A 12 May 1970 to Pfizer & Co] have been prepared from 4-chloro-6,7-dimethoxyquinazoline. The latter was obtained by the following sequence of reaction: veratole (4-hydroxy-3-methoxybenzaldehyde) + $Me_2SO_4 \rightarrow$ 3,4-dimethoxybenzaldehyde + $HNO_3 \rightarrow$ 4,5-dimethoxy-2-nitrobenzaldehyde + $KMnO_4 \rightarrow$ 4,5-dimethoxy-2-aminobenzoic acid + $HNCHO \rightarrow$ 6,7-dimethoxyquinazoline-4-one + $POCl_3/Me_2NPh \rightarrow$ 4-chloro-6,7-dimethoxyquinazoline, or some slight variations of this synthesis [Althuis & Hess *J Med Chem* **20** 146 1977, DOI: 10.1021/jm00211a031; see also Armarego & Reece *Aust J Chem* **34** 1561 1981, DOI: 10.1071/CH9811561]. The purity of the chloroquinazoline is important because impure material deteriorates rapidly. Stable material is best obtained by dissolving the residue, after evaporation of excess $POCl_3$, in CH_2Cl_2 , placing the solution on an Al_2O_3 column and eluting with CH_2Cl_2 followed by increasingly polar mixtures of CH_2Cl_2 and EtOAc. The condensation of 4-chloro-6,7-dimethoxyquinazoline with 3-chloroaniline can be achieved by fusing equimolar amounts at $\sim 100^\circ$ for 5min whereby the mixture melts and then solidifies, dissolving this in the minimum volume of hot EtOH, heat at boiling point for 30min and cool to crystallise out the quinazolinamine hydrochloride which is recrystallised from EtOH or *iso*PrOH. *Alternatively*, equimolar amounts of reagents are stirred with *iso*PrOH which has been heated to 80° and stirred at this

temperature for 30 minutes. On cooling, the product separates and is recrystallised from the same solvent. In another variant Et_3N is added to the *iso*PrOH solution which provides similar yields (65-88%), but in this case the **free base** is formed because Et_3N consumes the HCl liberated from the reaction. **AG 1478** is soluble in 0.1M HCl (0.4w/v%), in EtOH (1.0w/v%), in DMSO-MeOH (1:1, 1w/v%) and in CH_2Cl_2 , but insoluble in H_2O and in 0.1N NaOH. ***N*-(3-Chlorophenyl)-*N*-(6,7-dimethoxyquinazolin-4-yl)-*N*-methylamine hydrochloride** has **m 220-222° (235-237°)**, and ***N*-(3-chlorophenyl)-*N*-(6,7-dimethoxyquinazolin-4-yl)-*N*-ethylamine hydrochloride** has **m 261-263°** [See Myers et al. US Patent 6,645,969 B1, 11 Nov 2003, to Aventis Pharmaceuticals Inc, Armarego *Quinazolines, Fused Pyrimidines Part I* Brown Ed, Wiley-Interscience 1967, Brown *Quinazolines Supplement I* Taylor Ed, Wiley-Interscience 1996, ISBN 0-471-14565-3]

It is a strong and selective inhibitor of epidermal growth factor receptor kinase EGFR ($\text{IC}_{50} = 3\text{nM}$ for EGFR and $> 100\mu\text{M}$ for Her2-neu and EDGFR). [Levitzki & Gazit 'Tyrosine kinase inhibition: an approach to drug development: AG 1478' *Science* **267** 1782 1995, DOI: 10.1126/science.7892601; Han et al. 'Tyrphostin AG 1478 Preferentially Inhibits Human Glioma Cells Expressing Truncated Rather than Wild-Type Epidermal Growth Factor Receptors' *Cancer Res* **56** 3859-3861 1996, DOI: published 1 Sept 1996, not available, <http://cancerres.aacrjournals.org/content/56/17/3859>; Oshero & Levitzki 'Epidermal-Growth-Factor-Dependent Activation of the Src-Family Kinases' *Eur J Biochem (FEBS)* **225** 1047 1994, DOI: 10.1111/j.1432-1033.1994.1047b.x; Eguchi et al. 'Calcium-dependent Epidermal Growth Factor Receptor Transactivation Mediates the Angiotensin II-induced Mitogen-activated Protein Kinase Activation in Vascular Smooth Muscle Cell' *J Biol Chem* **273** 8890 1998, DOI: 10.1074/jbc.273.15.8890; Ward et al. 'Epidermal growth factor receptor tyrosine kinase: Investigation of catalytic mechanism, structure-based searching and discovery of a potent inhibitor' *Biochem Pharmacol* **48** 659 1994, DOI: 10.1016/0006-2952(94)90042-6]

An analogous compound, prepared in a similar manner except that 3-bromoaniline is used instead of 3-chloroaniline is ***N*-(3-bromophenyl)-6,7-dimethoxyquinazolinamine hydrochloride (Tyrphostin AG 1517, PD153035 HCl) [183322-45-4 hydrochloride]** $\text{C}_{16}\text{H}_{14}\text{BrN}_3\text{O}_2 \cdot \text{HCl}$, **M 396.7, decoposes $\geq 275-280^\circ$, pKa check w/la.** AG 1517 has chemical and physical properties similar to those of AG 1478, and is also a potent and selective ATP competitive inhibitor of the epidermal growth factor receptor tyrosine kinase EGFR.

9-Chloro-9-phenylxanthene (Pixyl chloride) [42506-03-6] $\text{C}_{19}\text{H}_{13}\text{ClO}$, **M 292.8, m 102-106°, 105-106°.** A possible impurity is 9-hydroxy-9-phenylxanthene. If the material contains a lot of the hydroxy product, then boil 10g of it in CHCl_3 (50ml) with redistilled acetyl chloride (1ml) until liberation of HCl is complete. Evaporation leaves the chlorophenylxanthene as the hydrochloride which on heating with *benzene loses HCl; and on adding petroleum ether prisms of chlorophenylxanthene separate and contain 0.5mol of *benzene. The *benzene-free compound is obtained on drying, and it melts to a colourless liquid. [Gomberg & Cone *Justus Liebigs Ann Chem* **370** 142 1909, DOI: 10.1002/jlac.19093700110.] The 9-phenylxanthyl group is called 'pixyl' and is a good protecting group [Chattopadhyaya & Reese *JCS Chem Commun* 639 1978, DOI: 10.1039/C39780000639; *Beilstein* **17** III/IV 1704.]

6-Chloropurine [87-42-3] $\text{C}_5\text{H}_3\text{ClN}_4$, **M 154.6, m 175-177°(dec), 179°(dec), $>300^\circ$ (dec), $\text{pK}_1^{20} 0.45$, $\text{pK}_2^{20} 7.88$.** 6-Chloropurine crystallises from water (solubility is 0.5% at $\sim 20^\circ$). The UV in water at pH 1 has λ_{max} 264nm ($\log \epsilon$ 3.94), at pH 5.2 has 56nm ($\log \epsilon$ 3.96), and at pH ~ 13 has 274nm ($\log \epsilon$ 3.94). [Lister *Purines Part II, Fused Pyrimidines* Brown Ed, Wiley-Interscience pp254-225 1971, ISBN 0-471-38205-1, *Beilstein* **26** III/IV 1742.]

2-Chloropyrazine [14508-49-7] $\text{C}_4\text{H}_3\text{ClN}_2$, **M 114.5, b 62-63°/31mm, 153-154°/atm, $d_4^{20} 1.302$, $n_D^{24} 1.535$, $\text{pK}_{\text{Est}} < 0$.** Fractionally distil it through a short column packed with glass helices. It has a penetrating, mildly pungent odour with a high vapour pressure at room temperature. [Erickson & Spoerri *J Am Chem Soc* **68** 400 1946, DOI: 10.1021/ja01207a018; Hetman & O'Donnell *J Org Chem* **28** 1682 1963, DOI: 10.1021/jo01041a064; *Beilstein* **23/5** V 366.]

2-Chloropyridine [109-09-1] $\text{C}_5\text{H}_4\text{ClN}$, **M 113.6, b 49.0°/7mm, 166°/714mm, $d_4^{20} 1.20$, $n_D^{20} 1.532$, $\text{pK}^{20} 0.49$ (0.72).** Dry 2-chloropyridine with NaOH for several days, then distil it from CaO under reduced pressure. [*Beilstein* **20/5** V 402.]

3-Chloropyridine [626-60-8] $\text{C}_5\text{H}_4\text{ClN}$, **M 113.6, b 148°/atm $d_4^{20} 1.194$, $n_D^{20} 1.533$, $\text{pK}^{20} 2.84$.** Distil 3-chloropyridine from KOH pellets. [*Beilstein* **20/5** V 406.]

4-Chloropyridine [626-61-9] C_5H_4ClN , M 113.6, b 85-86°/100mm, 147-148°/760mm, pK^{20} 3.84. Pour 4-chloropyridine into distilled water, and excess of 6M NaOH is added to give pH 12. The organic phase is separated and extracted with four volumes of diethyl ether. The combined extracts are filtered through paper to remove water, and the solvent is evaporated. The dark brown residual liquid is kept under high vacuum [Vaidya & Mathias *J Am Chem Soc* **108** 5514 1986, DOI: 10.1021/ja00278a024]. It can be distilled, but readily darkens and is best kept as the *hydrochloride* [7379-35-3] M 150.1, m 163-165°(dec). [Beilstein **20/5** V 410.]

2-Chloropyrimidine [1722-12-9] $C_4H_3ClN_2$, M 114.5, m 63-66°, 63-65°, 66°, b 91°/26mm, pK^{20} -1.90. It has been recrystallised from $*C_6H_6$, petroleum ether or a mixture of both. It sublimes at 50°/18mm and can be distilled in a vacuum. [IR: Short & Thompson *J Chem Soc* 168 1952, DOI: 10.1039/JR9520000168; Boardland & McOmie *J Chem Soc* 1218 1951, DOI: 10.1039/JR9510001218; Beilstein **23/5** V 343.]

2-Chloroquinoline [612-62-4] C_9H_6ClN , M 163.6, m 34°, 34-37°, b 147-148°/15mm, 266-267°/atm, d_4^{35} 1.235, n_D^{25} 1.629, pK_{Est} ~0.3. Purify it by crystallisation of its *picrate* to constant melting point (123-124°) from $*benzene$, regenerating the base and distilling it under vacuum [Cumper et al. *J Chem Soc* 1183 1962, DOI: 10.1039/JR9620001183]. 2-Chloroquinoline can be crystallised from EtOH. Its *picrate* has m 123-124° (from EtOH). [Beilstein **20** H 359, 20/7 V 312.]

4-Chloroquinoline [611-35-8] has m 28-31°, 29-32°, 31°, b 130°/15mm, 261°/744mm, pK^{25} 3.72. Possible impurities include the 2-isomer. It is best purified by converting to the *picrate* (m 212-213° dec) in EtOH and recrystallising it from EtOH (where the *picrate* of the 2-chloroquinoline remains in solution) or EtOAc. The *picrate* is decomposed with 5% aqueous NaOH, extracted in $CHCl_3$, washed with H_2O , dried ($MgSO_4$), evaporated and distilled in a vacuum. It can be steam distilled from slightly alkaline aqueous solutions, the aqueous distillate is extracted with Et_2O , evaporated and distilled. The distillate solidifies on cooling. [Bobránski *Chem Ber* **71** 578 1938, DOI: 10.1002/cber.19380710312; Beilstein **20/7** V 314.]

8-Chloroquinoline [611-33-6] C_9H_6ClN , M 163.6, b 171-171.5°/24mm, d_4^{20} 1.278, n_D^{20} 1.644, pK^{20} 3.12. Purify it by crystallisation of its $ZnCl_2$ complex (m 228°) from aqueous EtOH. The *free base* is then liberated with excess of cold aqueous NaOH, whereby the soluble zincate is formed, and the chloroquinoline is isolated by extraction into Et_2O , evaporated and distilled. [Beilstein **20** III/IV 3381, 20/7 V 315.]

5-Chloroquinolin-8-yl trifluoromethanesulfonate (5-chloro-8-quinoline triflate) [157437-38-2] $C_{10}H_5ClF_3NO_3S$, M 311.7, m 79-83°. If it is discoloured dissolve it in CH_2Cl_2 , wash it with N NaOH and half saturated K_2CO_3 , dry the organic layer over solid K_2CO_3 , filter, evaporate and dry the solid *in vacuo*, (cf. 2-methyl-5-pyridine triflate). [Matthew et al. *Tetrahedron Lett* **35** 5177 1994, DOI: 10.1016/S0040-4039(01)85088-7; Tilley & Zawoiski *J Org Chem* **53** 386 1988, DOI: 10.1021/jo00237a029; Ellingboe et al. *J Med Chem* **37** 542 1994, DOI: 10.1021/jm00030a013.]

8-Chlorotheophylline (8-chloro-1,3-dimethyl-2,6(1H,3H)-purinedione) [85-18-7] $C_7H_7ClN_4O_2$, M 214.6, m 290°(dec), 311°(dec), $pK_{Est(1)}$ ~5.4, $pK_{Est(2)}$ ~9.1. It crystallises from H_2O or EtOH (m 304° dec). The *choline salt* crystallises from H_2O with m 60-62° (2 H_2O) and m 97-99° (anhydrous). [Lister *Purines Part II, Fused Pyrimidines* Brown Ed, Wiley-Interscience pp254-225 1971, ISBN 0-471-38205-1, Beilstein **26** H 473, II 276, 26 III/IV 2442.]

2-Chlorothiophene (2-thienyl chloride) [96-43-5] C_4H_3ClS , M 118.6, b 126-128°/atm, 127-129°/atm, 128°~760mm, d_4^{20} 1.285, n_D^{20} 1.551. Purify it by fractional distillation at atmospheric pressure or by gas chromatography. [Conde et al. *Synthesis* 412 1976, DOI: 10.1055/s-1976-24065; Beilstein **17/1** V 303.]

5-Chlorouracil (5-chloro-2,4(6)-dihydroxypyrimidine) [1820-81-1] $C_4H_3ClN_2O_2$, M 146.5, m 314-418°(dec), 324-325°(dec), pK_1^{25} 7.95, pK_2^{25} >13. It recrystallises from hot H_2O (4g/500ml) using charcoal. [Chesterfield et al. *J Chem Soc* 3478 1955, DOI: 10.1039/JR9550003478; West & Barrett *J Am Chem Soc* **76** 3146 1954, DOI: 10.1021/ja01641a009; Beilstein **24** III/IV 1231.]

4-Chromanone (2,3-dihydro-4H-1-benzopyran-4-one) [491-37-2] $C_9H_8O_2$, M 148.2, m 35-37°, 39°, 41°, b 92-93°/3mm, 127-128°/13mm, 130-132°/15mm, 160°/50mm. It has been recrystallised from petroleum ether, or purified by dissolving in $*C_6H_6$ washing with H_2O , drying ($MgSO_4$), evaporating and distilling in a vacuum,

then recrystallising the residue. The liquid has a pleasant lemon-like odour. The *semicarbazone* has **m 227°**. [Loudon & Razdan *J Chem Soc* 4299 1954, DOI: 10.1039/JR9540004299.] The *oxime* is prepared from 3g of chromanone, 3g NH₂OH.HCl in EtOH (50ml), 6g K₂CO₃ and refluxed on a water bath for 6 hours. The solution is poured into H₂O, the solid is filtered off, dried and dissolved in hot *C₆H₆ which on addition of petroleum ether yields the *oxime* as glistening needles **m 140°**. Hydrolysis of this gives very pure chromanone. The *benzal derivative* is prepared from 3g of chromanone and 4g PhCHO in 50ml EtOH, heated to boiling, 10ml of conc HCl are added dropwise and set aside for several days. The derivative separates and is recrystallised from EtOH to give yellow needles, **m 112°** [Powell *J Am Chem Soc* 45 2708 1923, DOI: 10.1021/ja01664a033]. Reaction with Pb(OAc)₄ yields the *3-acetoxy derivative* **m 74°** (from petroleum ether + trace of EtOAc) [Cavill et al. *J Chem Soc* 4573 1954, DOI: 10.1039/JR9540004573; *Beilstein* 17/10 V 14].

Cinnoline [253-66-7] C₈H₆N₂, **M 130.2**, **m 38°, 40-41°, b 114°/0.35**, **pK²⁰ 2.37**. Distil at high vacuum, then recrystallised from petroleum ether. It forms an etherate **m 24-25°**. Keep under N₂ in sealed tubes in the dark at 0°. The *hydrochloride* [5949-24-6] **M 166.6** has **m 156-158°** (yellow *monohydrate*, from EtOH/Et₂O), and dehydrates on sublimation at 110-115°/3mm. The *picrate* has **m 196-196.5°**. [D.J. Brown *Cinnolines and Phthalazines* Suppl II 2005, Wiley & Sons NJ. ISBN-13: 978-0-471-48587-2; *Beilstein* 23 III.IV 1217.]

Citrazinic acid (2,6-dihydroxyisonicotinic acid) [99-11-6] C₆H₅NO₄, **M 155.1**, **m >300°, pK₁ 3.0, pK₂ 4.76**. The acid is normally a yellow powder with a greenish shade, but is white when ultra pure and turns blue on long standing. It is insoluble in H₂O but slightly soluble in hot HCl and soluble in alkali or carbonate solutions. It is purified by precipitation from alkaline solutions with dilute HCl, and dried in a vacuum over P₂O₅. The *ethyl ester* has **m 232°** (evacuated tube) and a pK_a of 4.81 in MeOCH₂CH₂OH [IR: Pitha *Coll Czech Chem Commun* 28 1408 1963, DOI: org/10.1135/cccc19631408]. [*Beilstein* 22/7 V 24.]

Clioquinol (5-chloro-8-hydroxy-7-iodoquinoline) [130-26-7] **M 305.5**, **m 181°, pK₁²⁵ 2.7, pK₂²⁵ 7.9**. It crystallises from AcOH or xylene, and dry it at 70° *in vacuo*. [*Beilstein* 21 III/IV 1190.]

Conessine [546-06-5] C₂₄H₄₀N₂, **M 356.6**, **m 125°, 127-128.5°, [α]_D²⁰ -1.9 (in CHCl₃) and +26 (c 3, EtOH), pK_{Est(1)} ~10.4, pK_{Est(2)} ~10.7**. It crystallises from acetone, sublimes at 95°/0.01mm and boils at 0.1mm with bath temperature at 220°. The *dihydrochloride* has **m >340°** (browns at 235° and decomposes at 338-240°) and has [α]_D²⁰ +9.3 (c 2, H₂O). The (+)-*dihydrobromide* [5913-82-6], **m ~340°**, crystallises from EtOH/Et₂O. [Marshall & Johnson *J Am Chem Soc* 84 1485 1962, DOI: 10.1021/ja00867a026; *Beilstein* 22 III/IV 4382.]

Coproporphyrin I [531-14-6] C₃₆H₃₈N₄O₈, **M 654.7**, **λ_{max} 591, 548, 401nm in 10% HCl**. It crystallises from pyridine/glacial acetic acid. The *dihydrochloride* [69477-27-6] has **M 727.6** and **λ_{max}** at 395nm in water. [*Beilstein* 26 III/IV 3094.]

Coumalic acid (2-pyrone-5-carboxylic acid) [500-05-0] C₆H₄O₄, **M 140.1**, **m 203-205°(dec), 205-210°(dec), 218°/120mm, pK_{Est} ~0**. The acid crystallises from MeOH. The *methyl ester* has **m 73-74°** (from petroleum ether) and **b 178-180°/60 mm**. [*Beilstein* 18/8 V 120.]

Coumarin [91-64-5] C₉H₆O₂, **M 146.2**, **m 68-69°, 68-70°, 68-73°, b 68-73°, 139°/5mm, pK²⁵ -4.97 (aqueous H₂SO₄)**. Coumarin crystallises from ethanol or water (solubility is 0.25w/v% at ~20°, and 2.0w/v% ~100°), and is soluble in organic solvents and alkaline solutions. It sublimes *in vacuo* at 43° [Srinivasan & De Levie *J Phys Chem* 91 2904 1987, DOI: 10.1021/j100295a050]. [*Beilstein* 17/10 V 143.] It is a flavouring (vanilla) agent. **Coumarin-3-carboxylic acid** [531-81-7] C₁₀H₆O₄, **M 190.2**, has **m 188°(dec), 189-192°(dec), pK_{Est} ~1.5**, and crystallises from water. [*Beilstein* 18/8 V 323.]

γ-Crotonolactone [2(5H)-furanone] [497-23-4] C₄H₄O₂, **M 84.1**, **m 3-4°, 76-77°/3.5mm, 90.5-91°/11.5mm, 92-93°/14mm, 107-109°/24mm, 212-214°/760mm, d₄²⁰ 1.197, n_D²⁰ 1.470**. Fractionally distil the lactone under reduced pressure. Its IR(CCl₄) has 1784 and 1742 cm⁻¹, UV no max above 205nm (ε 1160 cm⁻¹ M⁻¹) and ¹HNMR (CCl₃) has τ at 2.15 (pair of triplets 1H), 3.85 (pair of triplets 1H) and 5.03 (triplet 2H). [Price & Judge *Org Synth Coll Vol* 5 255 1973, DOI: 10.15227/orgsyn.045.0022; Jones et al. *Can J Chem* 37 2007, DOI: 10.1139/v59-293; 2092 1959, Smith & Jones *Can J Chem* 37 2092 1959, DOI: 10.1139/v59-306; *Beilstein* 17/9 V 112.]

15-Crown-5 [33100-27-5] $C_{10}H_{20}O_5$, M 220.3, b 93-96°/0.1mm, d_4^{20} 1.113, n_D^{20} 1.465. Dry it over 3A molecular sieves and distil it in a high vacuum. [Beilstein 19/12 V 252.]

18-Crown-6 [17455-13-9] $C_{12}H_{24}O_6$, M 264.3, has m 37-39°, 42-45°. Recrystallise it from acetonitrile and dry it in a vacuum. Purify it also by precipitating the 18-crown-6/nitromethane 1:2 complex with Et₂O/nitromethane (10:1 mixture). The complex is decomposed in vacuum whereby 18-crown-6 distils off under the reduced pressure. [Beilstein 19/12 V 601.] It is a complexing agent with cations which solubilises them in non-polar solvents, and is useful as a phase-transfer catalyst in aqueous-organic mixed solvents [Gokel *Crown Ethers and Cryptands* The Royal Society of Chemistry (Cambridge, England 1991), Gokel et al. *Org Synth Coll Vol* 6 301 1988, DOI: 10.15227/orgsyn.057.0030; cf. Kotha & Kashinath *Synthesis* 971 2005(6), DOI: 10.1055/s-2005-861840].

Cryptopine [482-74-6] $C_{21}H_{23}NO_5$, M 369.4, m 220-221°, 220-223°, pK²⁵ 8.09. It crystallises from *benzene, hot EtOH (0.25% cold, 1.2% at boiling), petroleum ether or methyl ethyl ketone. It is a relatively strong organic base — store in the absence of CO₂ in the dark. The *perchlorate* crystallises from aqueous MeOH with m 226-228°(dec). [Thomas et al. *Can J Chem* 33 570 1955, DOI: 10.1139/v55-067; Haworth & Perkin *J Chem Soc* 1769 1926, DOI: 10.1039/JR9262901769; Beilstein 27 III/IV 6652.]

Cupreine (6'-hydroxycinchonidine) [524-63-0] $C_{19}H_{22}N_2O_2$, M 310.4, m 202°(anhydrous), $[\alpha]_D^{17}$ -176 (c 0.5, MeOH), pK¹⁵ 7.63 (6.57). Cupreine crystallises from EtOH (*anhydrous* crystals) and wet Et₂O (as *dihydrate* crystals). It has K_b 2.7x10⁻⁷ [Kolthoff *Biochem Z* 162 323]. The *sulfate* forms needles, m 257°(dec), from MeOH, amyl alcohol or H₂O, with $[\alpha]_D^{20}$ -197.9 (c 1.2, H₂O). [Beilstein 22 I 165, 22 II 416.]

5-Cyanoindole [15861-24-2] $C_9H_6N_2$, M 142.2, m 106-108°, 107-108°, pK²⁵ <0. Dissolve the nitrile in 95% EtOH, boil it in the presence of charcoal, filter, evaporate to a small volume and add enough H₂O to cause crystallisation and cool. Recrystallise it directly from aqueous EtOH and dry it in a vacuum. Its UV has λ_{max} at 276 nm (log ϵ 3.6) in MeOH. [Lindwall & Mantell *J Org Chem* 18 345 1953, DOI: 10.1021/jo01132a001; Singer & Shive *J Org Chem* 1458 1955, DOI: 10.1021/jo01127a027; Thesing et al. *Chem Ber* 95 2205 1962, DOI: 10.1002/cber.19620950917; NMR: Lallemand & Bernath *Bull Soc Chim Fr* 4091 1970, Beilstein 22/3 V 45.]

2-Cyanopyridine [100-70-9] $C_6H_4N_2$, M 104.1, m 24-27°, 27°, 29°, b 212-215°/atm, pK²⁵ 0.26. Purify the 2-nitrile by distillation followed by recrystallisation to constant melting point from cyclohexane, or *o*-xylene/hexane. The UV has λ_{max} , ϵ_{max} (solvent): $\pi \rightarrow \pi$ bands at 265nm, 2730; $n \rightarrow \pi$ infl at 278nm, 340 (cyclohexane). [Mason *J Chem Soc* 1247 1959, DOI: 10.1039/JR9590001247, Beilstein 22/2 V 19.]

3-Cyanopyridine [100-54-9] has m 48-52°, 50°, b 201°/atm, pK²⁵ 1.26. Purify the 3-nitrile by distillation followed by recrystallisation to constant melting point from cyclohexane or *o*-xylene/hexane. The UV has λ_{max} , ϵ_{max} (solvent): $\pi \rightarrow \pi$ bands at 265nm, 2230; $n \rightarrow \pi$ infl at 279nm, 430 (cyclohexane). [Mason *J Chem Soc* 1247 1959, DOI: 10.1039/JR9590001247, Beilstein 22/2 V 115.]

4-Cyanopyridine [100-48-1] has m 76-79°, 79°, 80°, b 194-196°/atm, pK²⁵ 1.90. Purify the 4-nitrile by distillation followed by recrystallisation to constant melting point from cyclohexane or dichloromethane/diethyl ether mixture. The UV has λ_{max} , ϵ_{max} (solvent): $\pi \rightarrow \pi$ bands at 271nm, 2840; $n \rightarrow \pi$ infl at 290nm, 500 (cyclohexane). [Mason *J Chem Soc* 1247 1959, DOI: 10.1039/JR9590001247, Beilstein 22/2 V 214.]

Cyanuric acid (2,4,6-trihydroxy-1,3,5-triazine) [108-80-5] $C_3H_3N_3O_3$, M 129.1, m >300°, pK²⁵ 6.78. It crystallises from water. Dry it at room temperature in a desiccator in a vacuum. [Beilstein 26 III/IV 632] It is a disinfectant.

Cyanuric chloride (TCT, 2,4,6-trichloro-1,3,5-triazine) [108-77-0] $C_3Cl_3N_3$, M 184.4, has m 146-149°, 154°, b 190°. TCT crystallises from CCl₄ or petroleum ether (b 90-100°) and is dried under vacuum. It has also been recrystallised twice from anhydrous *benzene immediately before use [Abuchowski et al. *J Biol Chem* 252 3582 1977, <http://www.jbc.org/content/254/11/4856>]. [Beilstein 26 III/IV 66.]

Cyclohexene oxide (7-oxabicyclo[4.1.0]heptane) [286-20-4] $C_6H_{10}O$, M 98.1, b 131-133°/atm, d_4^{20} 0.971, n_D^{20} 1.452. Fractionate the oxide through an efficient column. The main impurity is probably H₂O. Dry the oxide over MgSO₄, filter it, and redistil it several times (b 129-134°/760mm). The residue can be hard to remove from the distilling flask. To avoid this difficulty, add a small amount of a mixture of ground NaCl

and Celite (1:1) to help break up the residue particularly if hot H₂O is added. [Osterberg *Org Synth Coll Vol I* 185 1948, DOI: 10.15227/orgsyn.005.0035; *Beilstein* 17 H 21, 17/I V 203.]

Cyclohexene sulfide [7-thiabicyclo[4.1.0]heptane] [286-28-2] C₆H₁₀S, M 114.2, b 55-58°/12mm, 67-68°/16mm, 69-71°/19mm, 71.5-73.5°/21mm, 73-74.5°/22mm, 80°/40mm, 83-87°/46mm, d₄²⁰ 0.971, n_D²⁰ 1.5292. Prepared by reaction of cyclohexeneoxide with thiourea, KCNS or NH₂CNS in EtOH at 60° for 1 to 3hrs, poured into H₂O extracted with CHCl₃, then evaporated and the residual oil distilled (short Vigreux column) to give 41 to 73% yields of the sulfide. It can be stored at 5° in a closed container for at least 1 month without apparent decomposition. Its UV (iso-octane) has λ_{max} 262.5nm (ε_{max} 39.6 L.mole⁻¹.cm⁻¹) and λ_{min} 237nm (ε_{min} 19.7 L.mole⁻¹.cm⁻¹). [Synth: Van Tamelen *Org Synth Coll Vol 4* 232 1963; DOI: 10.15227/orgsyn.032.0039; Synth and reactions: Culvenor et al. *J Chem Soc* 1050 1946, DOI: 10.1039/JR9460001050; UV: Davis *J Org Chem* 23 216 1958, DOI: 10.1021/jo01096a017; synth and reactions: van Tamelen *J Am Chem Soc* 73 3444 1951, DOI: 10.1021/ja01151a132; *Beilstein* 17 III 166, 17/I V 204.]

Cycloheximide (actidione) [68-81-9] C₁₅H₂₃NO₄, M 281.4, m 119.5-121°, [α]₅₄₆²⁰ +9.5 (c 2, H₂O), pK 11.2. Crystallise it from water/MeOH (4:1), amyl acetate, isopropyl acetate/isopropyl ether or water. The *acetate* has m 150-152° (from aqueous EtOH), the *p-nitrobenzoate* has m 215-220°(dec) (from aqueous dioxane) and the *oxime* has m 203-204° (from MeOH). [Kornfeld et al. *J Am Chem Soc* 71 150 1949, DOI: 10.1021/ja01169a041; *Beilstein* 21 IV 6632, 21/13 V 434.]

1-Cyclohexyl-5-methyl-1H-tetrazole [7707-57-5] M 166.2, m 124-124.5°. Crystallise it from absolute EtOH or H₂O (heavy needles), then sublime it at 115°/3mm. [Harvill et al. *J Org Chem* 15 662, 668 1950, DOI: 10.1021/jo01149a035; Billuber Inc USP 2507337 1946, *Beilstein* 26 III/IV 1661.]

cis-Cyclooctene oxide {(1r, 8c)-9-oxabicyclo[6.1.0]nonane} [286-62-4] C₈H₁₄O, M 126.2, m 53-57°, 56-57°, 57.5-57.8°, 50-60°, b 55°/5mm, 85-88°/17mm, 82.5°/22mm, 90-93°/37mm, 189-190°/atm. It can be distilled in a vacuum, and the solidified distillate can be sublimed in a vacuum below 50°. It has a characteristic odour. [IR: Cope et al. *J Am Chem Soc* 74 5884 1952, DOI: 10.1021/ja01143a018; cf. *trans-isomer*: Cope et al. *J Am Chem Soc* 79 3905 1957, DOI: 10.1021/ja01571a075; Reppe et al. *Justus Liebig's Ann Chem* 560 1 1948, DOI: 10.1002/jlac.19485600102; *Beilstein* 17 VII 234.]

Cyclotrimethylenetrinitramine (RDX, Cyclonite, 1,3,5-trinitrohexahydro-1,3,5-triazine) [121-82-4] C₃H₆N₆O₆, M 222.2, m 203.8°(dec), 205-206°(dec). RDX crystallises from acetone (solubility at ~20° is ~4w/v%), but is less soluble in MeOH. [Bachmann & Sheehan *J Am Chem Soc* 71 1842 1949, DOI: 10.1021/ja01173a092; *Beilstein* 26 II 5, 26 III/IV 22.] **EXPLOSIVE.**

Cytisine (7R,9S-7,9,10,11,12,13-hexahydro-7,9-methano-12H-pyrido[1,2-a][1,5]diazocin-8-one, Laburnine, Ulexine) [485-35-8] C₁₁H₁₄N₂O, M 190.3, m 152-153°, 155°, b 218°/2mm, [α]_D¹⁷ -120 (H₂O), [α]_D²⁵ -115 (c 1, H₂O), pK₁¹⁵ 1.20, pK₂¹⁵ 8.12 [also stated are pK₁ 6.11, pK₂ 13.08]. Crystallise cytisine from acetone and sublime it in a vacuum. Its solubilities are: 77% (H₂O), 7.7% (Me₂CO), 28.6% (EtOH), 3.3% (*C₆H₆), 50% (CHCl₃) but it is insoluble in petroleum ether. The *tartrate* has m 206-207° [α]_D²⁴ +45.9, the *N-tosylate* has m 206-207°, and the *N-acetate* has m 208°. [Bohlmann et al. *Angew Chem* 67 708 1955, DOI: 10.1002/ange.19550672214; van Tamelen & Baran *J Am Chem Soc* 77 4944 1955, DOI: 10.1021/ja01623a090; Isolation: Ing *J Chem Soc* 2195 1931, DOI: 10.1039/JR9310002195; Govindachari et al. *J Chem Soc* 3839 1957, DOI: 10.1039/JR9570003839; Abs config: Okuda et al. *Chem Ind (London)* 1751 1961, *Beilstein* 24 H 134, 24 I 244, 24 II 70, 24 III/IV 321.] **TOXIC.**

cis-Decahydroisoquinoline [2744-08-3] C₉H₁₇N, M 139.2, b 97-98°/15mm, 208-209°/730mm, pK²⁰ 11.32. The free base is treated with saturated aqueous picric acid, allowed to stand for 12 hours, filtered, washed with MeOH to remove the more soluble *trans isomer* and recrystallised from MeOH to give pure *cis-picrate* m 149-150°. The picrate (~5g) is shaken with 5M aqueous NaOH (50ml) and Et₂O (150ml) while H₂O is added to the aqueous phase to dissolve insoluble Na picrate. The Et₂O extract is dried over solid NaOH and then shaken with Al₂O₃ (Merck for chromatography) until the yellow color of traces of picric acid disappears (this color cannot be removed by repeated shaking with 5-10 M aqueous NaOH). The extract is

concentrated to 50ml and dry HCl is bubbled through until separation of the white crystals of the **cis-HCl** is complete. These are washed with Et₂O, dried at 100° and recrystallised from EtOH/EtOAc to yield pure **cis-hydrochloride m 182-183°** (dried in a vacuum desiccator over KOH) with IR (KBr) ν_{\max} 2920, 2820, 1582, 1470, 1445, 1410, 1395, 1313, 1135, 1080, 990, 870 cm⁻¹. The pure **free base** is prepared by dissolving the **hydrochloride** in 10 M aqueous NaOH, extracted with Et₂O, dried over solid KOH, filtered and distilled in a vacuum. It has IR (film) ν_{\max} 2920, 2820, 2720, 2560, 1584, 1470, 1445, 1415, 1395, 1315, 1300, 1135, 1080, 1020, 990, 873 cm⁻¹. The ¹HNMR in CDCl₃ is characteristically different from that of the **trans-isomer**. [Armarego *J Chem Soc (C)* 377 1967, DOI: 10.1039/J39670000377; Gray & Heitmeier *J Am Chem Soc* **80** 6274 1958 DOI: 10.1021/ja01556a028; Witkop *J Am Chem Soc* **70** 2617 1948, DOI: 10.1021/ja01188a001; Skita *Chem Ber* **57** 1977 1924, DOI: 10.1002/cber.19240571103; Helfer *Helv Chim Acta* **6** 785 1923, DOI: 10.1002/hlca.19230060186; Beilstein **20** II 73, **20** III/IV 2026.]

trans-Decahydroisoquinoline [2744-09-4] C₉H₁₇N, M **139.2**, b **106°/15mm**, pK²⁰ **11.32**. This is purified as the **cis-isomer** above. The **trans-picrate** has m **175-176°**, and the **trans-hydrochloride** has m **221-222°** and has IR (KBr) ν_{\max} at 2930, 3800, 1589, 1450, 1400, 1070, 952, 837 cm⁻¹. The pure **free base** is prepared as above and had IR (film) with ν_{\max} at 2920, 2820, 2720, 2560, 1584, 1470, 1445, 1415, 1395, 1315, 1300, 1135, 1080, 1020, 990, 873 cm⁻¹. The ¹HNMR in CDCl₃ is characteristically different from that of the **cis-isomer**. (references as above and Helfer *Helv Chim Acta* **9** 814 1926, DOI: 10.1002/hlca.192600901110). [Beilstein **20** II 73, **20** III/IV 2026.]

cis-Decahydroquinoline [10343-99-4] C₉H₁₇N, M **139.2**, b **207-208°/708mm**, pK²⁰ **11.29**. It is available as a **cis-trans-mixture** (b 70-73°/10mm, Aldrich, ~ 18% **cis-isomer** [2051-28-7]), but the isomers can be separated by fractionating in a spinning band column (1~1.5 metre, type E) at atmospheric pressure and collecting 2ml fractions with a distillation rate of 1 drop in 8-10 seconds. The lower boiling fraction solidifies and contains the **trans-isomer** (see below, m **48°**). The higher boiling fraction b 207-208°/708mm remains liquid and is mostly pure **cis-isomer**. This is reacted with PhCOCl and M aqueous NaOH to yield the **N-benzoyl derivative m 96°** after recrystallisation from petroleum ether (b 80-100°). It is hydrolysed with 20% aqueous HCl by refluxing overnight. PhCO₂H is filtered off, the filtrate is basified with 5M aqueous NaOH and extracted with Et₂O. The dried extract (Na₂SO₄) is saturated with dry HCl gas, and the **cis-decahydroquinoline hydrochloride** which separates has m **222-224°** after washing with Et₂O and drying at 100°; and has IR (KBr) with ν_{\max} at 2900, 2780, 2560, 1580, 1445, 1432, 1403, 1165, 1080, 1036, 990, 867 cm⁻¹. The **free base** is obtained by dissolving the **hydrochloride** salt in 5M aqueous NaOH, extracting with Et₂O and drying the extract (Na₂SO₄), evaporating and distilling the residue; it has IR (film) with ν_{\max} at 2900, 2840, 2770, 1445, 1357, 1330, 1305, 1140, 1125, 1109, 1068, 844 cm⁻¹. The ¹H NMR in CDCl₃ is characteristically different from that of the **trans-isomer**. [Armarego *J Chem Soc (C)* 377 1967, DOI: 10.1039/J39670000377; Hüchel & Stepf *Justus Liebigs Ann Chem* **453** 163 1927, DOI: 10.1002/jlac.19274530111; Bailey & McElvain *J Am Chem Soc* **52** 4013 1930, DOI: 10.1021/ja01373a037; Beilstein **20** H 157, **20** I 35, **20** II 72-73, **20** III/IV 2017.]

trans-Decahydroquinoline [767-92-0] has m **48°**, b **205-206°/708mm**, pK²⁰ **11.29**. The lower boiling fraction from the preceding spinning band column fractionation of the commercial **cis-trans-** mixture (~ 20:60; see the **cis-isomer** above) solidifies readily (m **48°**), and the receiver has to be kept hot with warm water. It is further purified by conversion to the **hydrochloride m 285-286°** after recrystallisation from EtOH/AcOEt. This has IR (KBr) with ν_{\max} at 2920, 2760, 2578, 2520, 1580, 1455, 1070, 1050, 975, 950, 833 cm⁻¹. The **free base** is prepared as for the **cis-isomer** above and distilled; and has IR (film, at ca 50°) with ν_{\max} at 2905, 2840, 2780, 1447, 1335, 1305, 1240, 1177, 1125, 987, 900, 835 cm⁻¹. The ¹HNMR in CDCl₃ is characteristically different from that of the **cis-isomer**. [Armarego *J Chem Soc (C)* 377 1967, DOI: 10.1039/J39670000377; Hüchel & Stepf *Justus Liebigs Ann Chem* **453** 163 1927, DOI: 10.1002/jlac.19274530111; Bailey & McElvain *J Am Chem Soc* **52** 4013 1930, DOI: 10.1021/ja01373a037; Prelog & Szpilfogel *Helv Chim Acta* **28** 1684 1945, DOI: 10.1002/hlca.6602801233; Beilstein **20** H 157, **20** I 35, **20** II 72-73, **20** III/IV 2017.]

Delphinine [561-07-9] C₃₃H₄₅NO₉, M **559.7**, m **197-199°**, [α]_D²⁰ **+26 (c 1, EtOH)**. It crystallises from EtOH with 198-200° uncorrected (187.5-188.5°) and from Et₂O or Me₂CO. Its solubility at ~25° in EtOH, CHCl₃ and Et₂O is 4%, 5% and 10% respectively. In c 1, EtOH, it has [α]_D²⁰ **+22** changing to **+19** in 4 hours. [Markwood *J Am Pharm Assoc* **16** 928 1927.] The **hydrochloride** has m **214°(dec)** from MeOH or MeOH/Et₂O. [Weisner et al. *Can J Chem* **50** 1925 1972, DOI: 10.1139/v72-308; Jacobs & Craig *J Biol Chem* **127** 361 1939, <http://www.jbc.org/content/127/2/361>; Beilstein **21** III/IV 2867.]

3,6-Diaminoacridine hydrochloride [952-23-8] $C_{13}H_{11}N_3 \cdot HCl$, M 245.7, m 270°(dec), ϵ_{456} 4.3×10^4 , pK_1^{20} 1.5, pK_2^{20} 9.60 (9.65 free base). It is first purified by precipitation of the *free base* by adding aqueous NH_3 solution to an aqueous solution of the hydrochloride or hydrogen sulfate (see below), drying the precipitate and subliming at 0.01mm Hg [Müller & Crothers *Eur J Biochem*, **54** 267 1975]. The *free base* in EtOH is treated with ethanolic HCl, and Et_2O is added to crystallise out the hydrochloride which is filtered off washed with Et_2O and dried *in vacuo* over solid KOH. [Beilstein **22** H 487.]

3,6-Diaminoacridine sulfate (proflavin sulfate) [1811-28-5] $C_{13}H_{11}N_3 \cdot 0.5H_2SO_4 \cdot xH_2O$, M 516.6, has m >300°(dec), λ_{max} 456nm. An aqueous solution, after treatment with charcoal, is concentrated, chilled overnight, filtered and the precipitate is rinsed with a little diethyl ether. The precipitate is dried in air, then overnight in a vacuum oven at 70°. [Beilstein **22** I 650, **22/11** V 323.]

4,5-Diamino-2,6-dihydroxypyrimidine (diamino uracil) sulfate [32014-70-3] $C_4H_6N_4O_2 \cdot xH_2SO_4 \cdot yH_2O$, M 382.3, m >260°(dec), >300°(dec), pK_1^{20} 1.7, pK_2^{20} 3.20, pK_3^{25} 4.56. The salt is quite insoluble in H_2O but can be converted to the *free base* which is recrystallised from H_2O and converted to the sulfate by addition of the required amount of H_2SO_4 . The *hydrochloride* has m 300-305°(dec) and can be used to prepare the sulfate by addition of H_2SO_4 ; it is more soluble than the sulfate. The *perchlorate* has m 252-254°. The *free base* has λ_{max} at 260nm (log ϵ 4.24) in 0.1M HCl. [Bogert & Davidson *J Am Chem Soc* **55** 1667 1933, DOI: 10.1021/ja01331a059; Bredereck et al. *Chem Ber* **86** 850 1953, DOI: 10.1002/cber.19240571103; Sherman & Taylor *Org Synth Coll Vol* **4** 247 1963, DOI: 10.15227/orgsyn.037.0015; Barlin & Pfeleiderer *J Chem Soc (B)* 1425 1971, DOI: 10.1039/J29710001425; Beilstein **25** II 382.]

5,6-Diamino-1,3-dimethyluracil hydrate (5,6-diamino-1,3-dimethyl-2-pyrimidine-2,4-dione hydrate) [5440-00-6] $C_6H_{10}N_4O_2 \cdot xH_2O$, M 188.2, m 205-208°(dec), 209°(dec), 210°dec, pK_1 1.7, pK_2 4.6. It recrystallises from EtOH. The *hydrochloride* has m 310° (from MeOH), and the *perchlorate* has m 246-248°. [UV: Bredereck et al. *Chem Ber* **92** 583 1959, DOI: 10.1002/cber.19590920310; Taylor et al. *J Am Chem Soc* **77** 2243 1955, DOI: 10.1021/ja01613a066; Beilstein **25** III/IV 4133.]

6,9-Diamino-2-ethoxyacridine (Ethacridine) [442-16-0] $C_{15}H_{15}N_3O$, M 257.3, m 226°, pK^{20} 11.6. It crystallises from 50% EtOH or EtOH (yellow-orange crystals). It also crystallises as a *monohydrate* m 116-118°. It has a pK^{20} of 11.04 in 50% aqueous EtOH. The *methiodide* is soluble in H_2O and has m 332-334° (dec) (from aqueous Me_2CO). It is an antiseptic and abortive agent. [Albert & Gledhill *J Soc Chem Ind* **61** 159 1942, Foye et al. *J Pharm Sci* **57** 1793 1968, DOI: 10.1002/jps.2600571040; Albert & Goldacre *J Chem Soc* 706 1946, DOI: 10.1039/JR9460000706; Beilstein **22** II 458, **22** III/IV 6679, **22/12** V 243.]

6,9-Diamino-2-ethoxyacridine dl-lactate monohydrate (Rivanol, Acrinol) [6402-23-9] $C_{15}H_{15}N_3O \cdot C_3H_6O_3 \cdot H_2O$, M 361.4, has m 235° (dark at ~200°), pK^{20} 11.6. It forms yellow crystals from 90% EtOH/ Et_2O . Its solubility in H_2O is ~15% at 25° and ~9% at 100°, and its solutions have a yellow fluorescence which is stable on boiling. It is an antiseptic and abortive agent. See ethacridine above, Beilstein **22** II 458, **22** III/IV 6680, **22/12** V 243.]

2,4-Diamino-6-hydroxypyrimidine [56-06-4] $C_4H_6N_4O$, M 126.1, m 260-270°(dec), 285-286°(dec), pK_1^{25} 1.34, pK_2^{25} 3.27, pK_3^{25} 10.83. It recrystallises from H_2O . [Beilstein **25** III/IV 3642.]

4,5-Diamino-6-hydroxypyrimidine hemisulfate [102783-18-6] $C_4H_6N_4O \cdot 0.5H_2SO_4$, M 350.3, has m 268°(dec), 270°(dec), pK_1^{25} 1.34, pK_2^{25} 3.57, pK_3^{25} 9.86. It crystallises from H_2O . The *free base*, obtained by basifying the salt with aqueous ammonia, also crystallises from H_2O (m 239°). [Mason *J Chem Soc* 2071 1954, DOI: 10.1039/JR9540002071; Elion et al. *J Am Chem Soc* **74** 411 1952, DOI: 10.1021/ja01122a037; Beilstein **25** III/IV 3645.]

2,4-Diamino-5-phenylthiazole (DAPT, Amiphenazole) [490-55-1] $C_9H_9N_3S$, M 191.3, m 163-164°(dec). The thiazole crystallises from aqueous EtOH or water. Store it in the dark under N_2 . It is a barbiturate and narcotic antagonist. The *hydrochloride* has m 273-274°(dec) (from MeOH/ $EtOAc$), and the *picrate* has m 189-191°(dec) (from H_2O). [Davies et al. *J Chem Soc* 3491 1950, DOI: 10.1039/JR9500003491; Dodson & Turner *J Am Chem Soc* **73** 4517 1951, DOI: 10.1021/ja01154a005; Beilstein **27** III/IV 5139.]

2,3-Diaminopyridine [452-58-4] $C_5H_7N_3$, M 109.1, m 110-115°, 116°, pK_1^{25} -0.50, pK_2^{25} 6.92. It crystallises from *benzene and sublimes *in vacuo*. It is a ligand for organometallic complexing. [cf. Fidalgo et

al. *J Organomet Chem* **447** 299 1993, DOI: 10.1016/0022-328X(93)80253-8; *Beilstein* **22/11** V 241.]

2,6-Diaminopyridine [141-86-6] has **m 117-122°**, **121.5°**, **b 285°/atm**, **pK_{Est(1)} <-6.0**, **pK_{Est(2)} ~7.3**. It crystallises from *benzene and sublimes *in vacuo*. [*Beilstein* **22** III/IV 255.]

3,4-Diaminopyridine [54-96-6] has **m 216-218°**, **218-219°**, **220°**, **pK₁²⁰ 0.49**, **pK₂²⁰ 9.14**. It crystallises from *benzene and is stored under N₂ because it is *deliquescent* and absorbs CO₂. [*Beilstein* **22/11** V 266.]

3,5-Diamino-1,2,4-triazole (Guanazole) [1455-77-2] C₂H₅N₅, **M 99.1**, **m 202-205°**, **206°**, **pK₁²⁰ 4.43**, **pK₂²⁰ 12.12**. The triazole crystallises from water or EtOH. It is a DNA synthesis inhibitor. [*Beilstein* **26** III/IV 1161.]

1,3-Diazaazulene (cycloheptimidazole) [275-94-5] C₈H₁₂N₂, **M 136.2**, **m 110-112°**, **120°**. It is recrystallised repeatedly from de-aerated cyclohexane in the dark or from petroleum ether/*C₆H₆ and forms yellow needles. It is soluble in H₂O, EtOH and *C₆H₆, and forms a *monohydrate* which loses H₂O at 60°. The *picrate* has **m 207°(dec)**. [Nozoe et al. *J Am Chem Soc* **76** 3352 1954, DOI: 10.1021/ja01641a086; Nukai et al. *Bull Chem Soc Jpn* **40** 1967 1967, *Beilstein* **23** III/IV 1216.]

1,5-Diazabicyclo[4.3.0]non-5-ene (DBN, 2,3,4,,6,7,8-hexahydropyrrolo[1,2-*a*]-pyrimidine) [3001-72-7] C₇H₁₂N₂, **M 124.2**, **b 95-98°/7.5mm**, **96-98°/11mm**, **100-102°/12mm**, **118-121°/32mm**, **d₄²⁰ 1.040**, **n_D²⁰ 1.520**, **pK²⁵ >13.0**. Distil this *strong base* from BaO. It forms a *hydroiodide* on addition of 47% HI; dry it and dissolve it in MeCN, evaporate and repeat; recrystallise from EtOH, dry at 25°/1mm for 5 hours, then at 80°/0.03mm for 12 hours and store and dispense it in a dry box, **m 154-156°** [Jaeger et al. *J Am Chem Soc* **101** 717 1979, DOI: 10.1021/ja00497a039]. The *methiodide* is recrystallised from CHCl₃/Et₂O, **m 248-250°**, and the *hydrogen fumarate* has **m 159-160°** and is crystallised from *iso*-PrOH [Rokach et al. *J Med Chem* **22** 237 1979, DOI: 10.1021/jm00189a004; Oediger et al. *Chem Ber* **99** 2012 1966, DOI: 10.1002/cber.19660990633; Reppe et al. *Justus Liebigs Ann Chem* **596** 158 1955, DOI: 10.1002/jlac.19555960109]. [*Beilstein* **23/5** V 239.]

1,4-Diazabicyclo[2.2.2]octane (DABCO, triethylenediamine, TED) [280-57-9] C₆H₁₂N₂, **M 112.2**, **m 156-157° (sealed tube)**, **pK₁²⁵ 2.97**, **pK₂²⁵ 8.82**. DABCO crystallises from 95% EtOH, petroleum ether or MeOH/diethyl ether (1:1). Dry it under vacuum over CaCl₂ and BaO. It can be sublimed *in vacuo*, and readily at room temperature. It has also been purified by removal of water during azeotropic distillation of a *benzene solution. It is then recrystallised twice from anhydrous diethyl ether under argon, and stored under argon [Blackstock et al. *J Org Chem* **52** 1451 1987, DOI: 10.1021/jo00384a013]. [*Beilstein* **23/3** V 487.]

1,8-Diazabicyclo[5.4.0]undec-7-ene (DBU, 2,3,4,6,7,8,9,10-octahydropyrimidino[1,2-*a*]-azepine) [6674-22-2] C₉H₁₆N₂, **M 152.2**, **b 80-83°/06mm**, **115°/11mm**, **d₄²⁰ 1.023**, **n_D²⁰ 1.522**, **pK_{Est} ~ >13**. Fractionally distil DBU under vacuum. Also purify it by chromatography on Kieselgel and eluting with CHCl₃/EtOH/25% aqueous NH₃ (15:5:2) and checking by IR and MS. [Oediger et al. *Chem Ber* **99** 2012 1962, DOI: 10.1002/cber.19660990633; *Angew Chem Int Ed* **6** 76 1967, DOI: 10.1002/anie.196700761; Guggisberg et al. *Helv Chim Acta* **61** 1050 1978, DOI: 10.1002/hlca.19780610315; *Beilstein* **23/5** V 271.] It is a very useful *strong hindered base* in organic syntheses [Bella et al. *Tetrahedron* **60** 4821 2004, DOI: 10.1016/j.tet.2004.04.007; Ghosh *Synlett* **3** 574 2004, DOI: 10.1055/s-2004-815436].

1,8-Diazabiphenylene [259-84-7] C₁₀H₆N₂, **M 154.2**, **m 156-158.5°**, **pK_{Est} ~4.4**. Recrystallise it from cyclohexane, then sublime it in a vacuum. [Barton & Walker *Tetrahedron Lett* **16** 569 1975, DOI: 10.1016/S0040-4039(00)71923-X; Deroski et al. *Can J Chem* **62** 2235 1984, DOI: 10.1139/v84-381.]

2,7-Diazabiphenylene [31857-42-8] C₁₀H₆N₂, has **m 192-192.5°**, **pK_{Est} ~4.5**. It forms yellow crystals from cyclohexane, and sublimes in a vacuum. [MacBride *JCS Chem Commun* 359 1974, DOI: 10.1039/C3974000359B; Kramer & Berry *J Am Chem Soc* **94** 8336 1972, DOI: 10.1021/ja00779a010.]

1,8-Diazafluoren-9-one (cyclopenta[1.2-*b*:4,3-*b'*]dipyridin-9-one) [54078-29-4] C₁₁H₆N₂O, **M 182.2**, **m 205°**, **229-231°**, **pK_{Est} ~2.6**. Recrystallise it from Me₂CO, and has λ_{ex} 470nm, λ_{em} 570nm. The *oxime* has **m 119-200°**. [Druey & Schmidt *Helv Chim Acta* **33** 1080 1950, DOI: 10.1002/hlca.19500330438; *Beilstein* **24** III/IV 622.]

5H-Dibenz[*b,f*]azepine (iminostilbene, 2,2-iminostilbene) [256-96-2] C₁₄H₁₁N, **M 193.2**, **m 196-199°**, **204-206°**, **pK_{Est} ~12.3 (acidic NH)**. The dibenzazepine has been synthesised in a variety of ways [Kricka &

Ledwith *Chem Rev* **74**101 1974, DOI: 10.1021/cr60287a006]. One in particular is from 2,2'-diaminodihydrostibene (**m 86°** from Na/amyl alcohol reduction of the stilbene) by heating it and HCl at 265-275°/26-30hrs to give a dark brown product which is dissolved in warm EtOH, then treated with aqueous NaOH and steam distilled (6 hours). The distillate is filtered, and the solid is dissolved in Et₂O, washed with HCl (to remove aniline and starting amine), and then evaporated to give the colourless **10,11-dihydro5H-dibenz[b,f]azepine (iminodibenzyl)** [494-19-9] **C₁₄H₁₃N**, **M 195.3**, **m 110°** (from ligroin), **108°** (*C₆H₆), whose solubility (g/100ml at ~25°) is Et₂O (15.4), 96% EtOH (3.59) and petroleum ether (0.64) [the **5-nitroso derivative** has **m 108°** (Et₂O)] [Thiele & Häfliger *Justus Liebigs Ann Chem* **305** 96 1899, DOI: 10.1002/jlac.18993050108; cf. Jørgensen et al. *J Heterocycl Chem* **36** 57 1999, DOI: 10.1002/jhet.5570360110]. The **5-acetyl-10,11-dihydrodibenz[b,f]azepine** has **m 96-97°** (EtOH). Bromination of this acetyl-dihydro compound with N-bromosuccinimide in CCl₄ under UV light gave **5-acetyl-10-bromo-10,11-dihydrodibenz[b,f]azepine** with **m 118-119°** (EtOH) [UV has λ_{\max} (MeOH) at 280nm (log ϵ 3.20) and IR has 5.97 μ (nujol, *tert*-amido gp), 6.24, 6.30, 6.33, 6.72 and 13.02 μ (1,2-disubstit-phenyl)] which, when 31g are dissolved in EtOH (68ml) and treated with 50% aqueous KOH (13g, exothermic reaction) followed by cooling (60°), allowing to stand for 1hr then diluting with H₂O (104ml), precipitated **5-acetyldibenz[b,f]azepine** (80% yield) with **m 121-122°** (after recrystallisation from Et₂O). Hydrolysis of this acetate (23.5g) with KOH (14g) in EtOH (70ml) by refluxing for 18 hours, and strong cooling precipitated yellow-orange needles, which were collected, washed with cold EtOH then H₂O until the filtrate pH was ~7, recrystallised first from EtOH then from 10 parts of Et₂O to give pure **5H-dibenz[b,f]azepine** (17.3g, 90% yield) with **m 204-206°** [Schindler & Plattner *Helv Chim Acta* **44** 753 1961, DOI: 10.1002/hlca.19610440319].

Dehydrogenation of **10,11-dihydrodibenz[b,f]azepine** to **dibenz[b,f]azepine** has also been achieved by (a) boiling in PhNO₂ for 8-10 hours, evaporate *in vacuo*, dissolve residue in *C₆H₆ and filter through Al₂O₃, and evaporate, (b) heating with stoichiometric amount of sulfur at 200°/2hrs the 230-240°/1hr, cool, dissolve in *C₆H₆ and filter through Al₂O₃, and evaporate, (c) treat a solution in *C₆H₆ with excess Pb(OAc) overnight, filter add *C₆H₆ and filter through Al₂O₃, and evaporate, and (d) reflux a toluene solution with chloranil (turns green) for 12hrs, filter off the dark solid and subject to chromatography [Teuber & Schmidtke *Chem Ber* **93** 1257 1960, DOI: 10.1002/cber.19600930602]. Dehydrogenation has also been achieved with sulfur as stated and also by boiling the dihydrodibenzepine (1.95g) in diethyl maleate (40ml) containing Pd (Mohr, 2g) under N₂ for 5hrs. The crystals that separated were treated with warm EtOH and the Pd was filtered off. After removal of EtOH and ester at 145-150° (bath temp)/0.001mm, the crude dibenzo-azepine, m192-194°, was obtained in 48-60% yield. **5-Methyldibenz[b,f]azepine (m 143-144.5°**, yellow needles from petroleum ether b 40-80°) was obtained by dehydrogenation of **5-methyl-10,11-dihydrodibenz[b,f]azepine (m 107-108°**, colourless needles from EtOH or MeOH) with S by the previous procedure, as well as by methylation of **10,11-dihydrodibenz[b,f]azepine** with dimethylsulfate/Na₂CO₃ at 100° (56% yield, **m 106-107°**), and with PhLi/Et₂O then MeI (63% yield, **m 107-108°**, colourless needles from MeOH) [Huisgen et al. *Chem Ber* **93** 392 1960, DOI: 10.1002/cber.19600930222].

10,11-Dihydrodibenz[b,f]azepine has UV with λ_{\max} (EtOH) (nm, log ϵ) at (206, 4.54) and (287, 4.29); the ¹H NMR (250MHz, CDCl₃, TMS) has δ at 3.06 (4H, s, H-10, H-11), 7.03 (2H, d, $J_{1,2}$ = 7.4Hz, H-1 and 9), 6.76 (2H, d, $J_{2,3}$ = 7.4Hz, H-2 and 8), 7.06 (2H, d, $J_{3,4}$ = 8.1Hz, H-3 and 7), 6.70 (2H, d, $J_{1,3}$ = 1.6Hz, H-4 and 6), and 5.94 (1H, br, NH); **5-Me** has δ at 3.00 (4H, s, H-10, H-11), 7.10-6.6 (8H, m, H-1,2,3,4,5,6,7,8) and 3.14 (1H, br, NMe); ¹³C NMR (63MHz, CDCl₃, TMS) has δ at 130.51 (C-1 and 9), 123.00 (C-2 and 8), 129.45 (C-3 and 7), 119.3 (C-4 and 6), 132.12 (C-10 and 11), 129.75 (C-9a and 11a) and 148.39 (C-4a and 5a).

Dibenz[b,f]azepine has UV with λ_{\max} (MeOH) (nm, log ϵ) at (258, 4.62), (292, 3.45) and (355sh, 2.86); (C₆H₁₂) (nm, log ϵ) at (258, 4.65), (293, 3.43) and (365, 2.89); ¹H NMR (250MHz, CDCl₃, TMS) has δ at 6.32 (2H, s, H-10, H-11), 6.85 (1H, d, $J_{1,2}$ = 7.3Hz, H-1 and 9), 6.81 (2H, d, $J_{2,3}$ = 6.6Hz, H-2 and 8), 7.02 (2H, d, $J_{3,4}$ = 7.7Hz, H-3 and 7), 6.47 (2H, d, $J_{1,3}$ = 2.3Hz, H-4 and 6), and 4.92 (1H, br, NH); ¹³C NMR (63MHz, CDCl₃, TMS) has δ at 130.51 (C-1 and 9), 123.00 (C-2 and 8), 129.45 (C-3 and 7), 119.3 (C-4 and 6), 132.12 (C-10 and 11), 129.75 (C-9a and 11a) and 148.39 (C-4a and 5a). [for further NMR data see Hallberg et al. *J Heterocycl Chem* **21** 197 1984, DOI: 10.1002/jhet.5570210139.]

A series of **5-(phenolic-anilino-3-propanoyl)-dibenz[b,f]azepines** which may have possibilities as therapeutic antioxidants, were found to have antioxidant properties when evaluated using the following studies: DPPH (2,2-diphenyl-1-picrylhydrazine) free radical scavenging activity, inhibition of human LDL (low density lipoprotein) oxidation, reducing power assay and inhibition of lipid peroxidation in a β -carotene linoleate system, [Kumar & Naik *Eur Med Chem* **45** 2 2010, DOI: 10.1016/j.ejmech.2009.09.016].

Dibenzo-18-crown-6 [14187-32-7] $C_{20}H_{24}O_6$, **M 360.4**, **m 162-164°, 163-164°**. Crystallise it from *benzene, *n*-heptane or toluene and dry it under vacuum at room temperature for several days. Useful phase-transfer catalyst and ion shield. [Szczygiel *J Phys Chem* **91** 1252 1987, DOI: 10.1021/j100289a042; Vögtle ed. *Top Corr Chem* (Host Guest Complex Chemistry) **98** 1981, DOI: 10.1007/BFb0111244.]

Dibenzo-24-crown-8 [14174-09-5] $C_{24}H_{32}O_8$, **M 448.5**, has **m 103-105°, 103-106°**. Recrystallise it from EtOH, and dry in a vacuum at 60° over P_2O_5 for 16 hours. Useful phase-transfer catalyst and ion shield. [Delville et al. *J Am Chem Soc* **109** 7293 1987, DOI: 10.1021/ja00258a008; Vögtle ed. *Top Corr Chem* (Host Guest Complex Chemistry) **98** 1981, DOI: 10.1007/BFb0111244.]

Dibenzofuran (Dipheylene oxide) [132-64-9] $C_{12}H_8O$, **M 168.2**, **m 80-82°, 82.4°, b 154-155°/20mm**. Dissolve dibenzofuran in diethyl ether, then shake it with two portions of aqueous NaOH (2M), wash it with water, separate and dry ($MgSO_4$) it. After evaporating the ether, dibenzofuran is crystallised from aqueous 80% EtOH and then dried under vacuum. [Cass et al. *J Chem Soc* 1406 1958, DOI: 10.1039/JR9580001406.] High purity material is obtained by zone refining. [Beilstein **17** V 234.]

Dibenzothiophene (Dipheylene sulfide) [132-65-0] $C_{12}H_8S$, **M 184.3**, **m 97-100°, 99°, b 332-333°/atm**. Purify dibenzothiophene by chromatography on alumina with petroleum ether, in a darkened room. Recrystallise it from water or EtOH (white crystals). [Beilstein **17/2** V 239.]

1,3-Dibromo-5,5-dimethylhydantoin [77-48-5] $C_5H_6Br_2N_2O_2$, **M 285.9**, **m 190-192°(dec), 190-193°(dec), 197-199°(dec)**. Recrystallise it from H_2O . Its solubility in CCl_4 is 0.003 mol/L at 25° and 0.024 mol/L at 76.5°. It is a brominating agent. [Fieser **1** 208, **2** 108; Beilstein **24** III/IV 1101.]

4',5'-Dibromofluorescein [596-03-2] $C_{20}H_{10}Br_2O_5$, **M 490.1**, **m 285°**. Crystallise this fluorescent dye from aqueous 30% EtOH (red plates). It dissolves in concentrated H_2SO_4 to give an orange solution which becomes yellow-brown on dilution and precipitates an orange solid (of the sulfonate?). [Beilstein **19/6** V 462.]

5,7-Dibromo-8-hydroxyquinoline (Broxyquinoline) [521-74-4] $C_9H_5Br_2NO$, **M 303.0**, **m 196°, pK_I²⁵ 5.84, pK₂²⁵ 9.56**. Crystallise it from acetone/EtOH. It can be sublimed. It complexes with Cu, Fe and Ti ions, is a disinfectant and an antiseptic. [Beilstein **21/3** V 290.]

2,6-Dibromopyridine [626-05-1] $C_5H_3Br_2N$, **M 236.9**, **m 117-119°, 118.5-119°, b 249°/757.5mm, pK_{Est} <0**. Purify 2,6-dibromopyridine by steam distillation, then recrystallise it twice from EtOH. It does not form an $HgCl_2$ salt. [den Hertog & Wibaut *Recl Trav Chim Pays-Bas* **51** 381 1932, DOI: 10.1002/recl.19320510412; Beilstein **20/5** V 435.]

3,4-Dibromothiophene [3141-26-2] $C_4H_2Br_2S$, **M 241.9**, **m 4-5°, b 94-95°/12mm, 212-213°/atm, 221-222°/atm, d₄²⁵ 2.188, n_D²⁰ 1.640**. Distil it in a vacuum, but if it is discoloured then dissolve it in Et_2O , wash it with 2M NaOH solution, separate the layers, filter, dry the Et_2O layer over $CaCl_2$, filter, evaporate, and distil the residual oil in a vacuum. Nitration with Ac_2O/HNO_3 at 60° yields **3,4-dibromo-2-nitrothiophene m 115-116°** (from EtOH). [Steinkopf et al. *Justus Liebigs Ann Chem* **512** 136 1934, DOI: 10.1002/jlac.19345120113; Beilstein **17** III/IV 248, **17/1** V 308.] It is a useful thiophene synthon [cf. He et al. *J Org Chem* **72** 442 2007, DOI: 10.1021/jo061853y].

2,6-Di-tert-butyl-4-methylpyridine [38222-83-2] $C_{14}H_{23}N$, **M 205.4**, **m 31-32°, 33-36°, b 148-153°/95mm, 223°/760mm, n_D²⁰ 1.476, pK_{Est} ~5.7**. A possible impurity is 2,6-di-tert-butyl-4-neopentylpyridine. Attempts to remove coloured impurities directly by distillation, acid-base extraction or treatment with activated charcoal were unsuccessful. Pure material is obtained by dissolving 0.3mole of the alkylpyridine in pentane (150ml) and introducing it at the top of a cold water jacketed chromatographic column (40 x 4.5cm) (cooling is necessary because the base in pentane reacts exothermically with alumina) containing activated and acidic alumina (300g). The column is eluted with pentane using a 1L constant pressure funnel fitted at the top of the column to provide slight pressure. All the pyridine is obtained in the first two litres of eluent (the progress of elution is monitored by spotting a fluorescent TLC plate and examining under short wave UV light—a dark blue spot is evidence for the presence of the alkylpyridine). Elution is complete in 1 hour. Pentane is removed on a rotovap with 90-93% recovery yielding a liquid which solidifies on cooling, **m 31-32°**, and the **free base** can be distilled. The

***H₂PtCl₆* salt** has **m 213-314°(dec)**, and the ***CF₃SO₃H* salt** has **m 202.5-203.5°** (from CH₂Cl₂). [Anderson & Stang *Org Synth Coll Vol* **7** 144 1981, DOI: 10.15227/orgsyn.060.0034; *Beilstein* **20/6** V 190.]

2,6-Di-*tert*-butylpyridine [585-48-8] C₁₃H₂₁N, **M 191.3**, **b 100-101°/23mm**, **d₄²⁵ 0.852**, **n_D²⁰ 1.473**, **pK²⁵ 5.02**. Redistil it from KOH pellets. Used for α-enolisation of aldehydes (in presence of ceric ammonium nitrate) [Jang et al. *J Am Chem Soc* **129** 7004 2007, DOI: 10.1021/ja0719428], and as a proton trap [cf. Gyor et al. *J Macromol Sci, Pure Appl Chem* **A29** 639 1992, DOI: 10.1080/10601329208052189]. [*Beilstein* **20** III/IV 2868.] § Polystyrene supported version is commercially available [107054-29-5] 200-400 mesh, crosslinked with 1% divinylbenzene containing ~1.8mmol N/g polymer.

3,5-Dicarbethoxy-1,4-dihydro-2,4,6-collidine [632-93-9] C₁₄H₂₁NO₄, **M 267.3**, **m 130-132°**, **131-132°**. Crystallise the ester from hot EtOH/water mixture. [*Beilstein* **22** H 147, **22** I 529, **22** II 100, **22** III/IV 1594.]

1S-(+)-(8,8-Dichlorocamphorsulfonyl)oxaziridine {(7S)-8,8-dichloro-10,10-dimethyl-5-thia-4-azatricyclo-[5.2.1.0^{3,7}]dec-3-ene-5,5-dioxide} [127184-05-8] C₁₀H₁₃Cl₂NO₃S, **M 298.2**, **m 178-180°**, **181-186°**, **[α]_D²⁰ +89.4** (c 1, CHCl₃). This dichlorooxaziridine was prepared by Davis and Weismiller [*J Org Chem* **55** 3715 1990, DOI: 10.1021/jo00299a007; supporting data jo00299a007_si_001.pdf] from **1S-(-)-Camphorsulfonylimine** [see 60886-80-8 above] by treatment with **sodium bis(trimethylsilyl)amide (NHMDs)** at -78° in THF to form the azaenolate and then chlorinated with *N*-chlorosuccinimide (NCS, 2.5 equivalents) at -78° to form the **1S-(+)-8,8-dichlorocamphorsulfonylimine** (note inversion of rotation) which they purified by flash chromatography (silica gel, eluting with CH₂Cl₂/hexane 6:4) in 64% yield and had **m 174°**, **[α]_D²⁰ +7.9** (c 1, CHCl₃), TLC R_F = 0.42 [silica gel plate (250μm thick), eluted with CH₂Cl₂ and developed with 10% molybdophosphoric acid in EtOH and heated for a few minutes in a 100° oven]; the ¹H NMR (CDCl₃) had δ at 1.19 (s, 3H, CH₃), 1.22 (s, 3H, CH₃), 1.85-2.45 (m, 4H), 2.80 (d, 1H, *J* = 3.6Hz), 3.28 (AB q, *J* = 13.5, 2H). Oxidation of the **(+)-8,8-dichloroimine** in CH₂Cl₂ was achieved by stirring vigorously with *m*-chloroperbenzoic acid (1.5equiv) in saturated K₂CO₃ for 18 hours to give **(+)-(8,8-dichlorocamphorsulfonyl)oxaziridine** in >90% yield after recrystallisation from EtOH which had **m 178-180°**, **[α]_D²⁰ +89.4** (c 1, CHCl₃), TLC R_F = 0.68 (same conditions as above) and the ¹H NMR (CDCl₃) had δ at 1.18 (s, 3H, CH₃), 1.49 (s, 3H, CH₃), 1.85-2.50 (m, 4H), 2.77 (d, 1H, *J* = 3.5Hz), 3.36 (AB q, *J* = 14.1, 2H).

1R-(-)-8,8-dichlorocamphorsulfonylimine was prepared from **1R-(+)-camphorsulfonylimine** by chlorination in the same manner (NHMDs/NCS), and oxidised to **(-)-(8,8-dichlorocamphorsulfonyl)oxaziridine** [139628-16-3] in similar yields, and their properties were similar to those of the enantiomers except for the optical rotations which were exactly of opposite sign.

1S-(+)-(8,8-Dimethoxycamphorsulfonyl)oxaziridine has **m 189°(dec)** after crystallisation from absolute EtOH, with **[α]_D²⁰ +91.3** (c 3.39, CHCl₃); the IR (KBr) has ν_{\max} at 1356 and 1165 cm⁻¹; the ¹H NMR (CDCl₃) δ at 3.35 (s, 3H), 3.28 (s, 3H), 3.20 (AB q, 2H *J* = 12Hz), 2.30-1.75 (m, 5H), 1.32 (s, 3H), 1.05 (s, 3H); ¹³C NMR (CDCl₃) δ at 102.7, 97.6, 54.6, 52.9, 50.8, 47.5, 45.2, 28.2, 21.7 and 20.6. It was prepared in three steps from **1S-(-)-camphorsulfonylimine** by oxidation with SeO₂/AcOH/reflux for 20 hours to **1S-(-)-3-oxocamphorsulfonylimine** (72% yield), converted in 95% yield by boiling under reflux with (MeO)₃CH/MeOH/H₂SO₄ (as catalyst)/Amberlist-15 ion-exchange resin (as catalyst) to **1S-(+)-8,8-dimethoxycamphorsulfonylimine** which has **m 186-187°** (after recrystallisation from absolute EtOH), **[α]_D²⁰ +7.2** (c 3.6, CHCl₃); IR (KBr) with ν_{\max} at 1620, 1340 and 1160 cm⁻¹; the ¹H NMR (CDCl₃) has δ at 3.46 (s, 3H), 3.38 (s, 3H), 3.09 (AB q, 2H *J* = 12Hz), 2.38-1.80 (m, 5H), 1.10 (s, 3H), 1.00 (s, 3H); and the ¹³C NMR (CDCl₃) has δ at 188.6, 102.9, 64.2, 52.0, 50.5, 48.8, 46.0, 29.2, 20.6, 20.5 and 20.4; followed by oxidation with *m*-chloroperbenzoic acid in CH₂Cl₂/saturated aqueous K₂CO₃ to the **oxaziridine** in 96% yield. [Davis et al. *J Org Chem* **56** 1143 1991, DOI: 10.1021/jo00003a042.]

1R-(-)-(8,8-Dimethoxycamphorsulfonyl)oxaziridine was prepared in similar yields from the respective enantiomeric intermediates and the products differed only in the signs of their optical rotations.

In very elegant work, Davis and coworkers have shown that the above three camphorsulfonyloxaziridines are highly stereospecific asymmetric enolate oxidants (**Davis oxidation**) to produce α-hydroxyketones from enolisable ketones in high to very high chemical yields and with high enantioselective hydroxylations of 90 to ≥95 %ee. The order of stereoselectivity increases with chiral **camphorsulfonyl)oxaziridine** < **8,8-dichlorocamphor-sulfonyl)oxaziridine** < **8,8-methoxycamphorsulfonyl)oxaziridine** **clearly the effects of going from 8,8-H,H to 8,8-Cl,Cl to 8,8-OMe,OMe**. They have used these stereospecific enolate hydroxylations successfully to prepare synthons for various anthracycline antibiotics among other compounds. [Davis &

Kumar *Tetrahedron Lett* **32** 7671 1991, DOI: 10.1016/0040-4039(91)80561-J; Davis & Chen *Tetrahedron Lett* **31** 6823 1990, DOI: 10.1016/S0040-4039(00)97181-8; Davis et al. *J Org Chem* **56** 1143 1991, DOI: 10.1021/jo00003a042; Davis et al. *Tetrahedron Lett* **32** 867 1991, DOI: 10.1016/S0040-4039(00)92107-5.]

1,3-Dichloro-5,5'-dimethylhydantoin [118-52-5] $C_5H_6Cl_2N_2O_2$, M 197.0, m 132-134°, 136°. Purify it by dissolving in concentrated H_2SO_4 and diluting with ice H_2O , collect the solid, dry it in a vacuum and recrystallise it from $CHCl_3$. It sublimes at 100° in a vacuum. It exhibits time-dependent hydrolysis at pH 9. It is a chlorinating agent and a useful disinfectant. [Petterson & Grzeskowiak *J Org Chem* **24** 1414 1959, DOI: 10.1021/jo01092a006; *Beilstein* **24** III/IV 1100.]

4,5-Dichloro-3H-1,2-dithiol-3-one [1192-52-5] M 187.1, m 52-56°, 61°, b 87°/0.5mm, 125°/11mm. Dissolve it in CH_2Cl_2 (1g in 250ml), filter, wash it twice with H_2O , evaporate and distil the residue *in vacuo* and then recrystallise it from petroleum ether. Its IR has ν_{max} at 1650 cm^{-1} . [Boberg *Justus Liebigs Ann Chem* **679** 109 1964, DOI: 10.1002/jlac.19646790115; Boberg *Justus Liebigs Ann Chem* **693** 212 1966, DOI: 10.1002/jlac.19666930122; *Beilstein* **19/4** V 72.]

5,7-Dichloro-8-hydroxyquinoline [773-76-2] $C_9H_5Cl_2NO$, M 214.1, m 178-180°, 180-181°, pK_1 1.89, pK_2 7.62. Crystallise the dichloro-oxine from acetone/EtOH. It is a chelator that is useful in analytical chemistry, and is also used medicinally to clear sebaceous glands in the skin. [*Beilstein* **21** H 95.]

6,9-Dichloro-2-methoxyacridine (3,9-Dichloro-7-methoxyacridine) [86-38-4] $C_{14}H_9Cl_2NO$, M 278.1, m 160-161°, 164°, 163-165°. Crystallise it from *benzene or 1,2-dichloroethane (m 162-163°). [Hall & Turner *J Chem Soc* 694 1945, DOI: 10.1039/JR9450000694; *Beilstein* **21** III/IV 1553.]

5,7-Dichloro-2-methyl-8-hydroxyquinoline (5,7-dichloro-8-hydroxyquinaldine) [72-80-0] M 228.1, m 114-115°, $pK_{Est(1)} \sim 2.0$, $pK_{Est(2)} \sim 8.4$. Crystallise it from EtOH. [*Beilstein* **21** III/IV 1180, **21/3** V 346.]

4,6-Dichloro-5-nitropyrimidine [4316-93-2] $C_4HCl_2N_3O_2$, M 194.0, m 100-103°, 101-102°, $pK_{Est} < 0$. If too impure, then dissolve it in Et_2O , wash it with H_2O , dry it over $MgSO_4$, evaporate it to dryness and recrystallise it from petroleum ether (b 85-105°) to give a light tan solid. It is soluble in *ca* 8 parts of MeOH [Boon et al. *J Chem Soc* 96 1951, DOI: 10.1039/JR9510000096; Montgomery et al. in *Synthetic Procedures in Nucleic Acid Chemistry* Zorbach & Tipson eds, Wiley & Sons, NY, p76 1968]. [*Beilstein* **23** III/IV 899.]

2,6-Dichloropurine [5451-40-1] $C_5H_2Cl_2N_4$, M 189.0, m 180-181.5°, 181°, 185-195°(dec), 188-189°, pK_1^{20} 1.16 (aqueous H_2SO_4), pK_2^{20} 7.06. It can be recrystallised from 150 parts of boiling H_2O and dried at 100° to constant weight. It is soluble in EtOAc. The $HgCl_2$ salt separates from EtOH solution. It has UV with λ_{max} at 275nm (ϵ 8.9K) at pH 1; and 280nm (ϵ 8.5K) at pH 11 [Elion & Hitchings *J Am Chem Soc* **78** 3508 1956, DOI: 10.1021/ja01595a065; Schaeffer & Thomas *J Am Chem Soc* **80** 3738 1958, DOI: 10.1021/ja01547a068; Beaman & Robins *J Appl Chem (London)* **12** 432 1962, Montgomery *J Am Chem Soc* **78** 1928 1956, DOI: 10.1021/ja01590a043]. [Lister *Purines Part II, Fused Pyrimidines* Brown Ed, Wiley-Interscience pp254-225 1971, ISBN 0-471-38205-1, *Beilstein* **26** III/IV 1747.] Substrate in Suzuki-Miyaura coupling with arylboronic acids [cf. *Synthesis* 3515 2006].

2,6-Dichloropyridine [2402-78-0] $C_5H_3Cl_2N$, M 148.0, m 83-86°, 87-88°, pK^{25} -2.86 (aqueous H_2SO_4). It crystallises from EtOH. [*Beilstein* **20/5** V 416.]

3,5-Dichloropyridine [2457-47-8] has m 64-65°, 66-67°, b 178-178°/~760mm, pK^{25} 0.67. Crystallise 3,5-dichloropyridine from EtOH, or distil it. [den Hertog et al. *Recl Trav Chim Pays-Bas* **69** 673 1950, DOI: 10.1002/recl.19500690604; *Beilstein* **20** H 231, **20** III/IV 2502, **20** V 417.]

2,6-Dichloropyridine-3-carboxylic acid (2,6-dichloronicotinic acid) [38496-18-3] $C_6H_3Cl_2NO_2$, M 192.0, m 140-143°, 146-148°, 147°, $pK_{Est(1)} \sim -1$. Purify the acid by recrystallisation from H_2O or Et_2O /petroleum ether (colourless needles), and the IR(KBr) has ν_{max} at 1738 cm^{-1} , and λ_{max} (ϵ) at 228.5(4700) and 273.5(8300)nm. The *acid chloride* [58584-83-1] M 210.4, b 72-74°/0.01mm, 117-118°/45mm, n_D^{20} 1.5944, was obtained by treatment with $SOCl_2$ /DMF/* C_6H_6 and converted into the *amide* [70445-62-4] M 191.0, m 148-148.5°, or the *hydrazide* m 171-173° which crystallised from a large volume of MeCN. [Mutterer & Weis

Helv Chim Acta **59** 222 1976, DOI: 10.1002/hlca.19760590124; Newkome et al. *J Org Chem* **44** 2693 1979, DOI: 10.1021/jo01329a020; *Beilstein* **22** H 44.] A pharmaceutical research building block acid of 90% grade is also available commercially.

2,6-Dichloropyridine-4-carboxylic acid (2,6-dichloroisonicotinic acid) [5398-44-7] has **m 210°, 209-212°, pK_{Est(1)} ~ -1**. Purify the acid by crystallisation from H₂O, and it forms needles or platelets from EtOH. The **methyl ester** crystallises from aqueous MeOH with **m 82°**, the **ethyl ester** from aqueous EtOH has **m 66°**, and the **amide** from H₂O has **m 207-208°**. [Meyer et al. *Monatsh Chem* **36** 731 1915, DOI: 10.1007/BF01519430; Levelt & Wibaut *Recl Trav Chim Pays-Bas* **48** 466 1929, DOI: 10.1002/recl.19290480506.]

5,6-Dichloropyridine-3-carboxylic acid (5,6-dichloronicotinic acid) [41667-95-2] has **m 160-161°, 164-168°, 168°, pK_{Est(1)} ~ 0.5**. Recrystallisation of the acid from H₂O provides the **monohydrate** which becomes **anhydrous (m 161-162°)** on heating at 100°. The **acid chloride** distils at **125°/~24mm**, solidifies on cooling, and crystallises from petroleum ether with **m 48-49°**. The **methyl ester** has **m 67-68°** (from aqueous MeOH), and the **amide** forms colourless plates from aqueous EtOH with **m 218-220°**. [Räth et al. *Justus Liebigs Ann Chem* **487** 127 1931, DOI: 10.1002/jlac.19314870109; Meyer & Graf *Chem Ber* **61** 2202 1928, DOI: 10.1002/cber.19280610925; *Beilstein* **22** H 44, **22** II 36, **22** III/IV 511, **22/2** V 80.]

4,7-Dichloroquinoline [86-98-6] **M 198.1, C₉H₅Cl₂N, m 81-83°, 86.4-87.4°, b 148°/10mm, pK²⁵ 2.80**. Crystallise the dichloroquinoline from MeOH or 95% EtOH. [*Beilstein* **20/7** V 316.]

2,3-Dichloroquinoxaline [2213-63-0] **C₈H₄Cl₂N₂, M 199.0, m 152-153°, 152-154°, pK_{Est} <0**. Recrystallise it from *C₆H₆ and dry it in a vacuum [Cheeseman *J Chem Soc* 1804 1955, DOI: 10.1039/JR9550001804; *Beilstein* **23/7** V 144].

cis-Dicyclohexyl-18-crown-6 [16069-36-6] **M 372.5, m 47-50°**. Purify it by chromatography on neutral alumina and elute with an ether/hexane mixture [see Izatt et al. *Inorg Chem* **14** 3132 1975]. Dissolve it in ether at ca 40°, and spectroscopic grade MeCN is added to the solution, which is then chilled. The crown ether precipitates and is filtered off. It is dried *in vacuo* at room temperature [Wallace *J Phys Chem* **89** 1357 1985]. [Vögtle ed. *Top Corr Chem* (Host Guest Complex Chemistry) **98** 1981.] **SKIN IRRITANT**.

1,1'-Diethyl-2,2'-cyanine iodide [977-96-8] **M 454.4, m 274°(dec)**. It crystallises from EtOH and is dried in a vacuum oven at 80° for 4 hours. [*Beilstein* **23** II 267.]

5,5-Diethylbarbituric acid (Barbital) [57-44-3] **M 184.2, m 188-192°, pK₁²⁵ 8.02, pK₂²⁵ 12.7**. Crystallise barbital from water or EtOH and dry it in a vacuum over P₂O₅. [*Beilstein* **24** III/IV 1901.]

2,3-Dihydrobenzofuran (coumaran) [496-16-2] **C₈H₈O, M 120.2, m -21.5°, 72-73°/12mm, 84°/17mm, 188°/atm, d₄²⁰ 1.065, n_D²⁰ 1.5524**. Suspend coumaran in aqueous NaOH and steam distil, saturate distillate with NaCl, extract with Et₂O, dry extract (MgSO₄), filter, evaporate and distil residue. It gives a violet colour with FeCl₃/H₂SO₄. Its yellow **picrate**, **m 76°** (from EtOH or *C₆H₆) loses coumaran *in vacuo* [Bennett & Hafez *J Chem Soc* 287 1941, DOI: 10.1039/JR9410000287; Baddeley et al. *J Chem Soc* 2455 1956, DOI: 10.1039/JR9560002455]. [*Beilstein* **17/1** V 581.]

Dihydropyran (3,4-dihydro-2H-pyran) [110-87-2] **C₅H₈O, M 84.1, m -70°, b 84.4°/742mm, 85.4-85.6°/760mm, d₄²⁰ 0.9261, n_D²⁰ 1.4423, pK_{Est} ~ 4.2**. Dry dihydropyran with Na₂CO₃, then fractionally distil it. Fraction **b** 84-85° is refluxed with Na until H₂ no longer evolves; add fresh Na then distil again through a 60 x 1.2cm column packed with glass rings [Brandon et al. *J Am Chem Soc* **72** 2120 1950, DOI: 10.1021/ja01161a069; UV: Eglinton et al. *J Chem Soc* 2873 1952, DOI: 10.1039/JR9520002873; NMR: Bushweller & O'Neil *Tetrahedron Lett* **10** 4713 1969, DOI: 10.1016/S0040-4039(01)88791-8]. Characterise it as the **2-(3,5-dinitrobenzoyloxy)tetrahydropyrane** derivative, **m 103°** [made by dissolving 3,4-dinitrobenzoic acid (3g) in 50% excess of dihydropyran with warming, cooling, adding Et₂O (5ml), when the derivative crystallises out quantitatively] which forms pale yellow crystals from 80% dihydropyran/Et₂O [Woods & Kramer *J Am Chem Soc* **69** 2246 1947, DOI: 10.1021/ja01201a520]. [*Beilstein* **17/1** V 181.] It is an OH protecting group [Greene & Wuts *Protecting Groups in Organic Synthesis* 2nd Edn, J. Wiley (NY) p 31 1991].

3,4-Dihydro-2H-pyrido[1,2a]-pyrimidin-2-one [5439-14-5] **M 148.2, m 185-187°, 187-188°, 191-191.5°**. Dissolve it in CHCl₃, filter, evaporate, then recrystallise the residue from EtOH/Me₂CO (needles) which can be

washed with Et₂O and dried. It can also be recrystallised from CHCl₃/petroleum ether or CHCl₃/hexane. The *hydrochloride* has **m 295-295°** (dec, from EtOH or MeOH/Et₂O), the *hydrobromide* has **m 299-300°**(dec) (from MeOH/Et₂O) and the *picrate* has **m 224-226°**(corr), **m 219-220°** from EtOH. [Adams & Pachter *J Am Chem Soc* **74** 4906 1952, DOI: 10.1021/ja01139a051; Lappin *J Org Chem* **23** 1358 1958, DOI: 10.1021/jo01103a034; Hurd & Hayao *J Am Chem Soc* **77** 117 1955, DOI: 10.1021/ja01606a037; *Beilstein* **24** III/IV 299.]

7,8-Dihydroxycoumarin (Daphnetin) [486-35-1] C₉H₆O₄, **M 178.2**, **m 256°**(dec), **pK_{Est(1)} ~8.5**, **pK_{Est(2)} ~12.3**. Crystallise it from aqueous EtOH. It can be sublimed. It inhibits protein kinases. [*Beilstein* **18/3** V 202.].

trans-2,3-Dihydroxy-1,4-dioxane [4845-50-5] C₄H₈O₄, **M 120.1**, **m 91-95°, 100°**. Recrystallise it from Me₂CO. With phenylhydrazine it gives *glyoxal phenylhydrazone* **m 175°** (from Me₂CO/petroleum ether). The *diacetyl* derivative has **m 105-106°** [Head *J Chem Soc* 1030 1955, DOI: 10.1039/JR9550001030; Raudnitz *Chem Ind (London)* 166 1956]. [*Beilstein* **1** IV 3627.]

2,5-Dihydroxy-1,4-dithiane [40018-26-6] C₄H₈O₂S₂, **M 152.2**, **m (130° dec, 142-147°) 150-152°, 151°**. Recrystallise the dithiane from EtOH. **2,5-Diethoxy-dithiane** has **m 91° (92-93°)**; it crystallises from petroleum ether and can be sublimed at 60°/0.001mm [Hromatka & Haberl *Monatsh Chem* **85** 1088 1954, DOI: 10.1007/BF00899857; Thiel et al. *Justus Liebigs Ann Chem* **611** 121 1958, DOI: 10.1002/jlac.19586110113; Hesse & Jørdet *Chem Be* **85** 924 1952, DOI: 10.1002/cber.19520850915]. [*Beilstein* **1** IV 3966.]

S(-)-4',7-Dihydroxyflavanone (3,4'-dihydroxyisoflavone, Liquiritigenin) [578-86-9] **M 256.3**, **m 203-205°** [α]_D²⁰ **-225 (c 0.95, 95% EtOH)**. It crystallises from aqueous 50% EtOH. It inhibits rat monoamine oxidase *in vitro* [Pan et al. *Acta Pharmacolo Sinica* **21** 949 2000, PMID:11501051] [*Beilstein* **18** III/IV 1780, **18/4** V 82.]

5,7-Dihydroxy-4'-methoxyflavone (Acacetin) [480-44-4] C₁₆H₁₂O₅, **M 284.3**, **m 261°, 260-265°, 265°**. Acacetin crystallises from 95% EtOH or acetic acid as pale yellow needles (**m 263°**). It is an anti-inflammatory. [Zemplen & Bognar *Chem Ber* **76** 452 1943, DOI: 10.1002/cber.19430760503; Pillon *Bull Soc Chim Fr* 9 1954, UV: Gaydou & Bianchini *Bull Soc Chim Fr* II 43 1978, *Beilstein* **18** III/IV 2683, **18/4** V 575.]

(±)-7-(2,3-Dihydroxypropyl)theophylline (Diprophylline, Dyphylline) [479-18-5] C₁₀H₁₄N₄O₄, **M 254.3**, **m 158°, 160-164°, 161°, 161-162°, 161-164°, pK_{Est} ~8.7**. Recrystallise it from EtOH or H₂O. Its solubility in H₂O is 33% at 25°, in EtOH it is 2% and in CHCl₃ it is 1%. Its UV has λ_{\max} (H₂O) at 273nm (ϵ 8,855). [Roth *Arch Pharm* **292** 234 1959, DOI: 10.1002/ardp.19592920504.] The **4-nitrobenzoyl** derivative has **m 178°** [Ishay *J Chem Soc* 3975 1956, DOI: 10.1039/JR9560003975]. [Lister *Purines Part II, Fused Pyrimidines* Brown Ed, Wiley-Interscience pp254-225 1971, ISBN 0-471-38205-1, *Beilstein* **26** III/IV 2370.]

1,3-Diiminoisoindoline [3468-11-9] C₈H₇N₃, **M 145.2**, **m 193-194°(dec), 196°(dec), pK₂₅ 8.27**. It crystallises from H₂O, MeOH or MeOH/Et₂O (charcoal) in colourless prisms that become green on heating. [Elvidge & Linstead *J Chem Soc* 5000 1952, DOI: 10.1039/JR9520005000]. Its IR (nujol) has bands at 3150 and 690 cm⁻¹, and the UV has λ_{\max} at 251nm (ϵ 12,500), 256nm (ϵ 12,5000) and 303nm (ϵ 4,600) [Elvidge & Golden *J Chem Soc* 700 1957, DOI: 10.1039/JR9570000700; Clark et al. *J Chem Soc* 3593 1953, DOI: 10.1039/JR9530003593]. The *thiocyanate* has **m 250-255°** (dec), the *monohydrochloride* has **m 300-301°** (turns green), and the *dihydrochloride* has **m 326-328°** (turns green) and the *picrate* crystallises from EtOH with **m 299°** (dec). [*Beilstein* **22/13** V 5.]

5,7-Diiodo-8-hydroxyquinoline [83-73-8] C₉H₅INO, **M 397.0**, **m >200°(dec), 214-215°(dec), pK_{Est(1)} ~3.2, pK_{Est(2)} ~8.2**. It crystallises from xylene and is dried at 70° in a vacuum. It is a metal chelating agent. [*Beilstein* **21** II 58.]

6-Dimethylaminopurine [938-55-6] **M 163.1**, **m 257.5-258.5°, 259-262°, 263-264°, pK₁²⁵ 3.87, pK₂²⁵ 10.5**. It is purified by recrystallisation from H₂O, EtOH (0.32g in 10ml) or CHCl₃. [Albert & Brown *J Chem Soc* 2060 1954, DOI: 10.1039/JR9540002060; UV: Mason *J Chem Soc* 2071 1954, DOI: 10.1039/JR9540002071.] The *monohydrochloride* crystallises from EtOH/Et₂O, **m 253°(dec)** [Elion et al. *J Am Chem Soc* **74** 411 1952,

DOI: 10.1021/ja01122a037], the *dihydrochloride* has **m** 225°(dec) and the *picrate* has **m** 245° (235-236.5°) [Fryth et al. *J Am Chem Soc* **80** 2736 1958, DOI: 10.1021/ja01544a039]. [*Beilstein* **26** III/IV 3566.]

1,3-Dimethylbarbituric acid [769-42-6] $C_6H_8N_2O_3$, **M** 156.1, **m** 121-123°, 123°, **pK**₁²⁵ 4.56. Crystallise the acid from water and sublime it in a vacuum. Also purify it by dissolving 10g in 100ml of boiling $CCl_4/CHCl_3$ (8:2) (1g charcoal), filtering and cooling to 25°. Dry it *in vacuo* [Kohn et al. *Anal Chem* **58** 3184 1986, DOI: 10.1021/ac00127a058]. [*Beilstein* **24** III/IV 1875.]

5,6-Dimethylbenzimidazole [582-60-5] $C_9H_{10}N_2$, **M** 146.2, **m** 202-205°, 205-206°, **pK**₁²⁵ 5.96, **pK**₂²⁵ 12.52. Crystallise 5,6-dimethylbenzimidazole from diethyl ether. It sublimes at 140°/3mm. [*Beilstein* **23/6** V 454.]

2,3-Dimethylbenzothiophene [31317-17-6] **M** 212.3, **m** 69-70°, **b** 123-124°/10mm, **n**_D¹⁹ 1.6171. Fractionate it through a 90cm Monel spiral column, or other efficient fractionating or spinning band column and collect the middle fraction. It has also been purified by chromatography on basic alumina using pentane as eluent. [Tedjamulia et al. *J Heterocycl Chem* **20** 1485 1983, DOI: 10.1002/jhet.5570200610; *Beilstein* **17** III/IV 161.]

4,4'-Dimethyl-2,2'-bipyridine [1134-35-6] $C_{12}H_{12}N_2$, **M** 184.2, **m** 169-174°, 175-176°, **pK**_{Est(1)} ~0.2, **pK**_{Est(2)} ~4.9. Crystallise it from ethyl acetate. [Elliott et al. *J Am Chem Soc* **107** 4647 1985, DOI: 10.1021/ja00302a009; *Beilstein* **23/8** V 79.]

1,1'-Dimethyl-4,4'-bipyridylium dichloride (3H₂O; Methyl Viologen Dichloride, paraquat dichloride) [1910-42-5] $C_{12}H_{14}Cl_2N_2 \cdot xH_2O$, **M** 311.2, **m** >300°(dec). Recrystallise the dichloride from MeOH/acetone mixture. It has also been recrystallised three times from absolute EtOH [Bancroft et al. *Anal Chem* **53** 1390 1981, DOI: 10.1021/ac00232a021]. Dry it at 80° in a vacuum. It is an electron acceptor [cf. Kelly & Rodgers *J Phys Chem* **98** 6377 1994, DOI: 10.1021/j100076a023], and a transfer catalyst in redox reactions [cf. Koshechko et al. *Tetrahedron Lett* **33** 6677 1993, DOI: 10.1016/S0040-4039(00)61016-X]. [*Beilstein* **23/8** V 30.]

1,3-Dimethylbutadiene sulfone (1,3-dimethylsulfolene, 2,5-dihydro-2,4-dimethylthiophene) [10033-92-8] **M** 145.2, **m** 40.4-41.0°. Crystallise it from Et₂O, Et₂O/pentane or CCl_4 . [Grummitt et al. *J Am Chem Soc* **72** 5167 1950, DOI: 10.1021/ja01167a103; Bartlett et al. *J Org Chem* **32** 1290 1967, DOI: 10.1021/jo01280a601; *Beilstein* **17** III/IV 161.]

2,2-Dimethyl-1,3-dioxan-4,6-dione (Meldrum's Acid) [2033-24-1] $C_6H_8O_4$, **M** 144.1, **m** 92-96°, 94-95°, **pK**₂₅²⁵ 5.1, 7.32. Crystallise the dione from Me₂CO/H₂O. It is a useful synthon for the C3 malonic acid moiety. [Arnett et al. *J Am Chem Soc* **106** 6759 1984, DOI: 10.1021/ja00334a049; Bihlmayer et al. *Monatsh Chem* **98** 564 1967, DOI: 10.1007/BF00901364; Review: McNab *Chem Soc, Rev* **7** 345 1978, Chan & Huang *Synthesis* 452 1982, DOI: 10.1055/s-1982-29829; *Beilstein* **19/5** V 8.] It is synthesised by a modified procedure in which concentrated H₂SO₄ (1.5ml) is added to a stirred suspension of powdered malonic acid (52g, 0.5 mole) in acetic anhydride (60ml, 0.6 mole), whereby the malonic acid dissolves with spontaneous cooling. Acetone (40ml, 0.55 mole) is added to the mixture while keeping the temperature at 20-25°, then cooled in a refrigerator overnight, the crystals are filtered off, washed three times with ice-water (enough to cover the crystals) and air dried. The crude product (35g, 49%) is recrystallised without heating by dissolving it (10g) in Me₂CO (20ml), filtering, and adding H₂O (40ml), with ~70% recovery of material **m** 94-95°. [Pihlaja & Seilo *Acta Chem Scand* **22** 3053 1968, DOI: 10.3891/acta.chem.scand.22-3053.] *Alternatively*, concentrated H₂SO₄ (0.5ml) is added dropwise to a stirred suspension of powdered malonic acid (52g, 0.5 mole) in redistilled isoprenyl acetate (62ml, 55g, [108-22-5]), when the temperature rose from 23° to 31° in 45 minutes and all the solid dissolved within 1 hour. Treatment of the reaction as above gave Meldrum's acid (37g, 50%). [Davidson & Bernhard *J Am Chem Soc* **70** 3426 1948, DOI: 10.1021/ja01190a060]. Its ¹H NMR (CDCl₃, TMS) has δ at 1.73 (s, 2-Me₂) and 3.60 (s, 5-H) [Schuster & Schuster *Tetrahedron* **25** 199 1969, DOI: 10.1016/S0040-4020(01)99472-4]. A similar synthesis can be used with substituted malonic acids.

3,6-Dimethyl-1,4-dioxan-2,5-dione (Lactide) [*cis*-*RS,RS*-(±) 615-95-2, *cis*-*R,R*-(+) 95-96-5, *cis*-*S,S*-(-) 4511-42-6] $C_6H_8O_4$, **M** 144.1. This is the cyclic dilactone of lactic acid. The (±)-*cis*-*racemate* has been distilled with **b** 142°/8mm; the distillate which solidifies gives yellow needles on recrystallisation from EtOH with **m** 128°, from Et₂O with **m** 129°, or $CHCl_3$ with **m** 126°, with IR ν_{max} at 1720-1740 cm⁻¹. It hydrolyses in cold

H₂O. [Carothers et al. *J Am Chem Soc* **54** 761 1932, DOI: 10.1021/ja01341a046]. A *trans-form* (probably *RS,SR*) has been reported which crystallises from Et₂O with **m 42-43°** [Hummel et al. *Acta Cryst (Sect B)* **38** 1679 1982, DOI: 10.1107/S0567740882006840]. The *R,R-(+)-lactide* has **b 150°/2mm** and crystallises from Et₂O with **m 95°**, or **m 96.5-97.5°** (from CHCl₃) or **m 97.7°** (from EtOAc) and **[α]_D²² +297 (c 1.2, *C₆H₆)**. The *S,S-(-)-lactide* has **b 150°/2.5mm** and crystallises from EtOAc with **m 98.7°**, or **m 95°** (from CHCl₃) or **m 96.5-97.5°** (from CCl₄) and **[α]_D²² -297 (c 1.2, *C₆H₆)**. [Toniolo et al. *J Org Chem* **35** 6 1970, DOI: 10.1021/jo00826a002; *Beilstein* **19** H 154, **19** I 179, **19** II 176, **19** IV 1927, **19**/V 10.]

2,9-Dimethyl-4,7-diphenyl-1,10-phenanthroline (BCP, Bathocuproine) [4733-39-5] C₂₆H₂₀N₂, **M 360.5, m 279-283°, >280°, pK_{Est} ~5.6**. Purify it by recrystallisation from *benzene. It complexes with Cu. [Smith & Wilkins *Anal Chem* **25** 510 1953, DOI: 10.1021/ac60075a037; *Beilstein* **23** III/IV 2160.] For the *disulfonic acid disodium salt* and its analytical applications see ‘Miscellaneous As, B, P, Si, S, Se and Te Compounds’ in this Chapter.

2,2-Dimethylethyleneimine (2,2-dimethylaziridine) [2658-24-4] **M 71.1, b 70.5-71.0°, 72°, n_D²⁰ 1.405, pK²⁵ 8.64**. Dry the 2,2-dimethylaziridine over solid KOH, filter and freshly distil from sodium before use, and store it under dry nitrogen. The *N-phenylthiocarbamoyl* derivative crystallises from petroleum ether containing a trace of Me₂CO with **m 92.5-93.5°**. [Hassner et al. *J Am Chem Soc* **91** 5046 1969, DOI: 10.1021/ja01046a019; Cairns *J Am Chem Soc* **63** 871 1941, DOI: 10.1021/ja01848a504; Lamaty et al. *Justus Liebigs Ann Chem* **726** 77 1969, DOI: 10.1002/jlac.19697260112; *Beilstein* **20** III/IV 280.]

5,5-Dimethylhydantoin [77-71-4] C₅H₈N₂O₂, **M 128.1, m 174-177°, 177-178°, pK²⁴ 9.19**. Crystallise the hydantoin from EtOH and sublime it *in vacuo*. [*Beilstein* **24** III/IV 1097.]

4,6-Dimethyl-2-hydroxypyrimidine [108-79-2] **M 124.1, m 198-199°, 202-205°, pK₁²⁰ 3.77, pK₂²⁰ 10.50**. Crystallise the pyrimidine from absolute EtOH (charcoal). [*Beilstein* **24**/2 V 138.]

1,2-Dimethyl-1H-imidazole [1739-84-0] C₅H₈N₂, **M 96.1, m 37-39°, 38-40°, b 204°/atm, 206°/760mm, d₄²⁰ 1.084, pK_{Est} ~8.1**. Crystallise the imidazole from *benzene, dry and store it at 0-4°. The *picrate* crystallises from H₂O or EtOH with **m 181°**. [Balaban & Pymann *J Chem Soc* **125** 1564 1924, DOI: 10.1039/CT9242501564; Gorun et al. *J Am Chem Soc* **109** 4244 1987, DOI: 10.1021/ja00248a019; *Beilstein* **23** H 66, **23** II 56, **23** III/IV 594.]

1,3-Dimethyl-2-imidazolinone (DMI, N,N'-dimethylethyleneurea, DMEU) [80-73-9] C₅H₁₀N₂O, **M 114.2, m 8.2°, b 67-68°/2mm, 104°/5mm, 106-107°/17mm, 224-226°/atm, d₄²⁵ 1.056, n_D²⁰ 1.472**. After preparation by reaction of *N,N'*-dimethylethylenediamine in toluene with phosgene in toluene below 15°, excess phosgene is removed by blowing air through the mixture. The hydrochloride salt that separates is removed by filtration, and washed with CHCl₃. The combined filtrate and washings are evaporated off, and the residue is distilled (b 129°/39mm). The crude distillate is treated with H₂O; K₂CO₃ is added to saturate the solution, and extracted with CHCl₃, the extract is dried (K₂CO₃), filtered, and pure **1,3-dimethylimidazolin-2-one** distils at **b 104°/5mm**. It has been used as an alternative to HMPA [see 680-31-9] as a high dielectric solvent for reactions. [Boon *J Chem Soc* 307 1947, DOI: 10.1039/JR9470000307; Lien & Kumler *J Med Chem* **11** 214 1968, DOI: 10.1021/jm00308a005; Kohn et al. *J Org Chem* **42** 941 1977, DOI: 10.1021/jo00426a003; *Beilstein* **24** III/IV 9.]

Cis-2,3-dimethyloxirane (cis-epoxybutane) [1758-33-4] C₄H₈O, **M 72.1, m -84 to -83°, b 59.7°/742mm, 61°/atm, 59-61°/atm, d₂₅²⁵ 0.8226, n_D²⁰ 1.3830**. Dry the oxirane over CaCl₂ and redistil slowly through a short Vigreux column [Cornforth & Green *J Chem Soc C* 846 1970, DOI: 10.1039/J39700000846]. Its IR spectrum is described by H. van Risseghem [*Bull Soc Chim Fr* 1661 1959]. Crude *cis*-2,3-dimethyloxirane, obtained from the *m*-chloroperbenzoic acid oxidation of *cis*-2-butene, is distilled off and the fraction boiling up to 100° is collected and fractionated through a 2 ft helices-packed column to give a 52-60% yield of pure *cis*-dimethyloxirane **b 58-59°/748mm**. The stereochemical purity of the oxirane was shown to be greater than 99.5% by gas chromatography on a 30-ft, 20% Carbowax 20M on Chromosorb P column at 150°, with retention time 10.0 minutes (compare with 8.7 minutes for *trans*-oxirane). [Pasto & Cumbo *J Org Chem* **30** 1271 1965, DOI: 10.1021/jo01015a523.] [*Beilstein* **17** III/IV 45, 48, **17** V/1 61.]

Trans-2,3-dimethyloxirane (trans-epoxybutane) [*RS* 21490-63-1, 6189-41-9] C_4H_8O , *M* 72.1, *b* 54-55°/atm, d_4^{25} 0.804, n_D^{20} 1.3730. The *trans*-oxirane, obtained by the *m*-chloroperbenzoic acid oxidation of *trans*-2-butene, as in the preceding entry has *b* 52.0-53.0°/748mm and was shown to be 99.5% stereochemically pure by gas chromatography as for the *cis* isomer. [Pasto & Cumbo *J Org Chem* **30** 1271 1965, DOI: 10.1021/jo01015a523.] The chiral oxiranes obtained from the respective 2-bromo-3-acetoxbutanes [Mori & Tamada *Tetrahedron* **35** 1279 1979, DOI: 10.1016/0040-4020(79)80054-X] or butane-2,3-diol [Newman & Chen *J Org Chem* **38** 1173 1973, DOI: 10.1021/jo00946a023] with aqueous KOH are purified by distillation after drying with KOH pellets or $CaCl_2$. **2*R*,3*R*-(+)-2,3-dimethyloxirane** [1758-32-3] has *b* 53.5-53.7°/745mm, 56-58°/atm, d_4^{25} 0.7998, n_D^{20} 1.3729, n_D^{25} 1.3705, $[\alpha]_D^{20}$ +58.0 (c 2.31, Et_2O), +76.2 (c 0.0613 xylene), and **2*S*,3*S*-(-)-2,3-dimethyloxirane** [63864-69-7] has *b* 56-58°/atm, n_D^{20} 1.3728, $[\alpha]_D^{20}$ -61.5 (c 2.11, Et_2O), $[\alpha]_D^{25}$ -44.3 (neat, dcm) [Lucas & Garner *J Am Chem Soc* **70** 990 1948, DOI: 10.1021/ja01183a028; Pasto et al. *J Am Chem Soc* **88** 2194 1966, DOI: 10.1021/ja00962a022.] The oxiranes have similar IR (film), ν_{max} 2990s, 2940m, 1490m, 1450s, 1380s, 1336m, 1280w, 1260w, 1150w, 1110s, 1020vs, 950w, 880s, 810s, 735m and 720m cm^{-1} ; 1H NMR (60MHz, TMS, CCl_4), δ 1.20 (6H, d, *J* 5Hz) 2.51 (2H, dq, *J*₁ 1.5 Hz, *J*₂ 5 Hz), and MS: *m/z* 72 ($M^+ = C_4H_8O$). [Beilstein 17 V/1 61, 62.]

3,3-Dimethyloxetane [6921-35-3] $C_5H_{10}O$, *M* 86.1, *b* 79.2-80.3°/760mm, 81°/765mm, d_4^{20} 0.836, n_D^{20} 1.399, Purify 3,3-dimethyloxetane by gas chromatography using a 2m silicone oil column or distil it. Fractionate it at atmospheric pressure (preferably under N_2 or Ar. [Beilstein 17 II 21.]

2,9-Dimethyl-1,10-phenanthroline (neocuproine hemihydrate) [484-11-7] $C_{14}H_{12}N_2$, *M* 208.3, 217.3 (hemihydrate), *m* 162-164°, pK^{25} 5.85. Purify it as the *hemihydrate* (*m* 156-160°) by crystallisation from H_2O and as the *anhydrous* base from *benzene or ligroin. It also forms a *dihydrate* as needles from H_2O . The hydrates lose H_2O on drying at ~80° or over P_2O_5 *in vacuo*. [O'Reilly & Plowman *Aust J Chem* **13** 145 1960, DOI: 10.1071/CH9600145; Beilstein 23/8 V 527.]

4,4-Dimethyl-2,6-piperidinedione (4,4-dimethylglutarimide) [1123-40-6] $C_7H_{11}NO_2$, *M* 141.2, *m* 144-146°, pK_{Est} ~11.5. Recrystallise the imide from hot H_2O or EtOH [Arnett & Harrelson *J Am Chem Soc* **109** 809 1987, DOI: 10.1021/ja00237a028]. [Beilstein 21 H 391, 21 I 331, 21 II 309, 21 III/IV 4601, 21/9 V 592.]

2,5-Dimethylpyrazine [123-32-0] $C_6H_8N_2$, *M* 108.1, *b* 155°/atm, 156°/atm, d_4^{20} 0.990, n_D^{20} 1.502, pK_1^{25} -4.6 (aqueous H_2SO_4), pK_2^{25} 1.85. Purify it *via* its *picrate* (*m* 150°) which is decomposed with a base (e.g. KOH) and distilled. [Wiggins and Wise *J Chem Soc* 4780 1956, DOI: 10.1039/JR9560004780]. [Beilstein 23/5 V 403.]

3,5-Dimethylpyrazole [67-51-6] $C_6H_8N_2$, *M* 96.1, *m* 105-108°, 107-108°, *b* 218°/atm, pK^{20} 4.16. Recrystallise it from cyclohexane or water. [Barszcz et al. *JCS Dalton Trans* 2025 1986, DOI: 10.1039/DT9860002025; Beilstein 23/5 V 110.]

2,3-Dimethylquinoxaline [2379-55-7] $C_{10}H_{10}N_2$, *M* 158.2, *m* 104-108°, 106°, pK^{25} -3.84 (aqueous H_2SO_4). It has been purified by steam distillation with the base crystallising in the distillate. Recrystallise it from distilled water or aqueous EtOH. The *sulfate* crystallises from EtOH with *m* 151-152°(dec). [Gibson *J Chem Soc* 342 1927, DOI: 10.1039/JR9270000342; Beilstein 23 H 191, 23 II 197, 23 III/IV 1277.]

2,4-Dimethylsulfolane [1003-78-7] $C_6H_{12}O_2S$, *M* 148.2, *b* 123.3°/5mm, 128°/77mm, 280-281°/atm (with some dec), d_D^{25} 1.1314, n_D^{20} 1.474. Distil the yellow 2,4-dimethylsulfolane in a vacuum. It is a useful solvent. [Beilstein 17/1 V 82.]

1,3-Dimethyluracil [1,3-dimethyl-2,4(1*H*,3*H*)-pyrimidinedione] [874-14-6] $C_6H_8N_2O_2$, *M* 140.1, *m* 119-122°, 121-122°, pK^{25} -3.25 (aqueous H_2SO_4). Crystallise it from EtOH/ether. [Beilstein 24 III/IV 1196.]

9,9-Dimethylxanthene [4,4-dimethyl-2,3:5,6-dibenzopyran] [19814-75-6] $C_{15}H_{14}O$, *M* 210.3, *m* 35-38°, 36-38°, *b* 114-115°/0.6mm, n_D^{20} 1.5973. It is prepared under argon (Schlenk equipment) by adding Me_3Al solution (2.0M in toluene, 320ml, 0.64mmol), *via* a septum over 50 minutes, to an ice-water cooled stirred suspension of 9-xanthone (50.0g, 0.255mol) in toluene (300ml) (exothermic reaction, no gas evolved). The

dark red solution is allowed to warm to ~25° over 3 hours, and stirred for a further 14 hours. (*Alternatively*, it can be heated to 60° for ~0.5 hours.) The mixture is transferred (*via* a cannula under argon pressure) to a stirred mixture of concentrated HCl (250ml) and ice (4 L). The organic phase is separated, dried (MgSO₄), filtered, the solvent is evaporated *in vacuo* to give a yellow oil (51.5g, 96%) which distils at 114-115°/0.6mm, and solidifies on cooling. The ¹H NMR (CDCl₃, TMS) has δ at 1.63 (s, 6H, CH₃), 7.0-7.6 (m, 8H, aromatic-H); MS has *m/z* at 210 (M⁺), 195 (base peak, M⁺-CH₃). [Nowick et al. *J Am Chem Soc* **112** 8902 1990, DOI: 10.1021/ja00180a038; Meisters & Mole *Aust J Chem* **27** 1655 1974, DOI: 10.1071/CH9741655; *Beilstein* **17** II 287.]

1,3-Dioxolan-2-one (ethylene carbonate) [96-49-1] C₃H₄O₃, M 88.1, m 35-38°, 37°, 39°, 40°, b 65-67°/1mm, 126°/17mm, 238°/760mm, 243-244°/atm, d₄²⁰ 1.321, n_D⁴⁰ 1.4199. Dry 1,3-dioxolan-2-one over P₂O₅, then fractionally distil it at low or atmospheric pressure. Recrystallise it from dry Et₂O (plates, m 36.5°, 38.5-40° was also reported). It is soluble in H₂O. [*Beilstein* 19 II 135, 19 III/IV 1556, 19/4 V 6.]

1,3-Dioxane (formaldehyde trimethylene acetal) [505-22-6] C₄H₈O₂, M 88.1, m -45°, b 104.5°/751mm, 105-106°/atm, d₄²⁰ 1.040, n_D²⁰ 1.417. Dry the dioxane with Na and fractionally distil it. [*Beilstein* 19/1 V 11.]

1,4-Dioxane (Dioxane, diethylene oxide) [123-91-1] has m 10-12°, 11.8°, b 12°/20mm, 34°/60mm, 45°/100mm, 82°/400mm, 101.1°/760mm, 100-102°/atm, 101.3°/atm, d₄²⁵ 1.0292, n_D¹⁵ 1.4236, n_D²⁵ 1.42025. It is prepared commercially either by dehydration of ethylene glycol with H₂SO₄ and heating ethylene oxide or bis(β-chloroethyl)ether with NaOH. The usual impurities are acetaldehyde, ethylene acetal, acetic acid, water and peroxides. Peroxides can be removed (and the aldehyde content decreased) by percolation through a column of activated alumina (80g per 100-200ml solvent), by refluxing with NaBH₄ or anhydrous stannous chloride and distilling, or by acidification with concentrated HCl, shaking with ferrous sulfate and leaving in contact with it for 24 hours before filtering and purifying further. Hess and Frahm [*Chem Ber* **71** 2627 1938, DOI: 10.1002/cber.19380711234] refluxed 2L of dioxane with 27ml concentrated HCl and 200ml water for 12 hours with slow passage of nitrogen to remove acetaldehyde. After cooling the solution, KOH pellets were added slowly and with shaking until no more would dissolve and a second layer had separated. The dioxane was decanted, treated with fresh KOH pellets to remove any aqueous phase, then transferred to a clean flask where it was refluxed for 6-12 hours with sodium, then distilled from it. *Alternatively*, Kraus and Vingee [*J Am Chem Soc* **56** 511 1934, DOI: 10.1021/ja01318a004] heated it on a steam bath with solid KOH until fresh addition of KOH gave no more resin (due to acetaldehyde). After filtering through paper, the dioxane was refluxed over sodium until the surface of the metal was not further discoloured during several hours. It was then distilled from sodium.

The acetal (b 82.5°) is removed during fractional distillation. Traces of *benzene, if present, can be removed as the *benzene/MeOH azeotrope by distillation in the presence of MeOH. Distillation from LiAlH₄ removes aldehydes, peroxides and water. Dioxane can be dried using Linde type 4X molecular sieves. Other purification procedures include distillation from excess C₂H₅MgBr, refluxing with PbO₂ to remove peroxides, fractional crystallisation by partial freezing and the addition of KI to dioxane acidified with aqueous HCl. Dioxane should be stored out of contact with air, preferably under N₂.

A detailed purification procedure is as follows: Dioxane is stood over ferrous sulfate for at least 2 days, under nitrogen. Then water (100ml) and conc HCl (14ml)/ litre of dioxane are added (giving a pale yellow colour). After refluxing for 8-12 hours with vigorous N₂ bubbling, pellets of KOH are added to the warm solution to form two layers and to discharge the colour. The solution is cooled rapidly with more KOH pellets being added (magnetic stirring) until no more dissolved in the cooled solution. After 4-12 hours, if the lower phase is not black, the upper phase is decanted rapidly into a clean flask containing sodium, and refluxed over sodium (until freshly added sodium remained bright) for 1 hour. The middle fraction is collected (and checked for minimum absorbency below 250nm). The distillate is fractionally frozen three times by cooling in a refrigerator, with occasional shaking or stirring. This material is stored in a refrigerator. Before use it is thawed, refluxed over sodium for 48 hours, and distilled into a container. All joints are clad with Teflon tape.

Coetzee and Chang [*Pure Appl Chem* **57** 633 1985, DOI: org/10.1351/pac198557040633] dried the solvent by passing it slowly through a column (20g/L) of 3A molecular sieves activated by heating at 250° for 24 hours. Impurities (including peroxides) are removed by passing the effluent slowly through a column packed with type NaX zeolite (pellets ground to 0.1mm size) activated by heating at 400° for 24 hours or chromatographic grade basic Al₂O₃ activated by heating at 250° for 24 hours. After removal of peroxides the

effluent is refluxed for several hours over sodium wire, excluding moisture, distilled under nitrogen or argon and stored in the dark. One of the best tests of purity of dioxane is the formation of the purple **disodium benzophenone complex** during reflux and its persistence on cooling. (Benzophenone is better than fluorenone for this purpose and for storing the solvent.) [Carter et al. *Trans Faraday Soc* **56** 343 1960, DOI: 10.1039/TF9605600343; *Beilstein* **19** V 16.] **TOXIC — do not inhale vapour.**

Rapid purification: Check for peroxides (see Chapter 1 and Chapter 2 for test under ethers). Pre-dry with CaCl_2 or better over Na wire. Then reflux the pre-dried solvent over Na (1% w/v) and benzophenone (0.2% w/v) under an inert atmosphere until the blue colour of the benzophenone ketyl radical anion persists. Distil, and store it over 4A molecular sieves in the dark.

1,3-Dioxolane (formaldehyde ethylene acetal) [646-06-0] $\text{C}_3\text{H}_6\text{O}_2$, M **74.1**, m **-95°**, b **74-75°/atm 75.0-75.2°/atm** d_4^{20} **1.0600**, n_D^{21} **1.3997**. Dry it with solid NaOH, KOH or CaSO_4 , and distil it from sodium or sodium amalgam. Barker et al. [*J Chem Soc* 802 1959, DOI: 10.1039/JR9590000802] heated 34ml of dioxalane under reflux with 3g of PbO_2 for 2 hours, then cooled and filtered. After adding xylene (40ml) and PbO_2 (2g) to the filtrate, the mixture is fractionally distilled. Addition of xylene (20ml) and sodium wire to the main fraction (b 70-71°) led to a vigorous reaction, following which the mixture was again fractionally distilled. Xylene and sodium additions are made to the main fraction (b 73-74°) before it is finally distilled. [*Beilstein* **19/1** V 6.]

1,3-Diphenylisobenzofuran [5471-63-6] $\text{C}_{20}\text{H}_{14}\text{O}$, M **270.3**, m **128-130°, 129-130°**. Recrystallise it from EtOH or EtOH/ CHCl_3 (1:1) under red light (as in photographic dark rooms) or from *benzene in the dark. [*Beilstein* **17/2** V 503.]

2,5-Diphenyl-1,3,4-oxadiazole (PPD) [725-12-2] M **222.3**, m **70° (hydrate), 139-140° (anhydrous), b 231°/13mm, 248°/16mm**. Its solubility in CHCl_3 is 10%. Crystallise it from EtOH and sublime it *in vacuo*. [*Beilstein* **27** III/IV 2712.]

2,5-Diphenyloxazole (PPO) [92-71-7] $\text{C}_{15}\text{H}_{11}\text{NO}$, M **221.3**, m **72-74°, 74°**, b **360°/atm**. Distil it in steam and crystallise it from ligroin. Scintillation material for radioactive counting. [*Beilstein* **27** III/IV 1437.]

2RS,5RS-(±)-2,5-Diphenylpyrrolidine (± trans isomer) [22147-84-8] $\text{C}_{16}\text{H}_{17}\text{N}$, M **223.3**, b **136°/0.3mm**, n_D^{25} **1.5866**, $\text{pK}_{\text{Est}} \sim 8.0$. A mixture of *cis*-(*meso*) and *trans*-(±) pyrrolidines (b ~130-145°/0.15-0.2mm, ~138g, 0.6mol) in EtOH (210ml) and concentrated HCl (100ml) was treated dropwise, during ~2 hours, with a solution of NaNO_2 (61.5g, 0.9mol) in H_2O (100ml) at ~0° (ice-bath cooling). The mixture was poured into H_2O (1 L) and the precipitate (~160g) was collected, washed with H_2O and fractionally recrystallised from Me_2CO to give the *trans*-nitroso derivative (m **140-140.9°**, 51%) [22147-82-6], and the *cis*-nitroso derivative (m **97-98°**, 20%) [22147-81-5]. This *trans*-nitroso-pyrrolidine was de-nitrosated (HCl gas in Et_2O) to the *trans*-hydrochloride m **196.5-180°** which was recrystallised from EtOH/ Et_2O . The *trans*-free base was obtained by basifying an aqueous solution to pH 9-10 and extracting with Et_2O (3 x 50ml), drying the extract (Mg_2SO_4), filtering, evaporating and distilling the residual oil (b **136°/0.3mm**). The IR (film) has ν_{max} at 1110-1025 (three peaks, one is for C-N) and 3330 (N-H) cm^{-1} . Similarly the *cis* (*meso*) *N*-nitroso derivative gave the 2RS,5SR-diphenylpyrrolidine *meso*-hydrochloride which had m **224.7-225.5°**, and the *meso*-free base [22147-83-7] had b **121-123.4°/0.2mm**, n_D^{25} **1.5850**. The optically active 2R,5R-(+)-diphenylpyrrolidine enantiomer [155155-73-0] had m **47-52°, 49-53°**, $[\alpha]_D^{22}$ **+118** (c **1.0**, CHCl_3). [Overberger et al. *J Am Chem Soc* **91** 687 1969, DOI: 10.1021/ja01031a029; Breuer & Melumad *J Org Chem* **37** 3949 1972, DOI: 10.1021/jo00797a043.]

(±)-α, α-Diphenyl-2-pyrrolidinemethanol (α, α-diphenyl-DL-prolinol) [63401-04-7, 112022-88-5] $\text{C}_{17}\text{H}_{19}\text{NO}$, M **253.3**, m **82-83° (76° also reported)**, $\text{pK}_{\text{Est}} \sim 9.5$. It is prepared as the (±)-hydrochloride [16226-54-3] M **289.8**, m **267-269° (dec, 262-263° was also reported)** by dissolving the amine in Et_2O , $\text{Et}_2\text{O}/\text{MeOH}$ or CH_2Cl_2 , and dry HCl gas is bubbled through (avoid excess HCl as it may dissolve precipitated NH_4Cl), filtering, evaporating, and the residue is recrystallised from $\text{MeOH}/\text{Me}_2\text{CO}$; and from which the free base is obtained by treatment with aqueous NaOH and extraction with Et_2O . It has psychostimulating activity. [Enders et al. *Org Synth Coll Vol* **6** 542 1988, DOI: 10.15227/orgsyn.058.0113;

Enders et al. *Synthesis* 548 1976, DOI: 10.1055/s-1976-24121; Likhoshesterov et al. *Khim Farm Z* 1 30 1967, *Chem Abstr* 67 90642 1967. *Beilstein* 21 III/IV 1519.]

R-(+)- α , α -Diphenyl-2-pyrrolidinemethanol (α , α -Diphenyl-D-prolinol) [22348-32-9] $C_{17}H_{19}NO$, M 253.3, m 77-80°, 78-80°, $[\alpha]_D^{20}$ +69 (c 3, $CHCl_3$). It is purified by recrystallisation from EtOH or hexane. It was also purified via the (*R*)-hydrochloride [172152-19-1] as above.

S-(-)- α , α -Diphenyl-2-pyrrolidinemethanol (α , α -diphenyl-L-prolinol) [112068-01-6] has m 77-80°, 79-79.5°, 83°, $[\alpha]_D^{20}$ -68.1 (c 3.2, $CHCl_3$), $[\alpha]_D^{20}$ -87.5 (c 1.16, CH_2Cl_2). It is purified by recrystallisation from EtOH or hexane. The S-(-)-hydrochloride [148719-90-8] melts above 240°, and the benzoyl derivative m 183° crystallises from H_2O , MeOH or EtOH. [Enders et al. *Bull Soc Chim Belg* 97 691 1988, Kerrick & Beak *J Am Chem Soc* 113 9708 1991, DOI: 10.1021/ja00025a066; Corey & Bakshi *Tetrahedron Lett* 31 611 1990, DOI: 10.1016/S0040-4039(00)94581-7; Mathre *J Org Chem* 58 2880 1993, DOI: 10.1021/jo00062a037; *Beilstein* 21 III/IV 1519.]

Dipicolinic acid (pyridine-2,6-dicarboxylic acid) [499-83-2] $C_7H_5NO_4$, M 167.1, m 248-250°(dec), 255°(dec), λ_{max} 270nm, pK_1^{20} 2.10, pK_2^{20} 4.68. Recrystallise the acid from water, and sublime it in a vacuum. [*Beilstein* 22/4 V 128.]

Di-(4-pyridoyl)hydrazine (*N,N'*-di-isonicotinoyl hydrazine) [4329-75-3] M 246.2, m 254-255°, 259-260°. Crystallise it from water, aqueous EtOH or propan-1-ol. [Albert & Rees *Biochem J* 61 128 1955, DOI: 10.1042/bj0610128; *Beilstein* 22 III/IV 663.]

2,2'-Dipyridylamine [1202-34-2] $C_{10}H_9N_3$, M 171.2, m 84° and remelts at 95° after solidifying, 90-92°, b 176-178°/13mm, 222°/50mm, 307-308°/760mm, pK^{25} 6.69 (in 20% aqueous EtOH). Crystallise the amine from *benzene or toluene [Blakley & DeArmond *J Am Chem Soc* 109 4895 1987, DOI: 10.1021/ja00250a023]. The amine is also recrystallised from Me_2CO (m 95.1°) or distilled in a vacuum. [*Beilstein* 22 I 630, 22 II 331, 22 III/IV 3961, 22/8 V 415.]

2,2'-Dipyridyl disulfide (2,2'-dithiopyridine, Aldrithiol-2) [2127-03-9] $C_{10}H_8N_2S_2$, M 220.3, m 53°, 56-58°, 57-58°, pK_1^{25} 0.35, pK_2^{25} 2.45. Recrystallise the disulfide H_2O from * C_6H_6 /petroleum ether (6:7), ligroin or * C_6H_6 . The picrate has m 119° (from EtOH). [Walter et al. *Justus Liebigs Ann Chem* 695 77 1966, DOI: 10.1002/jlac.19666950110; Marckwald et al. *Chem Ber* 33 1556 1900, DOI: 10.1002/cber.19000330226; Brocklehurst & Little *Biochem J* 133 67,78 1973, DOI: 10.1042/bj1330067; *Beilstein* 21 III/IV 48.] It has been used as a 1mM solution in EtOH for the spectrophotometric estimation of thiols. Essentially the thiol displaces half the disulfide molecule liberating the 2-mercaptopyridine anion, thereby shifting the λ_{max} from 340nm (of the disulfide) to 268nm (of the anion) at pH 9, or 278nm in H_2O . (Compare with 4,4'-dipyridyl disulfide (below) which has been used for the same purpose [Humphrey et al. *Anal Chem* 42 698 1970, DOI: 10.1021/ac60289a021; for review see Aldrich in *Aldrichimica Acta* 4 33 1971].

4,4'-Dipyridyl disulfide (4,4'-dithiopyridine, Aldrithiol-4) [2645-22-9] has m 74-76°, 76-78°, $pK_{Est(1)}$ ~1.5, $pK_{Est(2)}$ ~4.5. Recrystallise the disulfide from H_2O , EtOH, Me_2CO , * C_6H_6 or petroleum ether. It has been used as a 1mM solution in EtOH for the spectrophotometric estimation of thiols. Essentially the thiol displaces half the disulfide molecule liberating the 4-mercaptopyridine (4-pyridinethiol) anion, thereby shifting the λ_{max} from 324nm (of the disulfide) to 285nm (of the anion) at pH 9. (Compare with 2,2'-dipyridyl disulfide above which has been used for the same purpose.) [Humphrey et al. *Anal Chem* 40 698 1970, DOI: 10.1021/ac60289a021; Cheng & Ritchie *Aust J Chem* 26 1785 1973, DOI: 10.1071/CH9731785; Hansen et al. *Anal Biochem* 363 77 2007, DOI: 10.1016/j.ab.2007.01.002.] [*Beilstein* 21 II 35, for review see Aldrich in *Aldrichimica Acta* 4 33 1971.]

1,2-Di-(4-pyridyl)-ethane [4916-57-8] $C_{12}H_{12}N_2$, M 184.2, m 110.9-111.2°, 114.5-116°, b 167-174°/3mm, $pK_{Est(1)}$ ~3.8, $pK_{Est(2)}$ ~5.4. Crystallise the ethane from cyclohexane/*benzene (3:1, solubility is ~7.5g/100ml). The dihydrochloride crystallises from EtOH with m 329-330°(dec). [Bergmann et al. *J Am Chem Soc* 74 5979 1952, DOI: 10.1021/ja01143a047; Thayer & Carson *J Am Chem Soc* 70 2330 1948, DOI: 10.1021/ja01187a007; Jampolsky et al. *J Am Chem Soc* 74 5222 1952, DOI: 10.1021/ja01140a516; Chow & Fuoss *J Am Chem Soc* 80 1095 1958, DOI: 10.1021/ja01538a020; *Beilstein* 23 III/IV 1389.]

trans-1,2-Di-(2-pyridyl)-ethylene [1437-15-6] $C_{12}H_{10}N_2$, M 182.2, m 118-119°, 150-160°/2mm, $pK_{Est(1)} \sim 2.0$, $pK_{Est(2)} \sim 4.9$. Crystallise the ethylene from water (1.6g/100ml at 100°). The *dihydrochloride* has m 240°, from EtOH. [Beilstein 23 I 54.]

trans-1,2-Di-(4-pyridyl)-ethylene [13362-78-2] has m 148-152°, 153-154°, 155.5-156.5°, $pK_1^{25} 3.65$, $pK_2^{25} 5.6$. Crystallise the ethylene from water (1.6g/100ml at 100°). The *dihydrochloride* has m 347°, from EtOH. [Beilstein 23/8 V 239.]

1,3-Di-(4-pyridyl)-propane [17252-51-6] $C_{13}H_{14}N_2$, M 198.3, m 60.5-61.5°, 62-65°, 65-68.5°, $pK_{Est(1)} \sim 4.5$, $pK_{Est(2)} \sim 5.5$. Crystallise the propane from *n*-hexane/*benzene (5:1) or Me₂CO. The *picrate* has m 185-185°. [Jampolsky et al. *J Am Chem Soc* 74 5222 1952, DOI: 10.1021/ja01140a516; Chow & Fuoss *J Am Chem Soc* 80 1095 1958, DOI: 10.1021/ja01538a020; Beilstein 23 III/IV 1400.]

2,5-Distyrylpyrazine [14990-02-4] $C_{20}H_{16}N_2$, M 284.3, m 219°. Recrystallise it from xylene; chromatograph it on basic silica gel (60-80 mesh) using CH₂Cl₂ as eluent, then sublime it in a vacuum on to a cold surface at 10⁻³ torr [Ebied et al. *JCS Faraday Trans 1* 78 3213 1982, DOI: 10.1039/F19827803213]. All operations should be carried out in the dark.

1,3-Dithiane [505-23-7] $C_4H_8S_2$, M 120.2, m 52-54°, 54°. Crystallise the 1,3-dithiane from 1.5 times its weight of MeOH at 0°, and sublime it at 40-50°/0.1mm. [Gröbel & Seebach *Synthesis* 357 1977, DOI: 10.1055/s-1977-24412; Beilstein 19/1 V 13.]

2,2'-Dithiobis(benzothiazole) [120-78-5] $C_{14}H_8N_2S_2$, M 332.2, m 180°, 182.5-183.5°, 186°. Recrystallise it from *benzene. [Beilstein 27 H 109, 27 III/IV 1862.]

4,4'-Dithiodimorpholine (S,S'-di-*N,N'*-dimorpholine, dimorpholine-*N,N'*-disulfide) [103-34-4] $C_8H_{16}N_2O_2S_2$, M 236.2, m 124-125°. Recrystallise it from hot aqueous dimethylformamide or EtOH. It is a fungicide. [Blake *J Am Chem Soc* 65 1267 1943, DOI: 10.1021/ja01247a004.]

1,3-Dithiole-2-thione (Vinylene trithiocarbonate, 1,3-dithiocyclopent-4-ene-2-thione, isotrithione) [930-35-8] $C_3H_2S_3$, M 134.2, m 47.5-48°, 48-50°, 50°, b 122-123°/0.8mm. Purify the thione by recrystallisation from EtOH and/or light petroleum (b 40-60°) which provides yellow needles. It has been distilled at high vacuum. The UV (cyclohexane) has λ_{max} nm (ϵ): 228.5 (8,700), 276 (1,440) and 362 (14,700). When treated with MeI in MeCN at room temperature the *S-methiodide* separates after a few hours and crystallises from MeOH with m 131-132° (dec). The same methiodide (m 132°) can also be obtained by treatment with Me₂SO₄ in *C₆H₆ or Me₂CO followed by KI. With CuCl₂ it forms the brick-red precipitate of $C_3H_2S_3 \cdot CuCl_2$, with HgCl₂ it gives yellow $C_3H_2S_3 \cdot HgCl_2$ [m 222-223° (dec)], and with AgNO₃ in EtOH it yields the yellow precipitate of $C_3H_2S_3 \cdot AgNO_3$ [m 124°(dec)]. Upon reducing the *methiodide* with NaBH₄ in MeOH it forms *tetrathiafulvalene* (TTF) [see below]. [Challanger et al. *J Chem Soc* 292 1953, DOI: 10.1039/JR9530000292; Melby et al. *J Org Chem* 39 2456 1974, DOI: 10.1021/jo00930a043; Wudl & Kaplan *Inorg Synth* 19 27 1979; DOI: 10.1002/9780470132500.ch7; Guzic Jr et al. *JCS Perkin Trans 1* 1068 1989, DOI: 10.1039/P19890001068; Beilstein 19/4 V 74.]

1-Dodecylpyridinium chloride [104-74-5] M 301.9, m 68-70°. Purify the chloride by repeated crystallisation from acetone (charcoal); then recrystallise it twice from EtOH [Chu & Thomas *J Am Chem Soc* 108 6270 1986, DOI: 10.1021/ja00280a026]. It is *hygroscopic* and should be stored with a desiccant. [Beilstein 20 III/IV 2314.]

Ellipticine (5,11-dimethylpyrido[4,3-*b*]carbazole) [519-23-3] $C_{17}H_{14}N_2$, M 246.3, m 311-315°(dec), 312-314°(dec), $pK^{25} 5.78$ (80% aqueous methoxyethanol). This DNA intercalator is purified by recrystallisation from CHCl₃ or MeOH and is dried *in vacuo*. The UV has λ_{max} values in aqueous EtOH/HCl at 241, 249, 307, 335 and 426nm. [Marini-Bettolo & Schmutz *Helv Chim Acta* 42 2146 1959, DOI: 10.1002/hlca.19590420643.] The *methiodide* has m 360°(dec), with UV λ_{max} (EtOH/KOH) at 223, 242, 251, 311, 362 and 432nm. [Goodwin et al. *J Am Chem Soc* 81 1903 1959, DOI: 10.1021/ja01517a031; Beilstein 23/9 V 417.]

Elymoclavine (8,9-didehydro-6-methylergoline-8-methanol) [548-43-6] $C_{14}H_{18}N_2O$, M 254.3, m 248-252°(dec), 249-253°(dec), 250-252°(dec), $[\alpha]_D^{20}$ -109 (c 0.4, EtOH), $[\alpha]_D^{20}$ -152 (c 0.9, pyridine). This Ergot alkaloid crystallises from MeOH, $CHCl_3$, Et_2O , Me_2CO or $*C_6H_6$. [Stoll et al. *Helv Chim Acta* **37** 1815 1954, DOI: 10.1002/hlca.19540370627; for structure and stereochemistry see Schreier *Helv Chim Acta* **41** 1984 1958, DOI: 10.1002/hlca.19580410708; *Beilstein* **23** III/IV 2716.] **TOXIC.**

Emetine hydrochloride [316-42-7] $C_{29}H_{40}N_2O_4 \cdot 2HCl$, M 553.6 + aq, m 235-240°, 235-250°, 240-250°, 248-250° (depending on H_2O content), $[\alpha]_D^{20}$ -49.2 (free base, c 4, $CHCl_3$), +18 (c 6, H_2O , dry salt), pK_1 5.77, pK_2 6.64. It crystallises from MeOH/ Et_2O , MeOH or Et_2O /EtOAc. The *free base* [483-18-1] has m 104-105°, and the (-)-*phenylthiourea derivative* has m 220-221° (from EtOAc/petroleum ether, $[\alpha]_D^{25}$ -29.3 ($CHCl_3$)). Its IR has ν_{max} at 3413 (OH) and 2611 (NH^+) cm^{-1} ; and UV λ_{max} at 230nm (ϵ 16 200) and 282nm (ϵ 6 890) [Brossi et al. *Helv Chim Acta* **42** 1515 1959, DOI: 10.1002/hlca.19590420513; Barash et al. *J Chem Soc* 3530 1959, DOI: 10.1039/JR9590003530]. This alkaloid is antiamoebic. [*Beilstein* **23** III/IV 3419.]

(±)-Epichlorohydrin (chloromethyloxirane, γ-chloromethylpropyleneoxide) [106-89-8] C_3H_5ClO , M 92.5, m -57°, b 16.5°/10mm, 42°/40mm, 115.5°/760mm, d_4^{25} 1.183, n_D^{20} 1.438. Distil epichlorohydrin at 760mm, heat it on a steam bath with one-quarter its weight of CaO, then decant and fractionally distil it. [Clarke & Hartman *Org Synth Coll Vol* **1** 233 1941, DOI: 10.15227/orgsyn.003.0047; Braun *Org Synth Coll Vol* **2** 256 1943, DOI: 10.15227/orgsyn.016.0030; *Beilstein* **17** H 6, **17** III/IV 20, **17** V/1 20.]

Ergocornine [564-36-3] $C_{31}H_{39}N_5O_5$, M 561.7, m 181°(dec), 182-184°, $[\alpha]_D^{20}$ -176 (c 0.5, $CHCl_3$). It crystallises with solvent of crystallisation from MeOH. [Stadler et al. *Helv Chim Acta* **52** 1549 1969, DOI: 10.1002/hlca.19690520616; *Beilstein* **25** III/IV 963, **27** II 860.]

Ergocristine [511-08-0] $C_{35}H_{39}N_5O_5$, M 609.7, m 155-157°, $[\alpha]_D^{20}$ -183 (c 0.5, $CHCl_3$). It crystallises with 2 molecules of solvent of crystallisation from *benzene. [Stadler et al. *Helv Chim Acta* **52** 1549 1969, DOI: 10.1002/hlca.19690520616; *Beilstein* **25** III/IV 966, **27** II 860.]

α-Ergocryptine [511-09-1] $C_{32}H_{41}N_5O_5$, M 575.7, m 212-214°, $[\alpha]_D^{20}$ -180 (c 0.5, $CHCl_3$). It crystallises with solvent of crystallisation, from acetone, *benzene or methanol. [Stadler et al. *Helv Chim Acta* **52** 1549 1969, DOI: 10.1002/hlca.19690520616; *Beilstein* **25** III/IV 964, **27** II 860.]

Ergotamine [113-15-5] $C_{33}H_{35}N_5O_5$, M 581.6, m 212-214°(dec), $[\alpha]_D^{20}$ -160 (c 0.5, $CHCl_3$), pK^{25} 6.40. Crystallise it from *benzene, then dry it by prolonged heating in high vacuum. It is very *hygroscopic*. [*Beilstein* **25** III/IV 964.] **Ergotamine tartrate** [379-79-3] $C_{33}H_{35}N_5O_5 \cdot 0.5 C_4H_6O_6$, M 657.1, has m ~195°(dec), 203°(dec). It crystallises from MeOH. [*Beilstein* **25** III/IV 964.]

Ergotaminine [639-81-6] $C_{33}H_{35}N_5O_5$, M 581.7, m 241-243°, $[\alpha]_D^{20}$ +369° (c 0.5, $CHCl_3$). It forms rhombic plates from MeOH which retain solvent unlike its isomer ergotamine (previous entry). It is less soluble than ergotamine, and its solubility is 0.1% in boiling ethanol and 0.07% in methanol. [Stoll *Helv Chim Acta* **28** 1283 1945, DOI: 10.1002/hlca.6602801182; *Beilstein* **25** II 860, 862, **25** III/IV 966.]

D-Erythronic acid γ-lactone (3R-3,4-dihydroxyfuran-2-one) [15667-21-7] $C_4H_6O_4$, M 118.1, m 98-100°, 100-102°, 103-104°, 104-105°, 105°, $[\alpha]_D^{20}$ -73.2 (c 0.5, H_2O), $[\alpha]_{546}^{20}$ -87.6 (c 4, H_2O). Recrystallise it from EtOAc (20 parts) or isoPrOH (3 parts). [Baker & MacDonald *J Am Chem Soc* **82** 2301 1960, DOI: 10.1021/ja01494a049; Glattfeld & Forbrich *J Am Chem Soc* **56** 1209 1934, DOI: 10.1021/ja01320a066; Weidenhagen & Wegner *Chem Ber* **72** 2010 1939, DOI: 10.1002/cber.19390721122; Musich & Rapoport *J Am Chem Soc* **100** 4865 1978, DOI: 10.1021/ja00483a037; *Beilstein* **18/2** V 457.]

Esculetin (cichorigenin, 6,7-dihydroxycoumarin) [305-01-1] $C_9H_6O_4$, M 178.2, m 271-273°, 272-275° (dec), 274° (dec), pK^{25} 8.60 (70% aqueous EtOH), $pK_{Est(1)}$ ~8.7, $pK_{Est(2)}$ ~12.4. It forms prisms from AcOH, aqueous EtOH or aqueous MeOH, and provides leaflets on sublimation in a vacuum. [Kagan *J Am Chem Soc* **88** 2617 1966, DOI: 10.1021/ja00963a064; Mabry et al. *Phytochemistry* **4** 487 1965, DOI: 10.1016/S0031-

9422(00)86201-9.] **Esculin (the 6-glucoside)** has **m 215°(dec)**, $[\alpha]_{\text{D}}^{20}$ -41 (c 5, pyridine). [*Beilstein* **18** III/IV 1322, **18/3** V 202.]

Eserine (Physostigmine, Physostol, [(3*aS-cis*)-1,2,3,3*a*,8,8*a*-hexahydro-1,3*a*,8-trimethyl-pyrrolo[2,3-*b*]indol-5-ol methylcarbamate ester] [57-47-6] $\text{C}_{15}\text{H}_{21}\text{N}_3\text{O}_2$, **M 275.4**, **m 102-104°**, **105-106°**, $[\alpha]_{\text{D}}^{17}$ -67 (c **1.3**, CHCl_3), $[\alpha]_{\text{D}}^{25}$ -120 (* C_6H_6), $\text{pK}_{\text{I}}^{25}$ **1.96**, $\text{pK}_{\text{2}}^{25}$ **8.08**. Eserine crystallises from Et_2O or * C_6H_6 and forms an unstable low melting form **m 86-87°** [Harley-Mason & Jackson *J Chem Soc* 3651 1954, DOI: 10.1039/JR9540003651; Wijberg & Speckamp *Tetrahedron* **34** 2399 1978I, DOI: 10.1016/0040-4020(78)89058-9]. It is an acetylcholinesterase inhibitor which can cross the blood-brain barrier. [*Beilstein* **23/11** V 401.]

2-Ethoxy-1-ethoxycarbonyl-1,2-dihydroquinoline (EEDQ) [16357-59-8] $\text{C}_{14}\text{H}_{17}\text{NO}_3$, **M 247.3**, **m 62-67°**, **63.5-65°**, **66-67°**. Dissolve EEDQ (~180g) in CHCl_3 , evaporate to dryness in a vacuum. Add dry Et_2O (20ml) and a white solid separates on standing. Set aside for a few hours, collect the solid, wash it thoroughly with cold Et_2O and dry it in a vacuum (~140g, **m 63.5-65°**). A further crop of solid (~25g) is obtained from the filtrate on standing overnight. [Fieser **2** 191, Belleau et al. *J Am Chem Soc* **90** 823 1968, DOI: 10.1021/ja01005a067; and **90** 1651 1968, DOI: 10.1021/ja01008a045; *Beilstein* **21/3** V 28.] It is an irreversible membrane-bound receptor antagonist [Gozlan et al. *Neuropharmacology* **33** 423 1994].

Ethoxyquin (1,2-dihydro-6-ethoxy-2,2,4-trimethylquinoline) [91-53-2] $\text{C}_{14}\text{H}_{19}\text{NO}$, **M 217.3**, **b 123-125°/0.5mm**, **169°/12-13mm**, d_4^{20} **1.000**, pK_{Est} ~ **5.8**. Purify Ethoxyquin by fractional distillation *in vacuo* whereby the distillate solidifies to a glass. [Knoevenagel **54** 1722, 1730 1921, DOI: 10.1002/cber.19210540807.] The *methiodide* has **m 179°** (from EtOH), and the *1-phenylcarbamoyl* derivative has **m 146-147°** (from EtOH). Used as a food antioxidant. [Beaver et al. *J Am Chem Soc* **79** 1236 1957, DOI: 10.1021/ja01562a053; *Beilstein* **21** III/IV 95.]

2-Ethyl-1,2-benzisoxazolium tetrafluoroborate [4611-62-5] **M 235.0**, **m 107-109°**, **109.5-110.2°**. Recrystallise it from MeCN/EtOAc to give magnificent crystals. It is not hygroscopic but on long exposure to moisture it etches glass. It is light-sensitive and should be stored in brown glass bottles and free from moisture. The UV (H_2O), has λ_{max} at 258nm (ϵ 13 100) and λ_{max} 297nm (ϵ 2 900); the IR (CH_2Cl_2) has ν_{max} at 1613 ($\text{C}=\text{N}$) and 1111-1000 (BF_4^-) [UV, IR, NMR: Kemp & Woodward *Tetrahedron* **21** 3019 1965, DOI: 10.1016/S0040-4020(01)96921-2].

N-Ethylcarbazole [86-28-2] $\text{C}_{14}\text{H}_{13}\text{N}$, **M 195.3**, **m 68-70°**, **69-70°**, **69-71°**, **b 199-200°/15mm**. Recrystallise it from MeOH, EtOH, EtOH/water or isopropanol and dry it below 55°. [*Beilstein* **20** H 436, **20** II 282, **20** III/IV 3829.]

Ethyl 1,3-dithiane-2-carboxylate [20462-00-4] $\text{C}_7\text{H}_{12}\text{O}_2\text{S}_2$, **M 192.3**, **b 75-77°/0.2mm**, **96°/0.4mm**, d_4^{20} **1.220**, n_{D}^{25} **1.5379**. Dissolve the ester in CHCl_3 , wash with aqueous K_2CO_3 , twice with H_2O , dry over MgSO_4 , filter, evaporate and distil the residue. [Eliel & Hartmann *J Org Chem* **37** 505 1972, DOI: 10.1021/jo00968a043; Seebach *Synthesis* **1** 17 1969, DOI: 10.1055/s-1969-34190; *Beilstein* **19/7** V 227.]

Ethyl 1,3-dithiolane-2-carboxylate [20461-99-8] $\text{C}_6\text{H}_{10}\text{O}_2\text{S}_2$, **M 178.3**, **b 85°/0.1mm**, d_4^{20} **1.250**, n_{D}^{20} **1.538**. Dissolve the ester in CHCl_3 , wash it with aqueous K_2CO_3 , twice with H_2O , dry it over MgSO_4 , filter, evaporate and distil the residue *in vacuo*. [Hermann et al. *Tetrahedron Lett* **14** 2599 1973, DOI: 10.1016/S0040-4039(01)96155-6; Corey & Erickson *J Org Chem* **36** 3553 1971, DOI: 10.1021/jo00822a019]. [*Beilstein* **19/7** V 225.]

Ethylene oxide (oxirane) [75-21-8] $\text{C}_2\text{H}_4\text{O}$, **M 44.0**, **m -111°**, **b 10.7°/atm**, **13.5°/746mm**, d_4^{10} **0.882**, n_{D}^{17} **1.3597**. Dry oxirane with CaSO_4 , then distil it from crushed NaOH. It has also been purified by passage, as a gas, through towers containing solid NaOH. [*Beilstein* **17/1** V 3.]

Ethylene thiourea (2-imidazolidinethione) [96-45-7] $\text{C}_3\text{H}_6\text{N}_2\text{S}$, **M 102.2**, **m 196-200°**, **203-204°**. Crystallise it from EtOH, amyl alcohol or H_2O in which the solubility is ~1.5w/v% at ~20° and ~50w/v% at ~100°.

[Beilstein 24 III/IV 22.]

Ethylene urea (2-imidazolidone) [120-93-4] $C_3H_6N_2O$, M 86.1, m 129-132°, 131°. Crystallise it from MeOH (charcoal). [Beilstein 24 III/IV 6.]

(±)-2-Ethylethylenimine (2-ethylaziridine) [2549-67-9] C_4H_9N , M 71.1, b 88.5-89°, pK_{a1}^{25} 8.31 (K_b 5.70 x 10⁻⁷). Freshly distil the aziridine from sodium before use. The *picrate* has m 103-104°. TOXIC. [O'Rourke et al. *J Am Chem Soc* 78 2159 1956, DOI: 10.1021/ja01591a035; Beilstein 20 III/IV 280.]

Ethyl hydrocupreine hydrochloride (Optochin) [3413-58-9] M 376.9, m 249-251°, pK_{a1}^{25} 5.50, pK_{a2}^{25} 9.95. Recrystallise it from H₂O [UV: Heidt & Forbes *J Am Chem Soc* 55 2701 1933, DOI: 10.1021/ja01334a012]. [Beilstein 24 H 385, 24 III/IV 1446.]

2-Ethyl-isothionicotinamide (ethionamide, 3-ethyl 4-pyridinecarbothioamide) [536-33-4] $C_8H_{10}N_2S$, M 166.2, m 163-164°, 164-166°(dec). It crystallises from EtOH as lemon yellow needles. The *hydrochloride* crystallises from EtOH (+ few drops of HCl) as orange yellow needles with m 212-214°. [Kutscherowa et al. *J Gen Chem USSR* (English transl) 29 915 1959, Beilstein 22 III/IV 737.] It causes peripheral and ocular neuropathy, and is carcinogenic and teratogenic, but also has antibacterial properties.

Ethyloxirane (1,2-epoxybutane) [RS 106-88-7; 2R(+) 3760-95-0; 2S(-) 30608-62-9] C_4H_8O , M 72.1, b 63-65°/atm and n_D^{20} 1.3829 for (±)-oxirane; b 61-63°/atm, n_D^{20} 1.3865 and $[\alpha]_D^{20}$ +13.6 (c 1.14, Et₂O) for R(+)-enantiomer, and b 62-63°/atm, n_D^{20} 1.3822 and $[\alpha]_D^{20}$ -13.7 (c 1.48, Et₂O) for S(-)-enantiomer 66.4-66.6°, also with d_4^{20} 0.837, $[\alpha]_D^{20}$ +10 and $[\alpha]_D^{20}$ -10 respectively neat. All epoxides were prepared by adding the respective 1-bromo-2-acetoxybutanes dropwise to a solution of KOH in H₂O (60%w/v) at 100° during 15 minutes, then steam distilling the oxirane out, drying the distillate over solid KOH and distilling the oxirane (30-50% yield) at atmospheric pressure. [Mori et al. *Tetrahedron* 35 1601 1979, DOI: 10.1016/0040-4020(79)80022-8.] Alternatively, dry with CaSO₄, and fractionally distil through a long (126cm) glass helices-packed column. The first fraction contains a water azeotrope where the water can be removed with solid KOH. The oxiranes have similar IR (film), ν_{max} 3080m, 2980s, 2940s, 2830m, 1470m, 1280m, 900s, 830s and 800m cm⁻¹; ¹H NMR (60MHz, TMS): δ 0.98 (3H, t, J 7 Hz), ~1.25 to ~1.80 (3H, m, 1.22, 1.42, 1.51, 1.58, 1.66), 2.20 to ~2.40 (3H, m); and MS: m/e 72 (M⁺ = C₄H₈O). [Beilstein 17 II 17, 17 III/IV 45, 17 V/1 56.]

(±)-3-Ethyl-5-phenylhydantoin (Ethotoin) [86-35-1] $C_{11}H_{12}N_2O_2$, M 204.2, m 94°. Crystallise it from water. It is an anticonvulsant and is used in epilepsy. [Dudley & Bius *J Heterocycl Chem* 10 173 1973, DOI: 10.1002/jhet.5570100208; Pinner *Chem Ber* 21 2320 1888, DOI: 10.1002/cber.18880210226; Beilstein 25 III/IV 963, 27 II 860.]

N-Ethyl-5-phenylisoxazolinium-3'-sulfonate (Woodward's reagent K) [4156-16-5] $C_{11}H_{11}NO_4S$, M 253.3, m 220°(dec). Crystallise the reagent from diethyl ether or ethyl acetate/petroleum ether. [Llamas et al. *J Am Chem Soc* 108 5543 1986, DOI: 10.1021/ja00278a030.] It is best purified by dissolving in excess of aqueous N HCl and precipitating with Me₂CO to give a white fluffy solid. [Woodward et al. *Tetrahedron* 22 Suppl 8 321 1966, DOI: 10.1016/S0040-4020(01)82192-X; Fieser 1 385, 2 198.]

(±)-3-Ethyl-3-phenyl-2,6-piperidinedione (Glutethimide) [77-21-4] $C_{13}H_{15}NO_2$, M 217.3, m 84°. Crystallise glutethimide from diethyl ether or ethyl acetate/petroleum ether. It has m 91-92° (from aqueous EtOH), 87-87.5° (from Et₂O/petroleum ether), 84-87° (from isopropanol), and 83-84° (from Et₂O). [Penprase & Biles *J Am Pharm Assoc* 47 523 1958, Hoffmann et al. *Helv Chim Acta* 40 387, 393 1957, DOI: 10.1002/hlca.19570400217; Beilstein 21 III/IV 5493.] The R(+)-enantiomer crystallises from EtOAc/petroleum ether with m 103-104°, and $[\alpha]_D^{20}$ +184 (c 1, EtOH). The S(-)-enantiomer has similar properties except it has $[\alpha]_D^{20}$ -181 (c 1, EtOH). [Branchini *Ricerche Scientifiche* 29 2435 1959, Finch et al. *Experientia* 31 1002 1975.]

2-Ethylpyridine [100-71-0] C_7H_9N , M 107.2, b 148.6°/atm, d_4^{25} 0.942, pK_{a1}^{25} 5.89. Dry 2-ethylpyridine with BaO, and fractionally distil it. Purify it further by conversion to the *picrate*, recrystallisation of the picrate and

regeneration of the free base followed by distillation. [Beilstein 20/6 V 3.]

4-Ethylpyridine [536-75-4] has **b 168.2-168.3°/atm**, **d₄²⁵ 0.942**, **pK₂₅ 6.02**. Dry 4-ethylpyridine with BaO, and fractionally distil it. Also purified by converting to the *picrate*, recrystallising and the free base is regenerated and distilled. [Beilstein 20/6 V 10.] **4-Ethylpyridine-1-oxide** [14906-55-9] has **m 109-110°**, **pK_{Est}~1.1**. Crystallise the oxide from acetone/ether. [Beilstein 20/6 V 10.]

Flavone (2-phenyl-4H-1-benzopyran-4-one) [525-82-6] **C₁₅H₁₀O₂**, **M 222.3**, **m 94-97°, 99-100°, 100°**. Crystallise flavone from petroleum ether. [Wheeler *Org Synth Coll Vol* 4 478 1963, DOI: 10.15227/orgsyn.032.007; Beilstein 17/10 V 552.]

Fluorescein [9-(o-carboxyphenyl)-6-hydroxy-3H-xanthene-3-one] [2321-07-5] **C₂₀H₁₂O₅**, **M 332.3**, **m 320°, ε_{495nm} 7.84 x 10⁴ (in 10⁻³M NaOH)**, **pK₁ 2.2, pK₂ 4.4, pK₃ 6.7**. Dissolve it in dilute aqueous NaOH, filter and precipitate it by adding dilute (1:1) HCl. The process is repeated twice more, and the fluorescein is dried at 100°. *Alternatively*, it has been crystallised from acetone by allowing the solution to evaporate at 37° in an open beaker. It has also been recrystallised from EtOH and dried in a vacuum oven. [Beilstein 19 I 721, 19 II 248, 19 III/IV 2904, 19/8 V 456.]

Fluoresceinamine (mixture of 5- and 6-aminofluorescein) [27599-63-9] **C₂₀H₁₃NO₅**, **M 347.3**, **m 215-220°(dec) and 223°(dec), (for 5-amino) and m 314-316°(dec, for 6-amino)**. Dissolve it in EtOH, treat with charcoal, filter, evaporate and dry the residue in a vacuum at 100° overnight. Also recrystallise it from 6% HCl, then dissolve it in 0.5% aqueous NaOH and precipitate it by acidifying with acetic acid. The separate amines are made from the respective nitro compounds, which are best separated *via* their *acetate salts*. They have similar R_F of 0.26 on Silica Gel Merck F₂₅₄ in 5 ml MeOH + 150 ml Et₂O saturated with H₂O. The IR (Me₂SO) has a band at ν_{max} 1690 cm⁻¹ (CO₂⁻) and sometimes a weak band at ν_{max} 1750 cm⁻¹ due to the lactone. The UV (EtOH) of the 6-isomer has λ_{max} at 222nm (ε 60 000) and the 5-isomer at λ_{max} 222nm (ε 60 000) and 285nm (ε 20.600). [IR: McKinney & Churchill *J Chem Soc (C)* 654 1970, DOI: 10.1039/J39700000654; McKinney et al. *J Org Chem* 27 3986 1962, DOI: 10.1021/jo01058a055; UV: Verbiscar *J Org Chem* 29 490 1964, DOI: 10.1021/jo01025a508; Beilstein 19 III/IV 4337, 19/8 V 713.]

Fluorescein isothiocyanate isomer I (FTIC, 5-isothiocyanato isomer) [3326-32-7; 27072-45-3 *mixture of 5- and 6-isomers*] **C₂₁H₁₁NO₅S**, **M 389.4**, **m >160°(slow dec)**. It is made from the pure 5-amino isomer. Purify it by dissolving it in boiling Me₂CO, filtering and adding petroleum ether (b 60-70°) until it becomes turbid. If an oil separates, then decant it and add more petroleum ether to the supernatant and cool. Orange-yellow crystals separate, collect and dry them *in vacuo*. It should give one spot on TLC (silica gel) in EtOAc/pyridine/AcOH (50:1:1) and in Me₂NCHO/CHCl₃/28% NH₄OH (10:5:4). Its IR (Me₂SO) has ν_{max} at 2110 (NCS) and 1760 (C=O) cm⁻¹. The ¹HNMR spectra in Me₂CO-d₆ of the 5- and 6-isomers are distinctly different for the protons in the *benzene ring; the UV in phosphate buffer pH 8.0 shows a λ_{max} at ~490nm. [Sinsheimer et al. *Anal Biochem* 57 227 1974, DOI:10.1016/0003-2697(74)90068-2; McKinney et al. *Anal Biochem* 7 74 1964, DOI:10.1016/0003-2697(64)90121-6; Beilstein 19 III/IV 4337.] Used for the FTIC labeling of proteins and microsequencing of peptides and proteins (by HPLC).

3-Fluoro-4-iodopyridine [22282-75-3] **C₅H₃FIN**, **M 223.0**, **m 80-81°, 85-89°, pK_{Est} ~1.7**. Crystallise it from petroleum ether and/or sublime it *in vacuo* (**m 87°**). The *picrate* [22282-76-4] has **m 140°** (from EtOH). [Gribble & Saulnier *Tetrahedron Lett* 21 4137 1980, DOI: 10.1016/S0040-4020(01)82192-X.]

4-Fluoro-7-nitrobenzofurazan (4-fluoro-7-nitrobenzo-2-oxa-1,3-diazole) [29270-56-2] **C₆H₂FN₃O₃**, **M 183.1**, **m 52-54°, 52.5-53.5°, 53-56°, 53.5-54.5°**. Purify it by repeated recrystallisation from petroleum ether (b 40-60°). On treatment with MeONa in MeOH it gives **4-methoxy-7-nitrobenzo-2-oxa-1,3-diazole m 115-116°**. [Nunno et al. *J Chem Soc (C)* 1433 1970, DOI: 10.1039/J39700001433.] It is a very good fluorophore for amino acids [Imai & Watanabe *Analyt Chim Acta* 130 377 1981, DOI: 10.1016/S0003-2670(01)93016-8], as it reacts with primary and secondary amines to form fluorescent adducts with λ_{ex} 470nm and λ_{em} 530nm. It gives a *glycine* derivative with **m 185-187°** [Miyano et al. *Anal Chim Acta* 170 81 1985, DOI:10.1016/S0003-2670(98)00125-1]. Useful for the fluorescent labeling of amines and amino acids for HPLC analysis.

5-Fluorouracil (5-fluoropyrimidinedi-2,4-[1*H*,3*H*]-one, 5-FU) [51-21-8] $C_4H_3FN_2O_2$, M 130.1, m 282-283° (dec), 282-286° (dec), pK_1^{25} 8.04, pK_2^{25} 13.0. Recrystallise it from H_2O or $MeOH/Et_2O$ and sublime it at 190-200°/0.1mm or 210-230°/0.5mm. UV: λ_{max} at 265-266nm (ϵ 7070). [Hesse et al. *J Org Chem* **37** 329 1972, DOI: 10.1021/jo00967a037; Duschinsky & Plevin *J Am Chem Soc* **79** 4559 1957, DOI: 10.1021/ja01573a087; *Beilstein* **24** III/IV 1229.] A potent antitumour agent which inhibits thymidilate synthase and the consequent depletion of d-TTP resulting in hindering cell division.

Fluram (Fluorescamine, 4-phenyl-spiro[furan-2(3*H*)-1-phthalan]-3,3'-dione) [38183-12-9] $C_{17}H_{10}O_4$, M 278.3, m 153-155°, 153-157°, 154-155°. Fluram is a non-fluorescent reagent that reacts with primary amines to form highly fluorescent compounds. Purify it by dissolving (~1g) in Et_2O / C_6H_6 (1:1, 180 ml), washing with 1% aqueous $NaHCO_3$ (50ml), drying (Na_2SO_4), and evaporating in a vacuum. Dissolve the residue in warm CH_2Cl_2 (5ml), dilute with Et_2O (12ml) and refrigerate. Collect the solid and dry it in a vacuum. Its IR ($CHCl_3$) has ν_{max} at 1810, 1745, 1722, 1625 and 1600 cm^{-1} , and 1H NMR ($CDCl_3$) with δ at 8.71 (s, -OHC=). [Weigle et al. *J Am Chem Soc* **94** 5927 1972, DOI: 10.1021/ja00771a084; Weigle et al. *J Org Chem* **41** 388 1976, DOI: 10.1021/jo00864a051; Lai *Methods Enzymol* **47** 236 1977, DOI: 10.1016/0076-6879(77)47028-9.]

Forskolin (Colforsin, Coleonol, 5-[acetyloxy]-3-ethenyldodecahydro-6,10,10b-trihydroxy-3,4a,7,7, 10a-penta-methyl-[3*R*-(3 α -4 $\alpha\beta$, 5 β , 6 β , 6 α , 10 α , 10 $\alpha\beta$, 10b α)-1*H*-naphtho[2,1-*b*]pyran-1-one) [66575-29-9] $C_{22}H_{34}O_7$, M 410.5, m 230-232°, 228-233°, $[\alpha]_D^{25}$ -26.2 (c 1.7, $CHCl_3$). Recrystallise this diterpene from C_6H_6 /petroleum ether, $EtOAc$ /petroleum ether. It is an antihypertensive, a positive ionotropic agent, a platelet aggregation inhibitor, and it has adenylate cyclase activating properties [*Chem Abstr* **89** 244150 1978, de Souza et al. *Med Res Rev* **3** 201 1983, DOI: 10.1002/med.2610030205; X-ray: Tandon et al. *Indian J Chem* **15B** 880 1977]. [*Beilstein* **18/5** V 55.]

Fumagillin {2,4,6,8-decatetraene-1,10-dioic acid mono[4-(1,2-epoxy-1,5-dimethyl-4-hexenyl)-5-methoxy-1-oxaspiro[2.5]oct-6-yl] ester} [23110-15-8] $C_{26}H_{34}O_7$, M 458.5, m 194-195°, $[\alpha]_D^{20}$ -26.2 (in 95% $EtOH$), $pK_{Est} \sim 4.5$. Forty grams of a commercial sample containing 42% fumagillin, 45% sucrose, 10% antifoam agent and 3% of other impurities are digested with 150ml of $CHCl_3$. The insoluble sucrose is filtered off and washed with $CHCl_3$. The combined $CHCl_3$ extracts are evaporated almost to dryness at room temperature under reduced pressure. The residue is triturated with 20ml of $MeOH$, and the fumagillin is filtered off by suction. It is crystallised twice from 500ml of hot $MeOH$ by standing overnight in a refrigerator (yellow needles). (The long-chain fatty ester used as anti-foam agent is still present, but is then removed by repeated digestion, on a steam bath, with 100ml of diethyl ether.) For further purification, the fumagillin (10g) is dissolved in 150ml of 0.2M ammonia, and the insoluble residue is filtered off. The ammonia solution (cooled in running cold water) is then brought to pH 4 by careful addition of M HCl with constant shaking in the presence of 150ml of $CHCl_3$. (Fumagillin is acid-labile and must be removed rapidly from the aqueous acid solution.) The $CHCl_3$ extract is washed several times with distilled water, dried (Na_2SO_4) and evaporated under reduced pressure. The solid residue is washed with 20ml of $MeOH$. The fumagillin is filtered off by suction, then crystallised from 200ml of hot $MeOH$. [Tarbell et al. *J Am Chem Soc* **77** 5610 1955, DOI: 10.1021/ja01626a045.] **Alternatively**, 10g of fumagillin in 100ml $CHCl_3$ is passed through a silica gel (5g) column to remove tarry material, and the $CHCl_3$ is evaporated to leave an oil which gives fumagillin on crystallisation from amyl acetate. It recrystallises from $MeOH$ (charcoal) or $Me_2CO/MeOH$. The fumagillin is stored in dark bottles in the absence of oxygen and at low temperatures. [Schenck et al. *J Am Chem Soc* **77** 5606 1955, DOI: 10.1021/ja01626a044; *Beilstein* **19** III/IV 1012.] It is used as an antiprotozoal agent.

Furan [110-00-9] C_4H_4O , M 68.1, b 31.3°/atm, 32°/758mm, d_4^{20} 1.42, n_D^{20} 1.4214. Shake it with aqueous 5% KOH , dry it with $CaSO_4$ or Na_2SO_4 , then distil it under nitrogen, from KOH or sodium, immediately before use. A trace of hydroquinone could be added as an inhibitor of oxidation. [*Beilstein* **17** H 27, **17** I 16, **17** II 34, **17/1** V 291.]

Furan-2-carboxylic (2-furoic) acid [88-14-2] $C_5H_4O_3$, M 112.1, m 128-132°, 133-134°, b 141-144°/20mm, 230-232°/760mm, pK_1^{25} -7.3 (O-protonation), pK_2^{25} 3.32. Crystallise the acid from hot water (charcoal), dry it at 120° for 2 hours, then recrystallise it from $CHCl_3$, and again dry it at 120° for 2 hours. For use as a standard in volumetric analysis, good quality commercial acid should be crystallised from $CHCl_3$ and dried as above or sublimed at 130-140°/50-60mm or less. [*Beilstein* **18** I 438, **18** II 265, **18** III/IV 3914, **18/6** V 102.]

Furan-3-carboxylic (3-furoic) acid [488-93-7] $C_5H_4O_3$, M 112.1, m 120-122°, 122-123°, 123°/5, pK²⁵ 4.03. Crystallise the acid from water or aqueous EtOH, and sublime it in a vacuum. [Beilstein 18 I 439, 18 III/IV 4052, 18/6 V 196.]

Furan-3,4-dicarboxylic acid [3387-26-6] $C_6H_4O_5$, M 156.1, m 217-218°, 221.5-222.5°, pK₁²⁵ 1.44, pK₂²⁵ 7.84. Crystallise it from water or Et₂O/petroleum ether, and sublime it in a vacuum. [Beilstein 18 III/IV 4497.]

Furan-2,5-dione (maleic anhydride) [108-31-6] $C_4H_2O_3$, M 98.1, m 51-56°, 54°, b 94-96°/20mm, 199°/760mm. Crystallise it from *benzene, CHCl₃, CH₂Cl₂ or CCl₄. Sublime it under reduced pressure. [Skell et al. *J Am Chem Soc* 108 6300 1986, DOI: 10.1021/ja00280a030; Beilstein 17 III/IV 5897, 17/11 V 55.]

3-(2-Furanyl)acrylic acid [539-47-9] $C_7H_6O_3$, M 138.1, (*cis*-isomer) m 106-108°, pK²⁵ 3.5; (*trans*-isomer) m 141°, 143-144°, b 285°/atm, pK²⁵ 4.5 (H₂O), 5.76 (50% aqueous EtOH), 6.65 (ethoxyethanol/H₂O—80:20). Recrystallise the *cis*-isomer from *C₆H₆ and the *trans*-isomer from H₂O, *C₆H₆ or petroleum ether (b 80-100°)(charcoal). [Beilstein 18 H 301, 18 III/IV 4143, 18/6 V 306.]

Furfural (2-furfuraldehyde) [98-01-1] $C_5H_4O_2$, M 96.1, m -36°, b 54-56°/11mm, 59-60°/15mm, 67.8°/20mm, 90°/65mm, 161°/760mm, d₄²⁰ 1.159, n_D²⁰ 1.52608, pK²⁵ -6.5 (O-protonation). Furfural is unstable to air, light and acids. Impurities include formic acid, β-formylacrylic acid and furan-2-carboxylic acid. Distil it in an oil bath from 7% (w/w) Na₂CO₃ (added to neutralise acids, especially pyromucic acid). Redistil it from 2% (w/w) Na₂CO₃, and then, finally fractionally distil it under vacuum. It is stored in the dark. [Evans & Aylesworth *Ind Eng Chem (Anal ed)* 18 24 1926, DOI: 10.1021/ie50193a013.]

Impurities resulting from storage can be removed by passage through chromatographic grade alumina. Furfural can be separated from impurities other than carbonyl compounds by the bisulfite addition compound. The aldehyde is steam volatile. It has been purified by distillation (using a Claisen head) under reduced pressure. This is essential as is the use of an oil bath with temperatures of no higher than 130° which is highly recommended. When furfural is distilled at atmospheric pressure (in a stream of N₂), or under reduced pressure with a free flame (caution: because the aldehyde is flammable), an almost colourless oil is obtained. After a few days and sometimes a few hours, the oil gradually darkens and finally becomes black. This change is accelerated by light and occurs more slowly when it is kept in a brown bottle. However, when the aldehyde is distilled under vacuum and the bath temperature kept below 130° during the distillation, the oil develops only a slight colour when exposed to direct sunlight during several days. The distillation of very impure material should **NOT** be attempted at atmospheric pressure; otherwise the product darkens very rapidly. After one distillation under vacuum, a distillation at atmospheric pressure can be carried out without too much decomposition and darkening. The liquid **irritates mucous membranes**. Store it in dark containers under N₂, preferably in sealed ampoules. [Adams & Voorhees *Org Synth Coll Vol* 1 280 1941, DOI: 10.15227/orgsyn.001.0049; Beilstein 17/9 V 292.]

Furfuryl alcohol (2-furylmethanol) [98-00-0] $C_5H_6O_2$, M 98.1, m -29°, b 68-69°/20mm, 170.0°/750mm, d₄²⁰ 1.132, n_D²⁰ 1.4873, n_D³⁰ 1.4801, pK²⁵ 2.61. Distil it under reduced pressure to remove tarry material, shake with aqueous NaHCO₃, dry it with Na₂SO₄ and fractionally distil it under reduced pressure from Na₂CO₃. It can be further dried by shaking with Linde 5A molecular sieves. [Beilstein 17/3 V 338.]

Furfurylamine (2-aminomethylfuran) [617-89-0] $C_5H_6O_2$, M 97.1, m -70°, b 54-56°/17mm, 142.5-143°/735mm, 145-146°/atm, d₄²⁰ 1.059, n_D²⁰ 1.489, pK³⁰ 8.89. Distil it under nitrogen from KOH through a column packed with glass helices, preferably under vacuum. Store it away from CO₂, or better as a salt. The *picrate* has m 184-184°(dec), the *hydrochloride* has m 147-149°, and the *oxalate salt* has m 145-147°. [Beilstein 18 H 584, 18 II 416, 18 III/IV 3068, 18/9 V 541.]

6-Furfurylaminopurine (Kinetin) [525-79-1] $C_{10}H_9N_5O$, M 215.2, m 266-267°, 269-271°, 270-272°, 272° (sealed capillary), pK₁ <1, pK₂ 3.8, pK₃ 10. It forms platelets from EtOH and sublimes at 220°, but is best done at lower temperatures in a good vacuum. It has been extracted from neutral aqueous solutions with Et₂O. [Miller et al. *J Am Chem Soc* 78 1375 1956, DOI: 10.1021/ja01588a032; Bullock et al. *J Am Chem Soc* 78 3693 1956, DOI: 10.1021/ja01596a037; Beilstein 26 III/IV 3586.] It is a plant cytokine that promotes cell division.

Furil [492-94-4] $C_{10}H_6O_4$, M 190.2, m 163-165°, 165-166°. Furil crystallises from MeOH or *benzene (charcoal). [Beilstein 19 III/IV 2008.]

(±)-Furoin [1,2-di-(2-furyl)-2-hydroxyethanone] [552-86-3] $C_{10}H_8O_4$, M 192.2, m 134-137°, 135-136°, 138-139°, 158-162°/9mm. It crystallises from MeOH (charcoal) and distils in a vacuum. [Hartman & Dickey *J Am Chem Soc* 55 1228 1933, DOI: 10.1021/ja01330a063.] The (-)-*enantiomer* crystallises from toluene or EtOH with m 131-131°, and has $[\alpha]_D^{20}$ -4.9 (dioxane) [Neuberg et al. *Arch Biochem* 1 393 1943, Beilstein 19 H 204, 19 I 710, 19 II 224, 19 III/IV 2543.]

Fusaric acid (5-*n*-butylpyridine-2-carboxylic acid) [536-69-6] $C_{10}H_{13}NO_2$, M 179.2, m 96-98°, 98°, 98-100°, 101-103°, pK_1 5.7, pK_2 6.16 (80% aqueous methoxyethanol). Dissolve it in $CHCl_3$, dry (Na_2SO_4), filter, evaporate and recrystallise the residue from 50 parts of petroleum ether (b 40-60°), $CHCl_3$ /petroleum ether or EtOAc, then sublime it *in vacuo*. The *amide* crystallises from MeOH with m 128.2-129.0°. The *copper salt* forms bluish violet crystals from H_2O and has m 258-259°. [Hardegger & Nikles *Helv Chim Acta* 39 505 1956, DOI: 10.1002/hlca.19560390222; Schreiber & Adam *Chem Ber* 93 1848 1960, DOI: 10.1002/cber.19600930823; NMR and MS: Tschesche & Führer *Chem Ber* 111 3502 1978, DOI: 10.1002/cber.19781111024; Beilstein 22 III/IV 764, 22/2 V 384.]

Glycidol (oxirane-2-methanol) [*RS*-(±)- 556-52-5; *R*-(+)- 57044-25-4; *S*-(-)- 60456-23-7] $C_3H_6O_2$, M 74.1, (*R,S*) b 61-62°/15mm, d_4^{20} 1.117, n_D^{20} 1.433, [*S*-(*-*)-isomer, § also available on polymer support, has b 49-50°/7mm, 66-67°/19mm, $[\alpha]_D^{20}$ -15 (neat)], [*R*-(+)-isomer has b 56-56.5°/11mm, d_4^{20} 1.117, n_D^{20} 1.429, $[\alpha]_D^{20}$ +15 (neat)]. Purify glycidol by fractional distillation. The 4-nitrobenzoates have m 56° (±); m 60-62°, $[\alpha]_D^{20}$ -37.9 (c 3.38 $CHCl_3$) for *R*-(*-*)-isomer [106268-95-5]; m 60-62°, $[\alpha]_D^{20}$ +38 (c 1 $CHCl_3$) for the *S*-(+)-isomer [115459-65-9] and are recrystallised from Et_2O or Et_2O /petroleum ether (b 40-60°) [*S*-isomer: Burgos et al. *J Org Chem* 52 4973 1987, DOI: 10.1021/jo00231a025; Sowden & Fischer *J Am Chem Soc* 64 1291 1942, DOI: 10.1021/ja01258a017.] [Beilstein 17 I 50, 17 III/IV 985, 17/3 V 9.]

Gramine (3-dimethylaminoethylindole) [87-52-5] $C_{11}H_{14}N_2$, M 174.3, m 132-134°, 134°, pK^{25} 16.00 (NH acidic), basic pK^{25} 9.2 (50% aqueous EtOH). Crystallise gramine from diethyl ether, ethanol or acetone. It sublimes at 59°/0.001mm. The *hydrochloride* crystallises from EtOH/ Et_2O with m 190.5-191.0°(dec). [Culvenor et al. *Aust J Chem* 17 1301 1964, DOI: 10.1071/CH9641301; Beilstein 22 III/IV 4302, 22/10 V 25.]

(2*S*,6'*R*)(+)-Griseofulvin [126-07-8] $C_{17}H_{17}ClO_6$, M 352.8, m 220°, $[\alpha]_D^{22}$ +365 (c 1, acetone). Crystallise it from *benzene or EtOH. Purify 2g of griseofulvin by chromatography on Alumina (40 x 1.5cm) and elute with * C_6H_6 /MeOH (199:1) and follow the UV blue fluorescent band. [MacMillan *J Chem Soc* 1823 1959, DOI: 10.1039/JR9590001823; Beilstein 18 III/IV 3160, 18/5 V 150.]

Guanosine (H_2O) [118-00-3] $C_{10}H_{13}N_5O_5 \cdot xH_2O$, M 283.2, m 237-237.5°(dec), 239°(dec), 250°(dec), $[\alpha]_D^{20}$ -86 (c 1, 0.1M NaOH), pK_1^{25} 1.9, pK_2^{25} 9.24, pK_3^{25} 12.33. It crystallises from water as a *dihydrate*. Dry it at 110°. [Beilstein 26/18 V 81.]

Guanylic acid (guanosine-5'-monophosphoric acid) [85-32-5] $C_{10}H_{14}N_5O_8P$, M 363.2, m 190-200°(dec), 208°(dec), pK_2^{25} 2.4, pK_3^{25} 6.66 (6.1), pK_4^{25} 9.4. Crystallise it from water and dry it at 110°. [Beilstein 26 III/IV 3910.]

Harmaline (7-methoxy-1-methyl-4,9-dihydro-3*H*- β -carboline, 4,9-dihydro-7-methoxy-1-methyl-3*H*-pyrido[3,4-*b*]indole) [304-21-2] $C_{13}H_{14}N_2O$, M 214.3, m 229-230°, 229-231°, 232-234°, 235-237° (after distillation at 120-140°/10⁻³), pK_1 4.2. Recrystallise harmaline from MeOH and sublime it at high vacuum. It has UV in MeOH with λ_{max} at 218, 260 and 376nm (log ϵ 4.27, 3.90 and 4.02, respectively); IR (Nujol) with ν_{max} at 1620, 1600, 1570 and 1535cm⁻¹ and in $CHCl_3$ ν_{max} at 1470 and 1629cm⁻¹. [Spenser *Can J Chem* 37 1851 1959, DOI: 10.1139/v59-272; Marion et al. *J Am Chem Soc* 73 305 1951, DOI: 10.1021/ja01145a100; UV Pruckner & Witkop *Justus Liebigs Ann Chem* 554 127 1943, DOI: 10.1002/jlac.19435540109.] The *hydro-*

chloride dihydrate has **m 234-236°(dec)**, the *picrate* has **m 228-229°** (sinters at 215°) from aqueous EtOH, and the *N-acetate* forms needles **m 204-205°**. [Beilstein 23 H 396, 23 I 119, 23 II 345, 23 III/IV 2666, 23/12 V 148.] Harmaline is a CNS stimulant.

Harmane (2-methyl- β -carboline, 1-methyl-9H-pyrido[3,4-*b*]indole, Aribine) [486-84-0] $C_{12}H_{10}N_2$, **M 182.2, m 235-238°, 237-238°, pK₁ 7.37 (basic, Pry N), pK₂ 14.7 (acidic, NH)**. Crystallise it from heptane/cyclohexane. It is insoluble in H₂O, but soluble in dilute HCl or H₂SO₄. Solutions show a blue fluorescence. Its UV (MeOH) has λ_{max} nm (log ϵ) at 234 (4.57), 287 (4.21) and 347 (3.66). The *hydrochloride* forms needles from EtOH/dil HCl which sublimes at **m 120-130°**. It is an imidazoline binding site agonist. [Wolfbeis et al. *Monatsh Chem* 113 509 1982, DOI: 10.1007/BF00799926; Beilstein 23/12 237.] Harmane is an I₁ imidazoline binding site agonist.

Harmine (7-methoxy-1-methyl-9H-pyrido[3,4-*b*]indole) [442-51-3] $C_{13}H_{12}N_2O$, **M 212.3, m 261°(dec), 262-264°, 265°, pK²⁰ 7.61**. Crystallise harmine from MeOH and sublime it in a vacuum. Its UV (MeOH) has λ_{max} nm (log ϵ) at 241 (4.61), 301 (4.21) and 336 (3.69). [Beilstein 23 II 348, 23 III/IV 2702, 23/12 V 237.] **Harmine hydrochloride (hydrate)** [343-27-1] has **M 248.7, m 262°(dec, hydrate), 280°(dec)**. The *hydrate* crystallises as fluorescent crystals from water. Freely soluble in hot water, but only 2.5% in cold water. The *anhydrous* salt has **m 319°**. [Beilstein 23/12 V 237.] It is a CNS stimulant.

Hesperetin (3',5,7-trihydroxy-4'-methoxyflavanone) [520-33-2] $C_{16}H_{14}O_6$, **M 302.3, (R,S) m 226-228°, 227-228°, pK_{Est} ~8.5-10.5 (phenolic)**. Crystallise it from EtOAc or ethanol. The natural S(-) form crystallises from EtOH and has **m 216-218°** and $[\alpha]_D^{20}$ -37.6 (c 2, EtOH). **Note** that C2 is chiral. It is a weak phenolic acid which precipitates at pH 5-7 from alkaline solutions (see pK_a values). [Beilstein 18 III/IV 3215, 18/5 V 214.]

Hesperidin (hesperetin 7-rhamnoside) [520-26-3] $C_{28}H_{34}O_{15}$, **M 610.6, m 258-262° (soft at ~250°), 261-263°(dec), $[\alpha]_{546}^{20}$ -82 (c 2, pyridine)**. Dissolve hesperidine [the natural glycoside of hesperetin (above) from lemons and sweet oranges] in dilute aqueous alkali and crystallises out as fine needles on adjusting the pH to 6-7. [Beilstein 18 III/IV 3219, 18/5 V 218.]

Hexahydro-1H-azepine (hexamethyleneimine, Azepane) [111-49-9] $C_6H_{13}N$, **M 99.2, b 70-72°/30mm, 135-138°/atm, 138°/749mm, d_4^{20} 0.879, n_D^{20} 1.466, pK²⁵ 11.10 (pK⁰ 9.71, pK⁷⁵ 9.71)**. Purify azepane by dissolving in Et₂O and adding ethanolic HCl until all the base separates as the white *hydrochloride*, filter, wash with Et₂O and dry it (**m 236°**). The salt is dissolved in the minimum volume of H₂O and basified to pH ~ 14 with 10N KOH. The solution is extracted with Et₂O, the extract is dried over KOH, evaporated and distilled. The *free base* is a **FLAMMABLE** and **TOXIC** liquid, and best kept as the salt. The *nitrate* has **m 120-123°**, the *picrate* has **m 145-147°**, and the *tosylate salt* has **m 76.5°** (ligroin). [Müller & Sauerwald *Monatsh Chem* 48 727 1927, DOI: 10.1007/BF01526517; Hjelt & Agback *Acta Chem Scand* 18 191 1964, DOI: 10.3891/acta.chem.scand.18-0191; Beilstein 20 II 1406, 20 III/IV 1406, 20/4 V 3.]

Hexamethylenetetramine (Urotropine, hexamine, HMTA) [100-97-0] $C_6H_{12}N_4$, **M 140.1, m 280° (subl), 290-292° (sealed tube, CARE), d_4^{20} 1.331, pK²⁵ 4.85 (6.30)**. It is soluble in H₂O (67%), CHCl₃ (10%), EtOH (8%) and Et₂O (0.3%), and a 0.2M solution has a pH of 8.4. Dissolve it in hot absolute EtOH (reflux, Norit), filter using a heated funnel, cool at room temperature first, then in ice. Wash the crystals with cold Et₂O, dry them in air or under a vacuum. A further crop can be obtained by adding Et₂O to the filtrate. It sublimes above 260° without melting. It forms salts readily with organic and inorganic acids. The *hydrobromide perbromide* [149261-40-5] $C_6H_{12}N_4 \cdot HBr \cdot Br_2$, **M 380.9**, has **m 220°(dec)**, and the *picrate* has **m 179°(dec)**. [pK²⁰ 4.85: Reilly & Schmid *Anal Chem* 30 947 1958, DOI: 10.1021/ac60137a021; pK²⁰ 6.30: Pummerer & Hofmann *Chem Ber* 56 1255 1923, DOI: 10.1002/cber.19230560603.] [Beilstein 26 I 306, 26 II 200, 26 III/IV 1680.] It is a urinary antibacterial and an antimicrobial food additive.

R-(+)-2-Hexyloxirane [R-(+)-1,2-epoxyoctane, R-(+)-octene oxide] [77495-66-0] $C_8H_{16}O$, **M 128.2, b 51-54°/10mm, 60-62°/15mm, 157°/atm, d_4^{25} 0.839, n_D^{20} 1.418, $[\alpha]_D^{20}$ +14 (neat), the S-(-)- enantiomer has [50418-68-3] $[\alpha]_D^{20}$ -14.5 (c 2.5, EtOH), n_D^{20} 1.412**. The enantiomeric oxiranes have been purified by passage through a silica gel column in pentane and eluted with pentane/Et₂O. The eluate is evaporated and subjected to bulb-to-bulb distillation in a vacuum. The enantiomeric purity is checked by HPLC. [White & Emmons *Tetra-*

hedron **17** 31 1962, DOI:10.1016/S0040-4020(01)99002-7; Johnson & Rogers *J Org Chem* **38** 1793 1973, DOI: 10.1021/jo00950a005; *Beilstein* **17** H 17, **17** I 11, **17** III/IV 111, **17**/I V 138.]

Histamine [4-(2-aminoethyl)imidazole, β -aminoethylglyoxaline, 1*H*-imidazole-4-ethanamine] [51-45-6] $\text{C}_5\text{H}_9\text{N}_3$, **M** 111.2, **m** 83-84°, 86° (sealed tube), **b** 167°/0.8mm, 209°/18mm, **pK**₁²⁵ 6.02, **pK**₂²⁵ 9.70. It was prepared by decarboxylation of histidine monohydrochloride with concentrated HCl at 265-270° in a sealed tube for 3 hours, cooled, evaporated, neutralised and treated with picric acid to isolate *histamine dipicrate* **m** 233-235° which crystallised from H₂O. Repeated recrystallisation from H₂O gave the *monopicrate* which also had **m** 233-234°. Similar treatment of *histidine monohydrochloride* with 20% aqueous H₂SO₄ at 265-270°/sealed tube/3hrs, cooled, basified with Na₂CO₃ until no further precipitation (Na₂SO₄), filtered, and the filtrate evaporated to a small volume and mixed with saturated picric acid to isolate histamine as its picrate. Alternatively, *N*-benzoylhistidine (prepared by the Schotten-Baumann method) was heated in a vacuum at 240° until frothing ceased, the dark tarry mass was dissolved in concentrated HCl and hydrolysed at 180°, cooled, treated with H₂O, benzoic acid was filtered off. The filtrate was extracted with Et₂O (to remove all the benzoic acid), neutralised, treated with aqueous picrolonic acid and *histamine picrolonate* was isolated after recrystallising from EtOH (bunched yellow needles) **m** 262-264° (**m** 233-234°, 266° also reported). The yield of histamine, as calculated from the isolated picrates and picrolonate, from the three procedures were ~20% to 25%. [Ewins & Pyman *J Chem Soc* 339 1911, DOI: 10.1039/CT9119900339.]

In an alternative procedure Windaus & Vogt [*Chem Ber* **40** 3691 1907, DOI: 10.1002/cber.190704003164] converted *ethyl 3-(imidazol-4-yl)propionate* into its *hydrazide* (**m** 142°) (3.08g) which on treatment with amyl nitrite (2.34g) in EtOH (15ml) followed by ethanolic HCl (0.73g), allowing to stand for 1 hour at ~25°, then heating until evolution of N₂ ceased, and heating in a boiling water-bath for 8 hours, was evaporated to dryness. The resulting urethane was hydrolysed with boiling concentrated HCl. Evaporation to dryness provided *histamine dihydrochloride* (see below) in 55% yield which crystallised in prisms from EtOH.

Pyman [*J Chem Soc* 530 1912, DOI: 10.1039/CT9120100530] isolated the *free base* by mixing an aqueous solution of a pure salt (*dihydrobromide* or *diphosphate*, and presumably *dihydrochloride* see below) with excess Na₂CO₃, evaporating completely dry in a vacuum, and extracting with cold CHCl₃. The extract was dried with NaOH, evaporated to a small volume and on keeping, the base crystallised in clear, colourless, wedge-shaped plates which melted at 83-84° (corr), after softening a few degrees earlier. An analytical sample was obtained by drying in a vacuum over H₂SO₄. The *free base* is soluble in H₂O or EtOH, readily soluble in hot CHCl₃, but sparingly so in cold CHCl₃, and practically insoluble in Et₂O. It distils at 209-210°/18mm, into a very viscous colourless oil which solidifies on seeding into a colourless crystalline mass. It also crystallises from *benzene. It is best stored as the dihydrochloride or preferably as the diphosphate (see below) because the *free base* is strongly basic (see pK_a values above) and likely to absorb CO₂ from the atmosphere to form carbonate-carbamate salts. It forms an anhydrous *monohydrobromide* **m** 182-183° (from absolute EtOH in prismatic rods) which is soluble in H₂O forming a strongly alkaline solution, but is sparingly soluble in cold EtOH. It is a potent endogenous vasodilator as it is a histamine receptor agonist and activates nitric oxide synthase. [*Beilstein* **25** I 628, **25** II 302, **25** III/IV 2049.]

Histamine dihydrochloride (Maxamine) [56-92-8] $\text{C}_5\text{H}_9\text{N}_3 \cdot 2\text{HCl}$, **M** 184.1, has **m** 249-252° (244-245°). The dihydrochloride crystallises from aqueous EtOH, Et₂O-Me₂CO (plates) or H₂O (prisms). It is freely soluble in excess H₂O and MeOH. [*Beilstein* **25** III/IV 2049.] See previous entry for biological activity.

Histamine diphosphate [51-74-1] $\text{C}_5\text{H}_9\text{N}_3 \cdot 2 \text{H}_3\text{PO}_4$, **M** 307.2, has **m** 132-133° (corr). The phosphate is particularly suitable for characterising and purifying histamine on account of its crystalline form, sparing solubility in H₂O, and in almost treble the weight of the free base. It is soluble in only four parts of H₂O, but freely soluble in hot H₂O from which it separates in magnificent *anhydrous*, clear, colourless, quadrihedral prisms, belonging to the monoclinic system. The crystals are almost completely capped by a pair of pyramid faces, with a diminutive second pair present and usually with a basal plane. Crystals resembling *augite*, with half inch lengths and quarter inch breadth can be deposited from only a few grams of phosphate. [Isolation: Pyman *J Chem Soc* 530 1912, DOI: 10.1039/CT9120100530; *Beilstein* **25** I 630.] See above for biological activity.

Homopiperazine (1,4-diazepane) [505-66-8] $\text{C}_5\text{H}_{12}\text{N}_2$, **M** 100.2, **m** 38-40°, 43°, **b** 60°/10mm, 92°/50mm, 169°/atm, **pK**₁²⁰ 6.70, **pK**₂²⁰ 10.41. Purify it by fractionation through a column of 10 theoretical plates with a reflux ratio of 3:1. It boils at 169°, and the cool distillate crystallises in plates **m** 43°. [Poppelsdorf and Myerly *J Org Chem* **26** 131 1961, DOI: 10.1021/jo01060a031] Its pK_a values are 6.67 and 10.09 at 29.7°, and 6.28 and

9.86 at 40° [Pagano et al. *J Phys Chem* **65** 1062 1961, DOI: 10.1021/j100824a513]. The **1,4-bis(4-bromobenzoyl) derivative** has **m 194-198°** (from EtOH); the **hydrochloride** has **m 270-290°** (from EtOH) and the **picrate** has **m 265°(dec)** [Lloyd et al. *J Chem Soc (C)* 780 1966, DOI: 10.1039/J39660000780]. [Beilstein **23** III/IV 388, **23/3** V 240.] It is an antihypertensive.

1-Hydrazinophthalazine (hydralazine, Alphapress) [86-54-4] $C_8H_8N_4$, **M 160.1**, **m 172-173°(dec)**, **pK₁²⁵ 2.90**, **pK₂²⁵ 7.25** (NNH₂). It crystallises from MeOH. Its UV has λ_{max} at 656nm at pH ~11. It complexes with Bi³⁺, Zn²⁺, Fe²⁺ and Co²⁺. The **hydrochloride (hydralazine hydrochloride)** [304-20-1] $C_8H_8N_4 \cdot HCl$, **M 196.6**, also crystallises from MeOH and has **m 172-173°(dec)**. [Druey & Ringier *Helv Chim Acta* **34** 195 1951, DOI: 10.1002/hlca.19510340122; Beilstein **25** III/IV 4552.] It is an antihypertensive.

2-Hydrazinopyridine [4930-98-7] $C_5H_7N_3$, **M 109.1**, **m 41-44°, 46-47°, 49-50°, b 90-92°/1mm, 105°/5mm, 128-135°/13mm**. Purify it by distillation under a vacuum and by recrystallisation from Et₂O/hexane. [Kauffmann et al. *Justus Liebigs Ann Chem* **656** 103 1962, DOI: 10.1002/jlac.19626560116; Potts & Burton *J Org Chem* **31** 251 1966, DOI: 10.1021/jo01339a057.] The **mono-hydrochloride** has **m 183°(dec)** from aqueous HCl, and the **di-hydrochloride** [62437-99-4] $C_5H_7N_3 \cdot 2HCl$, **M 182.1**, has **m 214-215°**. [Beilstein **22** II 487, **22** III/IV 7025, **22/14** V 486.]

4-Hydroxyacridine (4-acridinol, neo-oxine) [18123-20-1] $C_{13}H_9NO$, **M 195.2**, **m 116.5°, 122-123°, pK₁¹⁵ 5.28**, **pK₂¹⁵ 9.75**. Crystallise neo-oxine from EtOH or aqueous EtOH. It complexes with Zn²⁺, Cd²⁺, Ga²⁺, Pb²⁺, Cr²⁺, Mn²⁺, Fe²⁺, Co²⁺ and Ni²⁺. The **hydrochloride** crystallises from EtOH with **m 242°**. [Beilstein **21** II 78, **21** III/IV 1562, **21/4** V 90.]

1-Hydroxy-1,2-benziodoxol-3(1H)-one (IBX, 2-iodoxybenzoic acid) [61717-82-6] $C_7H_5IO_4$, **M 280.0**, **m 224-225°, 226-234°, 232-233°(dec), 233°(dec)**, **pK <4**. IBX prepared by the Dess-Martin procedure (KBrO₃/H₂SO₄) [Dess & Martin *J Am Chem Soc* **113** 7277 1991, DOI: 10.1021/ja00019a027] has been reported as being explosive and comparable to TNT [Hartmann & Meyer *Chem Ber* **26** 1727 1893, DOI: 10.1002/cber.189302602109; Plumb & Harper *Chem Eng News* (July 16) 3 1990] when apparently the iodine content was below 43.5% (theoretical value is 45.32%). In an attempt to avoid this, **IBX** has been washed with H₂O and EtOH to render it non-explosive. Although deliberate attempts to detonate it on several occasions have been unsuccessful; operators should exercise CAUTION however when working with **IBX** particularly with large scale preparations. This is among the substances for which it may be difficult to obtain permission to transport it, so detailed preparations are described here.

Dess-Martin method: To a vigorously stirred mixture of 2-iodobenzoic acid (85.2g, 340mmol, see [88-67-5]) and 0.73 M H₂SO₄ (730ml) in a bath at 55° is added KBrO₃ (76.0g, 450mmol) over 1 hour. The mixture is stirred further for 3.6 hours at 65°, cooled in an ice bath, and the solid is filtered off, washed with H₂O (1000ml) then EtOH (2 x 50ml) and dried *in vacuo* to give BTX (89.1g, 320mmol, 93%) of analytical purity. [Dess & Martin *J Am Chem Soc* **113** 7277 1991, DOI: 10.1021/ja00019a027]

Santagostino et al.'s method: To **oxone** (181.0g, 290mmol, see [70693-62-8] 2KHSO₅—KHSO₄—K₂SO₄ triple salt) in deionised H₂O (650ml, 450mmol) is added rapidly 2-iodobenzoic acid (50.0g, 200mmol), and kept at 70-73° for 20 minutes then stirred mechanically at 70-73° for 3 hours. Initially a thick slurry coating along the walls of the container is formed which gradually becomes finely dispersed and the now easily stirred suspension readily sediments if stirring is interrupted. The mixture is then stirred slowly at 5° for 1.5 hours, the white crystalline solid is collected onto a medium porosity sintered-glass funnel, washed with H₂O (6 x 100ml) and Me₂CO (2 x 100ml) then Et₂O, and left in dry air for 16 hours to give IBX (44.8 to 45.7g, ~80%). The filtrates are disposed of by treating with solid Na₂SO₃ (70g, 55mmol) and neutralising with M aqueous NaOH. Analysis indicated that the purity was ≥95% (judging by the ¹H NMR integrals of the triplets at 7.84 and 7.47 ppm, and elemental analysis), and it contained 2-iodosobenzoic acid (~4%) and 2-iodobenzoic acid (~0.5%). **IBX** of analytical purity (≥99%) is similarly obtained on a small scale by adding 2-iodobenzoic acid (5.0g, 20mmol) to a solution of oxone in deionised H₂O (37.2g, 61mmol, in 200ml), and the suspension is kept at 70° for 1 hour when it becomes clear. After 0.5 hours at 0-5° the white crystals that separate are collected, washed and dried as before to give analytically pure **IBX** (4.4g, 77%), **m 233° (dec)**. This procedure is more environmentally friendly than the Dess-Martin procedure. [Frigerio et al. *J Org Chem* **64** 4537 1999, DOI: 10.1021/jo9824596.] It has IR (film) with a ν_{max} at 1640 cm⁻¹; the ¹H NMR (DMSO-d₆, TMS) has δ at 8.15 (d,

1H), 8.01 (d, 1H), 7.89 (t, 1H), and 7.84 (t, 1H); and for ^{13}C NMR see reference see Frigerio and Santagostino [*Tetrahedron Lett* **35** 8019 1994, DOI: 10.1016/0040-4039(94)80038-3].

SIBX is an even more stabilised, non-explosive formulation that has all the oxidative characteristics of **IBX**. It is prepared as follows: 2-iodobenzoic acid (200g) and isophthalic acid (133g, see [121-91-5]) are added into a solution of oxone (625g) in H_2O (2000ml) and kept at 70° for 3 hours, then sodium benzoate (128g) in H_2O (500ml) is added at 40° . The precipitate that is formed on cooling to 20° is filtered off, washed with H_2O (700ml) and dried in a ventilated oven at 60° to give **SBTX** (420g, 90%) containing 49% w/w of active **IBX**. [Ozanne et al. *Org Lett* **5** 2903 2003, DOI: 10.1021/ol0349965.]

SIBX and/or **IBX** are water tolerant, inexpensive to prepare and easy to use. They oxidise primary and secondary alcohols smoothly to aldehydes and ketones, and vicinal diols are oxidised to mono and/or dicarbonyl compounds without C-C cleavage [Frigerio & Santagostino *Tetrahedron Lett* **35** 8019 1994, DOI: 10.1016/0040-4039(94)80038-3], as well as allylic and benzylic alcohols to respective aldehydes [Ozanne et al. *Org Lett* **5** 2903 2003, DOI: 10.1021/ol0349965], amino alcohols to amino carbonyls without amino protection [Chen & Aduda *Synth Commun* **37** 3493 2007, DOI: 10.1080/00397910701555469], and sensitive heterocyclic substituents are not affected [Frigeio et al. *J Org Chem* **60** 7272 1995, DOI: 10.1021/jo00127a036]. The structure and kinetics of the reactive intermediates in the oxidation have been studied in detail [Munari et al. *J Org Chem* **61** 9272 1961, DOI: 10.1021/jo961044m]. The reagents are generally sparingly soluble in organic solvents such as EtOAc, THF, Me_2CO , MeCN, toluene, DMSO and *N*-methylpyrrolidine, but the reactions usually proceed in these solvents, or mixtures of them, with ease and in good to excellent yields either at room temperature or at as high as reflux temperatures.

2-Hydroxybenzothiazole (benzothiazol-2(3H)-one) [934-34-9] $\text{C}_7\text{H}_5\text{NOS}$, **M 151.1**, **m 140-141°**, Crystallise it from aqueous EtOH or water. [Hoggarth *J Chem Soc* 3311 1949, DOI: 10.1039/JR9490003311; Hunter *J Chem Soc* 125 1930, DOI: 10.1039/JR9300000125; *Beilstein* **27** H 182, **27** I 270, **27** II 225, **27** III/IV 2693.]

1-Hydroxybenzotriazole hydrate (HOBt) [2592-95-2; 123333-53-9 (H_2O)] $\text{C}_6\text{H}_5\text{N}_3\text{O} \cdot x\text{H}_2\text{O}$, **M 135.1**, **m 155-158°, 157°, 159-160°, pK²⁰ 7.88**. Crystallise HOBt from hot aqueous EtOH or water (charcoal). It is prepared from *o*-nitrophenylhydrazine or its hydrochloride dissolved in a small volume of H_2O by treating with 25% of aqueous KOH or aqueous ammonia whereby the mixture warms up immediately and discolours. After the mixture cools to room temperature it is acidified with hydrochloric acid and the **hydroxybenzotriazole** separates as colourless needles. It is soluble in hot H_2O , EtOH and AcOH, but much less soluble in Et_2O , petroleum ether, $^*\text{C}_6\text{H}_6$ and CHCl_3 . It is an acid and forms metal salts such the **Pb salt** which crystallises from hot H_2O in glistening leaflets with **m 270°**. In aqueous solution it exists mainly as the zwitterionic N-oxide tautomer with H^+ on N-2. [Nietzki & Braunschweig *Chem Ber* **27** 3381 1894, DOI: 10.1002/cber.189402703148; Zincke & Schwartz *Justus Liebigs Ann Chem* **311** 329 1900, DOI: 10.1002/jlac.19003110304; Boyle & Jones *JCS Perkin Trans II* 160 1973, DOI: 10.1039/P29730000160; Tomita & Ikawa *J Pharm Soc Jpn* **75** 449 1955, *Beilstein* **26** III/IV 95.] It is a useful reagent for peptide synthesis [Heusel et al. *Angew Chem Int Ed* **16** 642 1977, DOI: 10.1002/anie.197706421].

§ A polystyrene supported version is commercially available. For use in solid phase peptide synthesis, see Dryland & Sheppard *JCS Perkin Trans I* 125 1986, DOI: 10.1039/P19860000125.

4-Hydroxycoumarin (4-hydroxy-1-benzopyran-2-one) [1076-38-6] $\text{C}_9\text{H}_6\text{O}_3$, **M 162.1**, **m 206°, 211-213°, pK_{Est} ~9.0**. Crystallise 4-hydroxycoumarin from water and dry it in a vacuum desiccator over Sicapent. [*Beilstein* **18**/I V 378.]

N-2-Hydroxyethylpiperazine-N'-2-ethanesulfonic acid (HEPES) [7365-45-9] $\text{C}_8\text{H}_{18}\text{N}_2\text{O}_4\text{S}$, **M 238.3**, **pK²⁰ 7.55**. Crystallise the acid from hot EtOH and water. The **disodium salt** [75277-39-3] $\text{C}_8\text{H}_{17}\text{N}_2\text{NaO}_4\text{S}$, **M 260.3**, is purified by dissolving in the minimum volume of H_2O and adding EtOH, then drying *in vacuo*. They are useful buffers in the pH range 6.8–8.2. [*Beilstein* **23** V 376.]

3-Hydroxyflavone (Flavanol) [577-85-5] $\text{C}_{15}\text{H}_{10}\text{O}_3$, **M 238.2**, **m 169-170°, 171-172°**. Recrystallise it from MeOH (**m 169.5-170°**), EtOH, aqueous EtOH (**m 167°**) or hexane. It has also been purified by repeated sublimation under high vacuum, and dried at high vacuum pumping for at least one hour [Brucker & Kelley *J Phys Chem* **91** 2856 1987, DOI: 10.1021/j100295a040]. [*Beilstein* **17** H 527, **17** I 268, **17** II 498, **17** III/IV 6428.]

7-Hydroxy-4-methylcoumarin (4-methylumbelliferone(β) hydrate) [90-33-5] $\text{C}_{10}\text{H}_8\text{O}_3 \cdot \text{H}_2\text{O}$, **M 194.2, m 185-186°, 188-188.5°, 194-195°, $\text{pK}^{25}_{\text{a}}$ 7.80 (phenolic OH).** Purify it by recrystallisation from EtOH. It is very slightly soluble in cold H_2O (solubility at 37° is 0.22%), slightly soluble in Et_2O and CHCl_3 , but soluble in MeOH and AcOH. It has a blue fluorescence in aqueous EtOH and has UV with λ_{max} at 221, 251 and 322.5nm (MeOH). The IR has ν_{max} at 3077 br, 1667, 1592, 1385, 1267, 1156, 1130 and 1066 cm^{-1} . The *acetate* has **m 153-154°**. It is highly fluorescent in alkaline solution. [Woods & Sapp *J Org Chem* **27** 3703 1962, DOI: 10.1021/jo01057a519; *Beilstein* **18** III/IV 332, **18/1** V 439.]

2-Hydroxymethyl-12-crown-4 [75507-26-5] $\text{C}_9\text{H}_{18}\text{O}_5$, **M 206.2, b 115°/0.04mm, d_4^{20} 1.186, n_D^{20} 1.480.** Purify it by chromatography on Al_2O_3 with EtOAc as eluent to give a *hygroscopic* colourless oil, which is distilled under high vacuum and is distilled *in vacuo*. It has IR with ν_{max} at 3418 (OH) and 1103 (COC) cm^{-1} , and NMR: δ 3.70s. [Kimura et al. *JCS Chem Commun* 492 1983, DOI: 10.1039/C39830000492; Puglia et al. *J Org Chem* **52** 2617 1987, DOI: 10.1021/jo00388a057.]

S-(-)-5-Hydroxymethyl-2(5H)-furanone [78508-96-0] $\text{C}_5\text{H}_6\text{O}_3$, **M 114.1, 39-42°, 41-43°, 40-44°, b 130°/0.3mm, $[\alpha]_{\text{D}}^{20}$ -180, $[\alpha]_{\text{D}}^{20}$ -148 (c 1.4, H_2O).** It is purified by chromatography on Silica gel using hexane/EtOAc (1:1) to give a colourless oil which is distilled using a Kugelrohr apparatus, and the distillate crystallises on cooling. It has R_F 0.51 on Whatman No 1 paper using pentan-1-ol and 85% formic acid (1:1) and developing with ammoniacal AgNO_3 . [Boll *Acta Chem Scand* **22** 3245 1968, DOI: 10.3891/acta.chem.scand.22-3245; NMR: Oppolzer et al. *Helv Chim Acta* **68** 2100 1985, DOI: 10.1002/hlca.19850680803; *Beilstein* **18** III/IV 56, **18/1** V 54.]

5-(Hydroxymethyl)furfural [67-47-0] $\text{C}_6\text{H}_6\text{O}_3$, **M 126.1, m 28-34°, 33.5°, b 114-116°/1mm, d_{25}^{25} 1.2620, n_D^{25} 1.5533.** Crystallise it from diethyl ether/petroleum ether. [*Beilstein* **18** III/IV 100, **18/1** V 130.]

dl-3-Hydroxy-N-methylmorphinan (Racemorphan), see entry in ‘Physiologically Active……’, in Chapter 6.

8-Hydroxy-2-methylquinoline [826-81-3] $\text{C}_{10}\text{H}_9\text{NO}$, **M 159.2, m 71-73°, 74-75°, b 266-267°/atm, pK_1^{25} 5.61, pK_2^{25} 10.16.** Crystallise the quinoline from EtOH or aqueous EtOH. Its solubility at 20° in H_2O is 0.366g/L, and in CHCl_3 it is 466g/L. It complexes with many metals. [*Beilstein* **21** H 106, **21** III/IV 2132, **21/3** V 341.]

4-Hydroxy-2-n-nonylquinoline N-oxide [316-66-5] **M 287.4, m 148-149°, pK_{Est} ~6.0.** Crystallise the N-oxide from EtOH. Its UV has λ_{max} at 220-230nm in 0.001N NaOH. [Cornforth & James *Biochem J* **63** 124 1956, DOI: 10.1042/bj0630124; *Beilstein* **21** III/IV 3834.]

1-Hydroxyphenazine (Hemipyocyanine) [528-71-2] $\text{C}_{12}\text{H}_8\text{N}_2\text{O}$, **M 196.2, m 156-160°, 157-158°, 165-166°, pK_1^{15} 1.61, pK_2^{15} 8.33 (10% aqueous MeOH)** Purify the hydroxyphenazine by chromatography on acidic alumina with *benzene/ether, and recrystallise it from aqueous EtOH, *benzene/heptane, then sublime it at 140°/1.5mm. It forms a red *chloride* or *sulfate salt*. When dissolved in alkaline solution it gives a purple-red colour which turns yellow on dilution. [UV, IR: Badger et al. *J Chem Soc* 3204 1951, DOI: 10.1039/JR9510003204; Hegedüs *Helv Chim Acta* **33** 766 1950, DOI: 10.1002/hlca.19500330340; *Beilstein* **23** II 360, **23** III/IV 2753.]

2-(2-Hydroxyphenyl)benzothiazole [3411-95-8] $\text{C}_{13}\text{H}_9\text{NOS}$, **M 227.2, m 128-132°, 132-133°, b 173-179°/3mm.** Recrystallise it several times from aqueous EtOH or dilute AcOH and sublime it. [Itoh & Fujiwara *J Am Chem Soc* **107** 1561 1985, DOI: 10.1021/ja00292a018; Bogert & Corbitt *J Am Chem Soc* **48** 783 1926, DOI: 10.1021/ja01414a040; *Beilstein* **27** H 118, **27** II 91.]

2-(2-Hydroxyphenyl)benzoxazole [835-64-3] $\text{C}_{13}\text{H}_9\text{NO}_2$, **M 211.2, m 122-124°, 127°, b 338°/760mm.** Recrystallise it several times from aqueous EtOH or dilute AcOH and sublime it. An aqueous alkaline solution containing EtOH has a blue fluorescence. [Itoh & Fujiwara *J Am Chem Soc* **107** 1561 1985, DOI: 10.1021/ja00292a018; *Beilstein* **27** II 91.]

3-Hydroxy-2-phenylcinchoninic acid (Oxycinchophen) [485-89-2] $C_{16}H_{11}NO_3$, M 265.3, m 206-207°(dec). It is precipitated from alkaline solution on acidification, and crystallises from EtOH or AcOH in yellow prisms. It has antidiuretic properties. [Marshall & Blanchard *J Pharmacol Exp Ther* **95** 186 1949, PMID: 18124061; *Beilstein* **22** H 245, **22** II 183, **22** III/IV 2383.]

(±)-2-(α-Hydroxypropyl)piperidine [2-piperidinepropanol, *RS*-1-(*SR*-2-piperidyl)propan-1-ol, (±)-α-conhydrine] [8338-62-8, 63401-12-7, 24448-89-3] $C_8H_{17}NO$, M 143.2, m 99-100°, $pK_{Est} \sim 10.2$. Crystallise it from ether. The [*RS*-1-(*SR*-2)]-isomer also crystallises from ether and has m 98-98.5°. The *methiodide* crystallises from $Me_2CO/MeOH$ with m 127-130.5°. **POISONOUS** [Synth: Shono et al. *Tetrahedron Lett* **24** 4457 1983, DOI: 10.1016/S0040-4039(00)85960-2; Sicher & Tichy *Coll Czech Chem Commun* **23** 2081 1958, DOI: 10.1135/cccc19582081; Govindachari & Rajappa *J Chem Soc* 1297, 1306 1958, DOI: 10.1039/JR9580001297; Stereochemistry: Hill *J Am Chem Soc* **80** 1609 1958, DOI: 10.1021/ja01540a025; *Beilstein* **21** II 21, **21** III/IV 122.]

(+)-2-(α-Hydroxypropyl)piperidine [2-piperidinepropanol, *R*-1-(*S*-2-piperidyl)propan-1-ol, (+)-α-conhydrine] [495-20-5] has m 121°, b 224-5°/720mm, 226°/atm, $[\alpha]_D^{20} +9.8$ (c 4, EtOH), +10 (c 10, EtOH), $pK_{Est} \sim 10.2$. This very **POISONOUS** alkaloid from hemlock crystallises in leaflets from ether, or from H_2O . The *O,N*-dibenzoyl derivative has m 133-134° and $[\alpha]_D^{24} -13$ (c 3, $CHCl_3$). The *acetate* has b 133-135°/3mm. [Sicher & Tichy *Coll Czech Chem Commun* **23** 2081 1958, DOI: 10.1135/cccc19582081; Stereochemistry: Hill *J Am Chem Soc* **80** 1609 1958, DOI: 10.1021/ja01540a025; Absolute config & ORD: Fodor et al. *Can J Chem* **47** 4393 1969, DOI: 10.1139/v69-727; Short Review: Bhat et al. in *Synthesis* **46** 2551 2014, DOI: 10.1055/s-0034-1379023; *Beilstein* **21** I 191, **21** II 21, **21** III/IV 122.]

(-)-2-(α-Hydroxypropyl)piperidine [2-piperidinepropanol, *S*-1-(*R*-2-piperidyl)propan-1-ol, (-)-α-conhydrine] [18209-3-5] has m 121°, b 224-5°/720mm, $[\alpha]_D^{27} -8.6$ (c 068, EtOH), $pK_{Est} \sim 10.2$. Crystallise the piperidine from ether. **POISONOUS**. [Synth, 1H NMR NOE, abs. config. and X-Ray: Enders et al. *Tetrahedron: Asymmetry* **13** 285 2002, DOI: 10.1016/S0957-4166(02)00066-6; Galinovsky & Mulley *Monatsh Chem* **79** 426 1948, DOI: 10.1007/BF00918557; *Beilstein* **21** I 191, **21** II 21, **21** III/IV 122.]

(±)-7-(2-Hydroxypropyl)theophylline (Proxiphylline, 1,3-dimethyl-3*H*,7*H*-purine-2,6-dione) [603-00-9] $C_{10}H_{14}N_4O_3$, M 238.2, m 135-136°. Crystallise it from EtOH, aqueous MeOH or EtOAc. It is a vaso(broncho) dilator. [Roth *Archiv Pharmazie* **292** 234 1959, DOI: 10.1002/ardp.19592920504; Zelnik et al. *Bull Soc Chim Fr* 1733 1956, *Beilstein* **26** III/IV 2366.]

6-Hydroxypurine (hypoxanthine) [68-94-0] $C_5H_4N_4O$, M 136.1, m 150°(dec), >300°(dec), $pK_1^{20} 1.98$, $pK_2^{20} 8.96$, $pK_3^{20} 12.18$. Crystallise it from hot water (solubility (w/v) is 0.07% at ~20° and 1.42% at ~100°) and dry it at 105°. [Lister *Purines Part II, Fused Pyrimidines* Brown Ed, Wiley-Interscience pp254-225 1971, ISBN 0-471-38205-1, *Beilstein* **26** II 252, **26** III/IV 2081.]

2-Hydroxypyridine (2-pyridone) [142-08-5] C_5H_5NO , M 95.1, m 105-107°, 107.8°, b 181-185°/24mm, 280-281°/atm, $d^{20} 3.1g/cm^3$, $\epsilon_{293nm} 5900$ (H_2O) $pK_1^{25} 1.25$ (0.75), $pK_2^{25} 11.99$ (11.62). It has been prepared from 2-aminopyridine and nitrous acid [Adams & Jones *J Am Chem Soc* **69** 1803 1947, DOI: 10.1021/ja01199a067], from pyridine N-oxide and Ac_2O , pyridine and KOH at 300°, or 2-chloropyridine and dilute mineral acid. X-ray analysis by Penfold [*Acta Cryst* **6** 591 1953 DOI: 10.1107/S0365110X5300168X, **6** 707 1953, DOI: 10.1107/S0365110X5300199X] who showed that it has the *pyridone* structure in the solid state, and UV spectral comparisons support this structure in solution, rather than the 2-hydroxypyridine structure since the UV spectra are similar to those of 1-methyl-2-pyridone and not to those of 2-methoxypyridine. [Mason *J Chem Soc* 5010 1957, DOI: 10.1039/JR9570005010; Mason *J Chem Soc* 1253 1959, DOI: 10.1039/JR9590001253; UV: Armarego in *Physical Methods in Heterocyclic Chemistry* (Ed Katritzky, Academic Press, LCCC No 62-12037, Internet Public Domain) **Vol III** 79 1971]. Distil 2-pyridone under vacuum to remove coloured impurity, then recrystallise from *benzene, CCl_4 , Me_2CO or EtOH. It can be sublimed under high vacuum then recrystallised from $CHCl_3$ /diethyl ether. [DePue et al. *J Am Chem Soc* **107** 2131 1985, DOI: 10.1021/ja00293a050; *Beilstein* **21/7** V 106.]

1-Methyl-2-pyridone [694-85-9, 94071-56-4] C_6H_7NO , M 109.1, m 30-32°, b 250°/740mm, $d^{20} 1.11g/cm^3$, $pK^{20} 0.32$. It has been prepared in 65-70% yield by converting pyridine to *1-methylpyridinium methosulfate* with dimethyl sulfate followed by oxidation with $K_3Fe(CN)_6/NaOH/H_2O$, salting out with Na_2CO_3 , extracting with isoamyl alcohol and distilling under vacuum. It distils at b 121°/10mm, 126°/12.5mm and 130°/14.5mm.

[Prill & McElvain *Org Synth Coll Vol* **2** 418 1943, *Org Synth* **15** 41 1935, DOI: 10.15227/orgsyn.015.0041.]

2-Methoxypyridine [1628-89-3] C_6H_7NO , **M 109.1**, **b** 142°/760mm, **d**²⁵ 1.038g/ml, **n**_D²⁰ 1.503, **pK**²⁰ 3.28, is obtained by reaction of 2-chloropyridine with MeOH and is purified by distillation after washing with H₂O (at pH slightly higher than 7).

2-Pyridyl acetate (2-acetoxypyridine) [3847-19-6] $C_7H_7NO_2$, **M 137.1**, prepared by treating the sodium salt of 2-pyridone with acetyl chloride and purified by vacuum distillation, has **b** 110-112°/10mm. This *acetate* is a useful acetylating agent and in THF, xylene or neat, it will acetylate a variety of alcohols, phenols and amines in yields generally above 80%. It is more reactive than 3-pyridyl acetate (see below) in Friedel-Crafts reactions (in the presence of AlCl₃, or better BF₃) as in the acylation of *C₆H₆, toluene (mostly *p*-) and anisole (mostly *p*-) [Ueno et al. *Bull Chem Soc Jpn* **37** 864 1964, DOI: 10.1246/bcsj.37.864].

3-Hydroxypyridine [109-00-2] C_5H_5NO , **M 95.1**, has **m** 125-128°, 129°, 130°, **pK**₁²⁵ 5.10 (4.88), **pK**₂²⁵ 8.60 (8.72). 3-Hydroxypyridine was obtained by fusing pyridine-3-sulfonic acid with KOH, or by decomposing pyridine-3-diazonium salts with H₂O and purified by recrystallisation from *C₆H₆, water or EtOH. Unlike the 2- and 4- isomers it cannot tautomerise to an amide-like pyridine, and displays properties not unlike phenols, e.g. gives a colour with FeCl₃. It has zwitterionic properties and when alkylated, e.g. with MeI/*n*-PrOH, it forms the *N*-methyl zwitterion which can be converted into *N*-methyl-3-hydroxypyridinium iodide **m** 109-111° and the *picrate* **m** 201-202°. [Shapiro et al. *J Am Chem Soc* **81** 5140 1959, DOI: 10.1021/ja01528a030; Albert & Phillips *J Chem Soc* 1294 1956, DOI: 10.1039/JR9560001294; Metzler & Snell *J Am Chem Soc* **77** 2431 1955, DOI: 10.1021/ja01614a022; and for UV see Mason *J Chem Soc* 1253 1959, DOI: 10.1039/JR9590001253; and Armarego in *Physical Methods in Heterocyclic Chemistry* (Ed Katritzky, Academic Press, LCCC No 62-12037, Internet Public Domain) **Vol III** 79 1971.] Methylation of 3-hydroxy-pyridine with diazomethane normally provides the *N*-methyl compound, however diazomethane in *tert*-BuOH at -15° gives **3-methoxypyridine** [7295-76-3] C_6H_7NO , **M 109.1**, **b** 65°/15mm, **d**²⁵ 1.083g/ml, **n**_D²⁰ 1.518, **pK**²⁰ 4.88, in 70% yield and is purified by vacuum distillation [Prins *Recl Trav Chim Pays-Bas* **76** 58 1957, DOI: 10.1002/recl.19570760107]. **3-Pyridyl acetate (3-acetoxypyridine)** [17747-43-2] $C_7H_7NO_2$, **M 137.1**, **b** 92°/9mm, **d**²⁵ 3.1g/ml, **n**_D²⁰ 1.503, is obtained by acetylation with acetic anhydride, as with phenols and unlike the 2-pyridone (see above), is purified by vacuum distillation with **b** 92°/9mm. Like the **2-isomer** (see above) it is an acetylating agent and undergoes the Friedel-Crafts reaction but is marginally less active [Ueno et al. *Bull Chem Soc Jpn* **37** 864 1964, DOI: 10.1246/bcsj.37.864]. [Beilstein **21** III/IV 402, **21/2** V 68.] 3-Hydroxypyridine forms esters with the terminal carboxy groups of *N*-protected peptides, using dicyclohexylcarbodiimide in EtOAc, whereby the carboxy groups is activated to form a peptide bond with an unprotected terminal NH₂ [Taschner et al. *Angew Chem Int Ed* **4** 594 1965, DOI: 10.1002/anie.196505941].

4-Hydroxypyridine (4-pyridone) [626-64-2] has **m** 65°, 68° (hydrate), 148.5°, 150-151°, 151-152° (anhydr), **b** 230-235°/12mm, >350°/760mm, **pK**₁²⁰ 3.20, **pK**₂²⁰ 11.12. It has been prepared from pyridine by treatment with SOCl₂ which forms 4-chloropyridine that reacts with pyridine to give *N*(4-pyridyl)-pyridinium chloride followed by hydrolysis with H₂O at 15°. Crystallise 4-pyridone from H₂O or wet CHCl₃ as the *monohydrate*. It loses H₂O on drying *in vacuo* over H₂SO₄. Store it over KOH because it is *hygroscopic*. Like 2-pyridone (see above) it exists in the keto **4-one tautomer** as evidenced by IR and Raman spectra [Spinner *J Chem Soc* 1226, 1232 1960,], and **pKa** and UV spectral comparison with 4-methoxypyridine and 1-methyl-4-pyridone [Mason *J Chem Soc* 5010 1957, DOI: 10.1039/JR9570005010; Mason *J Chem Soc* 1253 1959, DOI: 10.1039/JR9590001253; UV: Armarego in *Physical Methods in Heterocyclic Chemistry* (Ed Katritzky, Academic Press, LCCC No 62-12037, Internet Public Domain) **Vol III** 79 1971]. For protonated species see Katritzky and Jones [*Proc Chem Soc* 297 1960, DOI: 10.1039/PS9600000297]. [Beilstein **21** III/IV 446, **21/7** V 152.] **4-Methoxypyridine** [620-08-6] C_6H_7NO , **M 109.1**, **b** 95°/45mm, 108-111°/65mm, 191°/738mm, **d**²⁵ 1.075g/ml, **n**_D²⁰ 1.516, **pK**²⁰ 6.62, like the **2-isomer**, can be prepared from 4-chloropyridine and MeOH, washing the extracted product with aqueous bicarbonate before drying and distilling preferably under vacuum. Care should be taken that it should be free of traces of acid as this causes serious deterioration by polymerisation. **1-Methyl-4-pyridone** [695-19-2] C_6H_7NO , **M 109.1**, has **m** 94.5-95.5°, **b** 188-192°/2.4mm, **d**²⁰ 1.11g/cm³, **pK**²⁰ 3.33. Prepared from 4-pyridone and MeI, then washed with H₂O and recrystallised from MeOH. The *hydrochloride* crystallises from isoPrOH with **m** 186-187°.

2(6)-Hydroxypyridine-5(3)-carboxylic acid (6-hydroxynicotinic acid) [5006-66-6] $C_6H_5NO_3$, **M 139.1**, **m** 304°(dec), **pK**₁²⁰ 3.82 (proton gain) and **pK**₂²⁰ 9.92 (proton loss). It crystallises from water (400 parts and dried at 110°) with **m** 303.4-303.7°(dec), or with **m** 325°(dec) from aqueous EtOH. The *methyl ester* crystallises from Me₂CO with **m** 166° and **pK**²⁰ 9.92 (proton loss). Used in the synthesis of retinoids [Torrado

et al. *Synthesis* 285 1995, DOI: 10.1055/s-1995-3905]. [Synthesis, pKa and UV spectra: Albert *J Chem Soc* 1020 1960, DOI: 10.1039/JR9600001020, *Beilstein* 22 III/IV 2147, 22/6 V 119.]

4-Hydroxypyridine-2,6-dicarboxylic acid (chelidamic acid) [138-60-3] $C_7H_5NO_5 \cdot xH_2O$, M 183.1, m 248°(dec, 1 H_2O), 254°(dec), 263°(dec), 268°(dec), pK₁²² 1.9, pK₂²² 3.18, pK₃²² 10.85; pK₁ 3.62, pK₂ 4.82 (80% aqueous methoxyethanol). It crystallises from water. The *dimethyl ester* crystallises from MeOH with m 167°(monohydrate, from H_2O), m 170-171°(anhydrous, from MeOH), and has pK₂₅ 6.25. [Bensaude et al. *J Am Chem Soc* 99 4438 1972, DOI: 10.1021/ja00455a037; *Beilstein* 22 III/IV 2583, 22/7 V 164.]

2-Hydroxypyrimidine [557-01-7] $C_4H_4N_2O$, M 96.1, m 179-180°, pK₁²⁰ 2.24, pK₂²⁰ 9.17. It crystallises from EtOH or ethyl acetate. Its solubility in H_2O at 20° is 1g/2.2ml. [Albert *J Chem Soc* 4219 1952, DOI: 10.1039/JR9600001020; *Beilstein* 24 III/IV 194.]. **2-Hydroxypyrimidine hydrochloride** [38353-09-2] M 132.5, has m 203-205.5°(dec) and crystallises from EtOH. [*Beilstein* 24 III/IV 173.]. The *picrate* has m 199° and crystallises from EtOH. [Brown et al. *J Chem Soc* 211 1955, DOI: 10.1039/JR9550000211.]

4-Hydroxypyrimidine [4562-27-0, 51953-17-4] has m 163-167°, 164-165°, 166-169°, pK₁²⁰ 1.66, pK₂²⁰ 8.63. It crystallises from *benzene or ethyl acetate. The *picrate* has m 164-166° and crystallises from EtOH [Brown et al. *J Chem Soc* 4035 1955, DOI: 10.1039/JR9550004035; *Beilstein* 24 III/IV 171.]

R-(+)-3-Hydroxypyrrolidine [R(+) 2799-21-5, (±) 40499-83-0, S(-) 100243-39-8] C_4H_9NO , M 87.1, b 215-216°/atm, d₄²⁰ 1.078, n_D²⁰ 1.490, [α]_D²⁰ +6.5 and -6.5 (c 1.5, MeOH), pK_{Est} ~10.1. The (±)-*base* is purified by repeated distillation (b 102-104°/12mm, 108-110°/18mm), and the (±)-*picrate* crystallises from EtOH with m 140-141°. The *R*(+)-*enantiomer* has b 70°/0.6mm and [α]_D²⁰ +5.6 (c 3.63, MeOH). Its *hydrochloride* [104706-47-0] M 123.6, m 104-107°, has a negative rotation [α]_D²⁰ -7.6 (c 3.5, MeOH), and its *dimethiodide* has m 230° and [α]_D²⁴ -8.02°. [Uno et al. *J Heterocycl Chem* 24 1025 1987, DOI: 10.1002/jhet.5570240423; Flanagan & Joullie *Heterocycles* 26 2247 1987, DOI: 10.3987/R-1987-08-2247; *Beilstein* 21 III/IV 44.]

2-Hydroxyquinoline (carbostyryl) [59-31-4] C_9H_7NO , M 145.2, m 198-199°, 199-200°, pK₁²⁰ -0.31, pK₂²⁰ 11.76. Crystallise it from MeOH. It has m 200-201° after sublimation in a vacuum. The *picrate* has m 132° after crystallisation from Et₂O. [Gibson et al. *J Chem Soc* 4340 1955, DOI: 10.1039/JR9550004340; *Beilstein* 21 III/IV 1057, 21/8 V 217.]

8-Hydroxyquinoline (oxine, 8-quinolinol) [148-24-3] has m 70-73°, 71-73°, 75-76°, 76°, b 122°/0.1mm, ~267°/752m, pK₁²⁵ 4.91, pK₂²⁵ 9.81. Crystallise oxine from hot EtOH, acetone, petroleum ether (b 60-80°) or water. Crude oxine can be purified by precipitation of *CuII oxinate* [13014-03-4] $Cu(C_9H_7NO)_2$, m 240°(dec), followed by liberation of *free oxine* with H_2S or by steam distillation after acidification with H_2SO_4 . Store it in the dark. It forms complexes with many metals or metal oxides on fusing or in solution. [Feigl & Heisig *Anal Chim Acta* 3 561 1949, DOI: 10.1016/S0003-2670(00)87382-1]. The *hemisulfate hemi-hydrate* [207386-91-2] $C_9H_7NO \cdot 0.5H_2SO_4 \cdot 0.5H_2O$, M 203.2 has m 176-179°. [Manske et al. *Can J Research* 27F 359 1949, DOI:10.1139/cjr49f-038; Phillips *Chem Rev* 56 271 1956, DOI: 10.1021/cr50008a003; *Beilstein* 21 III/IV 1135, 21/3 V 252.]

8-Hydroxyquinoline-5-sulfonic acid (H_2O) [84-88-8, 20946-17-2] $C_9H_7NO_4S \cdot H_2O$, M 243.3, has m 322-323°(sintering at ~305°), >310°, pK₁²⁵ 4.09, pK₂²⁵ 8.66. Crystallise the acid from water (as the *1.5 hydrate*, m 316-317°) or dilute HCl (ca 2% by weight). It is a water-soluble metal complexing agent. [*Beilstein* 22 I 620, 22 II 313, 22 III/IV 3493.]

4-Hydroxy-2,2,6,6-tetramethylpiperidine [2403-88-5] $C_9H_{19}NO$, M 157.3, m 129-131°, 130-131°, b 212-215°/atm, pK₂₀ 10.05. The piperidine crystallises from water as a *hydrate*, and crystallises from dry ether or * C_6H_6 as the *anhydrous base*. The *hydrochloride* has m 282-284° (from EtOH/ H_2O), and the *formate* has m 207°(dec, from EtOH/EtOAc). [Mailey & Day *J Org Chem* 22 1061 1957, DOI: 10.1021/jo01360a017; *Beilstein* 21 I 195, 21 III/IV 146, 21/1 V 159.]

4(6)-Hydroxy-2,5,6(2,4,5)-triaminopyrimidine sulfate [35011-47-3] $C_4H_7N_5O \cdot H_2SO_4$, M 239.2, m >~300°, >340°, pK₁ 2.0, pK₂ 5.1, pK₃ 10.1. This salt has very low solubility in H_2O . It is best purified by conversion into the dihydrochloride salt, which is then re-converted to the insoluble sulfate salt. The sulfate salt (2.57g, 10mmoles) is suspended in H_2O (20ml) containing $BaCl_2$ (10mmoles) and stirred in a boiling water bath for 15 minutes. After cooling, the insoluble $BaSO_4$ is filtered off and washed with boiling H_2O (10ml). The

combined filtrate and washings are made acidic with HCl and evaporated to dryness. The residual hydrochloride salt is recrystallised from H₂O by adding concentrated HCl whereby the *dihydrochloride salt* separates as clusters which darken at 260° and dec > 300° [darkening > 360°]. [Baugh & Shaw *J Org Chem* **29** 3610 1964, DOI: 10.1021/jo01035a042; King & Spengley *J Chem Soc* 2144 1952, DOI: 10.1039/JR9520002144]. The *hydrochloride* is then dissolved in H₂O, and while hot an equivalent of H₂SO₄ is added when the *sulfate* separates as a white microcrystalline solid which is filtered off washed liberally with H₂O and dried in vacuum over P₂O₅. [Albert & Wood *J Appl Chem London* **3** 521 1953, UV: Cavalieri et al. *J Am Chem Soc* **70** 3875 1948, DOI: 10.1021/ja01191a102; see also Pfeleiderer *Chem Ber* **90** 2272 1957, DOI: 10.1002/cber.19570901022; Traube *Chem Ber* **33** 1371 1900, DOI: 10.1002/cber.190003301236.] The *hydrochloride* has **m** > 300°. [Beilstein **25** III/IV 3648.] It is a useful synthon for purines and pteridines.

3-Hydroxyxanthone (3-hydroxy-9H-9-xanthenone) [3722-51-8] C₁₃H₈O₃, **M** 212.2, **m** 243°, 246°, 249-250°. Purify the xanthone by chromatography on SiO₂ gel with petroleum ether/*C₆H₆ as eluent. Recrystallise it from *C₆H₆, EtOH or aqueous EtOH (white needles). It exhibits interesting two-step laser-excitation fluorescence spectra. The 9 C=O and 3-OH form a tautomeric couple, and it forms a resonance-stabilised anion and radical anion. An alkaline solution has a blue fluorescence. The *acetate* has **m** 157-158°, and the *methyl ether* [3722-52-9] C₁₄H₁₀O₃, **M** 226.2, forms plates with **m** 129° from aqueous EtOH. [Davies et al. *J Org Chem* **23** 307 1958, DOI: 10.1021/jo01096a610; Itoh et al. *J Am Chem Soc* **107** 4819 1985, DOI: 10.1021/ja00303a003; Atkinson & Heilbron *J Chem Soc* 2688 1926, DOI: 10.1039/JR9262902688; for spectroscopic and theoretical studies of protonation see Mizutani et al. *Bull Chem Soc Jpn*, DOI: org/10.1246/bcsj.47.1596]. [Beilstein **18** H 46, **18** I 315, **18** II 29, **21** III/IV 601.]

Ibogaine (12-methoxybogamine) [83-74-9] C₂₀H₂₆N₂O, **M** 310.4, **m** 152-153°, [α]_D²⁰ -54° (EtOH), **pK**₂₅²⁵ 8.1 (80% aqueous MeOCH₂CH₂OH). Crystallise this alkaloid from EtOH or aqueous EtOH and sublime it at 150°/0.01mm. It is soluble in organic solvents but insoluble in H₂O. The *hydrochloride*, **m** 299-300°(dec), [α]_D²⁵ -63 (EtOH), is soluble in H₂O and alcohols. [Büchi et al. *J Am Chem Soc* **88** 3099 1866, DOI: 10.1021/ja00965a039; Rosenmund et al. *Chem Ber* **108** 1871 1975, DOI: 10.1002/cber.19751080611; Beilstein **23** III/IV 2742.]

Imidazole (glyoxaline) [288-32-4] C₃H₄N₂, **M** 68.1, **m** 89.5-91°, 89-90°, **b** 140-145°/15mm, 256°/atm, 262-264°/atm, **pK**₁²⁵ 6.99, **pK**₂²⁵ 14.44. Crystallise imidazole from *benzene, CCl₄, CH₂Cl₂, EtOH, petroleum ether, acetone/petroleum ether and distilled de-ionized water. Dry it at 40° under vacuum over P₂O₅. Distil it at low pressure. It is also purified by sublimation or by zone melting. [Snyder et al. *Org Synth Coll Vol* **3** 471 1955, DOI: 10.15227/orgsyn.022.0065; Brederick et al. *Chem Ber* **97** 827 1964, DOI: 10.1002/cber.19640970326; Caswell & Spiro *J Am Chem Soc* **108** 6470 1986, DOI: 10.1021/ja00281a004.] **¹⁵N-imidazole** crystallises from *benzene [Scholes et al. *J Am Chem Soc* **108** 1660 1986, DOI: 10.1021/ja00267a041]. [Beilstein **23** II 34, **23** III/IV 564, **23/4** V 191.] The *hydrochloride* [1467-16-9] C₃H₄N₂. HCl, **M** 104.5, has **m** 158-161° on crystallisation from EtOH/HCl/Et₂O, and the *sodium salt* [5587-42-8] C₃H₃N₂Na, **M** 90.1, has **m** 284°(dec). Imidazole is useful for buffers in the pH range 6.2–7.8.

1H-Indazole-3-carboxylic acid [4498-67-3] C₈H₆N₂O₂, **M** 162.2, **m** 265-265.5°, 266-270°(dec), 268-268.5°, **pK**_{Est(1)} ~4.5. Purify the acid by recrystallisation from glacial acetic acid (charcoal), and dry the yellow crystals *in vacuo*. Alternatively, dissolve the acid in boiling H₂O, concentrate the solution to one third its volume, cool and collect the yellow powder. Its UV has λ_{max} (H₂O) at 295 (log ε 3.88) nm. The *ethyl ester* (prepared *via* the acid chloride) has **m** 139° (from EtOH) and has λ_{max} (MeOH) at 393 nm; and the *N-methylamide* crystallises in tan needles from MeNO₂ with **m** 191-191° (187-188.5° was also reported). [Rousseau & Lindwall *J Am Chem Soc* **72** 3047 1950, DOI: 10.1021/ja01163a068; Snyder et al. *J Am Chem Soc* **74** 2009 1952, DOI: 10.1021/ja01128a042; Beilstein **25** H 129, **25** I 238, **25** II 128, **25** IV 808.]

1H, 2H-Indazol-3-one (3-hydroxy-1H-indazole) C₇H₆N₂O, [7364-25-2] **M** 134.1, **m** 250-252°, 253-254°, **pK**_{Est(1)} ~6.0. Purify indazol-3-one by recrystallisation from MeNO₂ or aqueous MeOH, and sublimation at 220°/0.1mm. In the UV it has λ_{max} in MeOH at 215 and 306 nm, and it is in the 3-OH form in ethanolic solution. There appears to be some controversy regarding the tautomeric form in the solid state, but the IR peak at ν_{max} 1627 cm⁻¹ (KBr) supports the keto form. [Ainsworth *J Am Chem Soc* **79** 5242 1957, DOI: 10.1021/

ja01576a047; Elguero et al. *Adv Heterocycl Chem*, **Suppl 1** 354 1976, Chernokal'skii et al. *Khim Geterotsikl Soedin* 96 1966, *Beilstein* **24** H 111, **24** II 59, **24** III/IV 270.]

4'-(Imidazol-1-yl)acetophenone [10041-06-2] $C_{11}H_{10}N_2O$, **M 186.2**, **m 104-107°**, **108-110°**, **pK²⁵ 4.54**. Recrystallise it twice from CH_2Cl_2 /hexane [Collman et al. *J Am Chem Soc* **108** 2588 1986, DOI: 10.1021/ja00270a016].

2-Iminothiolane hydrochloride (2-iminotetrahydrothiophene, Traut's reagent) [4781-83-3] C_4H_7NS . **HCl**, **M 137.6**, **m 187-192°**, **190-195°**, **192-193°**, **193-194°**, **190-201°**, **202-203°**, **pK²⁵ <2 (free base)**. Recrystallise the hydrochloride from MeOH/Et₂O (**m 187-192°**) or (MeOH/Me₂CO), but after sublimation at ~180°/0.2mm the melting point rises to 202-203°. It has ¹HNMR with δ 2.27 (2H, t), 3.25 (2H, t) and 3.52 (2H, t) in (CD₃)₂SO. [King et al. *Biochemistry* **17** 1499 1978, DOI: 10.1021/bi00601a022.] The **free base** is purified by vacuum distillation (**b 71-72°/6mm**), has IR (film) with ν_{max} at 1700 (C=N)cm⁻¹ and ¹HNMR (CDCl₃) with δ at 3.58 (2H, t) and 2.10-2.8 (4H, m). The **free base** is stable on storage but slowly hydrolyses in aqueous solutions with half-lives at 25° of 390 hours at pH 9.1, 210 hours at pH 10 and 18 hours at pH 11. It is a useful reagent for thiolating primary amines. [Traut et al. *Biochemistry* **12** 3266 1973, DOI: 10.1021/bi00741a019; Jue et al. *Biochemistry* **17** 5399 1978, DOI: 10.1021/bi00618a013; Alagon & King *Biochemistry* **19** 4341 1980, DOI: 10.1021/bi00559a030; *Beilstein* **17/9** V 12.]

Indanthrene (N,N'-dihydro-1,2,1',2'-anthraquinonazine) [81-77-6] $C_{28}H_{16}N_2O_4$, **M 442.4**, **m 470-500°(dec)**. Crystallise indanthrene repeatedly from 1,2,4-trichlorobenzene to give a blue powder. It is soluble in alkali and concentrated H₂SO₄. It has been used for dyeing cotton. [Weinstein & Merritt *J Am Chem Soc* **81** 3759 1959, DOI: 10.1021/ja01523a067; *Beilstein* **24** II 317, **24** III/IV 2193.]

Indazole (1,2-benzopyrazole, 1,2-benzodiazole) [271-44-3] $C_7H_6N_2$, **M 118.1**, **m 145-148°**, **147°**, **150°**, **b 270°/743mm**, **pK₁²⁰ 1.32**, **pK₂²⁰ 13.80 (acidic NH)**. Crystallise indazole from water, sublime it *in vacuo*, then recrystallise it from petroleum ether (**b 60-80°**). The **picrate** crystallises from Et₂O with **m 136°**. [Ainsworth *Org Synth Coll Vol* **4** 536 1963, DOI: 10.15227/orgsyn.039.0027; *Beilstein* **23** III/IV 1055, **23/6** V 156.]

Indigo (indigo blue) [482-89-3] $C_{16}H_{10}N_2O_2$, **M 262.3**, **sublimes at ~300°**, **m 390°(dec)**, **CI 73000**, and **halogen-substituted indigo dyes**. First reduce indigo in alkaline solution with sodium hydrosulfite, and filter. The filtrate is then oxidised by air, and the resulting precipitate is filtered off, dried at 65-70°, ground to a fine powder, and extracted with CHCl₃ in a Soxhlet extractor. Evaporation of the CHCl₃ extract gives the purified dye. It has λ_{max} at 602nm. [Brode et al. *J Am Chem Soc* **76** 1034 1954, DOI: 10.1021/ja01633a033; spectral characteristics are listed, *Beilstein* **24** II 233, **24** III/IV 1791.]

Indole [120-72-9] C_8H_7N , **M 117.2**, **m 51-54°**, **52°**, **54.5°**, **b 124°/5mm**, **253-254°/760mm**, **pK₁²⁵ -2.47 (H₀ scale)**, **pK₂²⁵ 16.97 (acidic NH)**. Crystallise indole from *benzene, hexane, petroleum ether, water or EtOH/water (1:10). It can be further purified by sublimation in a vacuum or by zone melting. The **picrate** forms orange crystals from EtOH and has **m 175°**. [*Beilstein* **20** II 196, **20** III/IV 3176, **20/7** V 5.]

Indole-3-acetic acid (heteroauxin) [87-51-4] $C_{10}H_9NO_2$, **M 175.2**, **m 165-169°**, **167-169°(dec)**, **pK₁²⁵ -6.13 (aqueous H₂SO₄)**, **pK₂²⁵ 4.54 (CO₂H)**. Recrystallise heteroauxin from EtOH/water [James & Ware *J Phys Chem* **89** 5450 1985, DOI: 10.1021/j100271a028]. [*Beilstein* **22** III/IV 65.] Alternatively, recrystallise 30g of the acid with 10g of charcoal in 1L of hot water, filter and cool when 22g of colourless acid separate. Dry it and store it in a dark bottle away from direct sunlight [Johnson & Crosby *Org Synth Coll Vol* **5** 654 1973, DOI: 10.15227/orgsyn.044.0064]. The **picrate** has **m 178-180°**. [*Beilstein* **22** H 66, **22** I 508, **22** II 50, **22** III/IV 1088.] It is a plant growth substance.

3-Indoleacetonitrile [771-51-7] $C_{10}H_8N_2$, **M 156.2**, **m 33-36°**, **36-38°**, **b 157°/0.2mm**, **158-160°/0.1mm**, **viscous oil n_D²⁰ 1.6097**. Distil the nitrile at very high vacuum, and the viscous distillate crystallises on standing after a few days; the **picrate** has **m 127-128°** (from EtOH) [Coker et al. *J Org Chem* **27** 850 1962, DOI: 10.1021/jo01050a038; Thesing & Schülde *Chem Ber* **85** 324 1952, DOI: 10.1002/cber.19520850409]. Store it away from light. The **N-acetate** has **m 118°** (from MeOH) and $R_F = 0.8$, on Silica Gel F₂₅₄ in CHCl₂/MeOH

19:1 [Buzas et al. *Synthesis* 129 1977, DOI: 10.1055/s-1977-24296]. [Beilstein 22 III/IV 1097, 22/3 V 74.] It is a plant growth substance.

Indole-2-carboxaldehyde [19005-93-7] C_9H_7NO , M 145.2, m 138°, 138-142°, 140-141°, 138-142°, $pK_{Est} <1$. Recrystallise the indole from aqueous MeOH (m 141-142°), Et₂O (m 138°) or sublime it *in vacuo* (m 138°). The *N,N*-dimethylhydrazone [127280-17-5] has m 105-106° (from hexane/*C₆H₆). The *thiosemicarbazone* has m 229° (dec) (from 50% aqueous EtOH) [Doyle et al. *J Chem Soc* 2853 1956, DOI: 10.1039/JR9560002853]. The *2,4-dinitrophenylhydrazone* has m 315-320° (dec) (from pyridine/MeOH). [Suzuki et al. *Chem Pharm Bull Jpn* 39 2170 1991, DOI: org/10.1248/cpb.39.2170; Beilstein 21 III/IV 3754.]

Indole-3-propionic acid (IPA) [830-96-6] $C_{11}H_{11}NO_2$, M 189.2, m ~133°, 134-135°, pK^{25} 4.95. Recrystallise it from EtOH/water [James & Ware *J Phys Chem* 89 5450 1985, DOI: 10.1021/j100271a028]. The *picrate* has m 143-144°, and the *methyl ester* crystallises from *C₆H₆ or MeOH with m 81-83°. [Beilstein 22 III/IV 1113, 22/3 V 114.]

(±)-Indoline-2-carboxylic acid (2,3-dihydro-1H-indole-2-carboxylic acid) [78348-24-0] $C_9H_9NO_2$, M 163.2, m 168°(dec), $pK_{Est(1)} \sim 2.1$, $pK_{Est(2)} \sim 3.8$. Dissolve the acid in hot EtOH, add excess of dry Et₂O and cool to yield colourless plates that decompose in the range of 120-150°. The *amide* (m 208-209°) crystallises as colourless plates which sublime at 150°/1.0mm and has ν_{max} at 1625cm⁻¹ (paraffin mull) [Hudson & Robertson *Aust J Chem* 20 1935 1967, DOI: 10.1071/CH9671935]. [Beilstein 22/2 V 421.] See Chapter 5, Catalysts-Part 1 for optical isomers.

Indolizine [pyrrocoline, pyrrolo(1,2-*a*)pyridine] [274-40-8] C_8H_7N , M 117.1, m 73-74°, 75°, pK^{20} 3.94 (C-1 and very little C-3 2protonation). Purify indolizine through an alumina column in *C₆H₆ and elute with *C₆H₆ (toluene could be used instead). The eluate contained in the fluorescent band (using UV light λ 365nm) is collected, evaporated and the crystalline residue is sublimed twice at 40-50°/0.2-0.5mm. The colourless crystals have a 'naphthalene' odour, darken on standing and should be stored in dark sealed containers. If the original sample is dark in colour then it should be covered with water and steam distilled. The colourless crystals in the distillate are collected and dried between filter paper and sublimed. It protonates mostly on C3 in aqueous acid. It should give one fluorescent spot on paper chromatography (Whatman 1) in 3% aqueous ammonia and in *n*-BuOH/AcOH/H₂O (4:1:1). The *picrate* has m 101° from EtOH. [Armarego *J Chem Soc* 4226 1964, DOI: 10.1039/JR9640004226; Armarego *J Chem Soc (B)* 191 1966, DOI: 10.1039/J29660000191; Scholtz *Chem Be* 45 734 1912, DOI: 10.1002/cber.191204501108; Beilstein 20 II 200, 20 III/IV 3195.]

trans-Indol-3-ylacrylic acid [1204-06-4] $C_{11}H_9NO_2$, M 187.2, m 190-195°(dec), 195°(dec), 196°(dec), 195-196°(dec), $pK_{Est} \sim 4.2$. Recrystallise the acid from AcOH, H₂O or EtOAc/cyclohexane. UV in MeOH has λ_{max} at 225, 274 and 325nm. [Shaw et al. *J Org Chem* 23 1171 1958, DOI: 10.1021/jo01102a025; constitution: Rappe *Acta Chem Scand* 18 818 1964, DOI: 10.3891/acta.chem.scand.18-0818a; Moffatt *J Chem Soc* 1432 1442 1957, DOI: 10.1039/JR9570001432; Kimming et al. *Hoppe Seyler's Z Physiol Chem* 371 234 1958, Beilstein 22 V 249.]

3-Indolylbutyric acid (IBA) [133-32-4] $C_{12}H_{13}NO_2$, M 203.2, m 120-123°, 123-125°, 124°, pK^{25} 4.84. Recrystallise the acid from H₂O. It is soluble in EtOH, Et₂O and Me₂CO but insoluble in CHCl₃. [Bowman & Islip *Chem Ind London* 154 1971, Jackson & Manske *J Am Chem Soc* 52 5029 1930, DOI: 10.1021/ja01375a056; Albaum & Kaiser *Am J Bot* 24 420 1937, <http://www.jstor.org.virtual.anu.edu.au/stable/2436425>.] It has also been recrystallised from EtOH/water [James & Ware *J Phys Chem* 89 5450 1985, DOI: 10.1021/j100271a028]. Its UV has λ_{max} at 278 and 320nm in isoPrOH [Elvidge *Quart J Pharm Pharmacol* 13 219 1940]. The *methyl ester* has m 73-74° (from *C₆H₆/petroleum ether) and b 230°/6mm [Bullock & Hand *J Am Chem Soc* 78 5854 1951, DOI: 10.1021/ja01603a040]. It stimulates root growth in plant clippings. [Beilstein 22 III/IV 1128, 22/3 V 140.]

3-Indolylpyruvic acid [392-12-1] $C_{11}H_9NO_3$, M 203.2, m ~210°(dec), 208-210°(dec), 215°(dec), 219°(dec), $pK_{Est} \sim 2.4$. Recrystallise the acid from Me₂CO/*C₆H₆, EtOAc/CHCl₃, Me₂CO/AcOH (crystals have 1 molecule of AcOH) and dioxane/*C₆H₆ (with 0.5 molecule of dioxane) [Shaw et al. *J Org Chem* 23 1171 1958,

DOI: 10.1021/jo01102a025; Kaper & Veldstra *Biochim Biophys Acta* **30** 401 1958, DOI:10.1016/0006-3002(58)90065-9]. The *ethyl ester* has **m 133°** (from Et₂O), and its **2,4-dinitro-phenylhydrazone** has **m 255°** (from Me₂CO). [Baker *J Chem Soc* 461 1946, DOI: 10.1039/JR9460000461.] The *oxime* has **m 157°(dec)**, from EtOAc/Et₂O and **pK²⁰ 3.40** [Ahmad & Spenser *Can J Chem* **39** 1340 1961, DOI: 10.1139/v61-169]. [Beilstein **22** II 250, **22** III/IV 3080, **22/6** V 324.]

7-Iodoindole [89976-15-8] **C₈H₆IN**, **M 243.0**, **m 52-56°**, **pK_{Est} <1**. Purify 7-iodoindole by chromatography through a silica gel column and eluting with CH₂Cl₂/hexane (1:3, v/v) followed by recrystallisation from hexane (colourless plates, **m 55-56°**). [Somei et al. *Chem Pharm Bull Jpn* **35** 3146 1987, DOI:org/10.1248/cpb.35.3146; Somei & Saida *Heterocycle* **23** 3113 1985, DOI: 10.3987/R-1985-12-3113.]

Iodinine (1,6-dihydroxyphenazine-5,10-dioxide) [68-81-5] **C₁₂H₈N₂O₄**, **M 244.1**, **m 236°(dec)**, **pK²⁵ 12.5**. Purify iodinine through a column of silica gel and elute with Me₂CO/CHCl₃, then recrystallise it from CHCl₃ to give purple crystals with a copper-coloured luster. Gives coloured solutions with acids and with alkali. [Clemo & Dalglish *J Chem Soc* 1481 1950, DOI: 10.1039/JR9500001481; Gerber & Lechevalier *Biochemistry* **3** 598 1964, DOI: 10.1021/bi00892a022; Beilstein **23** III/IV 3227.]

Iodonitrotetrazolium chloride (2[4-iodophenyl]-3-[4-nitrophenyl]-5-phenyl-2H-tetrazolium chloride) [146-68-9] **C₁₉H₁₃ClIN₅O₂**, **M 505.7**, **m 229°(dec)**, **240°(dec)**, **~245°(dec)**. Recrystallise the chloride from H₂O, aqueous EtOH or EtOH/Et₂O. Alternatively, dissolve it in the minimum volume of EtOH and add Et₂O; or dissolve it in hot H₂O (charcoal), filter and precipitate it by adding conc HCl. Filter the solid off and dry it at 100°. Its solubility in H₂O at 25° is 0.5%, and in hot MeOH/H₂O (1:1) it is 5%. [Fox & Atkinson *J Am Chem Soc* **72** 3629 1950, DOI: 10.1021/ja01164a086; Beilstein **26** III/IV 1776.]

Iodonitrotetrazolium violet-Formazan [7781-49-9] **C₁₉H₁₄IN₅O₂**, **M 471.3**, **m 185-186°**. Dissolve it in boiling dioxane (20g in 300ml), add H₂O (100ml) slowly, cool, filter and dry it *in vacuo* at 100°. Its solubility in CHCl₃ is ~1%. [UV: Fox & Atkinson *J Am Chem Soc* **72** 3629 1950, DOI: 10.1021/ja01164a086; Beilstein **16** III/IV 1776, **26** III/IV 1776.]

Isatin (indole-2,3-dione) [91-56-5] **C₈H₅NO₂**, **M 147.1**, **m 193-195°**, **201-203°**, **205°**, **pK²⁵ >12 (acidic NH)**. Crystallise isatin from amyl alcohol and sublime it at 180°/1mm. In aqueous NaOH the ring opens to yield sodium *o*-aminobenzoylformate. Used as a chromatographic spray reagent for amino acids. [Beilstein **21** II 327, 567, **21** IV 58, **21/10** V 221.]

Isatoic anhydride (3,1-benzoxazin-2,4[1-H]-dione) [118-48-9] **C₈H₅NO₃**, **M 163.1**, **m 235-240°**, **240-243°**, **243°**, **243-245°**. Recrystallise it from EtOH or 95% EtOH (30ml/g) or dioxane (10ml/g) and dry it in a vacuum. [Wagner & Fegley *Org Synth Coll Vol* **3** 488 1955, DOI: 10.15227/orgsyn.027.0045; Ben-Ishai & Katchalski *J Am Chem Soc* **74** 3688 1952, DOI: 10.1021/ja01134a501; UV: Zentmyer & Wagner *J Org Chem* **14** 967 1949, DOI: 10.1021/jo01158a006; Beilstein **27** II 299, **27** III/IV 3330.]

3-Isobutyl-1-methylxanthine (3-isobutyl-1-methylpurine-2,6-dione) [28822-58-4] **C₁₀H₁₄N₄O₂**, **M 222.3**, **m 199-210°**, **200-201°**, **202-203°**, **pK_{Est} ~ 6.7 (acidic NH)**. Recrystallise it from aqueous EtOH. It is a non-specific inhibitor of cAMP and cGMP phosphodiesterases. [Beilstein **26** III/IV 2350.]

Isonicotinic acid (pyridine-4-carboxylic acid) [55-22-1] **C₆H₅NO₂**, **M 123.1**, **m >300°**, **320°**, **323-325°(dec)**, **pK₁²⁵ 1.70**, **pK₂²⁵ 4.89**. Crystallise the acid repeatedly from water and dry it under vacuum at 110° or sublime it at 260°/15mm (**m 319°**). [Beilstein **22** III/IV 518, **22/2** V 188.] **Isonicotinamide (pyridine-4-carboxylic acid amide)** [1453-82-3] **C₆H₆N₂O**, **M 122.1**, has **m 155-157°**, **155.5-156°**, **pK₁²⁰ -1.0 (protonation of CONH₂)**, **pK₂²⁵ 3.61**, **pK₃²⁵ 11.47 (acidic CONH₂)**. Recrystallise isonicotinamide from hot water or isopropanol (**158.5-159°**), and dry it in a vacuum at 100°. The *picrate* crystallises from aqueous EtOH or H₂O and has **m 217-218° (214-215°)**. [Beilstein **22** III/IV 527, **22/2** V 195.] **Isonicotinic acid hydrazide (isoniazide)** [54-85-3] **C₆H₇N₃O**, **M 137.1**, has **m 171-173°**, **172°**, **pK₁ 1.75 (NHNH₂)**, **pK₂ 3.57 (=N-)**, **pK₃ 10.75 (-NH)**. Crystallise isoniazide from 95% EtOH and dry it in a vacuum. [Beilstein **22** III/IV 545, **22/2** V 219.] It is an antimicrobial which inhibits mycolic acid biosynthesis.

1-Isonicotinyl-2-isopropylhydrazide (Iproniazid) [54-92-2] $C_9H_{13}N_3O$, M 179.2, m 112.5-113.5°, 114-115°, $pK_{Est} \sim 3.5$. Crystallise it from *benzene or *benzene/petroleum ether and dry it in a vacuum. It is soluble in H_2O and EtOH. [Fox & Gibas *J Org Chem* 18 994 1953, DOI: 10.1021/jo50014a014.] The *dihydrochloride* has m 227-228° (from EtOH). [Beilstein 22 III/IV 551.] It is an antidepressant.

1-Isonicotinyl-2-isopropylhydrazide phosphate (Iproniazid phosphate, Marsilide) [305-33-9] $C_9H_{13}N_3O \cdot H_3PO_4$, M 277.2, has m 175-184°, 178-179°, 180-184°, $pK_{Est} \sim 3.5$ (free base). Crystallise it from H_2O and Me_2CO . The *free base* (see above) has m 113-114° from * C_6H_6 /petroleum ether and is soluble in H_2O and EtOH. [Fox & Gibas *J Org Chem* 18 994 1953, DOI: 10.1021/jo50014a014; Beilstein 22 III/IV 551.] It is an antidepressant.

1-Isonicotinyl-2-salicylidenehydrazide (Salinazide) [495-84-1] $C_{13}H_{11}N_3O_2$, M 241.2, has m 232-233° (also 251°). Crystallise it from EtOH (m 265-266° or 244-245°), aqueous EtOH or MeOH (252-253°) and dry it in a vacuum at 100°. [Beilstein 22 III/IV 584.] Tuberculostatic.

5-Isonitrosobarbituric acid (violuric acid) [87-39-8, 1*H*₂O 26851-19-9] $C_4H_3N_3O_4$, M 175.1, m 221-223°, 240-241°(dec), 245-250°(dec, mono hydrate), pK_1 4.41, pK_2 9.66 (10.1). Crystallise violuric acid from water or EtOH. **1,1-Dimethylvioluric acid**, m 144-147° has pK^{25} 4.72 [Taylor & Robinson *Talanta* 8 518 1961, DOI: 10.1016/0039-9140(63)80055-7]. [Beilstein 24 III/IV 2142.] It chelates with metals.

N-Isopropylcarbazole [1484-09-9] $C_{15}H_{15}N$, M 209.3, m 120°. Crystallise it from isopropanol. It sublimes under vacuum. It was also purified by zone refining. The *picrate* has m 143° after recrystallisation from EtOH. [Beilstein 20 I 164.]

Isoquinoline [119-65-3] C_9H_7N , M 129.2, m 24°, 25.5-26°, b 120°/18mm, 242-243°/atm, 243.25°/760mm, d₄²⁰ 1.0986, n_D²⁰ 1.6148, pK^{25} 5.40. Dry isoquinoline with Linde type 5A molecular sieves or Na_2SO_4 and fractionally distil at reduced pressure. *Alternatively*, it can be refluxed with, and distilled from, BaO. It is also purified by fractional crystallisation from the melt and distilled from zinc dust. It forms a *phosphate* (m 135°) and a *picrate* (m 223°), which are purified by crystallisation, and the free base can be recovered and distilled. [Packer et al. *J Am Chem Soc* 80 905 1958, DOI: 10.1021/ja01537a039.] The procedure for purification *via* the picrate comprises the addition of quinoline to picric acid dissolved in the minimum volume of 95% EtOH to yield yellow crystals which are washed with EtOH and air dried before recrystallising from acetonitrile. The crystals are dissolved in dimethyl sulfoxide (previously dried over 4A molecular sieves) and passed through a basic alumina column, on which picric acid is adsorbed. The *free base* in the effluent is extracted with *n*-pentane and distilled under vacuum. Traces of solvent from small quantities are removed by vapour phase chromatography. The *hydrochloride* crystallises from EtOH with m 193°. [Moomaw & Anton *J Phys Chem* 80 2243 1976, DOI: 10.1021/j100561a022; Beilstein 20 II 236, 20 III/IV 3410, 20/7 V 333.]

Isoxanthopterin (2-amino-4,7-dihydroxypteridine) [529-69-1] $C_6H_5N_5O_2$, M 179.1, m>300°, pK_1^{20} -0.5 (basic), pK_2^{20} 7.34 (acidic), pK_3^{20} 10.06 (acidic). Purify it by repeated precipitation from alkaline solution with acid (preferably AcOH or formic acid), filter, wash well with H_2O , then EtOH and dry at 100°. The purity is checked by paper chromatography [R_F 0.15 (*n*-BuOH/AcOH/ H_2O , 4:1:1); 0.33 (3% aqueous NH_4OH)]. [Goto et al. *Arch Biochem Biophys* 111 8 1965, DOI: 10.1016/0003-9861(65)90316-4.] [For biochemistry see Blakley *Biochemistry of Folic Acid and Related Pteridines* North Holland Publ Co, Amsterdam 1969.] [Beilstein 26 III/IV 3999.]

Janus Green B (3-dimethylamino-7-[4-dimethylaminoazo]-5-phenylphenazonium chloride) [2869-83-2] $C_{30}H_{31}ClN_6$, M 511.1, m >200°, CI 11050. The dye dissolves in H_2O to give a bluish violet solution which becomes colourless when made 10M in NaOH. It dissolves in EtOH to give a blue-violet colour, filter from insoluble material, then add dry Et_2O whereby the dye separates out leaving a small amount of blue colour in solution. Filter off the solid and dry it in a vacuum. Store it in a dark bottle. [Colour Index Vol 4, 3rd edn, 4015 1971.] It is a biological stain.

Jervine (3 β ,23 β -17,23-epoxy-2-hydroxyvertraman-11-one, a steroidal alkaloid) [469-59-0] $C_{27}H_{39}NO_3$, M 425.6, m 243-245°, 247-248°, [α]_D²⁰ -150 (in EtOH), $pK_{Est} \sim 9.4$. Crystallise Jervine from MeOH/ H_2O or Me_2O . The *hydrochloride* has m 300-302° (from MeOH/ Et_2O), and the *picrate* has m 262.5° (from aqueous

MeOH). [Kutney et al. *Can J Chem* **53** 1796 1975, DOI: 10.1139/v75-252; *Beilstein* **27** III/IV 3590.] It is **teratogenic**.

Julolidine (2,3,6,7-tetrahydro-1*H*,5*H*-benzo[*ij*]quinolizidine) [479-59-4] $C_{12}H_{15}N$, **M 173.3**, **m 34-36°, 40°, b 105-110°/1mm, 155-156°/17mm, 280°(dec)**, **pK_{Est} ~7.0**. Purify julolidine by dissolving it in dilute HCl, steam is bubbled through the solution and the residual acidic solution is basified with 10N NaOH, extracted with Et₂O, the extract is washed with H₂O, dried (NaOH pellets), filtered, evaporated and distilled *in vacuo*. The distillate crystallises on cooling (**m 39-40°**). It develops a red colour on standing in contact with air for several days. The colour can be removed by distilling or dissolving in 2-3 parts of hexane, adding charcoal, filtering and cooling in an Me₂CO/Dry-ice bath when **julolidine** crystallises out (85-90% yield **m 39-40°**). The **hydrobromide** [83646-41-7] has **m 218° (239-242°)**, the **picrate** has **m 174°(165°)** and the **methiodide** crystallises from MeOH, with **m 186°** [Glass & Weissberger *Org Synth Coll Vol* **3** 504 1955, DOI: 10.15227/orgsyn.026.0040; Smith & Yu *J Org Chem* **17** 1281 1952, DOI: 10.1021/jo50009a015; *Beilstein* **20** H 332, **20** I 133, **20** II 214, **20** III/IV 3281.] **Highly TOXIC**.

Kainic acid H₂O (2*S*,3*S*,4*S*-2-carboxy-4-isoprenyl-3-pyrrolidine- acetic acid) [487-79-6] $C_{10}H_{15}NO_4$, **M 231.4**, **m 235-245°(dec), 251°(dec)**, **[α]_D²⁰ -14.6 (c 1.46, H₂O)**, **pK₁ 2.09, pK₂ 4.58, pK₃ 10.21**. Purify the acid by adsorbing on to a strongly acidic ion-exchange resin (Merck), elute the diacid with aqueous M NaOH, the eluate is evaporated, H₂O is added, and filtered through a weakly acidic ion-exchange resin (Merck). The filtrate is then evaporated and recrystallised from EtOH. Its solubility is 0.1g in 1ml of 0.5N HCl. (**±**)-**α-Kainic acid** is recrystallised from H₂O with **m 230-260°**. Its UV (MeOH) has λ_{max} at 219 (log ε 3.9); the ¹HNMR (CCl₄, 100MHz, Me₄Si standard) has δ at 1.64 (s 1H), 1.70 (s 3H), 3.24 (d *J* = 7.5, 2H), 3.3-4.2 (1H), 3.70 (s 3H), 3.83 (s 3H), 4.35 (dd *J* = 7.5, 14.5, 1H), 5.21 (t *J* = 7.5, 1H), 7.26 (t *J* = 7.5, 1H). [Oppolzer & Andres *Helv Chim Acta* **62** 2282 1979, DOI: 10.1002/hlca.19790620724; *Beilstein* **22** III/IV 1523.] It is a useful neurobiology tool, and has anthelmintic properties.

Ketanserine [3(4-*p*-fluorobenzoyl)piperidinyl-*N*-ethyl)quinazolin-2,4-dione] [74050-98-9] $C_{22}H_{22}FN_3O_3$, **M 395.4**, **m 227-235°, pK₂₅ 7.5**. Its solubility is 0.001% in H₂O, 0.038% in EtOH and 2.34% in Me₂NCHO. It has been purified by recrystallisation from 4-methyl-3-pentanone [Peeters et al. *Cryst Structure Commun* **11** 375 1982, Kacprowicz et al. *J Chromatogr* **272** 417 1983, Davies et al. *J Chromatogr* **275** 232 1983]. The **Tartrate salt** [83846-83-7] $C_{22}H_{22}FN_3O_3 \cdot C_4H_6O_6$, **M 545.5**, is the antihypertensive drug **Serepress**.

Khellin (4,9-dimethoxy-7-methyl-5-oxofuro[3,2-*g*]-1,2-chromene) [82-02-0] **M 260.3**, **m 154-155°, b 180-200°/0.05mm**. Crystallise khellin from H₂O, MeOH, petroleum ether or Et₂O. The **hydrochloride** has **m 98°(dec)** (from EtOH/HCl). [*Beilstein* **19** II 236, **19** III/IV 2816, **19/6** V 320.]

Kojic acid [(5-hydroxy-2-hydroxymethyl)-4*H*-pyran-4-one] [501-30-4] $C_6H_6O_4$, **M 142.1**, **m 152°, 152-155°, 154-155°, pK₁²⁵ -1.38, pK₂²⁵ 7.66**. Crystallise the acid (yellow solid) from MeOH (charcoal) by adding Et₂O, or from H₂O (solubility is 0.9w/v% at boiling). It sublimes at 150-200°/0.1torr. It is a tyrosinase inhibitor. [*Beilstein* **18** II 57, **18** III/IV 1145, **18/2** V 516.]

Kynurenic acid (4-hydroxyquinoline-2-carboxylic acid) [492-27-3] $C_{10}H_7NO_3$, **M 189.1**, **m 282-283°, 285°, pK_{Est(1)} ~2, pK_{Est(2)} ~10**. Crystallise the acid from absolute EtOH. The **methyl ester** crystallises from MeOH with **m 224-226°**. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p2723 1961, *Beilstein* **22** II 174, **22** III/IV 2245, **22/6** V 280.]

L-Kynurenine [2-amino-3-(2-aminobenzoyl)-4-propionic acid] [343-65-7] $C_{10}H_{12}N_2O_3$, **M 208.2**, **m 190°(dec), 210°(dec)**, **[α]_D²⁰ -30 (c 0.4, H₂O)**, **pK_{Est(1)} ~2.3, pK_{Est(2)} ~3.5, pK_{Est(3)} ~9.2**. Crystallise it from H₂O or aqueous AcOH. The **picrate** has **m 188.5-189°(dec)** after crystallisation from H₂O. (**±**)-**Kynurenine** has **m 218° (~235° dec, also reported)**. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p2726 1961, *Beilstein* **14** III 1657, **14** IV 2562.] **L-Kynurenine sulfate** [16055-80-4] $C_{10}H_{12}N_2O_3 \cdot H_2SO_4 \cdot H_2O$, **M 306.3**, has **m 194°, monohydrate m 178°, [α]_D²⁵ +9.6 (H₂O)**, and crystallises from water by addition of EtOH. The (**±**)-**sulfate** has **m 173°** (darkening at 166° and decomposing at 194° is also reported). [*Beilstein* **14** IV 2562.]

dl- and l-Laudanosine {1-[(3,4-dimethoxyphenyl)methyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-2-methylisoquinoline} [(±) 1699-51-0, (-) 2688-77-9] **M 357.4, m 114-115°, 118°**. Crystallise these from EtOH. The (±)-*picrate* crystallises from EtOH with **m 177-178°**. The (-)-isomer has **m 83-85°** and $[\alpha]_D^{24}$ -110° (c 0.3, EtOH). The *hydrobromide tetrahydrate* [303136-74-5] **M 436.3** has **m 232-234°**. [Frydman et al. *Tetrahedron* **4** 342 1958, Elliott *J Heterocycl Chem* **9** 853 1972, *Beilstein* **21** II 183, **21** III/IV 2704.]

Luminol (5-aminophthalazin-1,4-dione) [521-31-3] **M 177.2, m 329-332°, pK₁ 3.37, pK₂ 6.35**. Dissolve luminol in KOH solution, treat with Norit (charcoal), filter and precipitate it with conc HCl. [Hardy et al. *Talanta* **24** 297 1977.] Store it in the dark in an inert atmosphere, because its structure changes during its luminescence. It has been recrystallised from 0.1M KOH [Merenyi et al. *J Am Chem Soc* **108** 7716 1986, DOI: 10.1021/ja00284a040]. [*Beilstein* **25** II 389, **25** III/IV 4192.]

dl-Lupinine (1-hydroxymethyloctahydroquinolizine, 1-hydroxymethylquinolizidine) [10248-30-3] **M 169.3, m 57-58°, 59°, 107°/1mm, pK_{Est} ~7.0**. It crystallises from Me₂CO, pentane and petroleum ether (b. 40-60°) and can be sublimed or distilled in high vacuum. The *picrate* has **m 127°**(from Et₂O), the *picrolonate* has **m 203-204°**, and the *methiodide* has **m 203°**(dec, from EtOH). [Clemo et al. *J Chem Soc* 965 1937, DOI: 10.1039/JR9370000965, Boekelheide & Lodge *J Am Chem Soc* **73** 3681 1951, DOI: 10.1021/ja01152a033; *Beilstein* **21** II 28, **21** III/IV 291.]

Lutidine (dimethylpyridine mixture). For the preparation of pure 2,3-, 2,4- and 2,5-lutidine from commercial '2,4- and 2,5-lutidine' see Coulson et al. *J Chem Soc* 1934 1959, DOI: 10.1039/JR9590001934; and Kyte et al. *J Chem Soc* 4454 1960, DOI: 10.1039/JR9600004454.

2,3-Lutidine (2,3-dimethylpyridine) [583-61-9] **C₇H₉N, M 107.2, m -14.8°, -15°, b 160.6°/760mm, 162-163°/atm, d₄²⁰ 0.9464, n_D²⁰ 1.50857, pK²⁵ 6.57**. Steam distil it from a solution containing about 1.2 equivalents of 20% H₂SO₄, until ca 10% of the base has been carried over with the non-basic impurities. The acidic solution is then made alkaline, and the base is separated, dried over NaOH or BaO, and fractionally distilled. The distilled lutidine is converted to its *urea complex* by stirring 100g with 40g of urea in 75ml of H₂O, cooling to 5°, filtering at the pump, and washing with 75ml of H₂O. The complex, dissolved in 300ml of H₂O, is steam distilled until the distillate gives no turbidity with a little solid NaOH. The distillate is then treated with excess solid NaOH, and the upper layer is removed: the aqueous layer is then extracted with Et₂O. The combined organic solutions are dried (K₂CO₃), and distilled through a short column. Final purification is by fractional crystallisation using partial freezing. The *picrate* crystallises from EtOH with **m 187-188°**. [Kyte et al. *J Chem Soc* 4454 1960, DOI: 10.1039/JR9600004454; **20** H 243, **20** II 159, **20** III/IV 2765, **20/6** V 15.]

2,4-Lutidine (2,4-dimethylpyridine) [108-47-4] has **m -60°, b 157.8°/760mm, 159°/atm, d₄²⁰ 0.9305, n_D²⁰ 1.50087, n_D²⁵ 1.4985, pK²⁵ 6.77**. Purify as described in *Purification of Laboratory Chemicals* 7th edn, 2013; ISBN: 978012382161-4. [see Kyte et al. *J Chem Soc* 4454 1960, DOI: 10.1039/JR9600004454.] *Alternative* purifications are via the *picrate* **m 183-184°** (from H₂O). [Clarke & Rothwell *J Chem Soc* 1885 1960, DOI: 10.1039/JR9600001885, or *hydrobromide* [Warnhoff *J Org Chem* **27** 4587 1962, DOI: 10.1021/jo01059a109]. The latter is precipitated from a solution of lutidine in *benzene by passing dry HBr gas, recrystallised from CHCl₃/MeCOEt, then decomposed with NaOH, and the free base is extracted into Et₂O, dried, evaporated and the residue is distilled. [*Beilstein* **20** II 180, **20** III/IV 2718, **20/6** V 19.]

2,5-Lutidine (2,5-dimethylpyridine) [589-93-5] has **m -15.3°, b 156.7°/759mm, 157°/atm, d₄²⁰ 0.927, n_D²⁵ 1.4982, pK²⁵ 6.40**. Purify as above, distil with Na then fractionally distil through a Todd column packed with glass helices. The *hydrochloride* has **m 219°**(dec), and the *picrate* has **m 170.5°** (from EtOH or H₂O). [*Beilstein* **20** H 244, **20** II 160, **20** III/IV 2774, **20/6** V 27.]

2,6-Lutidine (2,6-dimethylpyridine) [108-48-5] has **m -5.8°, b 79°/87mm, 144.0°/760mm, d₄²⁰ 0.92257, n_D²⁰ 1.49779, pK²⁵ 6.72**. Likely contaminants include 3- and 4-picoline (similar boiling points) which are removed with BF₃ (4ml for 100ml) or by distillation of commercial material from AlCl₃ (14g per 100ml). 2,6-Lutidine can be dried with KOH (do not use Na, see picolines below) or by refluxing with (and distilling from) BaO or CaH₂, prior to distillation. Also purified via its *picrate*, 2,6-lutidine **m 163-164.5° (166-167°)**, then partitioned between ammonia and CHCl₃/Et₂O. The organic layer, after washing with dilute aqueous KOH, is dried (Na₂SO₄) and fractionally distilled. [Warnhoff *J Org Chem* **27** 4587 1962, DOI: 10.1021/jo01059a109.] *Alternatively*, purify via its *urea complex*, as described above. Other purifications include azeotropic distillation with phenol [Coulson et al. *J Appl Chem* **2** 71 1952, DOI: 10.1002/jctb.5010020205], crystallisation by partial freezing, and vapour-phase chromatography using a 180-cm column of polyethylene glycol-400

(Shell, 5%) on Embacel (May and Baker) at 100°, with argon as carrier gas [Bamford & Block *J Chem Soc* 4989 1961, DOI: 10.1039/JR9610004989]. The **hydrochloride** has **m 235-237°, 239°** (from EtOH). [Beilstein 20 II 160, 20 III/IV 2776, 20/6 V 32.]

3,5-Lutidine (2,5-dimethylpyridine) [591-22-0] has **m -9°, -6.3°, b 169-170°/atm, 172.0°/767mm, d₄²⁰ 0.9419, n_D²⁰ 1.50613, n_D²⁵ 1.5035, pK²⁵ 6.15**. Dry 3,5-lutidine with CaH₂ and fractionally distil it through a Todd column packed with glass helices. Methods used for other isomers apply here too. The **hydrochloride** has **m 229°** (sublimes at 190-231°), and the **picrate** has **m 242-243°(dec, from H₂O), 249-250°(dec, from AcOH)**. [Beilstein 20 II 161, 20 III/IV 2788, 20/6 V 60.]

Lycorine [476-28-8] C₁₆H₁₇NO₄, **M 287.3, m 275-280°(dec), b 175-185°/0.002mm. [α]_D²⁰ -91 (c 0.16, EtOH), [α]_D¹⁶ -127 (c 0.16, absEtOH), pK²⁵ 6.9 (30% aqueous dimethylformamide)**. It crystallises as orange crystals from MeOH (**m 281-283°**), CHCl₃/EtOH (**m 272-274°**), pyridine or from EtOH (**m 277° dec, also 275-280° dec**). It has been distilled under high vacuum. The **hydrochloride** has **m 288°** (from MeOH/HCl), **m 217° (dec with prior sintering, needles from H₂O)**, [α]_D²⁰ +63 (EtOH), and the **picrate** has **m 196-197°(from EtOH)**. [Cook et al. *J Chem Soc* 4176 1954, DOI: 10.1039/JR9540004176; Martin & Tu *J Org Chem* 46 3763 1981, DOI: 10.1021/jo00331a049; Beilstein 27 II 547, 27 III/IV 6463.] Alkaloid.

(+)-Lysergic acid [82-58-6] **M 268.3, m 240°(dec), [α]_D²⁰ +40° (pyridine), pK₁²⁵ 3.32, pK₂²⁵ 8.66, pK₃²⁰ 8.50, pK₄⁴⁰ 8.27**. It crystallises from water as a *hydrate*. The **methyl ester** crystallises from *C₆H₆ and has **m 168°**; the **amide** [478-94-4] has **m 242°(dec)** (from MeOH) and [α]₅₄₆ +15° (c 0.5, pyridine). The **(-)-hydrochloride** has **m 208-210°(dec, from MeOH)**. [Kornfeld et al. *J Am Chem Soc* 76 5256 1954, Kornfeld et al. *J Am Chem Soc* 78 3087 1956, DOI: 10.1021/ja01594a039; Beilstein 25 III/IV 934.]

Maltol (3-hydroxy-2-methyl-4-pyrone) [118-71-8] C₆H₆O₃, **M 126.1, m 160-164°, 161-162°, 162-162.5°**. It crystallises from CHCl₃, toluene, aqueous 50% EtOH or H₂O, and is volatile in steam. It can be readily sublimed in a vacuum. It forms a Cu²⁺ complex. [Beilstein 18/1 V 114.] It is a food flavouring agent.

Meconic acid (3-hydroxy-γ-pyrone-2,6-dicarboxylic acid) [497-59-6] C₇H₄O₇, **M 200.1, m 100° (loses H₂O), pK₁²⁵ 1.83, pK₂²⁵ 2.3, pK₃²⁰ 10.10**. Crystallise the acid from water (0.25g/ml) and dry it at 100° for 20 minutes to dehydrate the *mono* or *dihydrate*. It decarboxylates above 120° or in boiling H₂O. It is soluble in MeOH (2%), EtOAc (2%) and Me₂CO (1%). The **picrate** has **m 206.5-208.5°(dec, from H₂O)**. [Wibaut & Kleinpools *Recl Trav Chim Pays-Bas* 66 24 1947, DOI: 10.1002/recl.19470660103; Beilstein 18 H 409, 18 I 523, 18 II 367, 18 III/IV 6136.]

Melamine (2,4,6-triamino-1,3,5-triazine) [108-78-1] C₃H₆N₆, **M 126.1, m >300°, 353°, pK²⁵ 5.00**. Crystallise Melamine from water or dilute aqueous NaOH. It sublimes at ~240° on prolonged heating. [Beilstein 26 I 74, 26 II 132, 26 III/IV 1253.]

(±)-Mellein [(±)-3,4-dihydro-8-hydroxy-3-methyl-2-benzopyran-1-one, 8-hydroxy-3-methylisochroman-1-one] [1200-93-7] **M 178.2, m 37-39°, 39°, pK_{Est} ~9.5**. Purify it by recrystallisation from H₂O or aqueous EtOH. It has UV with λ_{max} at 247 and 314nm. [Arakawa et al. *Justus Liebigs Ann Chem* 728 152 1969, DOI: 10.1002/jlac.19697280117; Blair & Newbold *Chem Ind (London)* 93 1955, *J Chem Soc* 2871 1955, DOI: 10.1039/JR9550002871.] The **methyl ether** has **m 66-67°** and UV with λ_{max} at 242nm (ε 7,400) and 305nm (ε 4,600). **R(-)-Mellein** has **m 56°(from aqueous Me₂CO), [α]_D²⁵ -102.5 (c 1, CHCl₃)** and **R(+)-mellein** has **m 56-57°(from hexane), [α]_D²⁵ +102 (c 1, CHCl₃) or [α]_D²⁵ +88 (c 1, MeOH)**. [Beilstein 18 III/IV 188, 18/1 V 274.]

2-Mercaptobenzimidazole [583-39-1] C₇H₆N₂S, **M 150.2, m 302-304°, 312°, pK²⁰ 10.24**. Crystallise it from aqueous EtOH, AcOH or aqueous ammonia. It complexes with many metals. [Brown *J Chem Soc* 1974 1958, DOI: 10.1039/JR9580001974; Beilstein 24 II 65, 24 III/IV 287.]

2-Mercaptobenzothiazole [149-30-4] C₇H₅NS₂, **M 167.2, m 177-181°, 182°, pK²⁵ 7.5 (50% aqueous AcOH)**. Crystallise it repeatedly from 95% EtOH, or purify it by incomplete precipitation by dilute H₂SO₄ from a basic solution, followed by several crystallisations from acetone/H₂O or *benzene. It complexes with Ag, Au, Bi, Cd, Hg, Ir, Pt, and Tl. [Beilstein 27 II 233, 27 III/IV 2709.]

2-Mercaptoimidazole [872-35-5] $\text{C}_3\text{H}_4\text{N}_2\text{S}$, M 100.1, m 221-222°, 226-228°(monohydrate), 228-231°, pK_1^{20} -1.6, pK_2^{20} 11.6. Crystallise 2-mercaptoimidazole from Me_2CO or H_2O . Its UV has λ_{max} at 208 and 252nm (H_2O). [Fox et al. *J Am Chem Soc* **67** 496 1945, DOI: 10.1021/ja01219a506; *Beilstein* **24** II 7, **24** III/IV 61.]

2-Mercapto-1-methylimidazole [60-56-0] $\text{C}_4\text{H}_6\text{N}_2\text{S}$, M 114.2, m 144-147°, 145-147°, 146-148°, pK_1^{20} -2.0, pK_2^{20} 11.9. Crystallise it from EtOH. Its UV has λ_{max} at 251nm (H_2O), 260nm (EtOH) and 267nm (CHCl_3). [Lawson & Morley *J Chem Soc* 1103 1956, DOI: 10.1039/JR9560001103; *Beilstein* **24** H 17, **24** III/IV 61.]

6-Mercaptopurine monohydrate [6112-76-1] $\text{C}_5\text{H}_4\text{N}_4\text{S} \cdot \text{H}_2\text{O}$, M 170.2, m 314-315°(dec), ~315°(dec), 313-315°(dec), pK_1^{20} 0.5, pK_2^{20} 7.77, pK_3^{20} 10.84. Crystallise 6-mercaptopurine from pyridine (30ml/g), wash it with pyridine, then triturate with water (25ml/g) and adjust to pH 5 by adding M HCl. Recrystallise it by heating, then cooling, the solution. Filter off the solid, wash it with water and dry it at 110°. It has also been crystallised from water (charcoal) as yellow crystals of the *monohydrate* which become *anhydrous* on drying at 140°. It has UV with λ_{max} at 230 and 312nm (ϵ 14,000 and 19,600) in 0.1N NaOH; 222 and 327nm (ϵ 9,2400 and 21,300), and 216 and 329nm (ϵ 8,740 and 19,300) in MeOH. It forms a **1:1 complex** with Zn^{2+} , Pb^{2+} , Co^{2+} , and Ni^{2+} in aqueous dioxane. It is an antineoplastic agent which inhibits *de novo* purine synthesis by incorporating thiopurine methyltransferase metabolites into DNA and RNA. [Albert & Brown *J Chem Soc* 2060 1954, DOI: 10.1039/JR9540002060; IR: Brown & Mason *J Chem Soc* 682 1957, DOI: 10.1039/JR9570000682; UV: Fox et al. *J Am Chem Soc* **80** 1669 1958, DOI: 10.1021/ja01540a041; UV: Mason *J Chem Soc* 2071 1954, DOI: 10.1039/JR9540002071; *Beilstein* **26** III/IV 2097.]

8-Mercaptoquinoline (2H₂O, thioxine) [491-33-8] $\text{C}_9\text{H}_7\text{NS} \cdot \text{H}_2\text{O}$, M 179.2, m 58-59°, pK_1^{25} 2.0, pK_2^{25} 8.40. Thioxine readily oxidises in air to give diquinolyl-8,8'-disulfide (which is stable). It is more convenient to make 8-mercaptoquinoline by reduction of the disulfide. [Nakamura & Sekido *Talanta* **17** 515 1970, DOI: 10.1016/0039-9140(63)80055-7.] The *hydrochloride* (see thioxine hydrochloride below) is more stable. [*Beilstein* **21** III/IV 1197, **21/3** V 30.]

3-Methoxycarbonyl-2,5-dihydrothiophen-1,1-dioxide (methyl 3-sulfolene-3-carboxylate) [67488-50-0] $\text{C}_6\text{H}_8\text{O}_4\text{S}$, M 176.2, m 57-58°, 60-62°. If the IR shows OH bands, then dissolve the dioxide in CHCl_3 , wash it with aqueous Na_2CO_3 and H_2O , dry it over MgSO_4 , filter, evaporate and wash the residue with cold Et_2O and dry *in vacuo*. Its ^1H NMR (CDCl_3) has δ at 7.00 (m 1H), 3.98 (bs 4H) and 3.80 (s Me). [McIntosh & Sieler *J Org Chem* **43** 4431 1978, DOI: 10.1021/jo00417a007; *Beilstein* **18/6** V 5.] It has been used as a stable precursor of 2-methoxycarbonyl-1,3-butadiene for cycloaddition reactions [McIntosh & Sieler *J Org Chem* **43** 4431 1978, DOI: 10.1021/jo00417a007].

5-Methoxyindole [1006-94-6] $\text{C}_9\text{H}_9\text{NO}$, M 147.2, m 52-55°, 55°, 57°, b 176-178°/17mm, pK_{Est} ~0. Crystallise 5-methoxyindole from cyclohexane petroleum ether or petroleum ether/ Et_2O . [Saito & Kikugawa *J Heterocycl Chem* **16** 1325 1979, DOI: 10.1002/jhet.5570160707; *Beilstein* **21** III/IV 765, **21/3** V 18.]

5-(p-Methoxyphenyl)-1,2-dithiole-3-thione [42766-10-9] M 240.2, m 111°. Crystallise the thione from EtOAc, BuOAc or EtOH. It sublimes at 90°/0.001mm and complexes with Sb^{3+} , Sb^{5+} , Bi^{3+} , Sn^{4+} , Ag^+ , Au^{3+} and Hg^{2+} . [*Beilstein* **19** III/IV 2538.]

6-Methylaminopurine [443-72-1] $\text{C}_6\text{H}_7\text{N}_5$, M 149.2, m >300°, 312-314° (dec), pK_1^{20} <1, pK_2^{20} 4.15, pK_3^{20} 10.02. The purine is best purified by recrystallising 2g from 50ml of H_2O and 1.2g of charcoal. [UV: Albert & Brown *J Chem Soc* 2060 1954; DOI: 10.1039/JR9540002060; UV: Mason *J Chem Soc* 2071 1954, DOI: 10.1039/JR9540002071; see also Elion et al. *J Am Chem Soc* **74** 411 1952, DOI: 10.1021/ja01122a037.] The *picrate* has m 265°(257°) [Bredereck et al. *Chem Ber* **81** 307 1948, DOI: 10.1002/cber.19480810406]. [*Beilstein* **26** III/IV 3565.]

Methyl 3-aminopyrazine-2-carboxylate [16298-03-6] $\text{C}_6\text{H}_7\text{N}_3\text{O}_2$, M 153.1, m 169-172°, 172°. The ester forms yellow needles from H_2O (100 parts using charcoal). If it contains the *free acid* (see IR), then dissolve it in CH_2Cl_2 , wash it with saturated aqueous Na_2CO_3 , brine, dry over MgSO_4 filter, evaporate and recrystallise the residue. The *free acid* has m 203-204° (dec) [UV: Brown & Mason *J Chem Soc* 3443 1956, DOI: 10.1039/JR9560003443] with pK_1 <1 and pK_2 3.70. The *ammonium salt* has m 232°(dec) (from aqueous

Me₂CO) and the *amide* has **m 239.2°** (from H₂O) [Ellingson et al. *J Am Chem Soc* **67** 1711 1945, DOI: 10.1021/ja01226a028]. [*Beilstein* **25** III/IV 4412.]

9-Methylcarbazole (N-methylcarbazole) [1484-12-4] C₁₃H₁₁N, **M 181.2, m 87°, 89°, 90-92°**. Purify *N*-methylcarbazole by chromatography on silica gel and eluting with CHCl₃/Me₂CO/Et₂O (100:5:1 v/v), or by flash chromatography using petroleum ether, or by zone melting followed by recrystallisation from petroleum ether or EtOH. [Flo & Pindur *Annalen* 509 1987, DOI: 10.1002/jlac.198719870366; Kashima et al. *J Heterocycl Chem* **24** 913 1987, DOI: 10.1002/jhet.5570240405; UV: Armarego in *Physical Methods in Heterocyclic Chemistry* (Ed Katritzky, Academic Press, LCCC No 62-12037, Internet Public Domain) **Vol III** 158 1971; *Beilstein* **20** H 436.]

5-Methylcytosine [4-amino-5-methylpyrimidin-2(1H)-one] [554-01-8] C₅H₇N₃O, **M 125.1, m 270°(dec), pK₁ 4.6, pK₂ 12.4**. Crystallise it from water (solubility is 3.4w/v% at ~20°). The *hydrochloride* has **m 299-301°** (sintering at 280°) (from aqueous HCl/Me₂CO). [Hitchings et al. *J Biol Chem* **177** 357 1949, <http://www.jbc.org/content/177/1/357>; Cohn *J Am Chem Soc* **73** 1539 1951, DOI: 10.1021/ja01148a039; *Beilstein* **25** II 183, **25** III/IV 3727.]

2-Methyl-1,3-dithiane [6007-26-7] C₅H₁₀S₂, **M 134.3, b 53-54°/1.1mm, 56-59°/3mm, 66°/5mm, 79-80°/8-10mm, 85°/12mm, d₄²⁰ 1.121, n_D²⁰ 1.560**. Wash the dithiane with H₂O, 2.5 M aqueous NaOH, H₂O, brine, dry over K₂CO₃ (use toluene as solvent if the volume of reagent is small), filter, evaporate and distil the colourless residue. Its IR (film) has ν_{max} at 1455, 1371 and 1060 (all medium and CH₃), 1451m, 1422s, 1412m, 1275m, 1236m, 1190m, 1171w, 918m and 866w (all dithiane) cm⁻¹ [Corey & Erickson *J Org Chem* **36** 3553 1971, DOI: 10.1021/jo00822a019; Seebach & Corey *J Org Chem* **40** 231 1975, DOI: 10.1021/jo00890a018]. [*Beilstein* **19** III/IV 49, **19**/I V 53.] It is a useful acyl anion equivalent [Smith & Xian *J Am Chem Soc* **128** 66 2006, DOI: 10.1021/jo00890a018].

Methylene Blue [3,7-bis-(dimethylamino)phenothiazin-5-ium chloride] [61-73-4] **M 319.9, CI 52015, ε₆₅₄ 94,000 (EtOH), ε₆₆₄ 81,000 (H₂O), pK²⁵ 3.8**. Crystallise the chloride from 0.1M HCl (16ml/g), the crystals are separated by centrifugation, washed with chilled EtOH and diethyl ether, and dried under vacuum. Crystallise it from 50% aqueous EtOH, wash it with absolute EtOH, and dry it at 50-55° for 24 hours. It has also been crystallised from *benzene/MeOH (3:1). It was salted out with NaCl from a commercial concentrated aqueous solution, then crystallised from water, and dried at 100° in an oven for 8-10 hours. [*Beilstein* **27** III/IV 5152.]

Methylene Green [3,7-bis-(dimethylamino)-4-nitrophenothiazin-5-ium chloride] [2679-01-8] **M 364.9, m >200°(dec), CI 52020, pK²⁵ 3.2**. Crystallise the dye three times from water (18ml/g). [*Beilstein* **27** H 399.] The *ZnCl₂ double salt* [6722-15-2] **M 866.0** has the same **CI**. [*Beilstein* **27** III/IV 5157.]

2-Methylene-oxetan-2-one (diketene) [674-82-8] **M 84.1, m -7°, b 41°/50.5mm, 66-68°/90mm, 127°/760mm, d₄²⁰ 1.440, n_D²⁰ 1.4376, n_D²⁵ 1.4348**. Diketene polymerises violently in the presence of alkali. Distil it under reduced pressure, then fractionally crystallise it by partial freezing (using as a cooling bath made from a 1:1 solution of Na₂S₂O₃ in water, cool with Dry-ice until slushy, and store it in a Dewar flask). Freezing proceeds slowly, and takes about a day for half completion. The crystals are separated and stored in a refrigerator under N₂. [Miller & Carlson *J Am Chem Soc* **79** 3995 1957, DOI: 10.1021/ja01572a010; Andreades & Carlson *Org Synth Coll Vol* **5** 679 1973, DOI: 10.15227/orgsyn.045.0050; *Beilstein* **7** I 309, **7** II 525, **1** III 2947, **17** III/IV 4297, **17**/9 V 115.] See ketene [463-51-4] in 'Aliphatic Compounds', in this Chapter.

2-Methylfuran (Silvan) [534-22-5] C₅H₆S, **M 82.1, m -90.19°, b 62.7-62.8°/731mm, 63-66°/atm, d₄²⁰ 0.917, n_D²⁰ 1.436**. Wash it with acidified saturated ferrous sulfate solution (to remove peroxides), separate, dry with CaSO₄ or CaCl₂, and fractionally distil it from KOH immediately before use. To reduce the possibility of spontaneous polymerisation, addition of about one-third of its volume of heavy mineral oil to 2-methylfuran prior to distillation has been recommended. [*Beilstein* **17** H 36, **17** I 18, **17** II 39, **17** III/IV 265, **17**/I V 322.]

1-Methylguanine [938-85-2] C₆H₇N₅O, **M 165.2, m >300°(dec), pK₁²⁰ ~0, pK₂²⁰ 3.13, pK₃²⁰ 10.54**. Crystallise it from H₂O or 50% aqueous acetic acid. [*Beilstein* **26** III/IV 3892.]

7-Methylguanine [578-76-7] $\text{C}_6\text{H}_7\text{N}_5\text{O}$, M 165.2, $\text{pK}_1^{20} \sim 0$, $\text{pK}_2^{20} 3.50$, $\text{pK}_3^{20} 9.95$. Crystallise it from water. It has UV with λ_{max} at 280nm (pH 2.1). The *picrate* has m 267° (270-272° dec, also reported). [Beilstein 26 H 455, 26 I 134, 26 II 263, 26 III/IV 3890.]

N-Methylimidazole [616-47-7] $\text{C}_4\text{H}_6\text{N}_2$, M 82.1, m -6°, b 81-84°/27mm, 197-198°/760mm, $d_4^{20} 1.032$, $n_D^{20} 1.496$, $\text{pK}^{25} 7.25$. Dry it with sodium metal and then distil it. Store it at 0° under dry argon. The *picrate* has m 159.5-160.5° (from H_2O). [Beilstein 23 III/IV 568, 23/4 V 256.]

2-Methylimidazole (2-methylglyoxaline) [693-98-1] has m 140-141°, 143-143°, 144.5-145.5°, b 267°/760mm, 267-268°/atm, $\text{pK}^{25} 7.86$. Recrystallise 2-methylimidazole from *benzene or petroleum ether. The *picrate* has m 215° (from H_2O). [Beilstein 23 III/IV 594, 23/5 V 35.]

4(5)-Methylimidazole [822-36-6] has m 44-47°, 47-48°, b 263°/760mm, $\text{pK}^{25} 7.61$. Recrystallise 4-methylimidazole from *benzene or petroleum ether. It has m 56° after sublimation. The *picrate* has m 162-163.5° (from EtOH). [Beilstein 23 II 60, 23 III/IV 597, 23/5 V 89.]

2-Methylindole [95-20-5] $\text{C}_9\text{H}_9\text{N}$, M 131.2, m 57-59°, 61°, b 273°/atm, $\text{pK}^{25} -0.28$ (C-3 protonation, aqueous H_2SO_4). Crystallise it from *benzene. It has also been purified by zone melting. The *picrate* has m 139° (from Et_2O or $\text{Et}_2\text{O}/\text{MeOH}$). [Cohen et al. *J Am Chem Soc* 82 2184 1960, DOI: 10.1021/ja01494a024; Beilstein 20 III/IV 3202, 20/7 V 59.]

3-Methylindole (skatole) [83-34-1] has m 92-97°, 95°, b 265-266°/atm, $\text{pK}^{25} -4.55$ (C-3-protonation, aqueous H_2SO_4). Crystallise skatole from *benzene or petroleum ether (m 96.5°). It has also been purified by zone melting. The *picrate* has m 182° (from Et_2O or $\text{Et}_2\text{O}/\text{MeOH}$). [Beilstein 20 III/IV 3206, 20/7 V 69.]

N-Methylmorpholine (4-methylmorpholine) [109-02-4] $\text{C}_5\text{H}_{11}\text{NO}$, M 101.2, m -66°, b 115-116°/750mm, 116-117°/764mm, $d_4^{20} 0.919$, $n_D^{20} 1.436$, $\text{pK}^{25} 7.41$. Dry it by refluxing with BaO or sodium, then fractionally distil it through a helices-packed column. The *picrate* has m 227°, the *thiocyanate salt* has m 103° (from butanone). [Hall *J Phys Chem* 60 63 1956, DOI: 10.1021/j150535a017; Beilstein 27 I 203, 27 III/IV 22.]

4-Methylmorpholine-4-oxide (NMO) [7529-22-8] $\text{C}_5\text{H}_{11}\text{NO}_2$, M 117.2, m 180-184° (anhydr). When the oxide is dried for 2-3 hours at high vacuum, it dehydrates. Add MeOH to the oxide and distil off the solvent under vacuum until the temperature is ca 95°. Then add Me_2CO at reflux and cool to 20°. The crystals are filtered off, washed with Me_2CO and dried. The degree of hydration may vary and may be important for the desired reactions. [van Rheeën et al. *Tetrahedron Lett* 17 1973 1976, DOI: 10.1016/S0040-4039(00)78093-2; Schneider & Hanze *US Pat* 2 769 823; see also Sharpless et al. *Tetrahedron Lett* 17 2503 1976, DOI: 10.1016/S0040-4039(00)78130-5.] It is a non-metallic catalyst for the cyanoethylation of ketones [Zhou et al. *Synlett* 6 1077 2004, DOI: 10.1055/s-2004-820045], and a co-oxidant for Sharpless asymmetric dihydroxylation in ionic liquids [Branco & Afonso *J Org Chem* 69 4381 2004, DOI: 10.1021/jo035588h]. **4-Methylmorpholine-4-oxide monohydrate** [70187-32-5] $\text{C}_5\text{H}_{11}\text{NO}_2 \cdot \text{H}_2\text{O}$, M 135.2, has m 71-73°, 71-75°. The *hydrate* is a reagent for an improved catalytic OsO_4 oxidation of olefins to *cis*-1,2-diols [Van Rheeën et al. *Tetrahedron Lett* 17 1973 1976, DOI: 10.1016/S0040-4039(00)78093-2; Priebe & Zamojski *Pol J Chem* 54 731 1980, see review by Schroeder *Chem Rev* 80 187 1980, DOI: 10.1021/cr60324a003], and for a ruthenium catalysed oxidation of alcohols to ketones see Sharpless et al. [*Tetrahedron Lett* 17 2503 1976, DOI: 10.1016/S0040-4039(00)78130-5]. A 50wt% solution of NMO is commercially available.

3-Methyl-2-oxazolidone [19836-78-3] $\text{C}_4\text{H}_7\text{NO}_2$, M 101.1, m 15°, b 88-91°/1mm, $d_4^{20} 1.172$, $n_D^{20} 1.455$. Purify the oxazolidone by successive fractional freezing, then dry it in a dry-box over 4A molecular sieves for 2 days. Distil it under high vacuum and store it dry as before. [Beilstein 27 III/IV 2517.]

3-Methyl-3-oxetanemethanol (3-hydroxymethyl-3-methyloxetane) [3143-02-0] $\text{C}_5\text{H}_{10}\text{O}_2$, M 102.1, b 80°/4mm, 92-93°/12mm, $d_4^{20} 1.033$, $n_D^{25} 1.4449$. Purify the oxetane by fractionation through a glass column [Pattison *J Am Chem Soc* 79 3455 1957, DOI: 10.1021/ja01570a038; Corey & De *J Am Chem Soc* 106 2735 1984, DOI: 10.1021/ja00321a063]. [Beilstein 17 III/IV 1128.] It has been used for protecting carbonyl groups in a synthesis of α -substituted γ -lactams [Raghavan & Johnson *J Org Chem* 71 2151 2006, DOI: 10.1021/jo035588h].

5-Methyl-1,10-phenanthroline [3002-78-6] $C_{13}H_{10}N_2$, M 194.2, m 67°(monohydrate), 113°, 113-114° (anhydrous), pK^{25} 5.28. Crystallise it from *benzene/petroleum ether or from H_2O as the *monohydrate*. It complexes with many metals. [Beilstein 23 III/IV 1714.]

5-Methylphenazinium methyl sulfate (PMS) [299-11-6] $C_{13}H_{11}N_2 \cdot CH_3SO_4$, M 306.3, has m 155-157°(dec), 158-160°(dec), (198°dec by rapid heating), pK^{25} -3.5. It forms yellow-brown prisms from EtOH (charcoal), or EtOH/Et₂O. Its solubility in H_2O at 20° is 10%. In the presence of aqueous KI it forms a *semiquinone* which crystallises as blue leaflets from EtOH. [Wieland & Roseau *Chem Ber* 48 1117 1915, DOI: 10.1002/cber.191504801155; Voriskova *Coll Czech Chem Commun* 12 607 1947, DOI: org/10.1135/cccc19470607; Bülow *Chem Ber* 57 1431 1924, DOI: 10.1002/cber.191504801155; Campbell et al. *J Chem Soc* 404 1938, DOI: 10.1039/JR9380000404; Morley *J Chem Soc* 4008 1952, DOI: 10.1039/JR9520004008; Beilstein 23 I 59, 23 II 234, 23 III/IV 1658, 23/8 V 395.] Together with ascorbic acid, it is used to determine nitric oxide reductase.

N-Methylphenothiazine (10-methylphenothiazine) [1207-72-3] $C_{13}H_{11}NS$, M 213.2, α -form m 99.3°, 99-101°, 100-102°, and b 360-365°/atm, β -form m 78-79°. Recrystallise it (three times) from EtOH to give the α -form (prisms). Recrystallisation from EtOH/*benzene gives the β -form (needles). It has also been purified by vacuum sublimation and is carefully dried in a vacuum line. It has been crystallised from toluene or MeOH and stored in the dark [Guarr et al. *J Am Chem Soc* 107 5104 1985, DOI: 10.1021/ja00304a015; Olmsted et al. *J Am Chem Soc* 109 3297 1987, DOI: 10.1021/ja00245a018]. Its solubility in H_2O (pH 6.5) is 0.0544mg/100ml, and in hexane it is 2083mg/100ml. Its UV has λ_{max} at 255nm (log ϵ 4.60) (pH 6.3) and 255nm in hexane. [Cymerman-Craig & Warburton *Aust J Chem* 9 294 1956, DOI:10.1071/CH9560294; Beilstein 27 H 65, 27 II 33, 27 III/IV 1215.]

3-Methyl-1-phenyl-5-pyrazolone [89-25-8] $C_{10}H_{10}N_2O$, M 174.2, m 126-128°, 127°, 129°, b 287°/265mm, pK^{25} 2.7. Crystallise the pyrazolone from hot H_2O , EtOH or EtOH/water (1:1). It complexes with metals. [Veibel et al. *Acta Chim Scand* 6 1066 1952, DOI: 10.3891/acta.chem.scand.06-1066; Beilstein 24 II 9, 24 III/IV 71.] This reagent has been used for detecting reducing carbohydrates by ESI/MALDI-MS [Lattová et al. *J Am Mass Spectrom* 16 683 2005, DOI: 10.1016/j.jasms.2005.01.021].

1-Methyl-4-phenyl-1,2,3,6-tetrahydropyridine hydrochloride (MPTP.HCl) [23007-85-4] $C_{12}H_{15}N \cdot HCl$, M 209.7, m 196-198°, $pK_{Est} \sim 9.3$, pK_a 8.07 (50% aqueous EtOH). Purify MPTP by recrystallisation from Me_2CO /isoPrOH. The *free base* has m 40-42°(from heptane), b 99-100°/1.3mm, 128-132°/12mm, (137-142°/0.8mm), n_D^{25} 1.5347. The *hydrochloride* has m 251-252°(from Me_2CO /isoPrOH) and its solubility in H_2O is 10mg/ml at ~20°. [Schmidle & Mansfield *J Am Chem Soc* 78 425 1956, DOI: 10.1021/ja01583a048; DeFeudis *Drug Dev Res* 15 1 1988, DOI: 10.1002/ddr.430150102; Beilstein 20 III/IV 3240, 20/7 V 121.] It is a dopaminergic neurotoxin and used in *in vitro* studies of model Parkinsonism.

(±)-2-Methylpiperazine [109-07-9] $C_5H_{12}N_2$, M 100.2, m 61-62°, 61-63°, 66°, b 147-150°/739mm, 155°/763mm, pK_1^{25} 5.46, pK_2^{25} 9.90. Purify it by zone melting and by distillation. It is *hygroscopic*. The *picrate* has m 275-276°. [Beilstein 23 II 16, 23 III/IV 393, 23/3 V 267.] *R(+)-2-Methylpiperazine* [75336-86-6] has m 91-93°, $[\alpha]_D^{20}$ -16.5 (c 5, * C_6H_6), and is obtained by optical resolution of (±)-2-methylpiperazine *via* separation of the diastereoisomeric (+)-base (+)-di-*O*-benzoyl tartrate and (-)-base (+)-di-*O*-benzoyl tartrate. *S(+)-2-methylpiperazine* [74879-18-8] with m 91-93°, $[\alpha]_D^{20}$ +6.8 (c 1, EtOH), is obtained *via* this resolution and its absolute configuration is deduced from its preparation by cyclising glycyl-*S*-alanine to *S(-)-3-methyl-2,5-dioxopiperazine* $[\alpha]_{546}^{20}$ -9.6 (c 1.35 H_2O) followed by reduction with $NaAl(OCH_2CH_2OMe)_2H_2$, and 4N-HCl to give *S(-)-2-methylpiperazinium dihydrochloride* which after sublimation at 190°/2mm and recrystallisation from MeOH/Et₂O has m 286-289° (effervescence and sublimation) and $[\alpha]_{546}^{20}$ -3.5 (c 1.2, 2N-HCl). [Armarego et al. *Journal of Chemical Research* 133 (S), 1951 (M) 1980, Armarego et al. *JCS Chem Commun* 334 1980, DOI: 10.1039/C39800000334.]

(±)-3-Methylpiperidine [626-56-2] $C_6H_{13}N$, M 99.2, b 125°/763mm, 125-126°/763mm, d_4^{20} 0.846, n_D^{25} 1.4448, pK^{25} 10.92. Purify it *via* the *hydrochloride* (m 172°). The *hydrobromide* has m 162-163°(from iso-PrOH). [Chapman et al. *J Chem Soc* 1925 1959, DOI: 10.1039/JR9590001925; Beilstein 20 III/IV 1499, 20/4 V 100.]

4-Methylpiperidine [626-58-4] $C_6H_{13}N$, M 99.2, b 124.4°/755mm, d_4^{20} 0.839, n_D^{25} 1.4430, pK²⁵ 10.78. Purify it via the *hydrochloride* (m 189°). It is freed from 3-methylpyridine by zone melting. The *hydrobromide* has m 173°(from butanone/* C_6H_6). [Beilstein 20 III/IV 1511, 20/4 V 116.]

1-Methyl-4-piperidone [1445-73-4] $C_6H_{11}NO$, M 113.2, b 53-56°/0.5mm, 54-56°/9mm, 68-71°/17mm, 85-87°/45mm, d_4^{20} 0.972, n_D^{25} 1.4588, pK²⁵ 7.9. It is best purified by fractional distillation. The *hydrochloride* of the hydrate (4-diol) has m 94.7-95.5°, but the *anhydrous hydrochloride* which crystallises from $CHCl_3/Et_2O$ has m 165-168° (164-167°), and can also be obtained by sublimation at 120°/2mm. The *oxime* has m 130-132° (from Me_2CO). The *methiodide* crystallises from MeOH, the crystals with 1MeOH have m 189-190°, and the solvent-free *iodide* has m 202-204°(dec). [Lyle et al. *J Org Chem* 24 342 1959, DOI: 10.1021/jo01085a015; Bowden & Green *J Chem Soc* 1164 1952, DOI: 10.1039/JR9520001164; Tomita *Yakugaku Zasshi (J Pharm Soc Japan)* 71 1053 1951, Beilstein 21 III/IV 3183, 21/6 V 419.]

2-Methylpyrazine [109-08-0] $C_5H_6N_2$, M 94.1, m -28.8°, -29°, b 135°/761mm, 136-137°/atm, d_4^{20} 1.025, n_D^{20} 1.505, pK₁²⁵ -5.25 (aqueous H_2SO_4), pK₂²⁵ 1.47. Purify it via the *picrate* and distil the *free base*. The *picrate* has m 133-134°(from EtOH). [Wiggins & Wise *J Chem Soc* 4780 1956, DOI: 10.1039/JR9560004780; Beilstein 23 III/IV 911, 23/5 V 386.]

2-Methylpyridine (2-picoline, α -picoline) [109-06-8] C_6H_7N , M 93.1, m -70°, b 128-129°/atm, 129.4°/760mm, d_4^{20} 0.9444, n_D^{20} 1.50102, pK²⁵ 5.96. Biddiscombe et al. [*J Chem Soc* 1957 1954, DOI: 10.1039/JR9540001957] steam distilled a boiling solution of the base in 1.2 equivalents of 20% H_2SO_4 until about 10% of the base had been carried over, along with non-basic impurities. Excess aqueous NaOH is then added to the residue, the free base is separated, dried with solid NaOH and fractionally distilled. 2-Methylpyridine can also be dried with BaO, CaO, CaH_2 , $LiAlH_4$, or Linde type 5A molecular sieves. An alternative purification is via the *ZnCl₂ adduct*, which is formed by adding 2-methylpyridine (90ml) to a solution of anhydrous $ZnCl_2$ (168g) and 42ml concentrated HCl in absolute EtOH (200ml). Crystals of the complex are filtered off, recrystallised twice from absolute EtOH (to give m 118.5-119.5°), and the *free base* is liberated by addition of excess aqueous NaOH. It is steam distilled, and solid NaOH is added to the distillate to form two layers, the upper one of which is then dried with KOH pellets, stored for several days with BaO and fractionally distilled. Instead of $ZnCl_2$, $HgCl_2$ (430g in 2.4L of hot water) can be used. The complex, which separates on cooling, can be dried at 110° and recrystallised from 1% HCl (to m 156-157°). *The alkali metals, Na, Li or Cs should NOT be used for drying pyridine, and pyridine derivatives, as they form coloured pyridine radical anions leading to bipyridyls.* [see Schmulbach et al. *J Am Chem Soc* 90 6600 1968, DOI: 10.1021/ja01026a006]. The *hydrochloride* has m 78-79°, and the *picrate* has m 165.5°(from EtOH) and 180°(from H_2O). [Beilstein 20 III/IV 2679, 20/5 V 464.] **2-Picoline Borane Complex**, [3999-38-0] C_6H_7N . M 107.0, m 44-46°, 48°, can be purified by recrystallisation from hexane and stored for long periods without noticeable decomposition. It is unchanged after heating to ~150°, and is consequently much more stable than *pyridine borane*. It is an alternative reagent for reductive aminations [Sato et al. *Tetrahedron* 60 7899 2004, DOI: 10.1016/j.tet.2004.06.045]. **α -Picoline hydrochloride** [14401-91-3] C_6H_7N . HCl, M 129.6, m 77-78°, 85°, b 229°/atm, is prepared from a 1:1 mixture of α -picoline and HCl is distilled at 275°, then sublimed in a vacuum at 91-91.5°. [Beilstein 20 H 236, 20 III/IV 2685, 20/5 V 474.]

3-Methylpyridine (3-picoline) [108-99-6] has m -19°, -18.5°, b 144°/767mm, d_4^{20} 0.957, n_D^{20} 1.5069, pK²⁵ 5.70. In general, the same methods of purification that are described for 2-methylpyridine can be used. However, 3-methylpyridine often contains 4-methylpyridine and 2,6-lutidine, neither of which can be removed satisfactorily by drying and fractionation, or by using the $ZnCl_2$ complex. Biddiscombe et al. [*J Chem Soc* 1957 1954, DOI: 10.1039/JR9540001957], after steam distillation as for 2-methylpyridine, treated the residue with urea to remove 2,6-lutidine, then azeotropically distilled with acetic acid (the azeotrope had b 114.5°/712mm), and recovered the base by adding excess of aqueous 30% NaOH, drying with solid NaOH and carefully fractionally distilling. The distillate is then fractionally crystallised by slow partial freezing. An alternative treatment [Reithoff et al. *Ind Eng Chem (Anal Edn)* 18 458 1946, DOI: 10.1021/i560155a023] is to reflux the crude base (500ml) for 20-24 hours with a mixture of acetic anhydride (125g) and phthalic anhydride (125g) followed by distillation until phthalic anhydride begins to pass over. The distillate is treated with NaOH (250g in 1.5L of water) and then steam distilled. Addition of solid NaOH (250g) to this distillate (ca 2L) led to the separation of 3-methylpyridine which is removed, dried (K_2CO_3 , then BaO) and fractionally distilled. (Subsequent fractional freezing would probably be advantageous.) *The alkali metals, Na, Li or Cs should*

NOT be used for drying pyridine, and pyridine derivatives, as they form coloured pyridine radical anions leading to bipyridyls. [see Schmulbach et al. *J Am Chem Soc* **90** 6600 1968, DOI: 10.1021/ja01026a006]. The *hydrochloride* has **m 85°**, and the *picrate* has **m 153°**(from Me₂CO, EtOH or H₂O). [*Beilstein* **20** III/IV 2710, 20/5 V 506.]

4-Methylpyridine (4-picoline) [108-89-4] has **m 2.4°, 4.25°, b 145.0°/765mm, d₄²⁰ 0.955, n_D²⁰ 1.5058, pK²⁵ 4.99**. It can be purified as for **2-methylpyridine**. Biddescombe and Handley's method (above) for 3-methylpyridine is also applicable. Lidstone [*J Chem Soc* 241 1940, DOI: 10.1039/JR9400000241] purified it via the *oxalate* (**m 137-138°**) by heating 100ml of 4-methylpyridine to 80° and adding slowly 110g of anhydrous oxalic acid, followed by 150ml of boiling EtOH. After cooling and filtering, the precipitate is washed with a little EtOH, then recrystallised from EtOH, dissolved in the minimum quantity of water and distilled with excess 50% KOH. The distillate is dried with solid KOH and again distilled. Hydrocarbons can be removed from 4-methylpyridine by converting the latter to its hydrochloride, crystallising from EtOH/diethyl ether, regenerating the free base by adding alkali and distilling. As a final purification step, 4-methylpyridine can be fractionally crystallised by partial freezing to effect a separation from 3-methylpyridine. Contamination with 2,6-lutidine is detected by its strong absorption at 270nm. **The alkali metals, Na, Li or Cs should NOT be used for drying pyridine, and pyridine derivatives, as they form coloured pyridine radical anions leading to bipyridyls.** [see Schmulbach et al. *J Am Chem Soc* **90** 6600 1968, DOI: 10.1021/ja01026a006]. The *hydrochloride* has **m 161°**, and the *picrate* has **m 167°**(from Me₂CO, EtOH or H₂O). [*Beilstein* **20** III/IV 2732, 20/5 V 543.]

2-Methylpyridin-5-yltrifluoromethanesulfonate (2-methyl-5-pyridine triflate) [111770-91-3] **C₇H₆F₃N O₃S, M 241.2, b 80-82°/1.9mm, d₄²⁵ 1.412, n_D²⁰ 1.442**. Distil the triflate under vacuum, or if it is discoloured dissolve it in CH₂Cl₂, wash it with N NaOH and half saturated K₂CO₃, dry it over solid K₂CO₃, filter, evaporate, and distil by bulb-to-bulb distillation at 65-70°/0.1mm, then redistil *in vacuo*. Its ¹H NMR [(CD₃)₂SO] has δ at 2.50 (s, 3H, Me), 7.45 (d, *J* = 9.2 Hz, 1H), 7.90 (dd, *J* = 9.2, 2.3 Hz, 1H) and 8.60 (d, *J* = 2.3 Hz, 1H). [Tilley & Zawoiski *J Org Chem* **53** 386 1988, DOI: 10.1021/jo00237a029; Ellingboe et al. *J Med Chem* **37** 542 1994, DOI: 10.1021/jm00030a013]. When stirred with *m*-chloroperbenzoic acid in CH₂Cl₂ (16 hours, 25°), filtered, concentrated and purified by flash chromatography (2% MeOH/ CH₂Cl₂), the triflate gave colourless crystals of the *triflate N-oxide* **m 47-48°** with ¹H NMR [(CD₃)₂SO] with δ at 2.36 (s, 3H, Me), 7.56 (dd, *J* = 8.9, 2.3 Hz, 1H), 7.69 (d, *J* = 8.9 Hz, 1H) and 8.84 (d, *J* = 2.3 Hz, 1H) [Ellingboe et al. *J Med Chem* **37** 542 1994, DOI: 10.1021/jm00030a013].

N-Methylpyrrole (NMP) [96-54-8] **C₅H₇N, M 81.1, m -57°, b 112-113°/atm, 115-116°/756mm, d₄²⁰ 0.908, n_D²⁰ 1.487, pK²⁵ -3.4 (-2.90)**. Dry *N*-methylpyrrole with CaSO₄, then fractionally distil it from KOH immediately before use. [*Beilstein* **20** III/IV 2080, 20/5 V 8.]

1-Methyl-2-pyrrolidinone (1-methyl-2-pyrrolidone) [872-50-4] **C₅H₉NO, M 99.1, m -24.4°, b 65-76°/1mm, 78-79°/12mm, 81-82°/10mm, 94-96°/20mm, 202°/760mm, d₄²⁰ 1.0328, n_D²⁰ 1.4678, pK²⁵ -0.17 (also -0.92, and 0.2)**. Dry the pyrrolidone by removing water as the *benzene azeotrope. Fractionally distil at 10 torr through a 100-cm column packed with glass helices. [Adelman *J Org Chem* **29** 1837 1964, DOI: 10.1021/jo01030a041; McElvain & Vozza *J Am Chem Soc* **71** 896 1949, DOI: 10.1021/ja01171a037.] It is a useful solvent for reactions. The *hydrochloride* has **m 86-88°** (from EtOH or Me₂CO/EtOH) [Reppe et al. *Justus Liebigs Ann Chem* **596** 1 1955, DOI: 10.1002/jlac.19555960102]. [*Beilstein* **21** II 213, **21** III/IV 3145, 21/6 V 321.]

2-Methylquinoline (quinaldine) [91-63-4] **C₁₀H₉N, M 143.2, m -9° to -3°, b 86-87°/1mm, 105-107°/10mm, 155°/14mm, 246-247°/760mm, d₄²⁰ 1.058, n_D²⁰ 1.6126, pK²⁵ 5.65**. Dry it with Na₂SO₄ or by refluxing with BaO, then fractionally distil it under reduced pressure and redistil it from zinc dust. Purify it further by conversion to its *phosphate* (**m 220°**) or *picrate* (**m 192°**) from which after recrystallisation, the *free base* is regenerated. [Packer et al. *J Am Chem Soc* **80** 905 1958, DOI: 10.1021/ja01537a039.] Its **ZnCl₂ complex** can be used for the same purpose. [*Beilstein* **20** III/IV 3454, 20/7 V 375.]

4-Methylquinoline (lepidine) [491-35-0] has **m 9-10°, b 261-263°/atm, 265.5°/760mm, d₄²⁰ 1.084, n_D²⁰ 1.61995, pK²⁵ 5.59**. Reflux lepidine with BaO, then fractionally distil it. Further purify it via its recrystallised *dichromate salt* (**m 138°**) (from H₂O). [Cumper et al. *J Chem Soc* 1176 1962, DOI: 10.1039/JR9620001176.] [*Beilstein* **20** III/IV 3477, 20/7 V 389.]

6-Methylquinoline (p-toluquinoline) [91-62-3] has **b** 256-260°/7atm, 258.6°/760mm, d_4^{20} 1.067, n_D^{20} 1.61606, pK^{25} 4.92. Reflux it with BaO, then fractionally distil it. Further purified *via* its recrystallised **ZnCl₂ complex** (**m** 190°). [Cumper et al. *J Chem Soc* 1176 1962, DOI: 10.1039/JR9620001176; **20** III/IV 3498, **20/7** V 400.]

7-Methylquinoline [612-60-2] has **m** **m** 35-39°, 38°, **b** 258°/atm, 255-260°/760mm, d_4^{20} 1.052, n_D^{20} 1.61481, pK^{25} 5.29. Purify it *via* its **dichromate complex** (**m** 149°, after five recrystallisations from water). [Cumper et al. *J Chem Soc* 1176 1962, DOI: 10.1039/JR9620001176; *Beilstein* **20** III/IV 3497, **20/7** V 402.]

8-Methylquinoline [611-32-5] has **m** -80°, **b** 122.5°/16mm, 143°/34mm, 247.8°/760mm, d_4^{20} 1.703, n_D^{20} 1.61631, pK^{25} 4.60. Purify it as for 2-methylquinoline. The **phosphate** and **picrate** have **m** 158° and **m** 201°, respectively. [*Beilstein* **20** III/IV 3500, **20/7** V 405.]

(±)-3-Methylsulfolane (3-methyltetrahydrothiophene-1,1-dioxide) [872-93-5] **C₅H₁₀O₂S**, **M** 134.2, **m** 0.5°, 1.0°, **b** 101°/2mm, 104°/3mm, 125-130°/12mm, 278-282°/763.5mm, d_4^{20} 1.1885, n_D^{20} 1.4770. Distil the sulfolane under vacuum and recrystallise it from Et₂O at -60° to -70°, if necessary. An IR film has strong bands at 570 and 500 cm⁻¹. [Eigenberger *J Prakt Chem* [2] **131** 289 1931, DOI: 10.1002/prac.19311310118; Freaiheller & Katon *Spectrochim Acta* **20** 1099 1964, DOI: 10.1016/0371-1951(64)80160-0; Whitehead et al. *J Am Chem Soc* **73** 3632 1951, DOI: 10.1021/ja01152a022; *Beilstein* **17** I 8, **17** III/IV 64.]

(±)-2-Methyltetrahydrofuran [96-47-9] **C₅H₁₀O**, **M** 86.1, **b** 78-80°/atm, 80.0°/760mm, d_4^{20} 0.856, n_D^{20} 1.4053. Likely impurities are 2-methylfuran, methyl-dihydrofurans and hydroquinone (stabiliser, which is removed by distillation under reduced pressures). It is washed with 10% aqueous NaOH, dried, vacuum distilled from CaH₂, passed through freshly activated alumina under nitrogen, and refluxed over sodium metal under vacuum. Store it over sodium. [Ling & Kevan *J Phys Chem* **80** 592 1976, DOI: 10.1021/j100547a008.] Distil it from sodium under vacuum, and store it with sodium-potassium alloy (this treatment removes water and prevents the formation of peroxides). *Alternatively*, it can be freed from peroxides by treatment with ferrous sulfate and sodium bisulfate, then solid KOH, followed by drying with, and distilling from, sodium, or type 4A molecular sieves under argon. It may be difficult to remove *benzene if it is present as an impurity (can be readily detected by its ultraviolet absorption in the 249-268nm region). [Ichikawa & Yoshida *J Phys Chem* **88** 3199 1984, DOI: 10.1021/j150659a013.] It has also been purified by percolating through Al₂O₃ and fractionated collecting fraction **b** 79.5-80°. After degassing, the material is distilled onto degassed molecular sieves, then distilled onto anthracene and a sodium mirror. The solvent is then distilled from the green solution onto potassium mirror or sodium-potassium alloy, from which it is distilled again. [Kosower & Mohammad *J Am Chem Soc* **93** 2713 1971, DOI: 10.1021/ja00740a022.] It should be stored in the presence of 0.1% of hydroquinone or 2,6-di-*tert*-butyl-*p*-cresol as stabiliser. The **R(+)-enantiomer** has **b** 78-80°/atm and $[\alpha]_D^{20}$ +27.5 (neat), and the **S(-)-enantiomer** has **b** 86°/atm and $[\alpha]_D^{20}$ -27.0 (neat) [Iffland & Davis *J Org Chem* **42** 4150 1977, DOI: 10.1021/jo00445a038, Gagnaire & Butt *Bull Soc Chim Fr* 312 1961, *Beilstein* **17** III/IV 60, **17/1** V 78.] **HARMFUL VAPOURS.**

3-Methylthiophene [616-44-4] **C₅H₆S**, **M** 98.2, **m** -69°, **b** 60°/116mm, 111-113°/atm, 115.5°/atm, d_4^{20} 1.024, n_D^{20} 1.531. Dry it with Na₂SO₄, then distil it from sodium. [*Beilstein* **17** III/IV 277, **17/1** V 331.]

6(4)-Methyl-2-thiouracil [56-04-2] **C₅H₆N₂OS**, **M** 142.2, **m** 330°(dec), 299-303°(dec), 323-324°(dec), **d** 1.36, pK^{25} 8.1. Crystallise the thiouracil from a large volume of H₂O. Purify it further by dissolving in base, adding charcoal, filtering and acidifying with AcOH. Suspend the wet solid (*ca* 100g) in boiling H₂O (1L), stir and add AcOH (20ml), stir and refrigerate. Collect the product, wash it with cold H₂O (4 x 200ml), drain it for several hours then place it in an oven at 70° to constant weight. It has antithyroid properties. [IR: Short & Thompson *J Chem Soc* 168 1952, DOI: 10.1039/JR9520000168; Foster & Snyder *Org Synth Coll Vol* **4** 638 1963, DOI: 10.15227/orgsyn.035.0080; *Beilstein* **24** III/IV 1289.]

4-Methyl-1,2,4-triazoline-3,5-dione (MTAD) [13274-43-6] **C₃H₃N₃O₂**, **M** 113.1, **m** 103-104°, **m** 107-109°. MTAD is obtained as pink (red crystals) needles by sublimation at 40-50°/0.1mm (see 4-phenyl-1,2,4-triazoline-3,5-dione, PTAD below). [Cookson et al. *Org Synth* **51** 121 1971, DOI: 10.15227/orgsyn.051.0121; Cheng et al. *J Org Chem* **49** 2910 1984, DOI: 10.1021/jo00190a014; *Beilstein* **26** III/IV 538.] It is one of the most reactive of dienophiles towards dienes [Cookson et al. *Tetrahedron Lett* **3** 615 1962, DOI: 10.1016/S0040-4039(00)70917-8; Gillis & Hagarty *J Org Chem* **32** 330 1967, DOI: 10.1021/jo01288a016,] and mono-olefines [Pirkle & Stickler *JCS Chem Commun* 760 1967, DOI: 10.1039/C19670000760].

2-Methyltricycloquinazoline [2642-52-6] $C_{22}H_{14}N_4$, M 334.4, m >300°. Purify it by crystallisation from C_6H_6 , toluene or xylene followed by vacuum sublimation. [cf. *Beilstein* 26 III/IV 1932.] **CARCINOGEN**.

5-Methyltryptamine hydrochloride (3-[2-aminoethyl]-5-methylindole hydrochloride) [1010-95-3] $C_{11}H_{14}N_2$. HCl, M 210.7, m 289-291°(dec), 290-292°, $pK_{Est(1)} \sim -3$ (protonation of ring NH), $pK_{Est(2)} \sim 9.0$ (CH_2NH_2), $pK_{Est(3)} \sim 10.9$ (acidic indole NH). Recrystallise the hydrochloride from H_2O . The *free base* has m 96-98° (from C_6H_6 /cyclohexane) or m 99-100° (from petroleum ether), and the *picrate* has m 243°(dec) (from EtOH). [Young *J Chem Soc* 3493 1958, DOI: 10.1039/JR9580003493; Gaddum et al. *Quart J Exp Physiol* 40 49 1955, Röhm *Hoppe Seyler's Z Physiol Chem* 297 229 1954, PMID: 14371990; *Beilstein* 22 III/IV 4364, 22/10 V 167.]

6-Methyluracil [626-48-2] $C_5H_6N_2O_2$, M 126.1, m 270-280°(dec), 318°(dec), λ_{max} 260nm log ϵ 3.97, $pK_1 \sim 1.1$, pK_2 9.8. Crystallise 6-methyluracil from EtOH or acetic acid. [*Beilstein* 24 III/IV 1281.]

1-Methyluric acid [708-79-2] $C_6H_6N_4O_3$, M 182.1, m >350°, pK_1 5.75 (basic), pK_2 10.6 (acidic). Recrystallise it from H_2O . Its solubility at 17.5° is 1g in 353ml of H_2O . [Bergmann & Dikstein *J Am Chem Soc* 77 691 1955, DOI: 10.1021/ja01608a044.] It has UV with λ_{max} at 231 and 283nm (pH 3), and 217.5 and 292.5nm (pH >12) [Johnson *Biochem J* 51 133 1952, DOI: 10.1042/bj0510133]. [*Beilstein* 26 II 299, 26 III/IV 2621.]

3-Methyluric acid [39717-48-1] has m >350°, pK_1 5.75 (6.2), pK_2 >12. Crystallise it from water. Its solubility at 17.5° is 1g in 19.7L of H_2O . It has UV with λ_{max} at 232 and 287 nm (pH 3), and 214 and 292.5nm (pH >12). [*Beilstein* 26 II 299, 26 III/IV 2621.]

7-Methyluric acid [612-37-3; 30409-21-3] has m >380°, pK_1 5.6, pK_2 10.3. Crystallise it from water. It has UV with λ_{max} at 234 and 286nm (pH 3), 237 and 293nm (8.5), and 222 and 296.5nm (pH >12) [*Beilstein* 26 H 525, 26 II 299, 26 III/IV 2622.]

9-Methyluric acid [30345-24-5, 55441-71-9] has m 385-400°(dec), >400°, d_{25}^{25} 1.73g/ml, n_D^{20} 1.70 (predicted), $pK_{Est} \sim 5.65$. Crystallise it from water (solubility is 2mg/ml at 25°). [*Beilstein* 26 III/IV 2622.]

1-Methylxanthine (1-methyl-purin-2,6(3-*H*,7-*H*)-dione) [6136-37-4] $C_6H_6N_4O_2$, M 166.1, m >360° pK_1^{20} 1.3, pK_2^{20} 7.9, pK_3^{20} 11.8. Crystallise it from water. It has UV with λ_{max} at 266nm (pH 2.08), 242.5 and 276nm (pH 9). [*Beilstein* 26 II 263, 26 III/IV 2329.]

3-Methylxanthine [1076-22-8] has m >360° pK_1^{20} 8.45, pK_2^{20} 11.92. Crystallise it from water. [*Beilstein* 26 II 263, 26 III/IV 2329.]

7-Methylxanthine [552-62-5] has m >380°(dec) pK_1^{20} 8.42, pK_2^{20} >13. Crystallise it from water. [*Beilstein* 26 II 263, 26 III/IV 2330.]

8-Methylxanthine [17338-96-4] has m 292-293°(dec). Crystallise it from water. [*Beilstein* 26 III/IV 2464.]

9-Methylxanthine [1198-33-0] has m 384°(dec), pK_1^{20} 2.0, pK_2^{20} 6.12, pK_3^{20} 10.5 (>13). Crystallise it from water. [*Beilstein* 26 II 263, 26 III/IV 2330.]

Morin (hydrate) (2',3,4',5,7-pentahydroxyflavone) [90-34-6, 480-16-0 (anhydr), 6472-38-4 (dihydrate)] $C_{15}H_{10}O_7 \cdot 2H_2O$, M 302.2 (338.3), m 289-292°, CI 75660, pK_1 5.3, pK_2 8.74. Stir morin at room temperature with ten times its weight of absolute EtOH, then leave overnight to settle. Filter it off, and evaporate under a heat lamp to one-tenth its volume. An equal volume of water is added, and the precipitated morin is filtered off, dissolved in the minimum amount of EtOH and again precipitated with an equal volume of water. The precipitate is filtered off, washed with water and dried at 110° for 1 hour (yield ca 2.5%). [Perkins & Kalkwarf *Anal Chem* 28 1989 1956, DOI: 10.1021/ac60120a051.] It complexes with W and Zr. [*Beilstein* 18 H 239, 18 III/IV 3468, 18/5 V 492.] It protects cells against γ -radiation induced oxidative state [Zhang et al. *Basic & Clin Pharmacol & Toxicol* 108 63 2001 DOI: 10.1111/j.1742-7843.2010.00629.x.]

Morpholine [110-91-8] C_4H_9NO , M 87.1, m -7° to -5°, -4.9°, b 128.9°/760mm, d_4^{20} 1.0007, n_D^{20} 1.4540, n_D^{25} 1.4533, pK^{25} 8.33. Dry morpholine with KOH, fractionally distil it, then reflux it with Na, and again fractionally distil it. Dermer & Dermer [*J Am Chem Soc* 59 1148 1937, DOI: 10.1021/ja01285a503] precipitated it as the *oxalate* by adding slowly to slightly more than 1 molar equivalent of oxalic acid in EtOH. The precipitate is filtered off and recrystallised twice from 60% EtOH [*1:1 salt* has m 190-195°(dec)]. Addition of the oxalate to concentrated aqueous NaOH regenerated the base, which is separated and dried with solid

KOH, then sodium, before being fractionally distilled. The *hydrochloride* has **m 178-179°** (from MeOH/Et₂O), and the *picrate* has **m 151.6°** (from aqueous EtOH). [Beilstein 27 II 3, 27 III/IV 15.]

§ A polystyrene supported morpholine is commercially available.

2-(N-Morpholino)ethanesulfonic acid (MES) [4432-31-9] C₆H₁₃NO₄S. xH₂O, **M 195.2** (anhyd), **m >300°(dec)**, **pK₁ 1.99**, **pK₂ 6.15**. Crystallise MES from hot EtOH containing a little water. Useful buffer in the pH range 5.5–6.7. The *picrate* crystallises from EtOH and has **m 178.8-182°**. [Malkiel & Mason *J Org Chem* 8 199 1943, DOI: 10.1021/jo01191a001; Beilstein 27 III/IV 370.]

Murexide (ammonium purpurate, 5,5'-nitrilobarbituric acid) [3051-09-0] C₈H₈N₆O₆, **M 284.2**, **m >300°**, **λ_{max} 520nm** (ε 12,000), **pK₂ 9.2**, **pK₃ 10.9**. The sample may be grossly contaminated with uramil, alloxanthine, etc., and may be difficult to purify. It is better to synthesise it from pure alloxanthine [Davidson *J Am Chem Soc* 58 1821 1936, DOI: 10.1021/ja01300a509]. Recrystallise a purer sample from hot water to give purple-red crystals with a green lustre which produce deep purple solutions in H₂O that turn deep blue on addition of NaOH. [Kuhn & Lyman *Chem Ber* 69 1547 1936, DOI: 10.1002/cber.19360690656; Beilstein 25 I 709, 25 III/IV 4236.] It is a complexometric titration indicator.

α-Naphthoflavone (7,8-benzoflavone) [604-59-1] C₁₉H₁₂O₂, **M 272.3**, **m 153-155°, 153-157°, 155°, pK²⁵ 8-9** (phenolic OH). Recrystallise the yellow flavone from EtOH or aqueous EtOH. [IR: Cramer & Windel *Chem Ber* 89 354 1956, DOI: 10.1002/cber.19560890227; UV Pilon & Massicot *Bull Soc Chim Fr* 26 1954, Smith et al. *J Chem Soc* 542 1946, DOI: 10.1039/JR9460000542; Mahal & Venkataraman *J Chem Soc* 1767 1934, DOI: 10.1039/JR9340001767.] It is a competitive inhibitor of human estrogen synthase (aromatase). [Kellis & Vickery *Science* 225 1032 1984, DOI: 10.1126/science.6474163; Beilstein 17 III/IV 5550.] **β-Naphthoflavone (5,6-benzoflavone)** [6051-87-2] C₁₉H₁₂O₂, **M 272.3**, **m 164-166°, pK²⁵ 8-9** (phenolic OH) is purified in the same manner.

Naphthol AS-acetate (3-acetoxynaphthoic acid anilide) [1163-67-3] C₁₉H₁₅NO₃, **M 305.3**, **m 152°, 160°, 160-162°**. Recrystallise it from hot MeOH and dry *in vacuo* over P₂O₅. It has **m 252°** after sublimation at 210–215°. It is slightly soluble in AcOH, EtOH, CHCl₃ or *C₆H₆. It is a fluorogenic substrate for albumin esterase activity. [Chen & Scott *Anal Lett* 17 857 1984, DOI: 10.1080/0003271811681085] At λ_{ex} 320nm it has fluorescence at λ_{em} 500nm. [Brass & Sommer *Chem Ber* 61 993 1928, DOI: 10.1002/cber.19280610518; Beilstein 12 II 260, 12 III 960, 12 IV 923.]

1,5-Naphthyridine [254-79-5] C₈H₆N₂, **M 130.1**, **m 69-75°, 75°, b 112°/15mm**, **pK²⁰ 2.84**. Recrystallise the base from light petroleum (60-70°) and sublime it at 40°/0.005mm colourless needles. It can also be distilled under a vacuum. The *picrate* crystallises from EtOH with **m 200°(dec)**. The *dihydrochloride* has **m 234°(dec)** after recrystallisation from EtOH/HCl/Et₂O. [Albert *J Chem Soc* 1790 1960, DOI: 10.1039/JR9600001790; UV: Armarego in *Physical Methods in Heterocyclic Chemistry* (Ed Katritzky, Academic Press, LCCC No 62-12037, Internet Public Domain) Vol III 133 1971, Beilstein 23 II 178, 23 III/IV 1235.]

1,8-Naphthyridine [254-60-4] has **m 99°, 98-99°, pK²⁰ 3.36**. Purify 1,8-naphthyridine through an Al₂O₃ column and elute with toluene and petroleum ether, evaporate the eluate, crystallise the residue from petroleum ether (b 60-80°), and sublime it at 80°/13mm. The *picrate* [15936-16-0] has **m 207-208°** (from EtOH), and the *methiodide* has **m 180-181°** (from EtOH). [Albert *J Chem Soc* 1790 1960, DOI: 10.1039/JR9600001790; Armarego *J Chem Soc (C)* 377 1967, DOI: 10.1039/J39670000377, Hawes & Wibberley *J Chem Soc (C)* 1564 1967, DOI: 10.1039/J39670001564; UV: Armarego in *Physical Methods in Heterocyclic Chemistry* (Ed Katritzky, Academic Press, LCCC No 62-12037, Internet Public Domain) Vol III 134 1971, Beilstein 23 II 178, 23 III/IV 1237.]

(±)-Naringenin (4',5,7-trihydroxyflavanone) [480-41-1, 67604-48-2] C₁₅H₁₂O₅, **M 272.3**, **m 247-254°, 251°** (phenolic pK_s~8-11). Crystallise it from EtOH or aqueous EtOH. It has UV with λ_{max} at 290nm (EtOH). The *S(-)-enantiomer* (natural form) has **m 255-256°** (from EtOH) and [α]_D²⁰ -28.0 (c 2, EtOH), [α]_D²⁰ -35.2 (c 1, pyridine). [Beilstein 18 H 503, 18 II 164, 18 III/IV 2630.] **Genistein (4',5,7-trihydroxyisoflavone)** [446-72-0] C₁₅H₁₂O₅, **M 270.2** crystallises from 60% aqueous EtOH or water with **m 297-298°** and [α]_D²⁰ -28 (c 0.6, 20mM NaOH). [Beilstein 18/4 V 594.]

Neutral Red (2-amino-8-dimethylamino-3-methylphenazine HCl, Basic Red 5, CI 50040) [553-24-2] $\text{C}_{15}\text{H}_{16}\text{N}_4 \cdot \text{HCl}$, **M 288.8, m 290°(dec), pK²⁵ 6.5**. Crystallise the dye from *benzene/MeOH (1:1). In aqueous solution the indicator is red at pH 6.8 and yellow at pH 8.0. Useful biological stain. [Beilstein 25 III/IV 3054.]

Nicotinaldehyde thiosemicarbazone [3608-75-1] $\text{C}_7\text{H}_8\text{N}_4\text{S}$, **M 180.2, m 222-223°, 225°**. Crystallise the derivative from EtOH, BuOH or water. Its *hydrochloride* [2104-92-9] crystallises from aqueous EtOH with **m 237-239° (dec)**. [Beilstein 21 III/IV 3542 21/7 V 342.]

Nicotinic acid (Niacin is also used for the acid, pyridine-3-carboxylic acid) [59-67-6] $\text{C}_6\text{H}_5\text{NO}_2$, **M 123.1, m 232-234°, pK₁²⁵ 2.00, pK₂²⁵ 4.82**. Crystallise it from *C₆H₆, EtOH or H₂O. It sublimes without decomposition. [McElvain *Org Synth Coll Vol* 1 385 1941, DOI: 10.15227/orgsyn.004.0049; Beilstein 22 III/IV 439, 22/2 V 57.] **Nicotinic acid hydrazide** [553-53-7] $\text{C}_6\text{H}_7\text{N}_3\text{O}$, **M 137.1, m 159-161°, pK₁²⁵ 2.2, pK₂²⁵ 3.63, pK₃²⁵ 11.49(NH)**. Crystallise it from aq EtOH or *C₆H₆. [Beilstein 22 III/IV 439, 22/2 V 121.]

Nile Blue A (a benzophenoxazinium sulfate dye) [3625-57-8] $(\text{C}_{20}\text{H}_{20}\text{N}_3\text{O})_2 \cdot \text{SO}_4$, **M 732.9, m >300°(dec), CI 51180, pK²⁵ 2.4**. Crystallise the dye from aqueous AcOH. It has UV with λ_{max} at 630nm (96% aqueous EtOH) and 635nm (H₂O). The *betaine* has UV with λ_{max} at 513nm (EtOH). [Crossley et al. *J Am Chem Soc* 74 578 1952, DOI: 10.1021/ja01123a002; Merrill & Spencer *J Am Chem Soc* 70 3683 1948, DOI: 10.1021/ja01191a043; Beilstein 27 II 457, 27 III/IV 5166.]

5-Nitrobarbituric acid (dilituric acid) [480-68-2] $\text{C}_4\text{H}_3\text{N}_3\text{O}_5$, **M 173.1, m 176°, 176-183°(dec), pK²⁰ 10.25**. Crystallise dilituric acid from water as the *trihydrate*, **m 180-181° (dec)**. Drying over 70% H₂SO₄ converts the trihydrate to the *dihydrate*. Used for detection of traces of potassium and for precipitation of alkaloids. [Loeffler & Moore *J Am Chem Soc* 70 3650 1948, DOI: 10.1021/ja01191a031]. [Beilstein 24 H 474, 24 II 273, 24 III/IV 1882.]

4'-Nitrobenzo-15-crown-5 [60835-69-0] $\text{C}_{14}\text{H}_{19}\text{NO}_8$, **M 313.3, m 84-85°, 93-95°**. Recrystallise the crown ether from EtOH, MeOH or *C₆H₆/hexane as for the 18-crown-6 compound below. It complexes with Na⁺, K⁺, NH₄⁺, Ca²⁺, Mg²⁺ and Cd²⁺. The ¹H NMR spectrum (CDCl₃) has δ (ppm) at 3.6-4.4 (m 16CH₂), 6.8 (d 1H arom), 7.65 (d 1H arom), 7.80 (dd 1H arom $J_{\text{ab}} = 9\text{Hz}$ and $J_{\text{bc}} = 3\text{Hz}$) [Schmid et al. *J Am Chem Soc* 98 5198 1976, DOI: 10.1021/ja00433a024; Kikukawa et al. *Bull Chem Soc Jpn* 50 2207 1977, DOI org/10.1246/bcsj.50.2207; Toke et al. *Justus Liebigs Ann Chem* 349, 1988, DOI: 10.1002/jlac.198819880408; 761 1988 DOI: 10.1002/jlac.198819880810; Lindner et al. *Z Anal Chem* 322 157 1985].

4'-Nitrobenzo-18-crown-6 [53408-96-1] $\text{C}_{16}\text{H}_{23}\text{NO}_8$, **M 357.4, has m 82-86°, 83-84°**. If impure and discoloured, then chromatograph it through Al₂O₃ and elute with *C₆H₆/hexane (1:1) containing 1% MeOH. The fractions are followed by TLC on Al₂O₃ (with Dragendorff's reagent for detection: R_F 0.6 in the above solvent system). Recrystallise the residues from the required fractions from *C₆H₆/hexane to give yellowish leaflets. It complexes with Na or K ions with logK_{Na} 3.95 and logK_K 4.71. [Petranek & Ryba *Coll Czech Chem Commun* 39 2033 1974, DOI: 10.1135/cccc19742033.]

4-(4-Nitrobenzyl)pyridine (PNBP) [1083-48-3] $\text{C}_{12}\text{H}_{10}\text{N}_2\text{O}_2$, **M 214.2, m 69-71°, 70-71°, 74°, pK_{Est} ~5.0**. Crystallise PNBP from aqueous EtOH or cyclohexane. The *hydrochloride* has **m 194-196°(dec, from EtOH)**, and the *picrate* has **m 168°(dec, from EtOH/EtOAc)**. [Beilstein 20 II 272, 20/7 V 564.]

5-Nitroindole [6146-52-7] $\text{C}_8\text{H}_6\text{N}_2\text{O}_2$, **M 162.1, m 140-142°, 141-142°, pK²⁵ -7.4 (aqueous H₂SO₄)**. Decolourise (charcoal) 5-nitroindole and recrystallise it twice from aqueous EtOH or recrystallise it from octane. It has UV has λ_{max} at 265 and 324nm (EtOH). [Beilstein 20 III/IV 3194, 20/7 V 41.]

Nitron [1,4-diphenyl-3-phenylamino-(1H)-1,2,4-triazolium (hydroxide) inner salt] [2218-94-2] $\text{C}_{20}\text{H}_{16}\text{N}_4$, **M 312.4, m 189°(dec), 189-190°(dec)**. Crystallise it from EtOH, chloroform or EtOH/*C₆H₆. [Beilstein 25 III/IV 1075.] It is used for the spectroscopic determination of nitrate and perchlorate.

5-Nitro-1,10-phenanthroline [4199-88-6] $\text{C}_{12}\text{H}_7\text{N}_3\text{O}_2$, **M 225.2, m 197-198°, 203°, 203-204°, pK²⁵ 3.33**. Crystallise the phenanthroline from *benzene/petroleum ether, until anhydrous. It also crystallises from H₂O with **m 202°**, and EtOAc with **m 203°**. Its pK²⁵ varies from 3.20 to 2.69 with varying MeOH/H₂O ratios from 0

to 0.95 moles/L, and from 3.20 to 1.95 in varying EtOH/H₂O ratios from 0 to 0.94 moles/L [Ram et al. *J Prakt Chem* **319** 719 1977, DOI: 10.1002/prac.19773190504]. It forms complexes with Cu²⁺, Zn²⁺, In²⁺, Fe²⁺, Co²⁺, Ni²⁺. [Beilstein **23** III/IV 1682, **23/8** V 425.]

3-Nitro-2-pyridinesulfonyl chloride [68206-45-1] C₅H₃ClN₂O₂S, M **190.2**, m **205°(dec)**, **217-222°(dec)**. The chloride crystallises as yellow needles from CH₂Cl₂. When pure, it is stable for several weeks at room temperature, and no decomposition was observed after 6 months at <0°. It is moisture sensitive. Its UV (MeCN) has λ_{max} at 231nm (ε 12,988), 264nm (ε 5,784) and 372nm (ε 3,117). [NMR and UV: Matsueda & Aiba *Chem Lett* 951 1978, DOI: org/10.1246/cl.1978.951; Wagner et al. *Chem Ber* **75** 935 1942, DOI: 10.1002/cber.19420750802.]

5-Nitroquinoline [607-34-1] C₉H₆N₂O₂, M **174.2**, m **70°, 71-73°, pK²⁰ 2.69**. Crystallise 5-nitroquinoline from pentane, then from *benzene. The *hydrochloride* has m **224°** and the *picrate* has m **206°, 214°**(from MeOH). [Beilstein **20** H 371, **20** II 235, **20** III/IV 3397.]

8-Nitroquinoline [607-35-2] has m **88-89°, 89-91°, 91-92°, pK²⁰ 2.55**. Crystallise 8-nitroquinoline from hot water, MeOH, EtOH or EtOH/diethyl ether (3:1). It sublimes at 70°/2mm. [Beilstein **20** H 373, **20** III/IV 3399.]

4-Nitroquinoline 1-oxide [56-57-5] C₉H₆N₂O₃, M **190.2**, m **154-155°, 154-156°, 157°**. The *N*-oxide recrystallises from aqueous acetone as yellow needles or platelets. [Ochiai *J Org Chem* **18** 534 1953, DOI: 10.1021/jo01133a010; Beilstein **20** III/IV 3396.]

5-Nitrouracil (2,4-dihydroxy-5-nitropyrimidine) [611-08-5] C₄H₃N₃O₄, M **157.1**, m **280-285°, >300°, pK²⁰ 0.03, pK²⁰ 5.55, pK²⁰ 11.3**. The uracil recrystallises in prisms from boiling H₂O as the *monohydrate* and loses H₂O on drying *in vacuo*. [UV: Brown *J Chem Soc* 3633(3647) 1959, DOI: 10.1039/JR9590003633, Note; Brown *J Appl Chem* **2** 239 1952, DOI: 10.1002/jctb.5010020502; Johnson *J Am Chem Soc* **63** 263 1941, DOI: 10.1021/ja01846a063; Beilstein **24** I 313, **24** II 171, **24** III/IV 1236.]

4-Nonadecylpyridine (hydrogen ionophore II [ETH 1907] - Proton ionophore) [70268-36-9] C₂₄H₄₃N, M **345.6**, b **180°/0.07mm**, pK_{Est} ~ **6.0**. Dissolve the waxy ionophore (ca 60g) in CHCl₃ (200ml), wash it with H₂O (3 x 200ml), dry it and evaporate it to dryness, then distil it in a vacuum. A waxy solid is formed on cooling the distillate. Its UV has λ_{max} at 257nm (ε 1.86 x 10³ M⁻¹cm⁻¹), 308nm (ε 1.7 x 10² M⁻¹cm⁻¹). [IR, NMR UV: Valenty et al. *Inorg Chem* **18** 2160 1979, DOI: 10.1021/ic50198a023.]

Octadecyl isonicotinate (hydrogen ionophore IV ETH 1778) [103225-02-1] C₂₄H₄₁NO₂, M **375.6**, m **57.5°, pK_{Est} ~ 3.5**. Dissolve it in Et₂O and wash it 3 times with H₂O. Dry the extract (MgSO₄), evaporate, and recrystallise the residue from EtOAc/hexane (4:1). Used as an amperometric proton selective sensor [Faisal, Pereira, Rho & Lee *Phys Chem Chem Phys* **12**(46) 15184 10/2010, DOI: 10.1039/c0cp00750a; Oesch et al. *Anal Chem* **58** 2285 1986, DOI: 10.1021/ac00124a037.]

Oxalylindigo [2533-00-8] C₁₈H₈N₂O₄, M **316.3**. It crystallises twice from nitrobenzene as small yellow needles and is dried by heating *in vacuo* for several hours. It is an example of a *cis*-indigo structure. [Schanze et al. *J Am Chem Soc* **108** 2646 1986, DOI: 10.1021/ja00270a024; Pummerer & Reuss *Chem Ber* **80** 242 1947, DOI: 10.1002/cber.19470800311; van Alphen *Recl Trav Chim Pays-Bas* **58** 378 1939, DOI: 10.1002/recl.19390580502.]

2-Oxazolidinone (ethylene carbamate) [497-25-6] C₃H₉NO₂, M **87.1**, m **83-87°, 88-90°, 89-90°, 91°, b 152°/0.4mm, 200°/12mm, 220°/48mm**. It is prepared by reaction of ethanolamine with phosgene or diethylcarbonate. It can be prepared from ethanolamine (2g) in CHCl₃ (200ml, EtOH free by passing through an Al₂O₃ column) by bubbling COCl₂ through the solution which is allowed to stand for 3 hours, the acid is neutralised with powdered PbCO₃, filtered, evaporated to dryness and the solid residue is recrystallised from CHCl₃. Alternatively, ethanolamine (61g), Et₂CO₃ (150ml) and NaOMe (0.5g) are heated in an oil bath; and after the EtOH (~112ml) has distilled off, the residue solidifies on cooling and is recrystallised from CHCl₃ (100ml) to give the *oxazolidinone* (57g, 65%), m **87-89°**. Its IR (film) has ν_{max} at 3000, 2920 (CH), 1766 (carbamate, C=O), 1690 (amide, C=O), 1465, 1382 and 1357 (CH₂), 1298 (C-N), 1212, 1140 and 1030 (COC,

C-O), 953 cm⁻¹; and ¹H NMR (CDCl₃) with δ at 6.68 (NH, br s, 1H), 4.47 (H₅, t, ³J = 8Hz, 2H), 3.63 (H₄, t, ³J = 8Hz, 2H); and for ¹³C NMR see references [Hammer et al. *J Org Chem* **46** 1521 1981, DOI: 10.1021/jo00321a002.] [Homeyer US Pat 2,399,118 1964, *Chem Abs* **40** 4084 1964.] It can be recrystallised from *C₆H₆, dichloroethane, CHCl₃ or EtOH. It is a cyclic urethane, and is not very stable in aqueous solvents. The *N*-acetyl derivative [1432-43-5] **M 129.1, m 69-70°**, is formed by boiling 2-oxazolidinone (3g) with Ac₂O (20ml) and NaOAc (1g) for 1.5 hours, and recrystallised from *C₆H₆/Et₂O or sublimed at 65° *in vacuo*. [Homeyer US Pat 2,399,118 1964, *Chem Abs* **40** 4084 1964.] The *N*-methyl derivative [19836-78-3] **M 101.1, b 120°/0.1mm**, is made from *N*-methyl ethanolamine with COCl₂/CHCl₃/PbCO₃ or as in the preparation of the parent compound [with (EtO)₂CO], and distilled in a vacuum. It is **TOXIC**. The yellow *picrolonate* has **m 137-138°**. [cf. Fränkel & Cornelius *Chem Ber* **51** 1654 1918, DOI: 10.1002/cber.19180510251; Ben-Ishai *J Am Chem Soc* **78** 4962 1956, DOI: 10.1021/ja01600a042; *Beilstein* **27** H 135, **27** I 259, **27** III/IV 2516.]

Oxetane (1,3-trimethylene oxide) [503-30-0] C₃H₆O, **M 58.1, m -97°, b 45-46°/736mm, 47-49°/atm, 48°/760mm, d₄²⁰ 0.892, n_D²⁰ 1.395**. Distil it twice from sodium and fractionate it through a small column at 47.0-47.2°/atm. Also purified by preparative GC using a 2m silica gel column. *Alternatively*, add KOH pellets (50g for 100g of oxetane) and distil through an efficient column packed with 1/4in Berl Saddles; and the main portion at b 45-50° is redistilled over fused KOH. [Noller *Org Synth Coll Vol* **3** 835 1955, DOI: 10.15227/orgsyn.029.0092; Dittmer et al. *J Am Chem Soc* **79** 4431 1957, DOI: 10.1021/ja01573a051.]

Oxetan-2-one (β-propiolactone, propan-3-olide) [57-57-8] C₃H₄O₂, **M 72.1, m -35°, -33°, -31.2°, b 51°/10mm, 83°/45mm, 162°/atm, d₄²⁰ 1.1460, n_D²⁵ 1.4117**. Fractionally distil the lactone from sodium under reduced pressure. It gives an acidic solution in H₂O due to hydrolysis. Produces insoluble polymers on storage. It irritates the skin and is a possible **carcinogen**. [*Beilstein* **17** I 130, **17** III/IV 4157.]

Oxine Blue (8-hydroxy-5-*p*-diethylaminophenylimino-5,8-dihydroquinoline) [3733-85-5] C₁₉H₁₉N₃O, **M 369.4, m 134-135°, pKa 4.70**. Recrystallise the dye from EtOH and dry it in a desiccator over H₂SO₄. Useful indicator (a few drops of 0.25% EtOH solution) for strong acids—strong base, and *vice versa*, titrations where it is orange below pH 3.9 and blue above pH 5.50 (see pKa). [Prepn details: Bishop *Indicators* pp 127-128, Oxford Pergamon Press 1972, Library of Congress Catalog Card No 78-171464, ISBN 0080166172, 9780080166179, Lev Zhur *Anal Khim* **11** 359 1956.]

Paraldehyde (acetaldehyde trimer, 2*r*,4*c*,6*c*-trimethyl-1,3,5-trioxane, all-*cis*) [123-63-7] C₆H₁₂O₃, **M 132.2, m 12.5°, b 124°/751mm, 123-124°/atm, d₄²⁰ 0.995, n_D²⁰ 1.407**. Wash paraldehyde with water and fractionally distil it. *Alternatively*, it is purified by drying with anhydrous Na₂SO₄, then cooled to 5°, and the frozen material is separated by decantation. The solid is distilled (**b 121-124°/atm**), the distillate is collected, stored over anhydrous Na₂SO₄ for several days and re-distilled at atmospheric pressure before use [Le Fevre et al. *J Chem Soc* 290 1950, DOI: 10.1039/JR9500000290]. The 2*r*,4*c*,6*t*-trimethyl-1,3,5-trioxane has **m 14.5°, b 125°/760mm**. [*Beilstein* **19** II 394, **19** III/IV 4715, **19** V 112.]

Patulin [hydroxyl-4*H*-furo(3,2-*c*)pyran-2(6*H*)-one, Clavatin] [149-29-1] C₇H₆O₄, **M 154.1, m 110°, 111-112°, [α]_D²⁰ -74 (CHCl₃)**. Recrystallises it from Et₂O, EtOH or *C₆H₆ (prisms or thick plates), and sublime it at 90°/high vacuum. The *acetate* has **m 118-120°** (from 50% aqueous EtOH). [Bergel et al. *J Chem Soc* 415 1944, DOI: 10.1039/JR9440000415; *Beilstein* **18** III/IV 1184, **18**/3 V 5.] Highly **TOXIC, CARCINOGENIC**.

Pentachloropyridine [2176-62-7] C₅Cl₅N, **M 251.3, m 122-124°, 123°, 123-127°, 124°, 124-125°, 125.5°, 125-126°, b 279-280°/atm, pK₂₀ -6.02 (aqueous H₂SO₄)**. Purify it by recrystallisation from EtOH or aqueous EtOH and sublime at 150°/3mm. [den Hertog et al. *Recl Trav Chim Pays-Bas* **69** 673 1950, DOI: 10.1002/recl.19500690604; Schickh et al. **69** 2593 1936, DOI: 10.1002/cber.19360691202; *Beilstein* **20**/5 V 422.]

Pentafluoropyridine [700-16-3] C₅F₅N, **M 169.1, m -41.5°, b 83.5°, 84°, 83-85°, d₄²⁰ 1.609, n_D²⁰ 1.3818, pK_{Est} ~<0**. Distil it through a concentric tube column; it has λ_{max} in cyclohexane at 256.8nm. [Chambers et al. *J Chem Soc* 3573 1964, DOI: 10.1039/JR9640003573; ¹⁹F NMR: Bell et al. *J Fluorine Chem* **1** 51 1971, DOI:10.1016/S0022-1139(00)82533-6.] The *hexafluoroantimonate* has **m 98-102°(dec)** after crystallisation from liquid SO₂. [*Beilstein* **20**/5 V 401.]

3,3,6,9,9-Pentamethyl-2,10-diazabicyclo[4.4.0]dec-1-ene (PMDBD, 1,2,3,4,4a,5,6,7-octahydro-2,2,4a,7,7-pentamethylnaphthyridine) [69340-58-5] $C_{13}H_{24}N_2$, M 208.3, m 15°, b 65°/0.07mm, d_4^{20} 0.924, n_D^{20} 1.4840, $pK > 11$ (MeOCH₂CH₂OH/H₂O, 1:1). The pentamethylhydronaphthyridine is obtained by intramolecular cyclisation of 5-amidino-2,5,8-trimethylnona-2-7-diene [190.0g, 0.914mol, prepared from 2mols BuLi, one mol of propionitrile and 2mols of 3-methylbut-2-enyl bromide (isoprenhydrobromide) in THF at -78°, followed by NaNH₂ in boiling *C₆H₆] in CH₂Cl₂ (500ml) solution at 0°, by bubbling dry HCl gas through it until the pH is 4. The solution is evaporated *in vacuo* until free from HCl vapour to give a dark hygroscopic hydrochloride which is heated in a flask (protected from air with a CaCl₂ tube) at 200°, whereby excess of HCl is released after 30 minutes (cyclisation begins at *ca* 125°), and heating is continued for 18 hours. The dark brown residue is treated with H₂O (1.0L), acidified with 2N H₂SO₄ (500ml) and the aqueous phase is extracted with Et₂O (2 x 100ml), the aqueous phase is basified (cooling is necessary) with NaOH (*ca* 100g) to pH >10. The brown oil that separates is extracted with Et₂O (2 x 500ml), dried over K₂CO₃, filtered, and evaporated to give a viscous oil (190g) which is distilled through a Vigreux column at 0.05mm, and after a small fore run (3.8g), a colourless viscous oil of the *naphthyridine* distils (156.6g, 80% yield, b 60-67°). An analytical sample has m ~15° and b 65°/0.07mm. Store it in an inert dry atmosphere as it absorbs CO₂ in the presence of moisture to form a *bicarbonate* (see below). Its UV/VIS has λ_{max} (ϵ) at 219 (11,600) nm; the IR (CHCl₃) of the *hydrochloride* has an immonium band at ν_{max} 2500 cm⁻¹; the ¹H NMR (CCl₄) of the *free base* has δ at 1.08-1.13 (2s, 12H, 4CH₃), 1.19 (s, 3H, CH₃), 1.2-2.0 (m, 8H), 3.38 (s, NH); see reference for full IR and MS

The *hydrochloride*, when recrystallised twice from CH₂Cl₂/Et₂O (15 hours at 0°), has m 195-196°, and the ¹H NMR (CCl₄) has δ at 1.34-1.13 (2s, 9H, 3CH₃), 1.42 (s, 6H, 2CH₃), 1.6-2.3 (m, 8H), 9.78 (br s, 2H NH).

The *hydrogen carbonate salt* is formed from the free base and moist CO₂ in 2 minutes, then by extraction with CH₂Cl₂, drying (K₂CO₃) and evaporating gives a labile salt decomposing at 122-125°, and for ¹H NMR see references. The *benzoate* crystallises from CH₂Cl₂ with m 183-185°, the *hydrogen sulfate* crystallises from CH₂Cl₂/hexane with m 208-209°, the *tosylate* has m 114°, and the *dihydrogen phosphate* crystallises from MeOH/Et₂O with m 225-228°. [Heinzer, Soukup and Eschenmoser *Helv Chim Acta* **61** 2851 1978, DOI: 10.1002/hlca.19780610812; Sternbach et al. *Angew Chem* **91** 670 1979, DOI: 10.1002/ange.19790910827; *Beilstein* **23/5** V 302.] The strong basicity, proximity of the nitrogen atoms, and the steric hinderance from methyl groups around them, confer remarkable coordination properties [Sternbach et al. *Angew Chem* **91** 670 1979, DOI: 10.1002/ange.19790910827; Boyle et al. *JCS Chem Commun* 239 1992, DOI: 10.1039/C39920000239; Denmark *J Org Chem* **46** 3144 1981, DOI: 10.1021/jo00328a038].

Pentaquine monophosphate (1,4-pentanediamine, *n*-[6-methoxy-8-quinoliny]-*N'*-[1-methylethyl] (1:1) phosphate) [5428-64-8] $C_{18}H_{27}N_3O_2 \cdot H_3PO_4$, M 399.4, m 189-189.5°, 189-190°, pK^{70} 8.22. Crystallise it from H₂O or 95% EtOH (pale yellow crystals), and dry at 50° *in vacuo*. The *free base* has b 165-170°/0.02mm, n_D^{25} 1.5785. The *picrate* crystallises from Me₂CO/EtOH with m 164.5-165.5°. [Drake et al. *J Am Chem Soc* **68** 1529 1946, DOI: 10.1021/ja01212a041; *Beilstein* **22** III/IV 5814.] It is an antimalarial.

Phenanthridine (benzo[c]quinoline, 3,4-benzoquinoline) [229-87-8] $C_{13}H_9N$, M 179.2, m 104-107°, 106.5°, 108-109°, b 349°/769mm, 350°/atm, pK^{20} 4.61 (4.48). Purify it *via* the HgCl₂ addition compound formed when phenanthridine (20g) in 1:1 HCl (100ml) is added to aqueous HgCl₂ (60g in 3L), and the mixture is heated to boiling. The *HgCl₂ complex* separates as yellow red crystals with m 195-198° [Arcus & Mesley *J Chem Soc* 178 1953, DOI: 10.1039/JR9530000178]. Concentrated HCl is then added until all of the solid has dissolved. The compound separates on cooling and is decomposed with aqueous NaOH (*ca* 5M). Phenanthridine is extracted into Et₂O, evaporated, and the residue is crystallised from petroleum ether (b 80-100°) or EtOAc. [Cumper et al. *J Chem Soc* 4518 1962, DOI: 10.1039/JR9620004518.] It is also purified by chromatography on activated alumina from *benzene solution, with diethyl ether as eluent. Evaporation of ether gives crystalline material which is freed from residual solvent under vacuum, then further purified by fractional crystallisation, under N₂, from its melt. It was purified by zone melting and sublimes in a vacuum. The *picrate* has m 218.5-219.5° (from *iso*-PrOH) (also reported are m 244-245° and 247-248°, from EtOH or H₂O). [Slough & Ubbelohde *J Chem Soc* 911 1957, DOI: 10.1039/JR9570000911.] [*Beilstein* **20** H 466, **20** III/IV 4016, **20/8** V 223.]

1,10-Phenanthroline (*o*-phenanthroline) [66-71-7 (*anhydr*); 5144-89-8 (*H₂O*)] $C_{12}H_8N_2$, M 198.2, m 98-101°, 100-104°, 108-110° (hydrate), 114-117°, 118° (*anhydrous*), b >300°, pK_1^{25} -0.7 (aqueous HClO₄), pK_2^{25} 4.86 (4.96). Crystallise its *picrate* (m 191°) from EtOH; then the *free base* is liberated with aqueous alkali,

dried at 78°/8mm over P₂O₅ and crystallised from petroleum ether (b 80-100°). [Cumper et al. *J Chem Soc* 1188 1962, DOI: 10.1039/JR9620001188.] It can be purified by zone melting. It has also been crystallised from hexane, *benzene/petroleum ether (b 40-60°) or sodium-dried *benzene, dried and stored over H₂SO₄. The **monohydrate** is obtained by crystallisation from aqueous EtOH or ethyl acetate. It has been crystallised from H₂O (300 parts) to give the **monohydrate m 102-103°** which sublimes at 10⁻³mm [Fielding & LeFevre *J Chem Soc* 1811 1951, DOI: 10.1039/JR9510001811.] The **anhydrous** compound has **m 118°** (after drying at high vacuum at 80°) and is also obtained by recrystallisation from petroleum ether or *C₆H₆ (70 parts) and drying at 78°/8mm. [UV: Badger et al. *J Chem Soc* 3199 1951, DOI: 10.1039/JR9510003199.] It has a pK_a in H₂O of 4.857 (25°) or 5.02 (20°) and 4.27 in 50% aqueous EtOH (20°). [Albert et al. *J Chem Soc* 2240 1948, DOI: 10.1039/JR9480002240]. [Beilstein **23** H 227, **23** II 235, **23/8** V 419.]

1,10-Phenanthroline hydrochloride (o-phenanthroline hydrochloride) [3829-86-5 (HCl.H₂O), 18851-33-7 (xHCl.H₂O)] C₁₂H₈N₂. HCl. H₂O, **M 243.7**, has **m 212-219°**. The hydrochloride crystallises from 95% EtOH, **m 212-219°** as the **monohydrate**; the **half hydrate** has **m 217°**. The **3HCl** has **m 143-145°** (sinters at 128°). [Thevenet et al. *Acta Cryst Sect B* **33** 2526 1977, DOI: 10.1107/S0567740877008838]. [Beilstein **23** II 235, **23/8** V 415, 421.]

4,7-Phenanthroline-5,6-dione [84-12-8] C₁₂H₆N₂O₂, **M 210.2**, **m 295°(dec)**. The dione crystallises from MeOH, and the white crystals are dried at 100°/0.1mm/6hrs. The **mono-oxime** forms yellow crystals from MeOH with **m 250°(dec)**, the **di-oxime** forms yellow crystals from MeOH with **m 300°(dec)** and the **mono-semihydrazone** forms yellow crystals from MeOH with **m 195°(dec)**. [Druey & Schmidt *Helv Chim Acta* **33** 1080 1950, DOI: 10.1002/hlca.19500330438; Shabir & Forrow *J Chromatogr Sci* **43** 207 2005, DOI:10.1093/chromsci/43.4.207; Beilstein **24** III/IV 1741.]

Phenazine (dibenzo[*b,e*]pyrazine) [92-82-0] C₁₂H₈N₂, **M 180.2**, **m 172-176°, 171°, pK₁²⁰ -4.9 (aqueous H₂SO₄), pK₂²⁰ 1.21**. Phenazine crystallises from EtOH, CHCl₃ or ethyl acetate, after pre-treatment with activated charcoal. It can be sublimed *in vacuo* and purified by zone refining. [Beilstein **23/8** V 389.] The **5-methylphenazinium methyl sulfate** [299-11-6] C₁₃H₁₁N₂. CH₃SO₄, **M 306.3**, **m 158-160°(dec)**, is used with ascorbic acid to determine nitric oxide reductase activity [Heiss et al. *J Bacteriol* **171**(6) 3288 1989, DOI: jb.asm.org/content/171/6/3288.full.pdf].

Phenosafranine (3,7-diamino-5-phenylphenazinium chloride, CI 50200) [81-93-6] C₁₈H₁₅ClN₄, **M 322.8**, **m >300°, λ_{max} 530nm (H₂O)**. Crystallise the dark green powder from dilute HCl. The red solution has UV with λ_{max} at 517-521nm in H₂O. The **picrate** decomposes on heating and has a solubility of 0.0048% in H₂O at 18°. [Beilstein **23** H 395, Beilstein **25** H 394, **25** I 654, **25** II 338, **25** III/IV 3050.] It is a biological stain.

Phenothiazine [92-84-2] C₁₂H₉NS, **M 199.3**, **m 182-187°, 184-185°, 185°, b 371°/atm**. Crystallise it from *benzene, toluene, hexane or Me₂CO (charcoal) after boiling for 10 minutes under reflux. Filter the crystals off and dry them in an oven at 100°, then in a vacuum desiccator over paraffin chips. Also recrystallise it twice from water and dry it in an oven at 100° for 8-10 hours. It sublimes at 130°/1mm and has UV with λ_{max} at 253nm in heptane. [Beilstein **27** I 225, **27** II 32, **27** III/IV 1214.] Insecticide and anthelmintic.

Phenoxazine [135-67-1] C₁₂H₉NO, **M 199.2**, **m 156°, 156-158°, 158-159°, b 215°/4mm**. Crystallise phenoxazine from EtOH and sublime it *in vacuo*. If too impure then extract it in a Soxhlet extractor using toluene. Evaporate the solvent and dissolve the residue (ca 100g) in *C₆H₆ (1L), **CARCINOGEN** (use an efficient fume cupboard) and chromatograph it through an Al₂O₃ column (50 x 450 mm) using *C₆H₆. The eluent (ca 3L) is evaporated to ca 150ml and cooled when ca 103g of phenoxazine **m 149-153°** are obtained. Sublimation yields platelets **m 158-159°**. It forms a green **picrate m 141.5-142°**. [Gilman & Moore *J Am Chem Soc* **79** 3485 1957, DOI: 10.1021/ja01570a048; Müller et al. *J Org Chem* **24** 37 1959, DOI: 10.1021/jo01083a011; Beilstein **27** I 223, **27** III/IV 1209.]

2-Phenyl-1-azaindolizine (-phenylimidazo[1,2-a]pyridine) [4105-21-9] C₁₃H₁₀N₂, **M 194.2**, **m 140°, pK_{Est} ~1.9**. Crystallise the indolizine from EtOH, *benzene/petroleum ether, hexane (**m 135-136°**) or cyclohexane (**m 136-137°**). The **hydrochloride 2H₂O** has (**m 114-116°**, from H₂O), and the **picrate** has **m 228-229°** (from AcOH) and **236-238°** (from Me₂CO). The **hydrobromide** has **m 161-163° (165°)**, after recrystallisation from

EtOH/EtOAc and drying at 56°/0.05mm. [Adams & Dix *J Am Chem Soc* **80** 4618 1958, DOI: 10.1021/ja01550a051; *Beilstein* **23** III/IV 1705.]

2-Phenyl-1,3-diazaheptahydroazulene [2161-31-1] $C_{13}H_{18}N_2$, **M 212.3**. Recrystallise the azulene three times from de-aerated cyclohexane in the dark. Dry it *in vacuo*.

9-Phenyl-3-fluorone (2,6,7-trihydroxy-9-phenylxanthen-3-one) [975-17-7] $C_{19}H_{12}O_5$, **M 320.3**, **m >300°(dec)**, **350°**, λ_{max} **462nm** (ϵ **4.06 x 10⁴**, in 1M HCl aqueous EtOH). Recrystallise it from warm, acidified EtOH by addition of ammonia. The crude material (1g) can be extracted with EtOH (50ml) in a Soxhlet apparatus for 10 hours to remove impurities. Impurities can be detected by paper electrophoresis. The *triacetate* forms yellow needles from EtOH (**m 230-233°**). [Petrova et al. *Anal Lett* **5** 695 1972, DOI: 10.1080/00032717208064350; *Beilstein* **18** H 199, **18** I 404, **18** III/IV 2824.] Possible eye, skin and respiratory tract irritant.

2-Phenylindolizine [25379-20-8] $C_{14}H_{11}N$, **M 193.2**, **m 211-214°**, **214°(dec)**, **pK_{Est} ~4.4**. Crystallise 2-phenylindolizine from EtOH. The *0.25HCl* crystallises from MeCN with **m 109°**, and the *picrate* has **m 161°** when crystallised from EtOAc. [*Beilstein* **20** II 304, **20** III/IV 4033, **20/8** V 244.]

1-Phenyl-5-mercaptopotetrazole [86-93-1] $C_7H_6N_4S$, **M 178.2**, **m 143-147° (dec)**, **145° (dec)**, **pK²⁵ 3.65 (5% aqueous EtOH)**. Purify the tetrazole by recrystallisation from EtOH or $CHCl_3$ (**m 152°**) [Tautomerism: Kauer & Sheppard *J Org Chem* **32** 3580 1967, DOI: 10.1021/jo01286a064; UV: Lieber et al. *Can J Chem* **37** 563 1959, DOI: 10.1139/v59-077]. The *ammonium salt* crystallises from EtOH and decomposes at 176°. The *sodium salt* [15052-19-4] **M 200.2**, crystallises from EtOH/* C_6H_6 , melts at 96° and decomposes at 145° [Stollé *J Prakt Chem* [2] **133** 60 1932, DOI: 10.1002/prac.19321330106]. It is used for the determination of Bi and Pd. [Fresenius *Z Anal Chem* **261** 151 1972, *Beilstein* **26** III/IV 2065.]

4-(3-Phenylpropyl)pyridine [1-phenyl-3(4-pyridyl)propane] [2057-49-0] $C_{14}H_{15}N$, **M 197.3**, **b 150-152°/5-6mm**, **322°/atm**, d_4^{25} **1.03**, n_D^{20} **1.563**, **pK_{Est} ~6.0**. This is prepared from the potassium salt of 4-picoline [obtained by dissolving K metal (1 mol) and a few mg of ferric oxide as catalyst in liquid NH_3 until the blue colour is discharged, indicating the formation of KNH_2 , to which is added 4-picoline (1 mol) rapidly forming a reddish-amber solution] and phenethyl bromide (1 mol, see [103-63-9]), allowing the NH_3 to evaporate (6-10 hours, in an efficient fume cupboard), adding H_2O , the mixture is extracted into Et_2O , evaporated to dryness and the residual oil is distilled, preferably under a vacuum to give the desired *phenylpropylpyridine*, **b 150-152°/5-6mm**, in 56% yield. The *hydrochloride*, obtained by evaporating a concentrated HCl solution of the base on a steam bath and the residual solid, recrystallised from EtOH/ Et_2O , has **m 143.5°** and is hygroscopic. The *N-oxide* [34122-28-6] $C_{14}H_{15}NO$, **M 213.3**, has **m 58-65°**. [Bergstrom et al. *J Org Chem* **10** 452 1945, DOI: 10.1021/jo01181a011; *Beilstein* **20** III/IV 3687.]

4-Phenylpyridine-2-carbonitrile [18714-16-4] $C_{12}H_8N_2$, **M 180.1**, **m 97-101°**, **b 342.4°/760mm**, **pK_{Est} <0**. Purify the nitrile by recrystallisation from petroleum ether (**m 99-100°**). [Case & Kasper *J Am Chem Soc* **78** 5842 1956, DOI: 10.1021/ja01603a036; *Beilstein* **22** III/IV 1261.]

Phenyl 2-pyridyl ketoxime [1826-28-4] $C_{12}H_{10}N_2O$, **M 198.2**, **m 151-152°**, **154-156°**, **pK₁²⁵ 3.84**, **pK₂²⁵ 10.71 for E-isomer**. The *E-isomer* crystallises from EtOH (charcoal). It isomerises to the *Z-isomer* on melting or in boiling *o*-xylene, and crystallises from EtOH or cyclohexanol with **m 166-168°**. [*Beilstein* **21** H 330, **21** III/IV 4120, **21/8** V 568.]

6-Phenylquinoline [612-95-3] $C_{15}H_{11}N$, **M 205.3**, **m 110.5-111.5°**, **pK_{Est} ~5.2**. Crystallise 6-phenylquinoline from EtOH (charcoal). The *picrate* has **m 105°** (from Me_2CO). It has UV with λ_{max} at 253nm in aqueous EtOH and 263 and 325nm in aqueous HCl. [*Beilstein* **20** H 483, **20** III/IV 4151.]

2-Phenylquinoline-4-carboxylic acid (Cinchophen, Atophen) [132-60-5] $C_{16}H_{11}NO_2$, **M 249.3**, **m 214-215°**, **215°**, **pK_{Est(1)} ~0.5 (CO₂H)**, **pK_{Est(2)} ~5.1 (N)**. It crystallises from EtOH (*ca* 20ml/g), in several modifications with **m 196°** and **216°**(subliming at 65°). [*Beilstein* **22** H 103, **22** II 518, **22** II 70, **22** III/IV 1358.]

1-Phenyl-5-sulfanilamidopyrazole [526-08-9] $C_{15}H_{14}N_4O_2S$, M 314.4, m 177-178°, 178-179°. Crystallise it from EtOH or aqueous EtOH. Methylation with $Me_2SO_4/NaOH$ occurred on the amido nitrogen atom to give the *N*¹-methyl derivative with m 212-213.5° (from EtOH). [Schmidt & Druey *Helv Chim Acta* **41** 306 1958, DOI: 10.1002/hlca.660410138; Eichenberger et al. *Helv Chim Acta* **48** 524 1963, DOI: 10.1002/hlca.19650480310; *Beilstein* **25** III/IV 2029.]

4-Phenyl-1,2,4-triazolidine-3,5-diol (4-phenylurazole) [15988-11-1] $C_8H_7N_3O_2$, M 175.2, m 207-209°. Crystallise 4-phenylurazole from water or 95% EtOH. Dissolve 35g in 80ml of boiling 95% EtOH and on cooling 90-95% is recovered with m 209-210°. It has IR with ν_{max} at 1685 and 3120 cm^{-1} . [Cookson et al. *Org Synth* **51** 121 1971, DOI: 10.15227/orgsyn.051.0121; *Beilstein* **26** I 64, **26** III/IV 540.]

4-Phenyl-1,2,4-triazole-3,5-dione (PTAD) [4233-33-4] $C_8H_5N_3O_2$, M 177.2, m 165°(dec), 165-170°(dec), 170-177°(dec). PTAD forms carmine red needles by sublimation (ice cold finger) at 100°/0.1mm, and/or by recrystallisation from EtOH. Its IR has ν_{max} at 1760 and 1780 cm^{-1} . [Cookson et al. *Org Synth* **51** 121 1971, DOI: 10.15227/orgsyn.051.0121; Moore et al. *J Org Chem* **39** 3799 1974, DOI: 10.1021/jo00939a049; *Beilstein* **26** I 57, **26** III/IV 540.] Useful reagent for selective oxidation of thiols to disulfides [Christoforou et al. *Tetrahedron Lett* **47** 9211 2006, DOI: 10.1016/j.tetlet.2006.10.134].

9-Phenyl-9-xanthenol (hydroxypixyl) [596-38-3] $C_{19}H_{14}O_2$, M 274.3, m 158-161°, 158.5-159°, 159°. Dissolve hydroxypixyl in AcOH and add H_2O whereby it separates as colourless prisms. It is slightly soluble in $CHCl_3$, soluble in $*C_6H_6$ but insoluble in petroleum ether. It sublimes on heating. Its UV in H_2SO_4 has λ_{max} at 450nm (ϵ 5620) and 370nm (ϵ 24,900) and the $HClO_4$ salt in $CHCl_3$ has λ_{max} at 450 (ϵ 404) and 375nm (ϵ 2420). [Sharp *J Chem Soc* 2558 1958, DOI: 10.1039/JR9580002558; Bünzly & Decker *Chem Ber* **37** 2931 1904, DOI: 10.1002/cber.19040370371; Chattopadhyaya & Reece *JCS Chem Commun* 639 1978, DOI: 10.1039/C39780000639; Gomberg & Cone *Justus Liebigs Ann Chem* **370** 142 1909, DOI: 10.1002/jlac.19093700110; *Beilstein* **17** I 80, **17** II 161, **17** III/IV 1704, **17/4** V 675.]

'Phosphine' [dye CI 793, Chrysaniline mononitrate, 3-amino-9-(4-aminophenyl)-acridinium mononitrate] [10181-37-0] $C_{19}H_{15}N_3$. HNO_3 , M 348.4, m >250°(dec), pK^{20} 7.71 (50% aqueous EtOH). Crystallise the dye from $*benzene/EtOH$. The *free base* crystallises from $*C_6H_6$ in yellow crystals m 229-230°. [Dunstan & Hewitt *J Chem Soc* **89** 483 1906, DOI: 10.1039/CT9068900482; pK ; Albert & Goldacre *J Chem Soc* 706 1946, DOI: 10.1039/JR9460000706; *Beilstein* **22** H 91, **22** I 651, **22** II 403, **22** III/IV 5513.]

Phthalazine [253-52-1] $C_8H_6N_2$, M 130.2, m 89-92°, 90-91°, b 175°/17mm, 189°/29mm, 190°/30mm, pK^{20} 3.47. Phthalazine crystallises from diethyl ether or $*benzene$, and sublimes under a vacuum. The *hydrochloride* forms needles from EtOH with m 235-236°(dec) and the *picrate* has m 208-210°. [Armarego *J Appl Chem* **11** 70 1961, DOI: 10.1002/jctb.5010110207; Gabriel & Eschenbach *Chem Ber* **30** 3022 1897, DOI: 10.1002/cber.189703003116; Stephenson *Chem & Ind* 174 1957, *Beilstein* **23** H 174, **23** III/IV 1233.]

Phthalazine-1,4-dione (phthalhydrazide) [1445-69-8] $C_8H_6N_2O_2$, M 162.2, m 330-333°, 336°, 346°, pK^{20}_1 - 3.29 pK^{20}_2 -0.99, pK^{20}_3 5.67, pK^{20}_4 13.0. Recrystallise it twice from 0.1M KOH [Merenyi et al. *J Am Chem Soc* **108** 7716 1986, DOI: 10.1021/ja00284a040], EtOH or dimethylformamide and it sublimes >300°. [*Beilstein* **24** H 371, **24** II 194.]

Phthalazone (1-hydroxyphthalazine) [119-39-1] $C_8H_6N_2O$, M 146.2, m 183-184°, 186-188°, b 337°/760mm, pK^{20}_1 -2.2, pK^{20}_2 -1.4, pK^{20}_3 11.99. Phthalazone crystallises from H_2O or EtOH and sublimes *in vacuo*. [*Beilstein* **24** H 142, **24** III/IV 400.]

2-Picoline-N-oxide (2-methylpyridine-1-oxide) [931-19-1] C_6H_7NO , M 109.1, m 41-45°, b 89-90°/0.8-0.9mm, 90-100°/1mm, 110°/4mm, 135°/5mm, 123°/9mm, 123-124°/15mm, 259-261°/atm, n_D^{25} 1.5854 (supercooled), pK^{25} 1.10. Purify the *N*-oxide by fractional distillation, and it can be recrystallised from $*C_6H_6$ /hexane but is *hygroscopic*. [Bullitt & Maynard *J Am Chem Soc* **76** 1370 1954, DOI: 10.1021/ja01634a054; Ross et al. *J Am Chem Soc* **78** 3625 1956, DOI: 10.1021/ja01596a019; IR: Wiley & Slaymaker *J Am Chem Soc* **79** 2233 1957, DOI: 10.1021/ja01566a054.] The *picrate* has m 125-126.5° (from EtOH) [Boekelheide & Linn *J Am Chem Soc* **76** 1286 1954, DOI: 10.1021/ja01634a026]. The *phthalate* has m

115-116° (from EtOH) [den Hertog et al. *Recl Trav Chim Pays-Bas* **70** 591 1951, DOI: 10.1002/recl.19510700705.]. [*Beilstein* **20** III/IV 2689, **20/5** V 479.]

3-Picoline-N-oxide (3-methylpyridine-1-oxide) [1003-73-2] has **m 37-39°, 37-38°** (evacuated capillary), **84-85°/0.3mm**, **101-103°/0.7-0.8mm**, **114-115°/1.5mm**, **118°/2mm**, **150°/15mm**, **pK²⁵ 1.08**. Purify the *N*-oxide by careful fractionation *in vacuo*. The distillate remains supercooled for several days before solidifying. It is a slightly *hygroscopic* solid which could melt in the hand. The *picrate* has **m 149-151°** (from EtOH). [Taylor & Croveti *Org Synth Coll Vol* **4** 654 1963, DOI: 10.15227/orgsyn.036.0053; IR: Katritzky et al. *J Chem Soc* 3680 1959, DOI: 10.1039/JR9590003680; Jaffé & Doak *J Am Chem Soc* **77** 4441, 4481 1955, DOI: 10.1021/ja01622a001; Boekelheide & Linn *J Am Chem Soc* **76** 1286 195, DOI: 10.1021/ja01634a026]. [*Beilstein* **20** III/IV 2719, **20/5** V 517.]

4-Picoline-N-oxide (4-methylpyridine-1-oxide) [1003-67-4] has **m 182-184°, 185-186°, 186-188°, pK²⁵ 1.29**. Recrystallise the *N*-oxide from EtOH/EtOAc, Me₂CO/Et₂O or *C₆H₆. [Bullitt & Maynard *J Am Chem Soc* **76** 1370 1954, DOI: 10.1021/ja01634a054; Boekelheide & Linn *J Am Chem Soc* **76** 1286 1954, DOI: 10.1021/ja01634a026]. [*Beilstein* **20** III/IV 2741, **20/5** V 558.]

Picolinic acid (pyridine-2-carboxylic acid) [98-98-6] C₆H₅NO₂, **M 123.1**, **m 138°, 138-142°, pK₁²⁵ 1.03 (1.36), pK₂²⁵ 5.30 (5.80)**. Crystallise the acid from water or *benzene. The *picrate* has **m 185-187°** (from MeOH). [*Beilstein* **22** H 33, **22** I 502, **22** II 30, **22** III/IV 303, **22/2** V 3.]

N-4-Picolinoylbenzimidazole [100312-29-6] C₁₃H₉N₃O, **M 173.3**, **m 105-107°**. Recrystallise the imidazole three times from hexane [Fife & Przysas *J Am Chem Soc* **108** 4631 1986, DOI: 10.1021/ja00275a059].

Picrolic acid [3-methyl-4-nitro-1-(4-nitrophenyl)-2-pyrazolin-5-one, picrolonic acid] [550-74-3] C₁₀H₈N₄O₅, **M 264.2**, **m 116.1-117.2°, 116.5°(dec at 125°) 120°(dec), 125°, b 548.8°/760mm, d 1.65**. Crystallise picrolic acid from water or EtOH (solubility is 0.123% at 15° and 1.203% at 100° in H₂O; and 1.107% at 0° and 11.68% at 81° in EtOH). It forms Ca, Cu Hg, Mg, Na, Sr, Pb and many other metal complexes, also used for analysis of alkaloids, tryptophan and phenylalanine as well as a precipitant for organic bases and alkaloids. [Maquestiau et al. *Bull Soc Chim Belg* **82** 233 1973, DOI: 10.1002/bscb.19730820306; Iseki et al. *Chem Ber* **74** 1420 1941, DOI: 10.1002/cber.19410740811]. [*Beilstein* **24** H 51, **24** I 218, **24** II 25, **22** III/IV 105.]

Pinacyanol chloride (Quinaldine Blue) [2768-90-3] C₂₅H₂₅ClN₂, **M 388.9**, **CI 808**, **m 270°(dec)**. Crystallise the chloride from EtOH (blue-green prisms) or EtOH/diethyl ether. The crystals lose EtOH at ~100° and decompose at ~270°. It is dichroic with λ_{max} at 560nm and 604nm. It is used as a histological stain for chromosomes. The *iodide* [605-91-4] crystallises from MeOH or EtOH with **m 298-299°(dec)**. [*Beilstein* **23** H 320, **23** I 90, **II** 282, **III/IV** 2064, **23/10** V 129.]

dl-Pipecolic acid (piperidine-2-carboxylic acid) [535-75-1, 4043-87-2] C₆H₁₁NO₂, **M 129.1**, **m 264°, 280°(dec), 282°(dec), pK₁²⁵ 2.29, pK₂²⁵ 10.77**. It crystallises from water. The (*±*)-*picrate* has **m 158-159°** (from EtOH or *C₆H₆). [*Beilstein* **22** H 7, **22** III/IV 97, **22/1** V 220.] The *R*(+)-*enantiomer* [1723-00-8] has **m 277°(dec)** and [α]_D²⁰ +27 (c 4, H₂O), and the *S*(-)-*enantiomer* [3105-95-1] has **m 277°(dec)** and [α]_D²⁰ -26 (c 4, H₂O). [cf. p 603, *Beilstein* **22** III/IV 96, **22/1** V 220.]

Piperazine [110-85-0] C₄H₁₀N₂, **M 86.1**, **m 44° (hexahydrate 142-63-2), 109-112°, 110-112°, b 125-130°/760mm, 145-146°/760mm, pK₁²⁵ 5.33, pK₂²⁵ 9.73**. Piperazine crystallises from EtOH or anhydrous *benzene and is dried at 0.01mm. It can be sublimed under vacuum and purified by zone melting. The *hydrochloride* has **m 172-174°** (from EtOH), and the *dihydrochloride* crystallises from aqueous EtOH and has **m 318-320° (dec, sublimes at 295-315°)**. The *picrate* has **m ~200°**, and the *picrolonate* crystallises from dimethylformamide (**m 259-261°**). [*Beilstein* **23** H 4, **23** I 4, **23** II 3, **23** III/IV 15, **23/1** V 30.]

§ Piperazine on polystyrene support is commercially available. **Piperazine dihydrochloride (H₂O)** [142-64-3 (2HCl); 6094-40-2 (xHCl), 207605-49-0] C₄H₁₀N₂. 2HCl, **M 177.1**, has **m 82.5-83.5°**. Crystallise the salt from aqueous EtOH and dry it at 110°. [*Beilstein* **23** III/IV 17, **23/1** V 30.] **Piperazine phosphate (H₂O)** [18534-18-4] C₄H₁₀N₂. H₃PO₄, **M 197.6**. Crystallise it twice from water, air-dry and store for several days over Drierite. The salt dehydrates slowly if heated at 70°. [*Beilstein* **23** III/IV 18, **23/1** V 30.]

Piperazine-*N,N'*-bis(2-ethanesulfonic acid) (PIPES) [5625-37-6] $C_8H_{18}N_2O_6S_2$, **M 302.4**, $pK_1^{25} < 3$, $pK_2^{25} 6.82$ (7.82). Purify PIPES from boiling water (maximum solubility is about 1 g/L) or as described for ADA (see [26239-55-4] in 'Aliphatic Compounds' in this Chapter). The **disodium salt** [76836-02-7] $C_8H_{16}N_2NaO_6S_2$, **M 346.3**, decomposes at high temperature, and is purified by dissolving in the minimum volume of H_2O and adding EtOH until crystallisation is complete, filter it off and dry it *in vacuo*. Apart from being a very good buffer in the natural pH range (~pH 7, see pKa) it is used as an organisms buffer, like the free acid. [Good et al. *Biochemistry* **5**(2) 467 1966, DOI: 10.1021/bi00866a011; *Beilstein* **23/12** V 380.]

Piperidine [110-89-4] $C_5H_{11}N$, **M 85.2**, **m** -13° , -9° , **b** $35.4^\circ/40mm$, $106^\circ/760mm$, $d_4^{20} 0.862$, $n_D^{20} 1.4535$, $n_D^{25} 1.4500$, $pK^{25} 11.20$ (basic). Dry piperidine with BaO, KOH, CaH_2 , or sodium, and fractionally distil (optionally from sodium, CaH_2 , or P_2O_5). Purify from pyridine by zone melting. [*Beilstein* **22** H 6, **22** I 5, **22** II 3, **22** III/IV 287, **22/2** V 3.] § Piperidine on polystyrene support is commercially available. **Piperidinium chloride** [6091-44-7] $C_5H_{11}N \cdot HCl$, **M 121.6**, has **m** $244-245^\circ$, $245-248^\circ$, $247-248^\circ$. Crystallise the salt from EtOH/diethyl ether in the presence of a small amount of HCl. [*Beilstein* **20** H 6, **20** II 10, **20** III/IV 295, **20/2** V 13.] **Piperidinium nitrate** [6091-45-8] $C_5H_{11}N \cdot HNO_3$, **M 145.2**, has **m** 141° , ($155-157^\circ$ from EtOH). The nitrate crystallises from acetone/ethyl acetate or EtOH. [*Beilstein* **20** H 12, **20** II 10, **20** III/IV 295, **20/2** V 14.]

Piperidine-2,6-dione (glutarimide) [1121-89-7] $C_5H_7NO_2$, **M 113.1**, **m** $163-165^\circ$, $155-157^\circ$, 154° , $152-154^\circ$, $pK^{25} 11.43$ (acidic). Purify it by dissolving 75g in 200ml of H_2O , boil for 30 minutes with 2g of charcoal, filter, evaporate to dryness and recrystallise the residue from 125m L of 95% EtOH to give 70g of white crystals, **m** $152-154^\circ$. It also crystallises from Me_2CO (**m** $163-165^\circ$) or EtOH (**m** $153-154^\circ$). The ***N*-bromo derivative** (a brominating agent) crystallises from H_2O with **m** $180-185^\circ$. [Paris et al. *Org Synth Coll Vol* **4** 496 1963, DOI: 10.15227/orgsyn.037.0047; *Beilstein* **21** H 382, **21** I 331, **21** II 307, **21** III/IV 4582.]

4-Piperidone [piperidin-4(1*H*)-one, γ -] [41661-47-6] C_5H_9NO , **M 99.1** (anhydrous), cannot be distilled without decomposition although a boiling point of 79° has been reported, $pK^{25} 8.6$ (acidic). It is a yellow irritating oil which is an alkaloid in the leaves and branches of *Dichilus* species (leguminosae), e.g. *D. strictus*, *D. gracilis*, *D. lebeckioides*, *D. pilosus* and *D. reflexus*. It decomposes on distillation and is purified and stored as the **monohydrate monohydrochloride** [anhydrous 41979-39-9; hydrate 40064-34-4], **M 135.6** (anhydrous salt), which is in fact **4,4-dihydroxy-piperidinium chloride** that recrystallises from H_2O with **m** $94-96^\circ$, $97-100^\circ$ (+ 1 H_2O), or from EtOH/Et₂O with **m** $139-141^\circ$ (+ 1.5 EtOH); and the **anhydrous salt** has **m** $147-149^\circ$. See above for the *N*-acetyl- *N*-benzyl and *N*-methyl derivatives. The ***N*-benzoyl derivative** [24686-78-0], **M 203.2**, has **m** $49-50^\circ$ and **b** $158-160^\circ/0.2mm$, and the **oxime** [79858-41-6], **M 114.1**, crystallises from dry $*C_6H_6$ in needles **m** $117-118^\circ$ (anhydrous), and is hygroscopic. [*Beilstein* **21/6** V 419.]

Piperine (1-piperoylpiperidine) [94-62-2] $C_{17}H_{19}NO_3$, **M 285.4**, **m** $129-129.5^\circ$, $131-135^\circ$, $pK^{15} 1.98$. Piperine crystallises as light yellow crystals from EtOH or EtOAc (**m** 132°), aqueous EtOH (**m** $128-129^\circ$), Et₂O (**m** 129°), or $*benzene$ /ligroin. [*Beilstein* **20** H 79, **20** I 23, **20** II 53, **20** III/IV 1341, **20/3** V 469.] It was originally isolated from black pepper, has no taste at first but produces a burning after-taste and has insecticidal properties.

Poly(*N*-vinylcarbazole) (PVK) [25067-59-8] $(C_{14}H_{11}N)_n$, M_{avr} 25,000-50,000, **m** $>300^\circ$, **d** 1.2g/ml. Precipitate it seven times from tetrahydrofuran with MeOH, with final freeze-drying from $*benzene$. Dry it under vacuum. Also available is polymer with M_{avr} 1,000,000, **powder m** 220° .

Poly(4-vinylpyridine, Reillex 402) [25232-41-1] $(C_7H_7N)_n$, **M** (105.1)_n, M_{avr} ~60,000 has **T_g**(onset of anneal) 137° ; M_{avr} ~160,000 has **T_g**(onset of anneal) 142° (all soluble in DMF, AcOH and lower alcohols). Purify them by repeated precipitation from solutions in EtOH with dioxane, and then EtOH with ethyl acetate. Finally, freeze-dry a *tert*-butanol solution.

Poly(*N*-vinylpyrrolidone) [9003-39-8] $(C_6H_9NO)_n$, **M** (111.1)_n, **crosslinked** [25249-54-1] has **m** $>300^\circ$. Purify it by dialysis, and freeze-drying. Also by precipitation from $CHCl_3$ solution by pouring into ether. Dry it in a vacuum over P_2O_5 . For the crosslinked polymer, purification is by boiling for 10 minutes in 10% HCl and then washing with glass-distilled water until free from Cl ions. Finally, Cl ions are removed more readily by neutralising with KOH and continued washing.

(±)-Primaquine diphosphate (*RS*- 8-[4-amino-1-methylbutylamino]-6-methoxyquinoline di-phosphate) [63-45-6] $C_{15}H_{21}N_3O \cdot 2H_3PO_4$, *M* 455.4, *m* 197-198°, 204-206°(dec), $pK_{Est(1)} \sim 3.38$ (ring N^+), $pK_{Est(2)} \sim 10.8$ (NH_3^+). It forms yellow crystals from 90% aqueous EtOH and is moderately soluble in H_2O . The *oxalate salt* has *m* 182.5-185° (from 80% aqueous EtOH), and the *free base* is a viscous liquid *b* 165-170°/0.002mm, 175-177°/2mm. [Elderfield et al. *J Am Chem Soc* 77 4816 1955, DOI: 10.1021/ja01623a038; Elderfield et al. *J Am Chem Soc* 77 4819 1955, DOI: 10.1021/ja01623a039; Elderfield et al. *J Am Chem Soc* 68 1524 1964, DOI: 10.1021/ja01212a040; *Beilstein* 22 III/IV 5817.]

Proclavine (3,6-diaminoacridine) [92-62-6] $C_{13}H_{11}N_3$, *M* 209.2, *m* 284-286°, $pK_1^{25} -2.7$, $pK_2^{25} 0.55$, $pK_3^{25} 9.49$. It crystallises from aqueous MeOH. The *picrate* crystallises from aqueous pyridine with *m* ~185°. [Beilstein 22 H 487, 22 I 649, 22 II 397, 22 III/IV 5487, 22/11 V 322, also see Elderfield references above.] For proflavin see 3,6-diaminoacridine hydrochloride.

Propidium iodide (3,8-diamino-5-(3-diethylaminopropyl)-6-phenylphenanthridinium iodide methiodide) [25535-16-4] $C_{27}H_{34}I_2N_4$, *M* 668.4, *m* 210-230°(dec), $pK_{Est(1)} \sim 4$ (aniline NH_2), $pK_{Est(2)} \sim 8.5$ (EtN_2). It crystallises as red crystals from H_2O containing a little KI. It fluoresces strongly and intercalates with nucleic acids; useful in flow cytometry. [Watkins *J Chem Soc* 3064 1952, DOI: 10.1039/JR9520003059, *Beilstein* 22 III/IV 5519.] **TOXIC.**

(±)-Propylene carbonate (4-methyl-1,3-dioxalan-2-one) [108-32-7] $C_4H_6O_3$, *M* 102.1, *m* -55°, *b* 79-80°/0.08mm, 110°/0.5-1mm, 112-114°/2mm, 241°/760mm, $d_4^{25} 1.187$, $n_D^{20} 1.421$. It is manufactured by reaction of 1,2-propylene oxide with CO_2 in the presence of a catalyst (quaternary ammonium halide). Contaminants include propylene oxide, carbon dioxide, 1,2- and 1,3-propanediols, allyl alcohol and ethylene carbonate. It can be purified by percolation through molecular sieves (Linde 5A, dried at 350° for 14 hours under a stream of argon), followed by distillation under a vacuum. [Jasinski & Kirkland *Anal Chem* 39 1663 1967, DOI: 10.1021/ac50156a051.] It can be stored over molecular sieves under an inert gas atmosphere. When purified in this way it contains less than 2 ppm of water. Activated alumina and dried CaO have also been used as drying agents prior to fractional distillation under reduced pressure. It has been dried with 3A molecular sieves and distilled under nitrogen in the presence of *p*-toluenesulfonic acid, then redistilled and the middle fraction collected. [Beilstein 19 III/IV 1564, 19/4 V 21.] *R*(+)-propylene carbonate (4-methyl-1,3-dioxalan-2-one) [16606-55-6] $C_4H_6O_3$, *M* 102.1, has *b* 240°/760mm, $d_4^{25} 1.189$, $n_D^{20} 1.422$, and $[\alpha]_D^{20} +2$ (neat), and *S*(-)-propylene carbonate (4-methyl-1,3-dioxalan-2-one) [51260-39-0] has $[\alpha]_D^{20} -2$ (neat) could be purified in the same way as the racemate.

dl-Propylene oxide (methyloxirane, 1,2-epoxypropane) [75-56-9] *M* 58.1, *m* -112°, *b* 34.5°/70mm $d_4^{20} 0.829$, $n_D^{20} 1.3664$. Dry the oxide with Na_2SO_4 or CaH_2 and fractionally distil it through a packed column (glass helices), after refluxing with Na, CaH_2 , or KOH pellets. [Beilstein 17 I 4, 17 II 131, 17 III/IV 17, 17/1 V 17.] The *R*(+)-enantiomer [15448-47-2] and the *S*(-)-enantiomer [16088-62-3] have *b* 33-34°/atm and $[\alpha]_D^{20}$ (+)14.6 and (-)14.6 (neat) respectively. [Beilstein 17/1 V 17.]

Protopine {Fumarine, Macleyine, 4,6,7,14-tetrahydro-5-methyl-bis[1,3]-benzodioxolo[4,5-*c*:5',6'-*g*]azecine-13(5*H*)-one} [130-86-9] $C_{20}H_{19}NO_5$, *M* 353.4, *m* 208°, 209°, 211°, $pK^{25} 5.99$. It crystallises from EtOH/ $CHCl_3$. The *picrate* has *m* ~240°(dec). [Beilstein 27 H 558, 17 I 568, 17 II 620, 17 III/IV 6881.] It is an analgesic, inhibits histamine H1 receptors and platelet aggregation. [For anti-thrombotic and anti-inflammatory activities see Saeed et al. *Pharmacol Research* 36 1 1997, DOI: 10.1006/phrs.1997.0195; for Anticholine-sterase and Antiamnesic Activities see Kim et al. *Planta Med* 65 218 1999, DOI: 10.1055/s-1999-13983.]

Pteridine [91-18-9] $C_6H_4N_4$, *M* 132.2, *m* 139.5-140°, $pK_1^{20} 4.05$ (equilibrium, hydrate), $pK_2^{20} 11.90$ (OH of hydrate). It crystallises from EtOH (5 parts with 80% recovery), *benzene, *n*-hexane, *n*-heptane or light petroleum (*b* 60°-80°, 300 parts with 80% recovery). It is best purified by sublimation at 120-130°/20mm. Store at 0°, in the dark. The yellow crystalline plates turn green in the presence of light and on long standing in the dark, and sublimation leaves some dark-coloured material behind. The crystals induce sneezing. [Albert et al. *J Chem Soc* 474 1951, DOI: 10.1039/JR9510000474; for hydrated species see Albert & Armarego *Adv Heterocycl Chem* 4 1 1965, DOI:10.1016/S0065-2725(08)60873-9; D. J. Brown *Fused Pyrimidines: Pteridines*

Wiley-Interscience NY 1988, ISBN 0-471-83041-1(part 3), *Beilstein* **26** III/IV 1770.]

2,4-(1*H*,3*H*)-Pteridinedione H₂O (lumazine) [487-21-8] C₆H₄N₄O₂, M 182.1, m >350°, pK₁²⁰ <1.0, pK₂²⁰ 7.94. Crystallise the dione from water. It has also been purified as for pterin [2236-60-4] below. [Pfleiderer & Hutzenlaub *Chem Ber* **106** 3149 1973, DOI: 10.1002/cber.19731061007; Dallacker & Steiner *Justus Liebigs Ann Chem* **660** 98 1962, DOI: 10.1002/jlac.19626600111; D.J. Brown *Fused Pyrimidines: Pteridines* Wiley-Interscience NY 1988, ISBN 0-471-83041-1(part 3); *Beilstein* **26** III/IV 2489.]

Pterin (2-aminopteridin-4(3*H*)-one) [2236-60-4] C₆H₅N₅O, M 163.1, m >300°, pK₁²⁰ 2.27 (basic), pK₂²⁰ 7.96 (acidic). It is dissolved in hot 1% aqueous ammonia, filtered, and an equal volume of hot 1M aqueous formic acid is added. The solution is allowed to cool at 0-2° overnight. The solid is collected and washed with distilled water several times by centrifugation and dried *in vacuo* over P₂O₅ overnight, and then at 100° overnight (any ammonium formate in the sample evaporates off). [D. J. Brown *Fused Pyrimidines: Pteridines* Wiley-Interscience NY 1988, ISBN 0-471-83041-1(part 3); *Beilstein* **26** III/IV 3936.]

(-)-Pterocarpin {(6*aR*-cis)-6*a*,12*a*-dihydro-3-methoxy-6*H*-[1,3]dioxolo[5,6]benzofuro[3,2*c*][1]-benzopyran} [524-97-0] C₁₇H₁₄O₅, M 298.3, m 165°, 165-166°, [α]_D²⁰ -215 (c 0.5, CHCl₃). Crystallise it from EtOH, or petroleum ether. [Fukui & Nakayama *Bull Chem Soc Jpn* **42** 1408 1969, DOI:org/10.1246/bcsj.42.1408; Pachler & Underwood *Tetrahedron* **23** 1817 1967, DOI: 10.1016/S0040-4020(01)82581-3; *Beilstein* **19** II 459, **19** III/IV 5789.]

Purine [120-73-0] C₅H₄N₄, M 120.1, m 214-217°, 216-217°, pK₁²⁰ 2.30, pK₂²⁰ 9.86. It crystallises from toluene or EtOH, and sublimates at 100-150°/0.1mm or 160°/10⁻⁴mm. The *picrate* has m 207-209° after crystallisation from 20 volumes of H₂O. [Lister *Purines Part II, Fused Pyrimidines* Brown Ed, Wiley-Interscience 1971, ISBN 0-471-38205-1, *Beilstein* **26** H 354, **26** III/IV 1736.]

Pyocyanine (1-hydroxy-5-methylphenazinium zwitterion) [85-66-5] C₁₃H₁₀N₂O, M 210.2, m 133° (sublimes and decomposes on further heating), pK₂₅ -3.5. It crystallises from H₂O as dark blue needles. The *picrate* has m 190° (dec). [*Beilstein* **23** H 395, **23** I 59, **23** II 234, **23/8** V 395.] This antibiotic from *Pseudomonas aeruginosa* can generate reactive oxygen species.

Pyrazine [290-37-9] C₄H₄N₂, M 80.1, m 47°, 50-56°, 57°, b 115.5-115.8°/760mm, pK₁²⁰ -6.25 (aqueous H₂SO₄), pK₂²⁰ 1.1 (0.51 at 20°). Distil pyrazine in steam and crystallise it from water. Purify also by zone melting. [G.B. Barlin *The Chemistry of Heterocyclic Compounds: The Pyrazines* Vol **41** pp 685 1982, Wiley-Interscience NY, ISBN 04771381195; D.J. Brown *The Chemistry of Heterocyclic Compounds: The Pyrazines* (Supplement 1) Vol **58** pp 557 2002, Wiley-Interscience NY, ISBN 047140822, *Beilstein* **23** H 91, **23** II 80, **23** III/IV 899, **23/5** V 351.]

Pyrazine-1-oxide [2423-65-6] C₄H₄N₂O, M 96.1, has m 112-116°, 113-114°, pK_{Est} <0. Recrystallise the oxide from *C₆H₆. It is soluble in hot petroleum ether (b 60-68°, see pyrazine-1,4-dioxide above). Dry it *in vacuo*. It has ν_{max} at 1305cm⁻¹. [Koelsch & Gumprecht *J Org Chem* **23** 1605 1958, DOI: 10.1021/jo01105a003.]

Pyrazine-1,4-dioxide [2433-84-9] C₄H₄N₂O₂, M 112.1, has m 285-286°, pK_{Est} <0. If the sample contains pyrazine-1-oxide, then place it in a Soxhlet extractor and extract it with hot petroleum ether (b 60-68°) in which the mono-oxide is soluble. Collect the dioxide from the thimble and recrystallise it from MeOH. It is dried *in vacuo*. It has ν_{max} at 1270cm⁻¹. [Koelsch & Gumprecht *J Org Chem* **23** 1605 1958, DOI: 10.1021/jo01105a003.]

Pyrazinecarboxylic acid [98-97-5] C₅H₄N₂O₂, M 124.1, m 222-225°(dec), 225-229°(dec), pK₁²⁵ -3.0, pK₂²⁵ -0.7, pK₃²⁵ 2.70. It crystallises from water. The *methyl ester* has m 62° (from petroleum ether). [Sausville & Spoerri *J Am Chem Soc* **63** 3153 1941, DOI: 10.1021/ja01856a075; *Beilstein* **25** III/IV 771.]

Pyrazinecarboxamide [98-96-4] C₅H₅N₃O, M 123.1, has m 189-191° (sublimes slowly at 159°), pK₂₅ -0.5. The amide crystallises from water, EtOH or 1:1 hexane/EtOH in four modifications *viz* α-, β-, δ- and γ-form. [Rø & Sørum *Acta Cryst* **28B** 1677 1972, DOI: 10.1107/S0567740872004856; *Beilstein* **25** III/IV 772.]

Pyrazine-2,3-dicarboxylic acid [89-01-0] $C_6H_4N_2O_4$, M 168.1, m 183-185°(dec), 187°(dec), 188°(dec), $pK_1 < 2.0$, pK_2 0.9, pK_3 2.77 (2.20). Crystallise the dicarboxylic acid from water and dry it at 100°. The *dimethyl ester* has m 62-63° (from Et₂O or Et₂O/petroleum ether). [Beilstein 25 H 168, 25 II 164, 25 III/IV 1064.]

Pyrazole [288-13-1] $C_3H_4N_2$, M 68.1, m 67-70°, 70°, b 96°/16mm, 186-188°/atm, pK_1^{25} 2.48 (protonation), pK_2^{25} 14.12 (acidic). Crystallise pyrazole from petroleum ether, cyclohexane, or water. Its solubility in H₂O at 9.6° is 2.7moles/L, and at 24.8° it is 19.4moles/L; in cyclohexane at 31.8° it is 0.577moles/L, and at 56.2° it is 5.86moles/L; and in benzene at 5.2° it is 0.31moles/L, and at 46.5° it is 16.8moles/1000ml. It forms complexes with metals, e.g. silver. [Barszcz et al. *JCS Dalton Trans* 2025 1986, DOI: 10.1039/DT9860002025; Beilstein 23 H 39, 23 I 15, 23 II 33, 23 III/IV 550, 23/4 V 122.]

Pyrazole-*N*-1-carboximidine hydrochloride (Praxadine, 1-amidinopyrazole hydrochloride, 1*H*-pyrazole-1-carboxamidinium hydrochloride) [4023-02-3] $C_4H_6N_4 \cdot HCl$, M 146.6, m 165-166°, 167-168°, 167-168.5°, 167-170°, pK_{Est} ~8.5. The white crystalline hydrochloride, which can be prepared from pyrazole and cyanamide in *p*-dioxane containing HCl, is purified by recrystallisation from dioxane/H₂O. It is a good guanylating agent for primary and secondary amines [Bernatowicz et al. *J Org Chem* 57 2497 1992, DOI: 10.1021/jo00034a059; Bernatowicz et al. *Tetrahedron Lett* 34 3389 1993, DOI: 10.1016/S0040-4039(00)79163-5]. The *free base* can be obtained by suspending the hydrochloride in CHCl₃ and bubbling NH₃ gas through, whereby NH₄Cl separates and is filtered off. CHCl₃ is then distilled off and the residue is washed with a little EtOH and dried *in vacuo*. It crystallises from *C₆H₆ and has m 93-96° (94-98.5° and 97-101° have also been reported). The *picrate* has m 202-203° (208° has also been reported). [Bredereck et al. *Chem Ber* 98 3178 1965, DOI: 10.1002/cber.19650981013; Jones et al. *J Org Chem* 19 1428 1954, DOI: 10.1021/jo01374a004; Beilstein 23/4 V131.] It is a competitive inhibitor of the three isoforms of nitric oxide synthase [Lee et al. *Bioorg Medicin Chem Lett* 10 2771 2000, DOI: 10.1016/S0960-894X(00)00573-4].

1*H*-Pyrazole-3-carboxylic acid [1621-91-6] $C_4H_4N_2O_2$, M 112.1, m 208-210°, 210-214° (decarboxylates), pK^{25} 3.74, $pK_{Est(2)}$ ~12 (acidic NH). Purify the acid by precipitation from an alkaline solution with mineral acid, and recrystallise it from H₂O. On heating, it decarboxylates (to pyrazole m 70°) more readily than the 4-carboxylic acid below. The *N*-2-methyl derivative has m 222° (from H₂O) and pK^{25} 3.27. [Habraken et al. *Recl Trav Chim Pays-Bas* 85 1194 1966, DOI: 10.1002/recl.19660851204; Jones et al. *J Org Chem* 19 1428 1954, DOI: 10.1021/jo01374a004; Alley & Shirley *J Am Chem Soc* 80 6271 1958, DOI: 10.1021/ja01556a027; Knorr *Justus Liebigs Ann Chem* 279 188 1894, DOI: 10.1002/jlac.18942790113.]

1*H*-Pyrazole-4-carboxylic acid [37718-11-9] $C_4H_4N_2O_2$, M 112.1, has m 282° (decarboxylates), $pK_{Est(1)}$ ~3.6, $pK_{Est(2)}$ ~12 (acidic NH). This acid, also obtained by decarboxylating the more water soluble 3,4,5-tricarboxylic acid, crystallises in yellow prisms from H₂O. [Buchner & Fritsch *Justus Liebigs Ann Chem* 273 252 1893, DOI: 10.1002/jlac.18932730211; Knorr & Rothenburg *Chem Ber* 28 688 1895, DOI: 10.1002/cber.189502801157; Beilstein 25 H 116.]

Pyrazole-3,5-dicarboxylic acid [3112-31-0] $C_5H_4N_2O_4$, M 174.1, m 287-289°(dec), 292-285°(dec), 295-297°(dec), $pK_{Est(1)}$ ~1.2 (CO₂H), $pK_{Est(2)}$ ~3.7 (CO₂H), $pK_{Est(3)}$ ~12 (NH). It crystallises from water as a *monohydrate*, or EtOH. Dry it in a vacuum at 100°. The *dimethyl ester* crystallises from Et₂O or *C₆H₆ with m 155°. [Buchner & Papendieck *Justus Liebigs Ann Chem* 273 246 1893, DOI: 10.1002/jlac.18932730210; Buchner & Papendieck *Justus Liebigs Ann Chem* 273 232 1893, DOI: 10.1002/jlac.18932730208; Beilstein 25 III/IV 1047.]

Pyridazine [289-80-5] $C_4H_4N_2O$, M 80.1, m -8°, b 87/14mm, 208°/atm, d_4^{20} 1.103, $n_D^{23.5}$ 1.5321, pK^{20} -7.1 (H₀ scale aq H₂SO₄). If it looks *dubious* then dissolve it in dry Et₂O, shake this with K₂CO₃, filter, and distil, preferably under a vacuum. The *hydrochloride*, yellow crystals m 161-163° (sealed tube), is obtained by crystallization (EtOH/Et₂O) and sublimation *in vacuo*. [Mizzoni & Spoerri *J Am Chem Soc* 73 1873 1951, DOI: 10.1021/ja01148a537; Beilstein 23/5 321.]

Pyridazine *N*-oxide (pyridazine 1-oxide) [1457-42-7] $C_4H_4N_2O$, M 96.1, has m 38-39°, b 138-140°/4mm, pK_{Est} <1. Purify the oxide by distillation in a vacuum and by sublimation *in vacuo*. When a solution of the oxide in MeOH is treated with an aqueous solution of CuCl₂, the [C₄H₄N₂O]₂—CuCl₂·2H₂O-complex (m 182-183°) is formed from which the oxide can be recovered. Its 1H NMR (60MHz, CDCl₃, TMS) has t 1.70(m H-3), 3.00(ddd, H-4), 2.43(ddd, H-5), 1.92(ddd, H-6) ($J_{4,5} = 7.5$, $J_{5,6} = 6.0$, $J_{4,6} = 1.0$, $J_{3,5} = 2.4$, $J_{3,6} = 0.6$ Hz).

[Pollak et al. *J Org Chem* **35** 2478 1970, DOI: 10.1021/jo00833a003; Klinge et al. *Recl Trav Chim Pays-Bas* **95** 21 1976, DOI: 10.1002/recl.19760950106; Ohsawa et al. *Tetrahedron Lett* **19** 1979 1978, DOI: 10.1016/S0040-4039(01)94726-4; Koelsch & Gumprecht *J Org Chem* **23** 1605 1958, DOI: 10.1021/jo01105a003; *Beilstein* **23** III/IV 890.]

Pyridazin-3,6-diol (pyridazine-(1*H*,2*H*-3,6-dione, maleic hydrazide) [123-33-1] C₄H₄N₂O₂, M 112.1, m 299-301°(dec), 305°(dec), 306°(dec), pK₁²⁵ 5.67, pK₂²⁵ 13.3. Crystallise the hydrazide from water. Dry it at ~100° over P₂O₅. [*Beilstein* **24** III/IV 1186.] Plant growth substance used in agriculture.

Pyridine [110-86-1] C₅H₅N, M 79.1, m -41.8°, b 115.6°/atm, d₄²⁰ 0.9831, n_D²⁰ 1.51021, pK_a²⁵ 5.23. Likely impurities are H₂O and amines such as the picolines and lutidines. Pyridine is *hygroscopic* and is miscible with H₂O and organic solvents. It can be dried with solid KOH, NaOH, CaO, or BaO, followed by fractional distillation. Other methods of drying include standing with Linde type 4A molecular sieves, CaH₂ or LiAlH₄, azeotropic distillation of the H₂O with toluene or *benzene, or treated with phenylmagnesium bromide in ether, followed by evaporation of the ether and distillation of the pyridine. A recommended [Lindauer & Mukherjee *Pure Appl Chem* **27** 265 1971, DOI: org/10.1351/pac197127010265] method dries pyridine over solid KOH (20g/Kg) for 2 weeks and fractionally distills the supernatant over Linde type 5A molecular sieves and solid KOH. The product is stored under CO₂-free nitrogen. Pyridine can be stored in contact with BaO, CaH₂ or molecular sieves. Non-basic materials can be removed by steam distilling a solution containing 1.2 equivalents of 20% H₂SO₄ or 17% HCl until about 10% of the base has been carried over along with the non-basic impurities. The residue is then made alkaline, and the base is separated, dried with NaOH and fractionally distilled. **The alkali metals, Na, Li or Cs should NOT be used for drying pyridine, and pyridine derivatives, as they form coloured pyridine radical anions leading to bipyridyls.** [see Schmulback et al. *J Am Chem Soc* **90** 6600 1968, DOI: 10.1021/ja01026a006]. **Alternatively**, pyridine can be treated with oxidising agents. Thus pyridine (800ml) has been stirred for 24 hours with a mixture of ceric sulfate (20g) and anhydrous K₂CO₃ (15g), then filtered and fractionally distilled. Hurd and Simon [*J Am Chem Soc* **84** 4519 1962, DOI: 10.1021/ja00882a030] stirred pyridine (135ml), water (2.5L) and KMnO₄ (90g) for 2 hours at 100°, then set aside for 15 hours before filtering off the precipitated manganese oxides. Addition of solid KOH (ca 500g) caused pyridine to separate. It was decanted, refluxed with CaO for 3 hours and distilled. Separation of pyridine from some of its homologues can be achieved by crystallisation of the oxalates. Pyridine is precipitated as its **oxalate** by adding it to the stirred solution of oxalic acid in acetone. The precipitate is filtered, washed with cold acetone, and pyridine is regenerated and isolated. Other methods are based on complex formation with ZnCl₂ or HgCl₂. Heap, Jones and Speakman [*J Am Chem Soc* **43** 1936 1921, DOI: 10.1021/ja01441a023] added crude pyridine (1L) to a solution of ZnCl₂ (848g) in 730ml of water, 346ml of concentrated HCl and 690ml of 95% EtOH. The crystalline precipitate of **ZnCl₂·(pyridine)₂** was filtered off, recrystallised twice from absolute EtOH, then treated with a concentrated NaOH solution, using 26.7g of solid NaOH to 100g of the complex. The precipitate was filtered off, and the pyridine was dried with NaOH pellets and distilled. Similarly, Kyte, Jeffery and Vogel [*J Chem Soc* 4454 1960, DOI: 10.1039/JR9600004454] added pyridine (60ml) in 300ml of 10% (v/v) HCl to a solution of HgCl₂ (405g) in hot water (2.3l). On cooling, crystals of **pyridine-HgCl₂ (1:1) complex** separated and were filtered off, crystallised from 1% HCl (to m 178.5-179°), washed with a little EtOH and dried at 110°. The free base was liberated by addition of excess aqueous NaOH and separated by steam distillation. The distillate was saturated with solid KOH, and the upper layer was removed, dried further with KOH, then BaO and distilled. Another possible purification step is fractional crystallisation by partial freezing. Small amounts of pyridine have been purified by vapour-phase chromatography, using a 180-cm column of polyethyleneglycol-400 (Shell 5%) on Embacel at 100°, with argon as carrier gas. The Karl Fischer titration can be used for determining water content. A colour test for pyrrole as a contaminant is described by Biddiscombe et al. [*J Chem Soc* 1957 1954, DOI: 10.1039/JR9540001957]. The **1:1-hydrochloride** crystallises from EtOH with m 144°, b 218-219°/760mm (see below) and is hygroscopic. The **1:2-hydrochloride** has m 46° [58888-58-7] and the **picrate** has m 165-166° [1152-90-5]. [*Beilstein* **20** H 181, **20** I 54, **20** II 96, **20** III/IV 2205, **20**/5 V 160.] § Polystyrene-supported pyridine is commercially available. **Pyridine hydrobromide perbromide (pyridinium bromide perbromide) [39416-48-3] C₅H₅N·HBr·Br₂, M 319.9, has m 130° (dec), 132-134°(dec), 135°(dec).** It is a very good stable brominating agent-liberating one mol of Br₂. Purify it by recrystallisation from glacial acetic acid (33g from 100ml of AcOH) to give orange-red crystals. Fieser prepared it by dissolving pyridine (15ml) in 48% HBr with cooling, then adding gradually bromine (25g) with cooling and swirling (fume cupboard). The solid that separated is collected, rinsed with

AcOH, and without drying is recrystallised from AcOH (100ml) to give red needles (33g, 69%). [Englert & McElvain *J Am Chem Soc* **51** 865 1929, DOI: 10.1021/ja01378a031; Fieser **1** 967; *Beilstein* **20/5** V 181.]

Pyridine hydrochloride [628-13-7] $C_5H_5N \cdot HCl$ **M 115.6**, has **m 144°**, **145-147°**, **b 218°/760mm**, **222-224°/atm**. Crystallise the salt from $CHCl_3/EtOAc$ and wash it with Et_2O . It is *hygroscopic*. [*Beilstein* **20** H 185, **20** I 57, **20** II 103, **20** III/IV 2230, **20/5** V 180.]

Pyridine-2-aldehyde (picolinaldehyde) [1121-60-4] C_6H_5NO , **M 107.1**, **b 81.5°/25mm**, **181°/atm**, d_4^{20} **1.121**, n_D^{20} **1.535**, pK_1^{25} **3.84**, pK_2^{25} **12.68**. Purification is achieved by bubbling sulfur dioxide into a solution of 50g of the aldehyde in 250ml of boiled water, under nitrogen, at 0°, until precipitation is complete. The *bisulfite addition compound* is filtered off rapidly and, after washing with a little water, is refluxed in 17% HCl (200ml) under nitrogen until a clear solution is obtained. Neutralisation with $NaHCO_3$ and extraction with ether separated the aldehyde which is recovered by drying the extract, then distilling twice, under nitrogen. [Kyte et al. *J Chem Soc* 4454 1960, DOI: 10.1039/JR9600004454; *Beilstein* **21** I 287, **21** III/IV 3495, **21/7** V 293.]

Pyridine-3-aldehyde (nicotinaldehyde) [500-22-1] has **b 78-81°/10mm**, **89.5°/14mm**, d_4^{20} **1.141**, n_D^{20} **1.549**, pK_1^{20} **3.80**, pK_2^{20} **13.10**. Purified as for pyridine-2-aldehyde. [*Beilstein* **21** I 288, **21** III/IV 3517, **21/7** V 334.]

Pyridine-4-aldehyde [872-85-5] has **b 71-73°/10mm**, **79.5°/12mm**, d_4^{20} **1.137**, n_D^{20} **1.544**, pK_1^{20} **4.77**, pK_2^{20} **12.20**. Purified as for pyridine-2-aldehyde. [*Beilstein* **21** III/IV 2529, **21/7** V 351.]

Pyridine-2-aldoxime (pyridine-2-carboxaldoxime) [873-69-8] $C_6H_6N_2O$, **M 122.1**, **m 110-112°**, **111-113°**, **114°**, pK_1^{25} **3.56**, pK_2^{25} **10.17**. Recrystallise it from Et_2O /petroleum ether or H_2O . The *picrate* has **m 169-171°** (from aqueous $EtOH$). It is used in peptide synthesis. [UV: Grammaticakis *Bull Chem Soc Fr* 109, 116 1956, Ginsburg & Wilson *J Am Chem Soc* **79** 481 1957, DOI: 10.1021/ja01559a067; Hanania & Irvine *Nature* **183** 40 1959, DOI: 10.1038/183040a0, Green & Saville *J Chem Soc* 3887 1956, DOI: 10.1039/JR9560003887; *Beilstein E-isomer* **21** I 288, **21** III/IV 3504, **21/7** V 305.] Its *methochloride* (Pralidoxime, 2-PAM chloride) [51-15-0] $C_7H_9N_2O \cdot Cl$, **M 172.6**, **m 226-227°(dec)** (from $EtOH/Et_2O$), is a prototypical reactivator of acetylcholine esterase that was inactivated by organophosphorus poisons, e.g. insecticides, or nerve gases [Kondritzer et al. *J Pharm Sci* **50** 109 1961, DOI: 10.1002/jps.2600500204]. The *methiodide* (Pralidoxime, 2-PAM iodide) [94-63-3] $C_7H_9N_2O \cdot I$, **M 266.1**, has **m 225-226°(dec)** (from $EtOH$),

Pyridine-3-aldoxime [1193-92-6] has **m 150°**, **150-153°**, pK_1^{20} **4.07**, pK_2^{20} **10.39**. Crystallise the oxime from water. [*Beilstein E-isomer* **21** III/IV 3521, **21/7** V 339.]

Pyridine-4-aldoxime [696-54-8] has **m 129°**, **130-133°**, pK_1^{20} **4.73**, pK_2^{20} **10.03**. Crystallise the oxime from water. [*Beilstein E-isomer* **21** III/IV 3533, **21/7** V 355.]

2,6-Pyridinedialdoxime [2851-68-5] $C_7H_7N_3O_2$, **M 165.1**, **m 212°**, **216°**, $pK_{Est(1)} \sim 3.0$, $pK_{Est(2)} \sim 10$. Crystallise it several times from water or $EtOH$ to give colourless needles. It is a tridentate chelate for metal ions, e.g. Fe^{2+} . [Lions & Martin *J Am Chem Soc* **79** 2733 1957, DOI: 10.1021/ja01568a018; *Beilstein* **21** III/IV 4746.]

Pyridine-2,5-dicarboxylic acid (isocinchomeronic acid) [100-26-5] $C_7H_5NO_4$, **M 167.1**, **m 242-247°**, **254°**, **(267° dec)**, pK_1^{25} **0.60**, pK_2^{25} **2.49**, pK_3^{25} **5.12**. Crystallise it from H_2O or dilute HCl. Attempted sublimation provides nicotinic acid via decarboxylation. [Napoli *J Inorg Nucl Chem* **32** 1907 1970, DOI:10.1016/0022-1902(70)80600-5; *Beilstein* **22** H 153, **22** I 533, **22** II 105, **22** III/IV 1632, **22/4** V 124.]

Pyridine-3,4-dicarboxylic acid (cinchomeronic acid) [490-11-9] has **m 253-255°**, **256°**, **262°(dec)**, pK_1^{25} **1.50** **(0.6)**, pK_2^{25} **2.43** **(2.95)**, pK_3^{25} **4.78** **(5.07)**. It has been prepared by the oxidation of isoquinoline with concentrated HNO_3 (44% yield) and crystallised from H_2O or dilute aqueous HCl. It has also been purified via the *dimethyl ester* which is distilled (**b 95-100°/1.5mm**) and hydrolysed with 3.5N HCl, evaporated and recrystallised. [Armarego & Evans *J Appl Chem* **12** 45 1962, DOI: 10.1002/jctb.5010120108; Foye et al. *J Med Chem* **9** 61 1966, DOI: 10.1021/jm00319a016; *Beilstein* **22** H 155, **22** I 534, **22** II 106, **22** III/IV 1641, **22/4** V 135.]

Pyridine N-oxide [694-59-7] C_5H_5NO , **M 95.1**, **m 62-67°**, **67°**, **68-69°**, **b 100-105°/1mm**, **270°/atm**, pK^{24} **0.79**. Purify the N-oxide by crystallisation from Et_2O and by vacuum sublimation. The *hydrochloride*, **m 179.5-181°**, crystallises from *i*-PrOH. The *picrolonate* has yellow needles with **m 182-184°**. [Katritzky *J Chem Soc* 2404 1956, DOI: 10.1039/JR9560002404, *Beilstein* **20** III/IV 2305, **20/5** V 217.]

Pyridine 3-sulfonic acid [636-73-7] $C_5H_5NO_3S$, M 159.2, m 365-366°(dec), 357°(sec), pK^{25} 2.89 (12% aqueous EtOH), 3.22 (H₂O)(protonation on N). Purify the acid by recrystallisation from H₂O or aqueous EtOH as needles or plates. [pKa: Evans & Brown *J Org Chem* 27 3127 1962, DOI: 10.1021/jo01056a034; IR: Arnett & Chawla *J Am Chem Soc* 100 214 1978, DOI: 10.1021/ja00469a037] Its UV in 50% aqueous EtOH has λ_{max} at 208 and 262nm. The **ammonium salt** has m 243° (from H₂O), the **sulfonyl chloride** has m 133-134° (from petroleum ether), the **amide** has m 110-111° (from H₂O), the **hydrochloride** has m >300°(dec), and the **N-methyl betaine** has m 130° (from H₂O). [Gastel & Wibaut *Recl Trav Chim Pays-Bas* 53 1031 1934, DOI: 10.1002/recl.19340531106; McIlvain & Goese *J Am Chem Soc* 65 2233 1943, DOI: 10.1021/ja01251a063; Machek *Monatsh Chem* 72 77 1939, DOI: 10.1007/BF02716118; *Beilstein* 22 I 616, 22 II 309, 22/7 V 552.]

2-Pyridinethiol (2-mercaptopyridine) [2637-34-5; 73018-10-7] C_5H_5NS , M 111.2, m 127.4°, 127-130°, 130-132°, pK_1^{20} -1.07, pK_2^{20} 9.97. If impure, dissolve it in CHCl₃, wash it with dilute AcOH, H₂O, dry (MgSO₄), evaporate under reduced pressure and recrystallise the residue from *C₆H₆ or H₂O. **2-Methylmercaptopyridine** (b 100-104°/33mm, pK^{20} 3.59) was formed by treatment with MeI/NaOH. [Albert & Barlin *J Chem Soc* 2384 1959, DOI: 10.1039/JR9590002384; *Beilstein* 21 H 45, 21 III/IV 373, 21/7 V 147.]

4-Pyridinethiol (4-mercaptopyridine) [4556-23-4] has m 177°, 179-189°, 186°, pK_1^{20} 1.43, pK_2^{20} 8.86. Purify the thiol by dissolving ~45g in boiling H₂O (100ml) (charcoal), filter and precipitate it by adding 50% aqueous NaOH (~80ml) to pH ~6. Dissolve the precipitate in EtOH, evaporate it to dryness, then crystallise it from boiling EtOH (~100ml, charcoal) to give yellow flat hexagonal plates (m 186°). It sublimes readily *in vacuo*. [King & Ware *J Chem Soc* 873 1939, DOI: 10.1039/JR9390000873.] The **picrate** forms yellow needles from H₂O with m 222°(dec). The **4-methylmercaptopyridine** derivative crystallises from petroleum ether (m 47°, also 44-45° was reported, with a pK^{20} of 5.97) and was prepared by treatment with MeI/NaOH. Its **picrate** has m 245° (from H₂O, MeOH or EtOH). The **N-methyl-4-pyridinethiol** derivative has m 168.5-170° (from EtOH), a pK^{20} of 1.30 and is soluble in CHCl₃. [Albert & Barlin *J Chem Soc* 2384 1959, DOI: 10.1039/JR9590002384; *Beilstein* 21 II 35, 22 III/IV 373, 22/7 V 147.]

1-(2-Pyridylazo)-2-naphthol (PAN) [85-85-8] $C_{15}H_{11}N_3O$, M 249.3, m 142°, 138-141°, 142°, pK_1^{30-36} 2.9, pK_2^{30-36} 11.2. Purify PAN by repeated crystallisation from EtOH or MeOH. It can also be purified by sublimation under vacuum. Purity can be checked by TLC using a mixed solvent (petroleum ether/Et₂O/EtOH; 10:10:1) on a silica gel plate. It has pK_1 1.9 and pK_2 12.2 in 20% aqueous ethoxyethanol. It chelates with copper [Pease & Williams *Anal Chem* 31 1044 1959, DOI: 10.1021/ac60150a027]. [*Beilstein* 22 I 694, 22 III/IV 7073, 22/4 V 618.]

4-(2-Pyridylazo)-resorcinol (PAR) [1141-59-9] $C_{11}H_9N_3O_2$, M 215.2, m 192-202°(dec), >195°(dec), λ_{max} 415nm, ϵ 2.59 x 10⁴ (pH 6-12), pK_1^{25} 2.69 (3.1), pK_2^{25} 5.50 (5.8), pK_3^{25} 12.5 (11.9). Purify PAR as the sodium salt by recrystallisation from 1:1 EtOH/water. Purity can be checked by TLC using a silica gel plate and a mixed solvent (*n*-BuOH:EtOH:2M NH₃; 6:2:2). [*Beilstein* 22 I 694, 22 III/IV 7074, 22/14 V 619.]

3-(2-Pyridyl)-5,6-diphenyl-1,2,4-triazine [1046-56-6] $C_{20}H_{14}N_4$, M 310.4, m 191-192°, 191-193°. Purify it by repeated recrystallisation from EtOH/dimethylformamide. It is a reagent for estimating Fe(II) and Ru(II). [Chriswell & Schilt *Anal Chem* 46 992 1974, DOI: 10.1021/ac60344a021; Karma & Ayers *Anal Chim Acta* 78 423 1975, DOI:10.1016/S0003-2670(00)00163-X.]

1-(4-Pyridyl)ethanol [*R*-(+) 27854-88-2; *S*-(-) 54656-96-1] C_7H_9NO , M 123.2, m 63-65°, 67-69°, b 138-140°/30mm, 254°/760mm, $[\alpha]_D^{20}$ *R*(+) +49.8 and *S*(-) -49.8 (c 0.5, EtOH), and *R*(+) +55 and *S*(-) -55 (c 1, CHCl₃), pK_{Est} ~5.4. Purify it by recrystallisation from petroleum ether. The m recorded after sublimation was 59.9-60.2°, and 55° after crystallisation from *C₆H₆/petroleum ether or petroleum ether/*C₆H₆. The (-)-**di-O-benzoyl tartrate salt** has m 146-148° (from EtOH). [UV, ORD: Gottarelli & Samori *JCS Perkin Trans* 2 1462 1974, DOI: 10.1039/P29740001462.] The **racemate** recrystallises from Et₂O with m 74-76°, b 90-94°/1mm. The **picrate** has m 125-126° (from *C₆H₆). [Ferles & Attia *Coll Czech Chem Commun* 38 611 1973, UV, NMR: Nielson et al. *J Org Chem* 29 2898 1964, DOI: 10.1021/jo01033a022; *Beilstein* 21/2 V 217.]

5-(3-Pyridyl)-4*H*-1,2,4-triazole-3-thiol [1,2-dihydro-5-(3-pyridinyl)-3*H*-1,2,4-triazole-3-thione] [32362-88-2] $C_7H_6N_4S$, M 178.2, m 285-286°, 296-302°, $pK_{Est(1)}$ ~3.0 (py N), $pK_{Est(2)}$ ~9 (SH). This thio-1,2,4-triazole

was prepared by fusing nicotinoylhydrazide with thiourea at $\sim 150^\circ$ then at $160\text{--}165^\circ$ (30 minutes), cooled, EtOH was added and the white solid was filtered off [Beyerman et al. *Recl Trav Chim Pays-Bas* **73** 109 1954, DOI: 10.1002/recl.19540730205]; or by heating 1-nicotinoyl-3-thiosemicarbazide in 5% aqueous NaOH at 100° (4 hours), cooled, acidified with AcOH, and filtered off [Yale & Pila *J Med Chem* **9** 42 1966, DOI: 10.1021/jm00319a010]. It was recrystallised from 95% EtOH, *n*-BuOH/H₂O, H₂O, and 0.5% aqueous HCl provides the *hydrochloride*, which is dried *in vacuo* [over P₂O₅ or H₂SO₄]. [Yoshida & Asai *J Pharm Soc Jpn* **74** 948 1954, *Chem Abstr* **49** 10937 1955, *Beilstein* **26** IV 2129].

5-(4-Pyridyl)-4H-1,2,4-triazole-3-thiol [1,2-Dihydro-5-(4-pyridinyl)-3H-1,2,4-triazole-3-thione] [14910-06-6] C₇H₆N₄S, M 178.2, m 301-303° (dec), 305-308° (dec), 308-313°, 320-322° (dec), 321-322° (dec), pK_{Est(1)} ~ 3.5 (py N), pK_{Est(2)} ~ 9 (SH). This thio-1,2,4-triazole was prepared and purified as for the 3-pyridyl-isomer above but by using isonicotinic acid derivatives. It has mild diuretic and natriuretic activity. [Blackman et al. *J Chem Soc C* 661 1967, DOI: 10.1039/J39670000661, Beyerman et al. *Recl Trav Chim Pays-Bas* **73** 109 1954, DOI: 10.1002/recl.19540730205; Yale & Pila *J Med Chem* **9** 42 1966, DOI: 10.1021/jm00319a010; Yoshida & Asai *J Pharm Soc Jpn* **74** 948 1954, *Chem Abstr* **49** 10937 1955, *Beilstein* **26** IV 2129]. **5-(4-Pyridyl)-4H-1,2,4-triazole-3-sulfonic acid hydrate** was obtained by oxidising the 3-thiol with saturated aqueous KMnO₄ at 25° (1 hour), decomposed with EtOH, filtered, precipitated as the Ag salt, filtered off, and decomposed with 5% HCl, then evaporated, and the residue was recrystallised from H₂O. It decomposed on heating. [Blackman et al. *J Chem Soc C* 661 1967, DOI: 10.1039/J39670000661.]

α -Pyrone (2H-pyran-2-one, coumalin) [504-31-4] C₇H₆N₄S, M 96.1, m 5°, 8-9°, b 102-103°/20mm, 110°/26mm, 104°/ 30mm, 115-118°/37mm, 206-207°/atm, d₄²⁰ 1.1972, n_D²⁰ 1.5298, pK²⁵ -1.14 (aqueous H₂SO₄). Dissolve α -pyrone in Et₂O, wash it with brine, dry (Na₂SO₄), filter, evaporate, distil the residue under vacuum and redistil it. It is a colourless liquid. Its IR has ν_{\max} at 1622 and 1752cm⁻¹ (CHCl₃). [Zimmermann et al. *Org Synth Coll Vol* **5** 982 1973, DOI: 10.15227/orgsyn.046.0101; Nakagawa et al. *Org Synth* **56** 49 1977, DOI: 10.15227/orgsyn.056.0049; Fried & Elderfield *J Org Chem* **6** 566 1941, DOI: 10.1021/jo01204a010.] The *picrate* has m 106-107° (from EtOH). [*Beilstein* **17** H 271, **17** II 305, **17** III/IV 4399, **17**/9 V 288.]

γ -Pyrone (4H-pyran-4-one) [108-97-4] has m 32.5-32.6°, 33°, 32-34°, b 88.5°/7mm, 91-91.5°/9mm, 95-97°/13mm, 105°/23mm, 215°/atm, pK²⁵ 0.10. Purify γ -pyrone by vacuum distillation; the distillate crystallises and is *hygroscopic*. It is non-steam volatile. The *hydrochloride* has m 139° (from EtOH), and the *picrate* has m 130.2-130.3° (from EtOH or H₂O). [Mayer *Chem Ber* **90** 2362 1957, DOI: 10.1002/cber.19570901035; IR: Jones et al. *Can J Chem* **37** 2007 1959, DOI: 10.1139/v59-293; Neelakatan *J Org Chem* **22** 1584 1957, DOI: 10.1021/jo01363a012; *Beilstein* **17** H 271, **17** I 145, **17** II 305, **17** III/IV 4399, **17**/9 V 290.]

Pyronin G [3,6-bis(dimethylamino)xanthylium chloride] [92-32-0] C₁₇H₁₉ClN₂O, M 302.8, m 250-260°, CI 45005, λ_{\max} 522nm, pK_{Est} ~ 7.6 . Commercial material may contain a large quantity of zinc. Purify it by dissolving 1g in 50ml of hot water containing 5g NaEDTA. Cool to 0° , filter, evaporate to dryness and the residue is extracted with EtOH. The solution is evaporated to 5-10ml, filtered, and the dye is precipitated by addition of excess of dry diethyl ether. It is centrifuged, and the crystals are washed with dry ether. The procedure is repeated, then the product is dissolved in CHCl₃, filtered and evaporated. The dye is stored in a vacuum. It forms lustruous green crystals of the FeCl₃ complex. [*Beilstein* **18** H 596, **18** III/IV 7361, **18**/10 V 181.] Bacterial and biological stain.

Pyrrol-2,5-dione (maleimide, 2,5-pyrroledione) [541-59-3] C₄H₃NO₂, M 97.1, m 91-93°, 92.6-93°, d_D^{105.5} 1.2493, n_D^{110.7} 1.49256. Purify it by sublimation in a vacuum. The UV has λ_{\max} at 216 and 280nm in EtOH. [de Wolf & van de Straete *Bull Soc Chim Belg* **44** 288 1935, UV: Rondestvedt et al. *J Am Chem Soc* **78** 6115 1956, DOI: 10.1021/ja01604a044; IR: Chiorboli & Mirone *Ann Chim (Rome)* **42** 681 1952.] Used in Rhodium-catalysed conjugate for arylation with arylboronic acids [Iyer et al. *Tetrahedron Lett* **48** 4413 2007, DOI: 10.1016/j.tetlet.2007.04.084]. [*Beilstein* **21**/10 V 3.]

Pyrrole [109-97-7] C₄H₅N, M 67.1, m -23.4°, b 66°/80mm, 129-130°/atm, d₄²⁰ 0.966, n_D²⁰ 1.5097, pK₁²⁵ -4.4 (Protonation on carbon), pK₂²⁵ 17.51 (aqueous KOH, H. scale). Dry pyrrole with NaOH, CaH₂ or CaSO₄. Fractionally distil it under reduced pressure from CaH₂. Store it under nitrogen as it turns brown in air. Redistil it immediately before use. It polymerises in the presence of acids, and with dilute HCl it gives 'pyrrole red'

which is an amorphous orange coloured solid. The *picrate* forms orange-red crystals with **m** 69°(dec). [Beilstein 20 H 4, 20 I 3, 20 II 3, 20 III/IV 61, 20/5 V 3.]

1H-Pyrrole-1-propanoic acid (3-[pyrrol-1-yl]propionic acid) [89059-06-3] $C_7H_9NO_2$, **M** 139.2, **m** 59-64°, 62°, 62.5°, **pK_{Est}** ~4.5. Recrystallise the acid from petroleum ether (b 80-100°) and dry it *in vacuo*. It readily forms the N-Na and N-K salts. The *ethyl ester* has **b** 122°/23mm. The *amide* forms colourless needles from *C₆H₆ with **m** 81° and is soluble in cold H₂O. [Clemo & Ramage *J Chem Soc* 49 1931, DOI: 10.1039/JR9310000049; Jefford & Johncock *Helv Chim Acta* 66 2666 1983, DOI: 10.1002/hlca.19830660835.]

Pyrrolidine (tetrahydropyrrole) [123-75-1] C_4H_9N , **M** 71.1, **b** 87.5-88.5°, **d**₄²⁰ 0.860, **n**_D²⁰ 1.443, **pK₂₅** 11.31. Dry pyrrolidine with BaO or sodium, then fractionally distil it, under N₂, through a Todd column packed with glass helices. It is a strong base, store away from CO₂. [Beilstein 20 H 159, 20 I 36, 20 II 79, 20 III/IV 2072, 20/1 V 162.]

Quercetin (2H₂O) (3,3',4',5,6-pentahydroxyflavone) [6151-25-3 (2H₂O); 117-39-3 (anhydrous)] $C_{15}H_{10}O_7 \cdot 2H_2O$, **M** 338.3, **m** ca 315°(dec), 317.4-317.9°(dec), (phenolic **pKs** 7–10). Crystallise the yellow flavone from aqueous EtOH and dry at 100°. It also sublimes in a vacuum. Its IR has ν_{max} at 2.5 μ and 16.7 μ (KBr). It complexes with Cu²⁺, Al³⁺, Ca²⁺, Ge⁴⁺, Zn²⁺, Ti⁴⁺, Zr⁴⁺, Th⁴⁺, UO⁴⁺. [Beilstein 18 H 242, 18 I 242, 18 II 236, 18 III/IV 3470, 18/5 V 494.] It exhibits anti-cancer activity.

Quinaldic (quinoline-2-carboxylic) acid [93-10-7] $C_{10}H_7NO_2$, **M** 173.2, **m** 155-157°, 156-157°, **pK₁²⁵** 1.45, **pK₂²⁵** 2.49 (2.97). Crystallise quinaldic acid from *C₆H₆ or AcOH. It is used for the estimation of many metals. The *methyl ester* has **m** 86-87° (from hexane) and **pK₂₅** 1.76. [Chauduri et al. *Frez Z Anal Chem* 281 361 1976, Beilstein 22 H 71, 22 II 55, 22 III/IV 1149, 22/3 V 183.] It forms insoluble salts with copper, uranium and zinc; and is used for their determination.

Quinazoline [253-82-7] $C_8H_6N_2$, **M** 130.2, **m** 46-48°, 48.0-48.5°, **b** 120-121°/17-18mm, 241°/764mm, 243°/772mm, **pK₁²⁰** -4.51 (aqueous H₂SO₄, anhydrous dication), **pK₂²⁰** 2.01 (anhydrous monocation), **pK₃²⁰** 4.3 (equilibrium with 3,4-hydrated species), **pK₄²⁰** 12.1 (hydrated anion). Purify quinazoline by passage through an activated alumina column in *C₆H₆ or petroleum ether (b 40-60°). Distil it under reduced pressure, sublime it under vacuum and recrystallise it from petroleum ether. The *hydrochloride hydrate* has **m** 127-128° (from EtOH/HCl/ Et₂O). The *picrate* has **m** 188-189° (from MeOH). [Armarego *J Appl Chem* 11 70 1961, DOI: 10.1002/jctb.5010110207; Armarego *Quinazolines, Fused Pyrimidines Part I* Brown Ed, Wiley-Interscience 1967, Brown *Quinazolines Supplement I* Taylor Ed, Wiley-Interscience 1996, ISBN 0-471-14565-3; for covalent hydration see Albert & Armarego *Adv Heterocycl Chem* 4 1 1965, DOI:10.1016/S0065-2725(08)60873-9; Elderfield et al. *J Org Chem* 12 405 1947, DOI: 10.1021/jo01167a007; Beilstein 23 H 175, 23 II 177, 23 III/IV 1221.]

Quinoline [91-22-5] C_9H_7N , **M** 129.2, **m** -20°, -16°, **b** 113-114°/17mm, 236°/758mm, **d**₄²⁰ 1.0937, **n**_D²⁰ 1.625, **pK₂₅** 4.80 (4.93). Dry quinoline with Na₂SO₄ and distil it from zinc dust in a vacuum. It has also been dried by boiling with acetic anhydride, then fractionally distilled. Calvin and Wilmarth [*J Am Chem Soc* 78 1301 1956, DOI: 10.1021/ja01588a009] cooled redistilled quinoline in ice and added enough HCl to form its hydrochloride. Diazotization removed aniline, the diazo compound being broken down by warming the solution to 60°. Non-basic impurities were removed by ether extraction. Quinoline was then liberated by neutralising the hydrochloride with NaOH, then dried with KOH and fractionally distilled at low pressure. Addition of cuprous acetate (7g/L of quinoline) and shaking under hydrogen for 12 hours at 100° removed impurities due to the nitrous acid treatment. Finally the hydrogen was pumped off, and the quinoline was distilled. Other purification procedures depend on conversion to the *phosphate* (**m** 159°, precipitated from MeOH solution, filtered, washed with MeOH, then dried at 55°) or the *picrate* (**m** 201°) which, after recrystallisation, were reconverted to quinoline. The method using the picrate [Packer et al. *J Am Chem Soc* 80 905 1958, DOI: 10.1021/ja01537a039] is as follows: quinoline is added to picric acid dissolved in the minimum volume of 95% EtOH, giving yellow crystals which were washed with EtOH, air-dried and crystallised from acetonitrile. These were dissolved in dimethyl sulfoxide (previously dried over 4A molecular sieves) and passed through a basic alumina column, onto which the picric acid is adsorbed. The *free base* in the effluent is extracted with *n*-

pentane and distilled under vacuum. Traces of solvent can be removed by vapour-phase chromatography. [Moomaw & Anton *J Phys Chem* **80** 2243 1976, DOI: 10.1021/j100561a022.] The ZnCl_2 and dichromate complexes have also been used [Cumper et al. *J Chem Soc* 1176 1962, DOI: 10.1039/JR9620001176]. [Beilstein **20** H 339, **20** I 134, **20** II 222, **20** III/IV 3334, **20/7** V 276.] **Quinoline ethiodide (1-ethylquinolinium iodide)** [634-35-5] $\text{C}_{11}\text{H}_{12}\text{IN}$, M **285.1**, has m **157-160°**, **158-159°**. Crystallise it from aqueous EtOH or EtOH/peroxide free Et_2O . [Beilstein **20** I 139, **20** II 231, **20** III/IV 3357, **20/7** V 276.]

2-Quinolinealdehyde [5470-96-2] $\text{C}_{10}\text{H}_7\text{NO}$, M **157.2**, m **70-72°**, **71°**, $\text{pK}_{\text{Est}} \sim 3.3$. Distil it in steam and recrystallise it from H_2O . Protect it from light. The *semicarbazone* has m **254°** (from aqueous EtOH), and the *picrate* has m **197-199°**. [Beilstein **21** H 322, **21** III/IV 4034, **21/8** V 442.]

8-Quinolinecarboxylic acid [86-59-9] $\text{C}_{10}\text{H}_7\text{NO}_2$, M **173.2**, m **183-185°**, **186-187.5°**, pK_1^{25} **1.82**, pK_2^{25} **6.87**. Crystallise the acid from water, aqueous EtOH, EtOH or $^*\text{C}_6\text{H}_6$. The *ethyl ester* has m **45°** and b **194-197°/13mm**. [Beilstein **22** H 81, **22** III/IV 1200, **22/3** V 217.] Forms complexes with Ag, Cd, Cu, Fe, Hg, Pb and Tl, and for estimation of Cu.

Quinoxaline [91-19-0] $\text{C}_8\text{H}_6\text{N}_2$, M **130.2**, m **28°** (anhydrous), **29-32°**, **37°(H₂O)**, b **108-110°/0.1mm**, **140°/40mm**, **220-223°/atm**, pK_1^{20} **-5.52** (-5.8, dication), pK_2^{20} **0.56** (c 0.5), **0.72** (c 1.0) (monocation). Crystallise quinoxaline from petroleum ether. It crystallises as the *monohydrate* on addition of water to a petroleum ether solution. It has UV with λ_{max} at 242 and 331nm ($\text{H}^0 -2$); 234 and 316nm (pH 7.1). The *picrate* has m **161-162°**. [Albert & Phillips *J Chem Soc* 1294 1956, DOI: 10.1039/JR9560001294; Brown *Chemistry of Heterocyclic Compounds: Quinoxaline (Supplement 1)* vol **61** pp 510 2004, Wiley-Interscience NY, ISBN: 0471264954; Beilstein **23** H 176, **23** II 177, **23** III/IV 1226, **23/7** V 135.]

Quinoxaline-2,3-dithiol [1199-03-7, 40158-04-1] $\text{C}_8\text{H}_6\text{N}_2\text{S}_2$, M **194.1**, m **345°(dec)**, pK_1 **6.9**, pK_2 **9.9**. Purify the dithiol by repeated dissolution in alkali and re-precipitation by acetic acid. It complexes with Ag^+ , Cd^{2+} , Pb^{2+} , Bi^{3+} and Ni^{2+} in aqueous NH_3 . It is used for the gravimetric analysis of Ni [Dalziel & Slawinski *Talanta* **15** 11385 1968, DOI: 10.1016/0039-9140(68)80198-5]. [Beilstein **24** III/IV 1428.]

Quinuclidine (1-azabicyclo[2.2.2]octane) [100-76-5] $\text{C}_7\text{H}_{13}\text{N}$, M **111.2**, m **157-160°**, **158°(sublimes)**, **158-159°**, pK^{25} **10.95**. Crystallise it from diethyl ether, or petroleum ether (b 40-50°) and cooling at 0°. It sublimes as colourless dendrites on heating during melting point determination in a sealed tube. It is a strong organic base. The *hydrochloride* [39896-06-5] has m **364-365°(dec)** (from EtOH or n-BuOH), and the *picrate* has m **275°** (also **254°**) (from aqueous EtOH). [Clemo & Metcalfe *J Chem Soc* 1989 1937, DOI: 10.1039/JR9370001989; Leonard & Elkin *J Org Chem* **27** 463 1962, DOI: 10.1021/jo01059a502; Beilstein **20** H 144, **20** II 71, **20** III/IV 1966, **20/4** V 335.]

Rhamnetin (3,3'-4',5-tetrahydroxy-7-methoxy flavone, 7-methyl quercetin) [90-19-7] $\text{C}_{16}\text{H}_{12}\text{O}_7$, M **316.3**, m **>300°(dec)**, several phenolic $\text{pKs} \sim 7-10.5$. Crystallise rhamnetin from EtOH (m **292-293°**), aqueous EtOH (m **294-296°**) or MeOH (m **290-294°**), or $\text{Me}_2\text{CO}/\text{MeOH}$. [Kuhn & Low *Chem Ber* **77** 211 1944, DOI: 10.1002/cber.19440770313; Jurd *J Am Chem Soc* **80** 5531 1958, DOI: 10.1021/ja01553a054.] The *tetraacetate* has m **189-190°** (from $\text{Me}_2\text{CO}/\text{MeOH}$). [Beilstein **18** H 245, **18** II 237, **18** III/IV 3474.]

Rhodamine 3B chloride [3,5-bis-(diethylamino)-9-(2-carboxyphenyl)xanthylum chloride] [81-88-9] $\text{C}_{28}\text{H}_{31}\text{ClN}_2\text{O}_3$, M **479.0**, m **210-211°(dec)**, CI **45170**, λ_{max} **543nm**, {Free base [509-34-2] CI **749**}, pK^{25} **5.53**. Major impurities are partially dealkylated compounds not removed by recrystallisation. Purify the dye by chromatography, using ethyl acetate/isopropanol/ammonia (880)(9:7:4, R_F 0.75 on Kieselgel G). It has also been crystallised from a concentrated solution in MeOH by slow addition of dry diethyl ether; or from EtOH containing a drop of concentrated HCl by slow addition of ten volumes of dry diethyl ether. The solid is washed with ether and air dried. The dried material has also been extracted with $^*\text{benzene}$ to remove oil-soluble material prior to recrystallisation. Store it in the dark. It is a colorimetric reagent for Cd, and suitable as a laser dye, and a powerful dye. [Beilstein **18** II 486, **18** III/IV 8246, **19/8** V 669.]

Rhodamine 6G [Basic Red 1, 3,5-bis-(ethylamino)-9-(2-ethoxycarbonylphenyl)-2,7-dimethylxanthylium chloride] [989-38-8] $C_{28}H_{31}ClN_2O_3$, M 479.3, CI 45160, λ_{max} 524nm, pK^{25} 5.58. Crystallise the dye from MeOH or EtOH, and dry it in a vacuum oven. It is a powerful dye, and has found use in identifying lipid materials on TLC plates, as it fluoresces under UV light. [Beilstein 18 III/IV 8244, 18/12 V 283.]

Rhodanine (2-mercaptothiazolidin-4-one) [141-84-4] $C_3H_3NOS_2$, M 133.2, has m 165-169°, 168.5° (capillary), pK^{20} 5.18. Crystallise rhodanine from glacial acetic acid or water. It is used to estimate Ag and gallic acid [Thies & Fischer *Mikrochimica Acta* 182 809 1973, DOI: 10.1007/BF01218140]. [Beilstein 27 H 242, 27 I 309, 27 II 288, 27 III/IV 3188.]

Saccharin (1,2-benzisothiazol-3(2H)-one 1,1-dioxide, o-benzoic acid sulfimide) [81-07-2] $C_7H_5NO_3S$, M 183.2, m 226-229°, 227-229°, 229°, 228.8-229.7°, pK^{25}_I 1.31, pK^{25}_2 12.8. Purify saccharin by recrystallisation from Me_2CO [solubility 7.14% at 0°, 14.4% at 50°], or aqueous isoPrOH to give a fluorescent solution. It sublimes *in vacuo*. It is an artificial sweetener which is 500 times sweeter than sucrose. [DeGarmo et al. *J Am Pharm Assoc (Sci Ed)* 41 17 1952, Beilstein 27 H 168, 870, 27 I 266, 27 II 214, 27 III/IV 2649.]

Safranine O (Safranine T, 3,7-diamino-2,8-dimethyl-5-phenylphenazinium chloride, cotton Red) [477-73-6] $C_{20}H_{19}ClN_4$, M 350.9, λ_{max} 530nm, pK^{25} 6.4. Crystallise it from *benzene/MeOH (1:1) or water. Dry it *in vacuo* over H_2SO_4 . It has UV: λ_{max} at 520nm (H_2O) and 530nm (EtOH). It is a certified dye for staining chromosomes, Gram negative bacteria and plant cells. [Beilstein 25 H 403, 25 I 657, 25 III/IV 3056.]

Safrole (5-allyl-1,3-benzodioxole, 4-allyl-1,2-methylenedioxybenzene) [94-59-7] $C_{10}H_{10}O_2$, M 162.1, m~11°, b 69-70°/1.5mm, 104-105°/6mm, 231.5-232°/atm, 235-237°/atm, d^{20}_4 1.0993, n^{20}_D 1.53738. Safrole has been purified by fractional distillation, although it has also been recrystallised from low boiling petroleum ether at low temperatures. [IR: Briggs et al. *Anal Chem* 29 904 1957, DOI: 10.1021/ac60126a014; UV: Patterson & Hibbert *J Am Chem Soc* 65 1862 1943, DOI: 10.1021/ja01250a021.] The *maleic anhydride adduct* forms yellow crystals from toluene m 257° [Hickey *J Org Chem* 13 443 1948, DOI: 10.1021/jo01161a020], and the *picrate* forms orange-red crystals from $CHCl_3$ [Baril & Megrdichian *J Am Chem Soc* 58 1415 1936, DOI: 10.1021/ja01299a031]. It is a sweet smelling oil. [Beilstein 19 I 617, 19 II 29, 19 III/IV 275, 19/I V 553.]

Scopoletin (7-hydroxy-6-methoxycoumarin) [92-61-5] $C_{10}H_8O_4$, M 192.2, m 203-205°, 206°, 208-209°, pK^{25} 8.96 (70% aqueous EtOH). Crystallise it from water, acetic acid or * C_6H_6 /MeOH. It is dimorphic with a second m at 193-195°. It sublimes at 120-130°/12mm. This dye is used for detecting release of oxygen reactive species, is a peroxynitrite scavenger and a cholinesterase inhibitor. [Beilstein 18 III/IV 1323, 18/3 V 203.]

Styrene oxide (phenyloxirane) [96-09-3] C_8H_8O M 120.2, m -37°, b 84-86°/16.5mm, 194°/atm, d^{20}_4 1.053, n^{25}_D 1.535. Fractional distillation under reduced pressure does not remove phenylacetaldehyde. If this material is present, the styrene oxide is treated with hydrogen under 3 atmospheres pressure in the presence of platinum oxide. The aldehyde, but not the oxide, is reduced to β -phenylethanol, and separation is now readily achieved by fractional distillation. [Schenck & Kaizermen *J Am Chem Soc* 75 1636 1953, DOI: 10.1021/ja01103a035; Beilstein 17/I V 577.]

2-Sulfobenzoic cyclic anhydride (2,1-benzoxathiazol-3-one-1,1-dioxide) [81-08-3] $C_7H_4O_4S$, M 184.2, m 126-127°, 129.5°, 130°, b 184-186°/18mm. The anhydride is purified by distillation in a vacuum and readily solidifies to a crystalline mass on cooling. [Heitman *J Am Chem Soc* 34 1591 1912, DOI: 10.1021/ja02212a019.] *Alternatively*, purify it by dissolving it in the minimum volume of toluene and refluxing for 2 hours using a Dean-Stark trap. Evaporate under reduced pressure and distil the anhydride at 18mm. It is then recrystallised three times from its own weight of dry * C_6H_6 . It is sensitive to moisture and should be stored in the dark in a dry atmosphere. The *O-methyloxime* has m 110-112° [Levy *Tetrahedron Lett* 13 3289 1972, DOI: 10.1016/S0040-4039(01)94025-0]. If the sample has hydrolysed extensively (presence of OH band in the IR) then treat with an equal bulk of $SOCl_2$, reflux it for 3 hours ($CaCl_2$ tube), evaporate and distil the residue in a vacuum, then recrystallise it from * C_6H_6 , Et_2O /* C_6H_6 or $CHCl_3$ (EtOH free by passing through Al_2O_3 , or standing over $CaCl_2$). [Clarke & Dreger *Org Synth Coll Vol* 1 495 1941, DOI: 10.15227/orgsyn.009.0080.] It

is used for modifying ζ -amino functions of lysyl residues in proteins [Bagree et al. *FEBS Lett* **120** 275 1980, DOI: 10.1016/0014-5793(80)80315-2]. [Beilstein **19** I 659, **19** II 137, **19** III/IV 1641, **19/4** V 215.]

Sulfolane (tetramethylenesulfone) [126-33-0] $C_4H_8O_2S$, **M 120.2**, **m 20-26°**, **28.5°**, **b 104°/0.2mm**, **153-154°/18mm**, **285°/760mm**, d_4^{20} **1.263**, n_D^{30} **1.4820**. It is prepared commercially by a Diels-Alder reaction of between 1,3-butadiene and sulfur dioxide, followed by Raney nickel hydrogenation. The principal impurities are water, 3-sulfolene, 2-sulfolene and 2-isopropyl sulfolanyl ether. It is dried by passage through a column of molecular sieves. Distil it under reduced pressure through a column packed with stainless steel helices. Again dry it with molecular sieves and distil. [Cram et al. *J Am Chem Soc* **83** 3678 1961, DOI: 10.1021/ja01478a029; Coetzee *Pure Appl Chem* **49** 211 1977, DOI: 10.1351/pac197749020211.] Alternatively, it is stirred at 50°, and small portions of solid $KMnO_4$ are added until the colour persists during 1 hour. Dropwise addition of MeOH then destroys the excess $KMnO_4$; the solution is filtered, freed from potassium ions by passage through an ion-exchange column and dried under vacuum. It has also been distilled in a vacuum from KOH pellets. It is *hygroscopic*. [See Sacco et al. *J Phys Chem* **80** 749 1976, DOI: 10.1021/j100548a018; Sacco et al. *JCS Faraday Trans 1* **73** 1936 1977, DOI: 10.1039/F19777301936; Petrella & Sacco *JCS Faraday Trans 1* **74** 2070 1978, DOI: 10.1039/F19787402070; Conway et al. *Trans Faraday Soc* **62** 2738 1966, DOI: 10.1039/TF9666202738.] Coetzee has reviewed the methods of purification of sulfolane, and also the removal of impurities. [Coetzee in *Recommended Methods of Purification of Solvents and Tests for Impurities*, Coetzee Ed. Pergamon Press, 1982, *Beilstein* **17** I 5, **17** III/IV 37, **17/1** V 39.]

2,2':6',2''-Terpyridyl [1148-79-4] $C_{15}H_{11}N_3$, **M 233.3**, **m 89-91°**, **91-92°**, pK_1^{23} **2.64**, pK_2^{23} **4.33**. Crystallise it from Et_2O , toluene or from petroleum ether, then aqueous MeOH, followed by sublimation in a vacuum at 90°. It is used for estimating Ag and Ru. [Kamra et al. *Anal Chim Acta* **81** 117 1976, DOI:10.1016/S0003-2670(00)89466-0; *Beilstein* **26** III/IV 258.]

Terthiophene (2,5-di[thienyl]thiophene; α -terthienyl) [1081-34-1] $C_{12}H_8S_3$, **M 248.4**, **m 93-95°**, **94-95.5°**, **94-96°**. Possible impurities are bithienyl and polythienyls. Suspend it in H_2O and steam distil it to remove bithienyl. The residue is cooled and extracted with $CHCl_3$, dried ($MgSO_4$), filtered, evaporated and the residue chromatographed on Al_2O_3 using petroleum ether/3% Me_2CO as eluent. The terphenyl zone is then eluted from the Al_2O_3 with Et_2O , the extract is evaporated and the residue is recrystallised from MeOH (40ml per g). The platelets are washed with cold MeOH and dried in air. [UV: Sease & Zechmeister *J Am Chem Soc* **69** 270 1947, DOI: 10.1021/ja01194a031; Uhlenbroek & Bijloo *Recl Trav Chim Pays-Bas* **79** 1181 1960, DOI: 10.1002/recl.19600791113.] It has also been recrystallised from MeOH, $*C_6H_6$, petroleum ether or AcOH. [UV: Zechmeister & Sease *J Am Chem Soc* **69** 273 1947, DOI: 10.1021/ja01194a032; Steinkopf et al. *Justus Liebigs Ann Chem* **546** 180 1941, DOI: 10.1002/jlac.19415460112.] It is a phototoxic nematocide [Cooper & Nitsche *Bioorg Chem* **13** 362 1985, DOI:10.1016/0045-2068(85)90036-7; Chan et al. *Phytochem* **14** 2295 1975, DOI:10.1016/S0031-9422(00)91121-X]. [*Beilstein* **19** III/IV 4763, **19/9** V 226.]

2,4,5,6-Tetraaminopyrimidine sulfate [5392-28-9] $C_4H_8N_6 \cdot H_2SO_4$, **M 238.2**, **m 255° (dec)**, **>300°**, **>350° (dec)**, pK^{20} **6.82**. Purify the salt by recrystallisation from H_2O , 2N H_2SO_4 (20 parts, 67% recovery) or 0.1N H_2SO_4 (40 parts, 62% recovery), and dried in air. [UV: Konrad & Pfeleiderer *Chem Ber* **103** 722 1970, DOI: 10.1002/cber.19701030311; Malletta et al. *J Am Chem Soc* **69** 1814 1947, DOI: 10.1021/ja01199a073; Cavalieri et al. *J Am Chem Soc* **70** 3875 1948, DOI: 10.1021/ja01191a102; *Beilstein* **25** H 423, **25** III/IV 3106.]

1,4,8,11-Tetraazacyclotetradecane (cyclam) [295-37-4] $C_{10}H_{24}N_4$, **M 200.33**, **m 173° (closed capillary and sublimates at 125°)**, **183-185°**, **184-186°**, **185°**, $pK_{Est(1)} \sim 3.8$, $pK_{Est(2)} \sim 6.0$, $pK_{Est(3)} \sim 9.0$, $pK_{Est(4)} \sim 9.6$. Purify cyclam by recrystallisation from dioxane (white needles), and it sublimates above 120°. It has been distilled, **b 132-140°/4-8mm**. It forms complexes with metals and gives a sparingly soluble *nitrate salt*, **m 205°(dec)**, which crystallises from H_2O and is dried at 150°. [UV: Bosnich et al. *Inorg Chem* **4** 1102 1963, DOI: 10.1021/ic50030a003; van Alphen *Recl Trav Chim Pays-Bas* **56** 343 1937, DOI: 10.1002/recl.19370560405; *Beilstein* **26** III/IV 1647.]

Tetrabenazine (\pm -2-oxo-3-isobutyl-9,10-dimethoxy-1,2,3,4,6,7-hexahydro-11bH-benzo[a]-quinolizine) [58-46-8] $C_{19}H_{27}NO_3$, **M 317.4**, **m 127-128°**, $pK_{Est} \sim 8$. Crystallise it from MeOH. The *hydrochloride* has **m 208-**

210°, and the *oxime* has **m 158°** (from EtOH). [Beilstein 21 III/IV 6488.] Drug for the treatment of hyperkinetic movement associated with Huntingdon's disease [Jankovic & Beach *Neurology* 48 358 1997, DOI: 10.1212/WNL.48.2.358; Robertson *Brain* 123(3) 425 2000, DOI: 10.1093/brain/123.3.425].

2,3,4,6-Tetrachloropyridine [14121-36-9] C_5HCl_4N , **M 216.9**, **m 74-75°**, (? 37-38°), **b 130-135°/16-20mm**, **248.5-249.5°/760mm**, **pK_{Est} ~ -5.7**. Crystallise it from 50% EtOH and/or distil it. The *N-oxide* has **m 210°** (from EtOH/CHCl₃). [Chivers & Suschitzky *J Chem Soc* 2867 1971, DOI: 10.1039/J39710002867; Beilstein 20 III/IV 6488, 20/5 V 421.]

Tetrahydrofuran (oxalane, THF) [109-99-9] C_4H_8O , **M 72.1**, **m -108°**, **b 25°/176mm**, **65-67°/atm**, **66°/760mm**, **d₄²⁰ 0.889**, **n_D²⁰ 1.4070**, **pK²⁵ -2.48 (aqueous H₂SO₄)**. It is obtained commercially by catalytic hydrogenation of furan from pentosan-containing agricultural residues. It was purified by refluxing with, and distilling from LiAlH₄ which removes water, peroxides, inhibitors and other impurities [Jaeger et al. *J Am Chem Soc* 101 717 1979, DOI: 10.1021/ja00497a039]. Peroxides can also be removed by passage through a column of activated alumina, or by treatment with aqueous ferrous sulfate and sodium bisulfate, followed by solid KOH. In both cases, the solvent is then dried and fractionally distilled from sodium. Lithium wire or vigorously stirred molten potassium have also been used for this purpose. CaH₂ has also been used as a drying agent. Several methods are available for obtaining the solvent almost anhydrous. Ward [*J Am Chem Soc* 83 1296 1961, DOI: 10.1021/ja01467a010] dried it vigorously with sodium-potassium alloy until a characteristic blue colour was evident in the solvent at Dry-ice/cellosolve temperatures. The solvent is kept in contact with the alloy until distilled for use. Worsfold and Bywater [*J Chem Soc* 5234 1960, DOI: 10.1039/JR9600005234], after refluxing and distilling from P₂O₅ and KOH, in turn, refluxed the solvent with sodium-potassium alloy and fluorenone until the green colour of the disodium salt of fluorenone was well established. [Alternatively, instead of fluorenone, benzophenone, which forms a blue ketyl, can be used.] The tetrahydrofuran was then fractionally distilled, degassed and stored above CaH₂. *p*-Cresol or hydroquinone inhibit peroxide formation. The method described by Coetzee and Chang [*Pure Appl Chem* 57 633 1985, DOI: 10.1351/pac198557040633] for 1,4-dioxane also applies here. Distillations should always be done in the presence of a reducing agent, e.g. FeSO₄. [Beilstein 17 H 10, 17 I 5, 17 II 15, 17 III/IV 24, 17/I V 27.] **It irritates the skin, eyes and mucous membranes, and the vapour should never be inhaled. It is HIGHLY FLAMMABLE, and the necessary precautions should be taken. Rapid purification:** Purification as for diethyl ether.

***l*-Tetrahydropalmatine (2,3,9,10-tetramethoxy-6*H*-dibenzo[*a,g*]quinolizidine)** [10097-84-4] $C_{21}H_{25}NO_4$, **M 355.4**, **m 147°**, **148-149°**, [α]_D²⁰ **-291 (EtOH)**. It crystallises from MeOH or EtOH (Norit) by addition of water as colourless prisms [see Kametani & Ihara *J Chem Soc (C)* 530 1967, DOI: 10.1039/J39670000530; Bradsher & Dutta *J Org Chem* 26 2231 1961, DOI: 10.1021/jo01351a018]. When crystallised from Me₂CO/Et₂O, it has **m 142°**. The *hydrate* has **m 115°**(effervescence). The (\pm)-*hydrochloride*, prepared by bubbling HCl through a dry ethereal solution of the base, is then recrystallised from MeOH to give colourless needles with **m 215-216°**. The *picrate* has **m 188°(dec)** (from aqueous EtOH). [Beilstein 21 III/IV 2769.]

Tetrahydropyran (oxane, THP) [142-68-7] $C_5H_{10}O$, **M 86.1**, **m -49.2°**, **-45°**, **b 88.0°/atm**, **d₄²⁰ 0.885**, **n_D²⁰ 1.4202**, **pK²⁵ -2.79 (aqueous H₂SO₄)**. Dry oxane with CaH₂, then pass it through a column of silica gel to remove olefinic impurities and fractionally distil it. Free it from peroxides and moisture by refluxing with sodium, then distil it from LiAlH₄. Alternatively, peroxides can be removed by treatment with aqueous ferrous sulfate and sodium bisulfate, followed by solid KOH, and fractional distillation from sodium. It forms an azeotrope with H₂O (b 71°, 8.5% H₂O). Add a stabiliser to avoid peroxide formation due to presence of air. [Beilstein 17 H 12, 17 I 6, 17 II 18, 17 III/IV 51, 17/I V 64.]

Tetrahydro-4*H*-pyran-4-one [29943-42-8] $C_5H_8O_2$, **M 100.1**, **b 57-59°/11mm**, **65-66°/15mm**, **67-68°/18mm**, **73°/20mm**, **164.7°/atm**, **166-166.5°/atm**, **d₄²⁰ 1.0844**, **n_D²⁰ 1.4551**. Purify the pyrone by repeated distillation, preferably in a vacuum. [Baker *J Chem Soc* 296 1944, DOI: 10.1039/JR9440000296; IR: Olsen & Bredoc *Chem Ber* 91 1589 1958, DOI: 10.1002/cber.19580910804.] The *oxime* has **m 87-88°** and **b 110-111°/13mm** [Cornubert et al. *Bull Soc Chim Fr* 36 1950]. The **4-nitrophenylhydrazone** forms orange-brown needles from EtOH, **m 186°** [Cawley & Plant *J Chem Soc* 1214 1938, DOI: 10.1039/JR9380001214]. [Beilstein 17 I 131, 17 II 287, 17 III/IV 4171, 17/9 V 21.]

Tetrahydrothiophene (thiophane) [110-01-0] C_4H_8S , **M 88.2**, **m -96°**, **b 14.5°/10mm**, **40.3°/39.7mm**, **120.9°/760mm**, d_4^{20} **0.997**, n_D^{20} **1.5289**. The crude material is purified by crystallisation of the mercuric chloride complex to a constant melting point. It is then regenerated, washed, dried, and fractionally distilled. [Whitehead et al. *J Am Chem Soc* **73** 3632 1951, DOI: 10.1021/ja01152a022.] It has been dried over Na_2SO_4 and distilled in a vacuum and the purity is checked by 1H NMR [Roberts & Friend *J Am Chem Soc* **108** 7204 1986, DOI: 10.1021/ja00283a011; for the C_4H_8S -AuCl complex see Uson et al. *Inorg Synth* **26** 85 2007; DOI: 10.1002/9780470132579.ch17]. [*Beilstein* **17** I 5, **17** II 15, **17** III/IV 34, **17**/I V 36.]

Tetrahydro-4H-thiopyran-4-one [1072-72-6] C_5H_8OS , **M 116.2**, **m 60-62°, 61-62°, 64-65°, 65-66°, 65-67°**. Purify it by recrystallisation from diisopropyl ether or petroleum ether and dry it in air. If too impure, then dissolve it in Et_2O , wash with aqueous $NaHCO_3$, then H_2O , dry ($MgSO_4$), filter, evaporate and the residue is recrystallised as before. [Cardwell *J Chem Soc* 715 1949, DOI: 10.1039/JR9490000715.] The **oxime** can be recrystallised from $CHCl_3$ /petroleum ether (at -20°) and has **m 84-85°** [Barkenbus et al. *J Org Chem* **20** 871 1955, DOI: 10.1021/jo01125a011]. The **2,4-dinitrophenylhydrazones** has **m 186°** (from $EtOAc$) [Barkenbus et al. *J Org Chem* **16** 232 1951, DOI: 10.1021/jo01142a011]. The **S-dioxide** is recrystallised from $AcOH$, **m 173-174°**, and **1,4-thiapyrone-1,1-dioxide oxime** has **m 197.8° dec** (from $MeOH$). [Fehnel & Carmack *J Am Chem Soc* **70** 1813 1948, DOI: 10.1021/ja01185a048; *Beilstein* **17** II 287, **17** III/IV 4172, **17**/I V 21].

Tetramethylene sulfoxide (tetrahydrothiophen 1-oxide) [1600-44-8] C_4H_8OS , **M 104.2**, **b 45°/3mm**, **235-237°/atm**, d_4^{20} **1.175**, n_D^{20} **1.525**. Shake the oxide with BaO for 4 days, then distil it from CaH_2 under reduced pressure. It can also be purified by distillation under reduced pressure over Lind 13X molecular sieves (conditioned by calcination at 400°/4hrs under N_2). Store below 5° as its shelf life at ~20° is *ca* 3 months. It is an excellent reagent for oxidising thiols and dithiols to the respective disulfides, requiring one mol of reagent per 2mols of SH [Wallace *J Am Chem Soc* **86** 2018 1964, DOI: 10.1021/ja01064a022; Wallace & Mahon *J Am Chem Soc* **86** 4099 1965, DOI: 10.1021/ja01073a039]. [*Beilstein* **17** III/IV 36, **17**/I V 38.]

2,2,6,6-Tetramethylpiperidiny-1-oxy (TEMPO) [2564-83-2] $C_9H_{18}NO$, **M 156.3**, **m 36-38°, 37-39°**. Purify TEMPO by sublimation (33°, water aspirator, **m 39°**). [Hay & Finke *J Am Chem Soc* **109** 8012 1987, DOI: 10.1021/ja00260a011; Rozantsev & Neiman *Tetrahedron* **20** 131 1964, DOI: 10.1016/S0040-4020(01)98404-2; Keana *Chem Rev* **78** 37 1978, DOI: 10.1021/cr60311a004].

2,2,6,6-Tetramethyl-4-piperidone hydrochloride (triacetoneamine HCl) [33973-59-0] $C_9H_{17}NO$. **HCl, M 191.7**, **m 190°(dec)**, **198-199°(dec)**, pK_{25}^{25} **7.90**. Purify the salt by recrystallisation from $EtOH/Et_2O$, $MeCN$ or $Me_2CO/MeOH$. The **free base** [826-36-8] $C_9H_{17}NO$, **M 155.2**, has **m 37-39°** (after sublimation, also **m 34-38°**), **b 102-105°/18mm**, and the **hydrate** has **m 56-58°** (wet Et_2O); the **hydrobromide** has **m 203°** (from $EtOH/Et_2O$), and the **picrate** has **m 196°** (from aqueous $EtOH$). [Sandris & Ourisson *Bull Soc Chim Fr* 345 1958, *Beilstein* **21** H 249, 246, **21** I 273, **21** II 222, **21** III/IV 3278, **21**/6 V 538.]

1,3,7,9-Tetramethyl uric acid [2309-49-1] $C_9H_{12}N_4O_3$, **M 224.2**, **m 225°, 226°, 228°, $pK_{Est} < 0$** . Crystallise the uric acid from H_2O or $MeOH$. [Lister *Purines Part II, Fused Pyrimidines* Brown Ed, Wiley-Interscience pp254-225 1971, ISBN 0-471-38205-1, *Beilstein* **26** H 532, **26** I 156, **26** II 302, **21** III/IV 2623.]

1,3,5,5-Tetranitrohexahydropyrimidine [81360-42-1] $C_4H_6N_6O_8$, **M 266.1**, **m 153-154°**. Crystallise the nitropyrimidine from $EtOH$ (5x) and sublime it (~65°/0.05mm). It has 1H NMR ($CDCl_3$ with 1drop of Me_2SO-d_6) with δ at 6.13 (s, NCH_2N), 5.23 (s, CCH_2N). [Cichra & Adolph *J Org Chem* **47** 2474 1982, DOI: 10.1021/jo00133a047; Shackelford *J Labelled Comp Radiopharm* **29** 1197 1991, DOI: 10.1002/jlcr.2580291104].

4,7,13,18-Tetraoxa-1,10-diazabicyclo[8.5.5]eicosane (Cryptand 211, Kryptofix 221) [31250-06-3] $C_{14}H_{28}N_2O_4$, **M 288.1**, **b 130°/0.002mm**, d_4^{20} **1.097**, n_D^{20} **1.505**, $pK_{Est} \sim 7.9$. Redistil Cryptand 211 under high vacuum, dry it under high vacuum over 24 hours, and store it under nitrogen. [Alberto et al. *J Am Chem Soc* **121** 3135 2001, DOI: 10.1021/ja003932b.]

1,7,10,16-Tetraoxa-4,13-diazacyclooctadecane (4,13-diaza-18-crown-6) [23978-55-4] $C_{12}H_{26}N_2O_4$, **M 262.3**, **m 118-116°, $pK_{Est} \sim 8.8$** . Twice recrystallise it from *benzene/*n*-heptane, and dry it for 24 hours under high vacuum to remove adhered solvent. It complexes with alkali metal ions in $MeCN$ and $MeOH$ at 25°.

[Weber & Vögtle *Top Curr Chem* (Springer Verlag, Berlin) **98** 1 1981, ISBN: 978-3-540-10793-4 (Print) 978-3-540-38706-0; D'Aprano & Sesta *J Phys Chem* **91** 2415 1987, DOI: 10.1021/j100293a042.]

5,10,15,20-Tetraphenylporphyrin (TPP) [917-23-7] $C_{44}H_{30}N_4$, **M 614.7**, **m 450°** (sublimes >400°), λ_{\max} **482nm**. Purify TPP by chromatography on neutral (Grade I) alumina, and recrystallisation from $CH_2Cl_2/MeOH$ or $*C_6H_6$. It forms complexes with metals. [Yamashita et al. *J Phys Chem* **91** 3055 1987, DOI: 10.1021/j100295a082; *Beilstein* **26** III/IV 1958.]

5,10,15,20-Tetra-4'-pyridinylporphyrin [16834-13-2] $C_{40}H_{26}N_8$, **M 618.7**, has **m >300°(dec)**, λ_{\max} **412nm**. Purify it by chromatography on alumina (neutral, Grade I), with $CHCl_3/MeOH$ (80:20) followed by recrystallisation from $CH_2Cl_2/MeOH$ [Yamashita et al. *J Phys Chem* **91** 3055 1987, DOI: 10.1021/j100295a082;]. It complexes with metals, e.g. Zn. [Kalyanasundaram *Inorg Chem* **23** 2453 1984, DOI: 10.1021/ic00184a019; Okuno et al. *Synthesis* 537 1980, DOI: 10.1055/s-1980-29111.]

Tetrathiafulvalene (TTF, 2,2'-bi-1,3-dithiole) [31366-25-3] $C_6H_4S_4$, **M 204.4**, **m 116-119°, 119.1-119.3°(corr), 122-124°**. Recrystallise TTF from cyclohexane (20%)/hexane (80%), (including Norite or Darco and drying with $MgSO_4$) which provides large (long) orange-yellow needles under an argon or N_2 atmosphere. TTF sublimes at 105°/0.1Torr. The 1H NMR ($CDCl_3$) has δ at 6.32ppm, and the IR (Nujol) has ν_{\max} at 1530, 1250, 1090, 795, 780 and 730 cm^{-1} . [Melby et al. *J Org Chem* **39** 2456 1974, DOI: 10.1021/jo00930a043]. TTF is produced by Zn/ $AcOH$ reduction of 1,3-dithiole-2-thione, or better by reduction of **1,3-dithiole-2-thione methiodide** (see above) with $NaBH_4$ in $MeOH$. On treatment with H_2O_2 and HBF_4 it forms $(TTF)_3(BF_4)_2$. Similar **tetrathiafulvalenium radical cation salts** are readily prepared in high yields, e.g. $(TTF)_{14}(NCSe)_8$, $(TTF)_{11}I_8$, $(TTF)_2[Pt(CN)_4]$, $(TTF)_2[Cu(mnt)_2]$, $(TTF)_2[Ni(mnt)_2]$, $(TTF)_2[Pt(mnt)_2]$, and $(TTF)[Pt(mnt)_2]$, {where mnt is 2,3-dimercapto-2-butanenitrilo(2 $^-$) (maleonitriledithiolato), i.e. $[N=C-C(S^-)=C(S^-)=CN]$ }, which have high compressed pellet resistances $>10^6$ (Ohms, Ω) [Wudl & Kaplan *Inorg Synth* **19** 27 1979; DOI: 10.1002/9780470132500.ch7]. It has electrical properties superior to other comparable electron donors which with organic acceptors, e.g. **TCQN** (tetracyanoquinodimethane [1518-16-7]), under oxidising conditions it forms charge transfer complexes with **metallic** properties. It is an organic electron donor which intercalates with $FeOCl$ producing solid-state low-dimensional conductive materials [Kauzlarich et al. *J Am Chem Soc* **109** 4561 1987, DOI:10.1021/ja00249a019]. With chloranil it forms a coloured charge transfer mixed stack organic semiconductor. [Girlando et al. *J Chem Phys* **79** 1075 1983, DOI: 10.1063/1.445833.] [*Beilstein* **19/11** V 380.]

1,2,3,4-(1H)Tetrazole [288-94-8] CH_2N_4 , **M 70.1**, **m 156°, 157.5-158°, pK²⁵ 4.89 (acidic)**. Crystallise the tetrazole from $EtOH$ and sublime it under high vacuum at *ca* 120° (*care should be taken due to possible EXPLOSION*). It is used in the phosphite trimer method of oligonucleotide synthesis [Barone et al. *Nucleic Acids Res* **12** 4051 1984, PMID:PMC318815]. [*Beilstein* **26** H 346, **26** I 108, **26** II 196, **26** III/IV 1652.]

2-Thenoyltrifluoroacetone [1-(2-thienyl)-4,4,4-trifluorobutan-1,3-dione] [326-91-0] $C_8H_5F_3O_2S$, **M 222.2**, **m 40-44°, 42-44°, b 96-98°/9mm, pK²⁵ 6.4**. Crystallise the dione from hexane or $*benzene$. (An aqueous solution slowly decomposes it). It has ν_{\max} at 1638($C=O$), 1657($C=C$) cm^{-1} . The **oxime** crystallises from H_2O or aqueous $EtOH$. It is used for the determination of Actinides and Lanthanides. [Chaston et al. *Aust J Chem* **18** 673 1965, DOI: 10.1071/CH9650673; Jeffrey et al. In *Vogel's Textbook of Quantitative Chemical Analysis* 5thedn J Wiley & Sons, p170 1989, *Beilstein* **17** III/IV 5989, **17/11** V 128.]

2-Thenylamine (2-thiophenemethylamine) [27757-85-3] C_5H_7NS , **M 113.2**, **b 78.5°/15mm, 95-99°/28mm, d₄²⁰ 1.137, n_D²⁰ 1.5643, pK³⁰ 8.92**. Distil the amine under reduced pressure (nitrogen), from BaO , through a column packed with glass helices. The **hydrochloride** has **m 193-194°** (from $EtOH/Me_2CO$), and the **picrate** has **m 181-18** [*Beilstein* **18** III/IV 7096.]

Thianthrene [92-85-3] $M C_{12}H_8S_2$, **216.3**, **m 151-155°, 158°, b 364-366°/atm**. Crystallise thianthrene from Me_2CO (charcoal), $AcOH$ or $EtOH$. It sublimes in a vacuum. [*Beilstein* **19** H 45, **19** I 619, **19** II 34, **19** III/IV 347, **19/2** V 49.]

5-Thiazolecarboxaldehyde [1003-32-3] C_4H_3NOS , **M 113.1**, **b 92-94°/16mm, d₄²⁰ 1.304, n_D²⁰ 1.5874, pK_{Est} ~0.6**. Dry the aldehyde over Na_2SO_4 and fractionate it in a vacuum. The **2,4-dinitrophenylhydrazone** forms

red crystals from MeOH with **m 238-240°**, and the *semicarbazone* has **m 210-212°** (from MeOH). [Erne et al. *Helv Chim Acta* **34** 143 1951, DOI: 10.1002/hlca.19510340117; *Beilstein* **27** III/IV 2615.]

Thiazoline-2-thiol [96-53-7] $C_3H_5NS_2$, **M 119.2**, **m 106-107°, 106-108°, pK_{Est} ~13.0**. Purify the thiol by dissolution in aqueous alkali, precipitation by addition of HCl and then recrystallisation from H₂O (as needles). [IR: Flett *J Chem Soc* 345(347) 1953, DOI: 10.1039/JR9530000345; Mecke et al. *Chem Ber* **90** 975 1957, DOI: 10.1002/cber.19570900618; Gabriel & Stelzner *Chem Ber* **28** 2929 1895, DOI: 10.1002/cber.189502803110; *Beilstein* **27** III/IV 2540.]

4-(2-Thiazolylazo)-resorcinol [2246-46-0] $C_9H_7N_3O_2S$, **M 221.2**, **m 200-202°(dec), 211°(dec), 218-219°, λ_{max} 500 nm, pK₁²⁵ 1.25, pK₂²⁵ 6.53, pK₃²⁵ 10.76**. Dissolve it in aqueous alkali, extract it with diethyl ether, and re-precipitate it with dilute HCl. The purity is checked by TLC on silica gel using petroleum ether/diethyl ether/EtOH (10:10:1) as the mobile phase. It complexes with Cu²⁺ (pH 3-4), Co²⁺ and Ni²⁺ (pH 7) and Zn²⁺, and Cd²⁺ (pH 8.4). [*Beilstein* **27** III/IV 5988.]

Thiazolyl blue tetrazolium bromide (MTT, 3-[4,5-dimethyl-2-thiazolyl]-2,5-diphenyl-2H-tetrazolium bromide) [298-93-1, 2348-71-2] $C_{18}H_{16}BrN_5S$, **M 414.3**, **m 171°, 195°(dec)**. It is recrystallised by dissolving in MeOH containing a few drops of HBr and then adding dry Et₂O to complete the crystallisation, wash the needles with Et₂O and dry them in a vacuum desiccator over KOH. [Beyer & Pyl *Chem Ber* **87** 1505 1954, DOI: 10.1002/cber.19540871020; *Beilstein* **27** III/IV 6045.]

Thietane (trimethylene sulfide) [287-27-4] C_3H_6S , **M 74.1**, **m -64°, -73.2°, b 93.8-94.2°/752mm, 94-94.5°/760mm, 95°/atm, d₄²⁰ 1.0200, n_D²⁰ 1.5020**. Purify thietane by preparative gas chromatography on a dinonyl phthalate column. It has also been purified by drying over anhydrous K₂CO₃, and distilling through a 25cm glass helices-packed column (for 14g of thietane), then drying over CaSO₄ before sealing it in a vacuum. [Haines et al. *J Phys Chem* **58** 270 1954, DOI: 10.1021/j150513a022.] It is characterised as the *dimethylsulfonium iodide m 97-98°* [Bennett & Hock *J Chem Soc* 2496 1927, DOI: 10.1039/JR9270002496]. The *S-oxide* has **b 102°/25mm, n_D²¹ 1.5075** [Tamres & Searles *J Am Chem Soc* **81** 2100 1959, DOI: 10.1021/ja01518a019]. [*Beilstein* **17** I 3, **17** II 12, **17** III/IV 14, **17**/I V 14.]

2-Thiobarbituric acid (TBA) [504-17-6] $C_4H_4N_2O_2S$, **M 144.2**, **m 235°(dec), 245°(dec), pK₁²⁵ 2.25, pK₂²⁵ 10.72 (2% aqueous EtOH)**. Crystallise it from water. It is used for the quantitative determination of lipopolysaccharides, carrageenan, sialic acid, and detecting lipid oxidation and lipid hydroperoxides. [*Beilstein* **24** H 476, **24** I 414, **24** II 275, **24** III/IV 1884.]

1,1'-Thiocarbonyldiimidazole [6160-65-2] $C_7H_6N_4S$, **M 178.1**, **m 98-102°, 100-102°, 105-106°**. It forms yellow crystals on recrystallisation from tetrahydrofuran or by sublimation at 10⁻³torr (bath temperature 70-80°). It is hydrolysed by H₂O and should be stored dry. Useful for thiating amides [Harpp & Mac Donald *Tetrahedron Lett* **24** 4927 1983, DOI:10.1016/S0040-4039(01)99813-2], in a modified Corey-Winter olefin synthesis [Vedejs et al. *Tetrahedron Lett* **14** 3793 1973, DOI:10.1016/S0040-4039(01)87038-6], and preparing thiocarbamates from alcohols for radical reactions with Bu₃SnH-AlBN [Hanessian et al. *Can J Chem* **65** 1859 1987, DOI: 10.1139/v87-312; RajanBabu et al. *J Am Chem Soc* **111** 1759 1989, DOI: 10.1021/ja00187a031]. [Staab & Walther *Justus Liebigs Ann Chem* **657** 98 1962, DOI: 10.1002/jlac.19626570113; Pullukat & Urry *Tetrahedron Lett* **8** 1953 1967, DOI:10.1016/S0040-4039(00)90762-7.]

Thiochrome {2,7-dimethyl-5H-thiachromine-8-ethanol; 3,8-dimethyl-2-hydroxyethyl-5H-thiazolo [2,3:1',2'] pyrimido[4',5'-d]pyrimidine} [92-35-3] $C_{12}H_{14}N_4OS$, **M 262.3**, **m 227-228°, pK₁²⁰ 8.11, pK₂²⁰ 12.6**. Crystallise thiochrome (found in Yeast) from chloroform. The *monohydrochloride* has **m 235-236°(dec)** (from EtOH) and the *dihydrochloride* has **m 237°(dec)**. [*Beilstein* **27** III/IV 9599.] Fluorescent compound used for detecting and estimating thiamine.

2-Thiocytosine (4-amino-2-mercaptopyrimidine) [333-49-3] $C_4H_4N_2O_2S$, **M 127.2**, **m 236-237°(dec), 285-290°(dec), pK₁²⁰ 3.90 (NH₂), pK₂²⁰ 11.10 (SH)**. It is recrystallised from hot H₂O and dried at 100° to constant weight. [Brown *J Appl Chem (London)* **9** 203 1959, DOI: 10.1002/jctb.5010090402; Russell et al. *J Am Chem*

Soc **71** 2279 1949, DOI: 10.1021/ja01175a001.] It is used in transcription and translation studies [Rachwitz & Scheit *Eur J Biochem* **72** 191 1977, DOI:10.1111/j.1432-1033.1977.tb11239.x].

Thioflavine T [2-(4-dimethylaminophenyl)-3,6-dimethylbenzothiazolium chloride] [2390-54-7] $C_{17}H_{19}ClN_2S$, **M 318.9**, **pK²⁵ 2.7**. Crystallise the chloride from *benzene/EtOH (1:1). Used for staining and quantifying amyloid protein aggregates [LeVine *Methods in Enzymology* **309** 274 1999, DOI:10.1016/S0076-6879(99)09020-5; Alavez et al *Nature* **472** 226 2011, DOI:10.1038/nature09873]. [Beilstein **27** III/IV 5052.]

1-Thioflavone (2-phenylthiochromen-4-one) [784-62-3] $C_{15}H_{10}OS$, **M 238.3**, **m 129-130°**. This yellow solid is purified by passage through a silica gel column, eluting with *C₆H₆/Me₆CO, evaporating and crystallising the residue from EtOH. The *sulfoxide* [65373-82-2] has **m 133-135°**, and the *sulfone* [22810-82-2] has **m 136.5-137°** (from EtOH). The *dimethylhydrazone* has **m 111-113°** (from BuOH). It forms easily hydrolysable salts. [Nakazumi et al. *J Heterocycl Chem* **21** 193 1984, DOI: 10.1002/jhet.5570210138; Chen et al. *J Org Chem* **51** 3282 1986, DOI: 10.1021/jo00367a007; Van Allan & Reynolds *J Heterocycl Chem* **8** 803 1971, DOI: 10.1002/jhet.5570080523; Beilstein **17** I 204, **17** III/IV 5420, **17/10** V 560.]

6-Thioguanine [154-42-7] $C_5H_5N_5S$, **M 167.2**, **m >300°**, **pK₁²³ 8.2 (SH)**, **pK₂²³ 11.6 (acidic, 9-NH)**. It crystallises from H₂O as needles. It has UV with λ_{max} at 258 and 347nm (H₂O, pH 1) and 242, 270 and 322nm (H₂O, pH 11). [Elion & Hitchings *J Am Chem Soc* **77** 1676 1955, DOI: 10.1021/ja01611a082; Fox et al. *J Am Chem Soc* **80** 1669 1958, DOI: 10.1021/ja01540a041.] It is an antineoplastic agent [Kataoka et al. *Cancer Res* **44** 519 1984, <http://cancerres.aacrjournals.org/content/44/2/519>]. [Beilstein **26** III/IV 3926.]

Thioindigo [522-75-8] $C_{16}H_8O_2S_2$, **M 296.2**, **m >280°, 280°, 359°**. Adsorb it on silica gel from CCl₄/*benzene (3:1), elute with *benzene, evaporate, crystallise the residue from CHCl₃ and dry it at 60-65° [Wyman & Brode *J Am Chem Soc* **73** 1487 1951, DOI: 10.1021/ja01148a023; this paper also gives details of purification of other thioindigo dyes]. [Beilstein **19** H 137, **19** I 690, **19** II 192, **19** III/IV 2091.]

Thiomorpholine (tetrahydro-2H-1,4-thiazine) [123-90-0] C_4H_9NS , **M 103.2**, **b 110°/100mm, 169°/atm, d₄²⁰ 1.026, n_D²⁰ 1.540, pK²⁵ 9.00**. Purify it by vacuum distillation. The *hydrochloride* [5967-90-8] has **m 179°** (from isoPrOH or EtOH/Et₂O/HCl), and the *picrolonate*, $C_4H_9NS \cdot C_{10}H_8N_4O_5$, **m 242°(dec)** forms orange prisms from EtOH. [Davies *J Chem Soc* 297 1920, DOI: 10.1039/CT9201700297; Beilstein **2** III/IV 636.]

Thionine (3,7-diaminophenothiazine, Lauth's violet) [135-59-1, 581-64-6 (HCl), 78338-22-4 (acetate)] $C_{12}H_9N_3S_2$, **M 263.7**, ϵ_{590} **6.2 x 10⁴ M⁻¹ cm⁻¹**, **pK¹⁵ 6.9**. The standard biological stain is usually highly pure. It can be crystallised from water or 50% EtOH, then chromatographed on alumina using CHCl₃ as eluent [Shepp et al. *J Phys Chem* **66** 2563 1962, DOI: 10.1021/j100818a055]. Dry it overnight at 100° and store it in a vacuum. The *hydrochloride* can be recrystallised from 50% EtOH or dilute HCl and aqueous *n*-butanol. Purify it also by column chromatography and washed with CHCl₃ and acetone. Dry it *in vacuo* at room temperature. Thionine derivatives are useful drugs for the treatment of Alzheimer and related diseases. [Synth and spectra: Pereteanu & Muller *Org Biomol Chem* **11** 5127 2013, DOI: 10.1039/C3OB40815A; Beilstein **27** H 391, **27** I 412, **27** II 447, **27** III/IV 5149.]

Thiooxine hydrochloride (8-mercaptoquinoline hydrochloride) [34006-16-1] $C_9H_7NS \cdot HCl$, **M 197.7**, **m 165° (dec), 170-175° (dec)**, **pK₁²⁵ 2.16, pK₂²⁵ 8.38**. It forms yellow crystals from EtOH. It has pK_a²⁰ values of 2.05 and 8.29 in H₂O. The salt is more stable than thiooxine. [UV: Albert & Barlin *J Chem Soc* 2384 1959, DOI: 10.1039/JR9590002384.] [Beilstein **21** H 99, **21** III/IV 1197, **21/3** V 30.]

Thiophene [110-02-1] C_4H_4S , **M 84.1**, **m -38.5°, b 84.2°/760mm, 20°/60mm, 46.5°/200mm, d₄²⁰ 1.525, n_D²⁰ 1.52890, n_D³⁰ 1.5223**. The simplest purification procedure is to dry thiophene with solid KOH, or reflux it with sodium, and fractionally distil it through a glass-helices-packed column. More extensive treatments include an initial wash with aqueous HCl, then water, drying with CaSO₄ or KOH, and passage through columns of activated silica gel or alumina. Fawcett and Rasmussen [*J Am Chem Soc* **67** 1705 1945, DOI: 10.1021/ja01226a026] washed thiophene successively with 7M HCl, 4M NaOH, and distilled water, dried with CaCl₂ and fractionally distilled it. *Benzene was removed by fractional crystallisation by partial freezing, and the thiophene was degassed and sealed in Pyrex flasks. [Also a method is described for recovering the thio-

phene from the *benzene-enriched portion.] [Phillips *Org Synth Coll Vol 2* 578 1943, DOI: 10.15227/orgsyn.012.0072; *Beilstein* 17 H 29, 17 I 17, 17 II 35, 17 III/IV 234, 17/I V 297.]

Thiophene-2-acetic acid [1918-77-0] $C_6H_6O_2S$, M 142.2, m 63-64°, 76°, b 160°/22mm, pK^{25} 3.89, pK^{25} 6.43 [MeO(CH₂)₂OH-H₂O/80:20]. Crystallise the acid from ligroin, hexane and/or distil it in a vacuum. The *acid chloride* [39098-97-0] C_6H_5ClOS , M 160.6, d^{25} 1.303/ml, n_D^{20} 1.551, has b 105-106°/22mm, 130-135°/90mm. The *amide* has m 148° (from H₂O or petroleum ether). [*Beilstein* 18 III/IV 4062, 18/6 V 207.] **Thiophene-3-acetic acid** [6964-21-2] has m 73-76°, 79-80°, $pK_{Est} \sim 3.1$. Crystallise the acid from ligroin or H₂O. [*Beilstein* 18 III/IV 4066.]

2-Thiophenecarboxaldehyde (2-thenaldehyde) [98-03-3] C_5H_4OS , M 112.2, b 75-77°/11mm, 106°/30mm, 198°/756mm, d^{20}_4 1.593, n_D^{20} 1.222. Wash it with 50% HCl and distil it under reduced pressure just before use. It has UV with λ_{max} 234nm (hexane). The *Z-oxime* has m 144°, 136-138° and 142° (H₂O). [*Beilstein* 17 H 285, 17 I 148, 17 II 313, 17 III/IV 4477, 17/9 V 349.]

3-Thiophenecarboxaldehyde (3-thenaldehyde) [498-62-4] has m -30°, b 80-81°/14mm, 194-196°/760mm, d^{25} 1.28, n_D^{20} 1.52660. Prepared by the reaction of 2-bromomethylthiophen and hexamine to give the respective hexaminium bromide (m 150°, dec), when heated with H₂O the aldehyde distils with steam. The distillate is acidified (HCl), extracted with Et₂O, dried (Drierite), filtered, evaporated and the residual oil is redistilled (preferably in a vacuum) to provide pure aldehyde [Campaigne et al. *Org Synth Coll Vol 4* 918 1963, DOI: 10.15227/orgsyn.033.0093].

Thiophene-2-carboxylic acid [527-72-0] $C_5H_4O_2S$, M 128.2, m 125-127°, 128.5°, 129-130°, b 260°/atm, pK^{25} 3.59. Crystallise the acid from water and dry it in a vacuum. The *acid chloride* [5271-67-0] has b 206-208°/atm, and the *amide* [5813-89-8] has m 181°(from H₂O) and pK^{25} 10.54 (50% aqueous dioxane). [*Beilstein* 18 H 289, 18 I 438, 18 II 269, 18 III/IV 4011, 18/6 V 158.]

Thiophene-3-carboxylic acid [88-31-1] has m 136-141°, 137-138°, 138-139°, pK^{25} 6.23(4.11). Obtained by the oxidation of the above 3-aldehyde with AgO, the acid is purified by recrystallisation from water and is dried in a vacuum. [Campaigne & LeSuer *Org Synth Coll Vol 4* 919 1963, DOI: 10.15227/orgsyn.033.0094; *Beilstein* 18 H 292, 18 III/IV 4053, 18/6 V 199.] The *amide* [51640-47-0] has m 179-180° (from H₂O, also 181-184°) [*Beilstein* 18 III/IV 4056].

Thiophene-2,5-dicarboxylic acid (2,5-dicarboxythiophene) [4282-31-9] $C_6H_4O_4S$, M 172.2, m 332-333°(sealed capillary), 358.5-359.5°(corrected, sealed tube), 360°, $pK_{Est(1)} \sim 3.3$, $pK_{Est(2)} \sim 7.4$. It can be precipitated from alkaline solution with acid, filtered and recrystallised from H₂O, dried and sublimed at 0.0001 mm. Its *mono-methyl ester* crystallises from aqueous MeOH (m 187-190°, $pK_{Est(1)} \sim 3.2$) or petroleum ether (m 192°), and sublimes in a vacuum unchanged. With diazomethane/Et₂O it provides the *di-methyl ester* (m 148.5-149.5°; m's 146-147° and 152° were also reported) which crystallises from MeOH or 1:1 aqueous MeOH in flattened needles, and sublimes at 80-95° in high vacuum. This ester has $\lambda_{max}(\log \epsilon)$ at 275(4.27) in EtOH. [Hartough & Kosak *J Am Chem Soc* 69 1012 1947, DOI: 10.1021/ja01197a010; Birkinshaw & Chaplen *Biochem J* 60 255 1955, DOI: 10.1042/bj0600255; Griffing & Salisbury *J Am Chem Soc* 70 3416 1948, DOI: 10.1021/ja01190a056; *Beilstein* 18 H 330, 18 III/IV 4496.]

Thiophene-2,5-dicarbonyl dichloride [2,5-bis(chlorocarbonyl)thiophene] [3857-36-1] $C_6H_2Cl_2O_2S$, M 209.1, has m 43-47°, 45-46°, b 102-103°/2mm, 150-152°/11mm. Purify it by distillation in a vacuum; or if discoloured, then heat it with SOCl₂ or oxalyl chloride, and distil it in a vacuum. It solidifies on cooling, and can be recrystallised from *C₆H₆/heptane. It provides the *di-phenyl ester* (m 136-137°) when treated with phenol at 200° or phenol/pyridine at 100°, followed by adding to cold H₂O, filtering, drying, and recrystallising from EtOH [Griffing & Salisbury *J Am Chem Soc* 70 3416 1948, DOI: 10.1021/ja01190a056]. A useful building block for synthesising chiral bis-oxazoline ligands for Cu- and Ru- catalyzed asymmetric cyclopropanations [Gao et al. *Synth Commun* 35 2665 2005, DOI:10.1080/00397910500213948; Gao et al. *Tetrahedron Lett* 45 5649 2004, DOI: 10.1016/j.tetlet.2004.05.120]. [*Beilstein* 18 H 330, 18 III/IV 4496.]

Thiopyronine (2,7-dimethylaminothiaxanthene chloride hydrochloride) [2412-14-8] $C_{17}H_{19}ClN_2S$, M 318.9, λ_{max} 564nm (ϵ 78,500) H₂O, $pK_{Est} \sim 7$. Purify it as the hydrochloride by recrystallisation from hydrochloric acid forming needles, m 245° (dec), and UV with λ_{max} at 564nm (ϵ 78,500, H₂O). [Fanghanel et

al. *J Phys Chem* **91** 3700 1987, DOI: 10.1021/j100297a048; *Beilstein* **18** H 596, **18** III/IV 7291.] It has photodynamic effects on cell growth, colony formation and RNA repair synthesis in *Saccharomyces* mutants with DNA repair deficiency [Roth et al. *Photochem Photobiol* **59**(S1) 627 1994, DOI: 10.1111/j.1751-1097.1994.tb08229.x].

Thiothienoyltrifluoroacetone [1-(2-thienyl)-4,4,4-trifluorobutan-3-one-1-thione] [4552-64-1] $C_8H_5F_3OS_2$, **M 228.2**, **m 61-62°, 64.5-65°, 73-74°, 74°**. It is easily oxidised and has to be purified before use. This is achieved by recrystallisation from *benzene or by dissolution in petroleum ether, extraction into 1M NaOH solution, acidification of the aqueous phase with 1-6M HCl solution, back extraction into petroleum ether and final evaporation of the solvent. The purity can be checked by TLC. It is stored in ampoules under nitrogen at 0° in the dark. It forms red crystals from petroleum ether (b 55-65°). Its IR has ν_{max} at 815m(C-S, C-H), 1260m(C-S), 1570sh(C=C) and 1612s(C=O)cm⁻¹. It yields a chocolate brown complex of $Ni[C_8H_5F_3OS_2]_2$ which has **m 237°**. [Müller & Rother *Anal Chim Acta* **66** 49 1973, DOI:10.1016/S0003-2670(00)89466-0; Chaston et al. *Aust J Chem* **18** 673 1965, DOI:10.1071/CH9650673.]

2-Thiouracil [141-90-2] $C_4H_4N_2OS$, **M 128.2**, **m 240°(dec), 315°(dec)**, **pK₁²⁵ 7.75, pK₂²⁵ 12.7**. Crystallise 2-thiouracil from water or EtOH. [*Beilstein* **24** H 323, **24** I 315, **24** II 171, **24** III/IV 1237.]

9H-Thioxanthene-9-one (thioxanthone, thionanthone) [492-22-8] $C_{13}H_8OS$, **M 212.3**, **m 200-202°, 209°, 210-213°, 212-214°, b 371-373°/712mm**. It forms yellow needles from $CHCl_3$ or EtOH and sublimes *in vacuo*. It is soluble in CS_2 , hot AcOH, and dissolves in concentrated H_2SO_4 to give a yellow colour with green fluorescence in VIS light. The *sulfone* has **m 187°** (from EtOH), and the *hydrazone* has **m 115°** (yellow leaflets from EtOH/* C_6H_6). The *oxime* has **m 194-196°** (from petroleum ether). [Szmant et al. *J Org Chem* **18** 745 1953, DOI: 10.1021/jo01134a022; Ullmann et al. *Chem Ber* **49** 2487 1916, DOI: 10.1002/cber.191604902110; NMR: Sharpless et al. *Org Magn Res* **6** 115 1974, DOI: 10.1002/mrc.1270060213; *Beilstein* **17** H 357, **17** I 191, **17** III/IV 5302, **17/10** V 437.]

Thymine (5-methylpyrimidin-2,4-dione, 5-methyluracil) [65-71-4] $C_5H_6N_2O_2$, **M 126.1**, **m 326°(dec), 335-337°(dec)**, **pK₁²⁵ 9.90 (9.82) pK₂²⁵ >13.0**. Crystallise thymine from EtOAc, 10% aqueous EtOH or water. It has **m 318-320°** after sublimation at 200°/12mm. Purify it by preparative (2mm thick) TLC plates of silica gel, eluting with ethyl acetate/isopropanol/water (75:16:9, v/v; R_F 0.75). The desired spot is located with a uv lamp, cut the band from the plate, place it in MeOH, shake and filter it through a millipore filter, then evaporate. It is an DNA building block. [Infante et al. *JCS Faraday Trans 1* **68** 1586 1973, DOI: 10.1039/F19736901586; *Beilstein* **24** H 353, **24** I 330, **24** II 183, **24** III/IV 1292.]

Tinuvin P (2-[2H-benzotriazol-2-yl]-p-cresol) [2440-22-4, 50936-05-5] $C_{13}H_{11}N_3O$, **M 225.3**, **m 125.5-129.5°, 128-132°, 131-133°, b 225°/10mm**, **pK_{Est(1)}~1.6** (N protonation), **pK_{Est(2)}~ 8** (phenolic OH). Recrystallise it from *n*-heptane or Me_2CO /pentane. It is a UV absorber of light (UVA) in screens and a stabiliser in plastics. [Woessner et al. *J Phys Chem* **89** 3629 1985, DOI: 10.1021/j100263a013.]

Toluidine Blue O [93-31-9] $C_{15}H_{16}N_3S^+ Cl^-$, **M 305.8**, **CI 52040**, **λ_{max} 626nm**, **pK²⁵ 7.5**. Crystallise this blue cationic dibenzothiazine dye from hot water (18ml/g) by adding one and a half volume of alcohol and chilling on ice. Dry it at 100° in an oven for 8-10 hours. [Merrill & Spencer *J Am Chem Soc* **70** 3683 1948, DOI: 10.1021/ja01191a043; *Beilstein* **27** I 417, **27** II 454, **27** III/IV 5161.] Used for selectively staining of pre-malignant lesions (in dysplasia).

1,5,7-Triazabicyclo[4.4.0]dec-5-ene (TBD, 1,3,4,6,7,8-hexahydro-2H-pyrimido[1,2-a]-pyrimidine) [5807-14-7] $C_7H_{13}N_3$, **M 139.2**, **m 125-130°, pK²⁵ ~ 16**. It crystallises from Et_2O but readily forms white crystals of the carbonate. It is a strong base (see pK, i.e. about 100 times more basic than tetramethylguanidine). The *picrate* has **m 220.5-222°** (from EtOH). It forms the *5-nitro* derivative **m 145-160°** that gives a *5-nitro nitrate salt* **m 100-101°** (from EtOH/ Et_2O) and a *5-nitro picrate* **m 144-145°** (from H_2O) [McKay & Kreling *Can J Chem* **35** 1438 1957, DOI: 10.1139/v57-189; Schwesinger *Chimia* **39** 369 1985, Hilpert et al. *J Chem Soc, Chem Commun* 1401 1983, DOI: 10.1039/C39830001401; Kampfen & Eschenmoser *Helv Chim Acta* **72** 185 1989, DOI: 10.1002/hlca.19890720202]. [*Beilstein* **26** III/IV 60.]

1,2,4(1H)-Triazole [288-88-0] $C_2H_3N_3$, M 69.1, m 119-121°, 121°, b 260°/760mm, pK_I^{25} 2.27 (basic), pK_2^{25} 10.26 (acidic). Crystallise 1,2,4-triazole from EtOH, H₂O, EtOAc (m 120.5-121°), or EtOH/*C₆H₆. The *hydrochloride* has m 170°, and the *picrate* has m 163-164° (from H₂O or CHCl₃). [Barszcz et al. *JCS Dalton Trans* 2025 1986, DOI: 10.1039/DT9860002025]. [Beilstein 26 H 13, 26 II 6, 26 III/IV 35.]

Tricycloquinazoline [195-84-6] $C_{21}H_{12}N_4$, M 230.4, m 322-323°. Crystallise it repeatedly from toluene, xylene or these solvents mixed with *C₆H₆. It can also be crystallised from CHCl₃ or cymene (m 308-310°), followed by sublimation at 210°/0.15-0.3 Torr in subdued light. [Beilstein 26 III/IV 1932.] It has carcinogenic properties [Balwin et al. *International J Cancer* 3 244 1968, DOI: 10.1002/ijc.2910030209].

Trifluoperazine dihydrochloride (10-[3-{4-methyl-1-piperazinyl}propyl]-2-trifluoro-methyl-phenothiazine 2HCl) [440-17-5] $C_{21}H_{24}F_3N_3S$. 2HCl, M 480.4, m 240-243°, 242-243°, pK_1 3.9, pK_2 8.1. Recrystallise the salt from absolute EtOH, filter the crystals, dry them *in vacuo* and store them in tightly stoppered bottles because it is *hygroscopic*. It is soluble in H₂O but insoluble in *C₆H₆, Et₂O and alkaline aqueous solution. It has UV with λ_{max} at 258 and 307.5nm (log ϵ 4.50 and 3.50) in EtOH (neutral species). [Craig et al. *J Org Chem* 22 709 1957, DOI: 10.1021/jo01357a618.] It is a calmodulin inhibitor [Levin & Weiss *J Pharmacol Exptl Ther* 208 454 1979, PMID: 34709] and is a psychotropic agent [Fowler *Arzneim.-Forsch* 27 866 1977]. [Beilstein 27 III/IV 1353.]

Trigonelline (1-methylnicotinic acid zwitterion) [535-83-1] $C_7H_7NO_2$, M 137.1, m 218°(dec), 230-233°, pK_I^{25} 2.10. Crystallise trigonelline (as *monohydrate*) from aqueous EtOH, then dry it at 100°. It also crystallises from H₂O as the monohydrate with m 230-233°(dec). It has been crystallised from EtOH with m 214-215°(dec). The *hydrochloride* [6138-41-6] M 173.6 has m 258-259°(dec) (from EtOH) [Smismann & Hite *J Am Chem Soc* 81 1201 1959, DOI: 10.1021/ja01514a047]. The *picrate* crystallises from EtOH with m 204-206°. [Green & Tong *J Am Chem Soc* 78 4896 1956, DOI: 10.1021/ja01600a023; Kosower & Patton *J Org Chem* 26 1319 1961, DOI: 10.1021/jo01063a623; Beilstein 22 I 504, 22 II 35, 22 III/IV 462, 22/2 V 143.]

4',5,7-Trihydroxyflavone (apigenin) [520-36-5] $C_{15}H_{10}O_5$, M 270.2, m 296-298°, 300-305°, 345-350° (pK's 7–10, for phenolic OH). Crystallise it from aqueous pyridine or aqueous EtOH. It dyes wool yellow when mixed with Cr ions. [Beilstein 18 H 181, 18 I 396, 18 II 178, 18 III/IV 2682, 18/4 V 574.]

2,2,5-Trimethyl-1,3-dioxane-4,6-dione (Methyl Meldrum's Acid, methylmalonic acid cyclic isopropylidene ester) [3709-18-0] $C_7H_{10}O_4$, M 158.2, m 111-114°, 113-114°, 115°, pK^{25} 4.77. This 'acid' is synthesised and purified in the same way as *Meldrum's acid* [2033-24-1], except that malonic acid is replaced by methylmalonic acid, in 58% yield. [Pihlaja & Seilo *Acta Chim Scand* 22 3053 1968, DOI: 10.3891/acta.chem.scand.22-3053; Davidson & Bernhard *J Am Chem Soc* 70 3426 1948, DOI: 10.1021/ja01190a060]. Its ¹H NMR (CDCl₃, TMS) has δ at 1.77 (m, isopropylidene Me, J = 0.6Hz), 1.85 (m, isopropylidene Me, J = 0.6Hz), 1.52 (d, 5-Me, J = 7Hz) and 3.81 (q, 5-H, J = 7.0Hz) ppm [Schuster & Schuster *Tetrahedron* 25 199 1969, DOI: 10.1016/S0040-4020(01)99472-4], and the proportion of *enol-form* is apparently not seriously altered by the polarity of the solvent, i.e. MeOH (55.7%), EtOH (51.2%), CDCl₃ (60.8%) and *C₆H₆ (58.9%) [Kabachnik et al. *Tetrahedron* 1 317 1967, DOI: 10.1016/S0040-4020(01)99472-4]. The un- and acid-catalysed aqueous hydrolysis has been studied in detail with the 5,5-dimethyl-dione (not described here) hydrolysing slightly faster than the 5-methyl-4,6-dione (described here) and the unmethylated acid [Meldrum's acid,] because, unlike the 5,5-dimethyl-4,6-dione, the latter two can enolise [Pihlaja & Seilo *Acta Chim Scand* 22 3053 1968, DOI: 10.3891/acta.chem.scand.22-3053; Beilstein 19 III/IV 1928, 19/5 V 11.]

2,2,6-Trimethyl-4H-1,3-dioxin-4-one (diketene acetone adduct) [5394-63-8] $C_7H_{10}O_3$, M 142.2, m 12-13°, b 40°/0.03mm, 65-67°/2mm, 275°/atm, d_4^{20} 1.0879, n_D^{20} 1.4678. The reactions of this dioxinone are very similar to those of diketene and it acts as a β -keto-ester synthon. Its purity can be easily assessed by ¹H NMR spectroscopy as it has only three characteristic peaks (see below). Purify it by fractional distillation, preferably under a vacuum. It is a pleasant smelling liquid that is quite stable in the **absence** of alkali. It is slightly soluble in H₂O that becomes faintly acidic, it gives a red colour with FeCl₃, and it rapidly reduces alkaline permanganate in alcoholic solution. It is readily prepared by refluxing a mixture of dry acetone (100ml), diketene (100ml, 109g, 1.3moles, see [674-82-8] **toxic**) and *p*-toluenesulfonic acid (0.5g, amount is critical) for 3 hours when the odour of diketene disappears. Excess of acetone is distilled off first followed by the *adduct*

(168g, 1.18moles, 91% based on diketene). This procedure should be carried out in an efficient fume cupboard as diketene is **TOXIC**. [Carroll & Bader *J Am Chem Soc* **75** 5400 1953, DOI: 10.1021/ja01117a076; Naylor (uses ZnCl_2 as catalyst) *J Chem Soc* 244 1945, DOI: 10.1039/JR9450000244; Dehmloew & Shamout (using quaternary ammonium salts as catalysts) *Justus Liebigs Ann Chem* 1753 1982, DOI: 10.1002/jlac.198219820917.] The structure of the **adduct** has been determined with certainty [Bader et al. *J Org Chem* **21** 821 1956, DOI: 10.1021/jo01113a619]. Its FT-IR (film) has ν_{max} at 1738.7, 1640.0, 1392.8, 1272.6, 1205.3, 1031.5, 901.2, 805.1 and 548.3 cm^{-1} ; the ^1H NMR (CDCl_3 , TMS) has δ at 1.69 [s, 6H, gem (CH_3)₂], 2.00 (s, but d at very high resolution with $J \sim 1$ Hz, 3H, allyl CH_3) and 5.21 (s, but q at very high resolution with $J \sim 1$ Hz, 1H, vinyl H); and the ^{13}C NMR (CDCl_3) has δ at 168.65, 161.06, 106.32, 93.81, 25.02 and 19.92. Labeling experiments with $(\text{CD}_3)_2\text{CO}$ showed that isoprenyl acetoacetate may be the key intermediate in the synthesis [Hyatt *J Org Chem* **49** 5102 1984, DOI: 10.1021/jo00200a017]. In addition to reacting as a diketene reagent, it has been functionalised, e.g. to 6-bromomethyl-2,2-dimethyl-4H-1,3-dioxin-4-one and 6-methylene-4-diethylphosphoryl-2,2-dimethyl-1,3-dioxinane, for the synthesis of natural products such as **ikarugamycin** and **tirandamycin** [Boeckman & Thomas *J Org Chem* **47** 2823 1982, DOI: 10.1021/jo00135a041]. [Beilstein **19** IV 1604]

1',3',3'-Trimethyl-6-nitrospiro(2H-benzopyran-2,2'-indoline) [1498-88-0] $\text{C}_{19}\text{H}_{18}\text{N}_2\text{O}_3$, **M 322.4**, **m 178°**, **179.5°**, **180°**. This photochromic dye crystallises from absolute EtOH [Hinnen et al. *Bull Soc Chim Fr* 2066 1968, Ramesh & Labes *J Am Chem Soc* **109** 3228 1987, DOI: 10.1021/ja00245a008; Berman et al. *J Am Chem Soc* **81** 5605 1959, DOI: 10.1021/ja01530a021]. [Beilstein **27** III/IV 1460.]

2,2,4-Trimethyl-6-phenyl-1,2-dihydroquinoline [3562-69-4] $\text{C}_{18}\text{H}_{19}\text{N}$, **M 249.3**, **m 102°**, **b 394°/atm**. It is the principal ingredient in Santoflex (a synthetic polymer stabiliser). Crystallise it three times from absolute EtOH. It is a glucocorticoid receptor agonist. The **1-phenylcarbamoyl** derivative has **m 148-149.6°** (from EtOH). [Hively et al. *Anal Chem* **27** 100 1955, DOI: 10.1021/ac60097a029; Beilstein **20** III/IV 4116.]

2,4,6-Trimethylpyridine (sym-collidine) [108-75-8] $\text{C}_8\text{H}_{11}\text{N}$, **M 121.2**, **m -46°**, **-43°**, **b 10°/2.7mm**, **36-37°/2mm**, **60.7°/13mm**, **65°/31mm**, **170.4°/760mm**, **175-178°/atm**, d_4^{25} **0.9100**, n_D^{20} **1.4939**, **1.4981**, n_D^{25} **1.4959**, pK^{25} **6.69(7.45)**. Commercial samples may be grossly impure. Likely contaminants include 3,5-dimethylpyridine, 2,3,6-trimethylpyridine and water. Brown, Johnson and Podall [*J Am Chem Soc* **76** 5556 1954, DOI: 10.1021/ja01650a085] fractionally distilled 2,4,6-trimethylpyridine under reduced pressure through a 40cm Vigreux column and added to 430ml of the distillate slowly, with cooling to 0°, 45g of BF_3 -diethyl etherate. The mixture was again distilled, and an equal volume of dry *benzene was added to the distillate. Dry HCl was passed into the solution, which was kept cold in an ice-bath, and the hydrochloride was filtered off. It was recrystallised from absolute EtOH (1.5ml/g) to **m 286-287°**[**m 256°**(sealed tube), also **m 293-294°** subliming slowly]. The **free base** was regenerated by treatment with aqueous NaOH, then extracted with *benzene, dried (MgSO_4) and distilled under reduced pressure. Sisler et al. [*J Am Chem Soc* **75** 446 1953, DOI: 10.1021/ja01098a055] precipitated trimethylpyridine as its phosphate salt from a solution of the base in MeOH by adding 85% H_3PO_4 , shaking and cooling. The **free base** was then regenerated as above. Garrett and Smythe [*J Chem Soc* 763 1903, DOI: 10.1039/CT9038300763] purified the trimethylpyridine via the **HgCl_2 complex**. It is more soluble in cold than hot H_2O [the solubility is 20.8% at 6°, 3.5% at 20°, 1.8% at 100°]. **Alternatively**, purify it by dissolving it in CHCl_3 , adding solid K_2CO_3 and Drierite, filtering and fractionally distilling through an 8in helix-packed column. **The alkali metals, Na, Li or Cs should NOT be used for drying pyridine, and pyridine derivatives, as they form coloured pyridine radical anions leading to bipyridyls.** [see Schmulbach et al. *J Am Chem Soc* **90** 6600 1968, DOI: 10.1021/ja01026a006]. The **sulfate** has **m 205°**, and the **picrate** (from hot H_2O) has **m 155-156°**. [Frank & Meikle *J Am Chem Soc* **72** 4184 1950, DOI: 10.1021/ja01165a097; Beilstein **20** H 250, **20** I 87, **20** II 164, **20** III/IV 2810, **20/6** V 93.]

1,3,7-Trimethyluric acid [5415-44-1] $\text{C}_8\text{H}_{10}\text{N}_4\text{O}_3$, **M 210.2**, **m 345°(dec)**, pK^{25} **6.0**. Crystallise it from water and dry it at 100° in a vacuum. It has UV with λ_{max} at 289nm (pH 2.5). [Beilstein **26** III/IV 2623.]

1,3,9-Trimethyluric acid [7464-93-9] has **m 340°(dec)**, **347°**, pK^{20} **9.39**. Crystallise it from water and dry it at 100° in a vacuum. [Beilstein **26** H 530, **26** II 301, **26** III/IV 2623.]

1,7,9-Trimethyluric acid [55441-82-2] $\text{C}_8\text{H}_{10}\text{N}_4\text{O}_3$, **M 210.2**, **m 316-318°**, **345°**, pK_{Est} **~9.0**. Crystallise the uric acid from water or EtOH, and sublime it *in vacuo*. [Beilstein **26** H 530, **26** II 302, **26** III/IV 2623.]

3,7,9-Trimethyluric acid [55441-72-0] $\text{C}_8\text{H}_{10}\text{N}_4\text{O}_3$, **M 210.2**, **m 373-375°(dec)**, **pK²⁰ 9.39**. Crystallise the uric acid from water and dry it at 100° in a vacuum. It has UV with λ_{max} at 294nm (pH 2.5). [Bergmann & Dikstein *J Am Chem Soc* **77** 691 1955, DOI: 10.1021/ja01608a044; *Beilstein* **26** H 530, **26** I 156, **26** II 301, **26** III/IV 2623.]

1,3,5-Trioxane (metaformaldehyde) [110-88-3] $\text{C}_3\text{H}_6\text{O}_3$, **M 90.1**, **m 59-62°, 64°, b 114.5°/759mm**. Crystallise 1,3,4-trioxane from sodium-dried diethyl ether or water, and dry it over CaCl_2 . It can also be purified by zone refining. Used as a source of formaldehyde. [*Beilstein* **19** II 392, **19** III/IV 4710, **19/9** V 103.]

Trioxsalen (2,5,9-trimethyl-7H-furo[3,2-g]benzopyran-7-one) [3902-71-4] $\text{C}_{14}\text{H}_{12}\text{O}_3$, **M 228.2**, **m 229-231°, 233-235°, 234.5-235°**. Purify trioxsalen by recrystallisation from CHCl_3 . If too impure, it is fractionally crystallised from CHCl_3 /petroleum ether (b 30-60°) using Norit, and finally crystallised from CHCl_3 alone to give colourless prisms, **m 234.5-235°**. It is a photosensitiser so it should be stored in the dark. [UV: Kaufmann *J Org Chem* **26** 117 1961, DOI: 10.1021/jo01060a028.] Used as a photochemical crosslinker of DNA and used to study nucleic acid structure and function [Higuchi et al. *Nucleic Acids Symposium Series* (2004) **49** 331, DOI: 10.1093/nass/49.1.331]. [*Beilstein* **19/4** V 472.]

2,3,5-Triphenyltetrazolium chloride (TTC, TTZ) [298-96-4] $\text{C}_{19}\text{H}_{15}\text{ClN}_4$, **M 334.8**, **m 243°(dec)**. Crystallise TTZ from EtOH or CHCl_3 , and dry it at 105°. Reagent used to distinguish between α -ketols and simple aldehydes, and used as a seed germination indicator. [*Beilstein* **26** H 363, **26** II 216, **26** III/IV 1774.]

Tripyridyl triazine [TPTZ, 2,4,6-tri(2-pyridyl)-s-triazine] [3682-35-7] $\text{C}_{18}\text{H}_{12}\text{N}_6$, **M 312.3**, **m 245-248°, 247-249°, 248-250°**. Purify it by repeated crystallisation from aqueous EtOH. It is a reagent for the determination of Fe(II) and total Fe [Collins et al. *Anal Chem* **31** 1862 1959, DOI: 10.1021/ac60155a056]. [*Beilstein* **26** III/IV 4192.]

1,1,1-Tris(acetyloxy)-1,1-dihydro-1,2-benziodoxol-3-(1H)-one (Dess-Martin periodinane, DMP) [87413-09-0] $\text{C}_{13}\text{H}_{13}\text{IO}_8$, **M 424.1**, **m 130-133°, 133-134° (dec)**. DMP is essentially the acetylated form of IBX (see above) that renders the latter more soluble in organic solvents. Unlike IBX, it is not explosive on melting or upon hard impact. However, it is subject to hydrolysis in the presence of moisture that may give rise to explosive impurities. It is important that it should be stored in a dry atmosphere and that care should be taken when using it. The original Dess-Martin preparation [Dess & Martin *J Am Chem Soc* **113** 7277 1991, DOI: 10.1021/ja00019a027] has been improved and is reported here. **1-Hydroxy-1,2-benziodoxol-3(1H)-one** (100g, 360mmol, see IBX [61717-82-6]) is added to Ac_2O (400ml, ~4mol) and $\text{TsOH} \cdot \text{H}_2\text{O}$ (0.5g), and stirred under N_2 at 80° for 2 hours, then cooled in an ice-water bath. The white solid that separates in the cold mixture is filtered off onto a fritted glass funnel, rinsed with anhydrous Et_2O (5 x 50ml), and the crystalline solid (138g, 91%, m 134°) is rapidly transferred to an argon filled amber-glass bottle and stored in a freezer. When exposed to light for several weeks some decomposition occurs so it should be kept away from light. [Ireland & Liu *J Org Chem* **58** 2899 1993, DOI: 10.1021/jo00062a040.] Its IR (CH_2Cl_2) has ν_{max} at 1726.9 (s) and 1707.5 cm^{-1} ; the ^1H NMR (CDCl_3 , TMS) has δ at 2.01 (s, 6H, COCH_3), 2.33 (s, 3H, COCH_3), 7.80 (t, 1H, $J_{\text{HH}} = 7.3$ and 8.5 Hz) and 8.07 (t, 1H, $J_{\text{HH}} = 7.3$ and 8.5 Hz) (C-4 and C-5), 8.29 (d, 1H, $J_{\text{HH}} = 8.5$ Hz) and 8.31 (d, 1H, $J_{\text{HH}} = 8.5$ Hz) (C-3 and C-6); and the ^{13}C NMR (CDCl_3) has δ at 20.29 (2 COCH_3), 20.43 (1 COCH_3), 126.01 (C-2), 126.51, 131.79, 133.81, 135.76, 142.36 (C-1), 166.08 (endocyclic C=O), 173.96 (1 acetate C=O), 175.66 and (2 acetate C=O's). A 0.3M solution of DMP in CH_2Cl_2 is available commercially.

DMP is an extremely useful reagent for the oxidation of primary and secondary alcohols to aldehydes and ketones to completion, without further oxidation to the acids, within ~2 hours at 20°; allylic and benzylic alcohols require ~30 minutes. The reaction is increased rapidly on addition of an alcohol. Furan rings or sulfides and vinyl ethers are unreactive. Geraniol is oxidised to geranial without rearrangement to nerol, and it oxidises *N*-benzylbenzamide to benzaldehyde. The reagent is usually added in an appropriate anhydrous solvent (e.g. CHCl_3 , CH_2Cl_2 , MeCN etc) followed by the substrate. Workup can be by adding Et_2O followed by aqueous NaOH which decomposes the reagent to iodobenzoate, or for base-sensitive substrates NaHCO_3 and sodium thiosulfate ($\text{Na}_2\text{S}_2\text{O}_3$) are added, which also remove iodine. A pyridine buffer can be used to keep near-neutral conditions throughout the entire oxidation followed by a thiosulfate work up procedure. [see Dess & Martin *J Am Chem Soc* **113** 7277 1991, DOI: 10.1021/ja00019a027 and papers cited in this reference] In the peptide field, substances like *N*-Fmoc phenylglycinol are oxidised to *N*-Fmoc phenylglycinal almost quantita-

tively in water-saturated CH_2Cl_2 with this reagent [Myers et al. *Tetrahedron Lett* **41** 1359 2000, DOI:10.1016/S0040-4039(99)02293-5].

1,3,5-Trithiane (thioformaldehyde trimer) [291-21-4] $\text{C}_3\text{H}_6\text{S}_3$, M 138.3, m 216-218°(dec), 215-220°(dec). Crystallise it from AcOH or toluene, after Soxhlet extraction with toluene (30g/300ml) [Beilstein **19** III/IV 4711, **19/9** V 105.]

3-Tropanol (Tropine) [120-29-6] $\text{C}_8\text{H}_{15}\text{NO}$, M 141.2, m 63°, 64-66°, b 229°/760mm, pK^{15} 3.80. Distil 3-tropanol in steam and crystallise it from Et_2O or toluene/petroleum ether. It is soluble in H_2O , EtOH, CHCl_3 , and sublimates at 60°/0.1mm. **Hygroscopic**. A 0.05M solution in H_2O has a pH of 11.5, and its solubility in H_2O is 0.1g/ml. The **hydrochloride** has m 280° (from EtOH/ Et_2O). [Beilstein **21** II 17, **21** III/IV 168, **21/1** V 219.]

Tryptamine [3-(2-aminoethyl)indole] [61-54-1] $\text{C}_{10}\text{H}_{12}\text{N}_2$, M 160.1, m 113-116°, 116°, 118°, pK_1^{25} -6.31 (aqueous H_2SO_4 , diprotonation), $\text{pK}_{\text{Est}(2)} \sim 4.9$, pK_3^{25} 16.60 (acidic indole NH). Crystallise tryptamine from *benzene, Et_2O (m 114°) or petroleum ether (m 118°). It has UV with λ_{max} at 222, 276, 282 and 291nm (EtOH) and 226, 275, 281 and 290nm (HCl). It is vasoactive agent, and a neuromodulator. [Beilstein **22** II 346, **22** III/IV 4319, **22/10** V 45.]

Tryptamine hydrochloride [343-94-2] $\text{C}_{10}\text{H}_{12}\text{N}_2 \cdot \text{HCl}$, M 196.7, has m 248°, 252-253°. Crystallise the salt from EtOH/water, EtOH/ Et_2O or EtOH/ EtOAc . See previous entry for UV. [Beilstein **22** II 347, **22** III/IV 4319, **22/10** V 46.]

Tryptophol [3-(2-hydroxyethyl)indole] [526-55-6] $\text{C}_{10}\text{H}_{11}\text{NO}$, M 161.2, m 56-59°, 59°, b 174°/2mm. Crystallise it from diethyl ether/petroleum ether, * C_6H_6 , * C_6H_6 /petroleum ether. The **picrate** has m 100-101° (from * C_6H_6). [Beilstein **21** I 218, **21** II 49, **21** III/IV 788, **21/3** V 61.]

Umbelliferone (7-hydroxycoumarin) [93-35-6] $\text{C}_9\text{H}_6\text{O}_3$, M 162.2, m 225-228°, 230-233°, $\text{pK}_{\text{Est}} \sim 8.0$. It crystallises from water (m 232-232.2°) or EtOH (m 232°). It sublimates at 160°/0.001mm. Fluorescence: Em_{max} 452nm/ Exc_{max} 325nm in 50% EtOH. [Beilstein **18** H 27, **18** I 306, **18** II 16, **18** III/IV 294, **18/1** V 386.]

Uracil (pyrimidine-2,4(1H)-dione) [66-22-8] $\text{C}_4\text{H}_4\text{N}_2\text{O}_2$, M 122.1, m 335°(dec), pK_1^{25} 9.43, pK_2^{25} 13.3-14.2. Uracil crystallises from water (m 339-341°) and m 338° after sublimation in high vacuum. Its solubility in H_2O at 20° is 1g/300ml. [Beilstein **24** H 312, **24** I 312, **24** II 169, **24** III/IV 1193.]

Uramil (5-aminobarbituric acid) [118-78-5] $\text{C}_4\text{H}_5\text{N}_3\text{O}_3$, M 143.1, m 310-312°, 320°, >400°(dec), $\text{pK}_{\text{Est}(1)} \sim 3.9$, $\text{pK}_{\text{Est}(2)} \sim 8.0$, $\text{pK}_{\text{Est}(3)} \sim 12.5$. It crystallises from water. It has also been purified by dissolving it in aqueous ammonia and precipitating it by dropwise addition of formic acid. The solid is collected and dried in a vacuum at 100°. [Hartman & Sheppard *Org Synth Coll Vol* **2** 617 1943, DOI: 10.15227/orgsyn.012.0084; Beilstein **25** H 492, **25** I 704, **25** III/IV 4228.]

Uric acid (2,6,8-trihydroxypurine) [69-93-2] $\text{C}_5\text{H}_4\text{N}_4\text{O}_3$, M 168.1, m >300° (dec) pK_1 5.75, pK_2 10.3. Crystallise uric acid from hot distilled H_2O (the solubility in H_2O is 1part/39,000parts at 18° and 1part/2,000parts at 100°). It is best purified by dissolving in an alkaline solution and acidifying with dilute HCl and drying it at 100° in a vacuum. [Bergmann & Dikstein *J Am Chem Soc* **77** 691 1955 DOI: 10.1021/ja01608a044; Lister *Purines Part II, Fused Pyrimidines* Brown Ed, Wiley-Interscience pp256-257 1971, ISBN 0-471-38205-1, Beilstein **26** H 513, **26** I 151, **26** II 293, **26** III/IV 2619.]

β -Uridine (uracil-1- β -D-ribofuranoside) [58-96-8] $\text{C}_9\text{H}_{12}\text{N}_2\text{O}_6$, M 244.2, m 163-167°, 165°, $[\alpha]_{\text{D}}^{20}$ +10.0 (c 1.6, H_2O), pK^{25} 9.51 (9.25). Crystallise β -uridine from aqueous 75% MeOH or EtOH (m 165-166°). [Beilstein **24** III/IV 1202.]

Urocanic acid (4-imidazolylacrylic acid) [104-98-3] $\text{C}_6\text{H}_6\text{N}_2\text{O}_2$, M 138.1, m 225°, 226-228°, $\text{pK}_{\text{Est}(1)} \sim 2.5$, $\text{pK}_{\text{Est}(2)} \sim 6$, $\text{pK}_{\text{Est}(3)} \sim 11$. Crystallise the acid from water and dry it at 100°. The **trans-isomer** [3465-72-3] has m 225° (229-230°, 230-231° or 231°(dec, from H_2O) and pK_1 3.5 and pK_2 5.6; and the **picrate** has m 225°(dec,

from H₂O). The *cis-isomer* [7699-35-6] has **m** 175-176° (178-179° or 180-184° dec, from H₂O) and pK₁ 3.0 and pK₂ 6.7, and the *picrate* has **m** 204° (from H₂O). [Beilstein 25 H 124, 25 I 536, 25 II 121, 25 III/IV 786.]

δ-Valerolactam (2-piperidone) [675-20-7] C₅H₉NO, **M** 99.1, **m** 38.5-39.5°, 39-40°, 40°, **b** 81-82°/0.1mm, 136-137°/15mm, 256°/atm, pK²⁵ 0.75 (in AcOH). Purify it by repeated fractional distillation. [Conley *J Org Chem* 23 1330 1958, DOI: 10.1021/jo01103a023; Reppe et al. *Justus Liebigs Ann Chem* 596 158 198 1955, DOI: 10.1002/jlac.19555960109; IR: Huisgen et al. *Chem Ber* 90 1437 1957, DOI: 10.1002/cber.19570900808.] The *hydrochloride* [5174-67-4] has **m** 183-184° (from isoPrOH or EtOH/Et₂O) [Hurd et al. *J Org Chem* 17 865 1952, DOI: 10.1021/jo01140a013], and the *oxime* has **m** 122.5° (from petroleum ether) [Behringer & Meier *Justus Liebigs Ann Chem* 607 67 1957, DOI: 10.1002/jlac.19576070109]. The *N-benzoyl derivative* [4252-56-6] has **m** 115° (from CHCl₃/petroleum ether), the *N-methyl derivative* [931-20-4] is a water soluble hygroscopic liquid with **b** 115°/14mm, and the *N-methyl hydrochloride* [87243-73-0] crystallises from EtOH/Et₂O with **m** 115°. The *picrate* has **m** 92-93°. [Beilstein 21/6 V 396.]

δ-Valerolactone (tetrahydro-2H-pyran-2-one) [542-28-9] C₅H₈O₂, **M** 100.1, **m** -13°, -12°, **b** 88°/4mm, 97°/10mm, 124°/24mm, 145-146°/40mm, 229-229.5°/atm, d₄²⁰ 1.1081, n_D²⁰ 1.4568. Purify the δ-lactone by repeated fractional distillation. Its IR has ν_{max} at 1750 (in CS₂), 1732 (in CHCl₃), 1748 (in CCl₄) and 1733 (in MeOH) cm⁻¹ [Huisgen & Ott *Tetrahedron* 6 253 1959, DOI:10.1016/0040-4020(59)80006-5; Linstead & Rydon *J Chem Soc* 580 1933, DOI: 10.1039/JR9330000580; Jones et al. *Can J Chem* 37 2007 1959, DOI: 10.1139/v59-293]. [Beilstein 17 H 235, 17 II 287, 17 III/IV 4169, 17/9 V 17.]

γ-Valerolactone (± 4,5-dihydro-5-methyl-2(3H)-furanone) [108-29-2] has **m** -37°, -36°, -31°, **b** 82-85°/10mm, 102-103°/28mm, 125.3°/68mm, 136°/100mm, 205.75-206.25°/754mm, d₄²⁰ 1.072, n_D²⁰ 1.4322. Purify the γ-lactone by repeated fractional distillation [Boorman & Linstead *J Chem Soc* 577, 580 1933, DOI: 10.1039/JR9330000577]. Its IR has ν_{max} at 1790 (CS₂), 1775 (CHCl₃) cm⁻¹ [Jones et al. *Can J Chem* 37 2007 1959, DOI: 10.1139/v59-293]. The **BF₃-complex** distils at 110-111°/20mm [Reppe et al. *Justus Liebigs Ann Chem* 596 179 1955, DOI: 10.1002/jlac.19555960109]. It is characterised by conversion to γ-hydroxy-*n*-valeramide on treatment with NH₃, which has **m** 51.5-52° (by slow evaporation of a CHCl₃ solution). [Beilstein 17 H 235, 17 I 131, 17 II 288, 17 III/IV 4176, 17/9 V 24.]

(±)-**Vinclozolin** [3-(3,5-dichlorophenyl)-5-methyl-5-vinyloxazolidine-2,4-dione] [50471-44-8] C₁₂H₉Cl₂NO₃, **M** 286.1, **m** 108°. Crystallise this fungicide (controlling plant diseases) from Me₂CO/H₂O. Its solubility at 20° (w/w%) is 44 (Me₂CO), 32 (CHCl₃), 25 (EtOAc) and 10 (H₂O). It irritates the eyes and skin. [GP 2,207,576 1973, *Chem Abstr* 79 137120 1973.]

N-Vinylcaprolactam [2235-00-9] C₈H₁₃NO, **M** 139.2, **m** 35-38°(polym), **b** 95-95.5°/4mm, 128°/21mm, d₄²⁰ 1.0287, n_D²⁰ 1.5133. Distil it under vacuum and with 0.0015% of 4-*tert*-butylcatechol as stabiliser. [Beilstein 21 III/IV 3207.]

N-Vinylcarbazole [1484-13-5] C₁₄H₁₁N, **M** 193.3, has **m** 60-65°, 66°, **b** 154-155°/3mm. Crystallise *N*-vinylcarbazole repeatedly from MeOH in amber glassware. It sublimes in a vacuum. [Beilstein 20 II 282, 20 III/IV 3830, 20/8 V 19.]

Vinylene carbonate (1,3-dioxol-2-one) [872-36-6] C₃H₂O₃, **M** 86.1, **m** 19-22°, 22°, **b** 76-78°/37mm, 165°/~760mm. Purify it by zone melting, or distillation, and stabilise it with 0.5% of 2,6-di-*tert*-butyl-*p*-cresol. [Beilstein 19 III/IV 1597, 19/4 V 72.]

2-Vinylpyridine monomer [100-69-6] C₇H₇N, **M** 105.1, **b** 79-82°/29mm, d 0.974, n 1.550, pK²⁵ 4.92(4.98). Steam distil it, then dry it with MgSO₄ and distil it in a vacuum. Stabilise it with 0.1w/w% of *p*-*tert*-butylcatechol if not used immediately. [Beilstein 20 H 256, 20 III/IV 2884, 20/6 V 211.]

4-Vinylpyridine monomer [100-43-6] has **b** 40-41°/1.4mm, 54°/5mm, 58-61°/12mm, 68°/18mm, 79°/33mm, d₄²⁰ 0.9836, n_D²⁰ 1.5486, pK²⁵ 5.62. Purify the monomer by fractional distillation under a good vacuum and in a N₂ atmosphere; store it in sealed ampoules under N₂, and keep it in the dark at -20°. Stabilise it with 0.1w/w% of *p*-*tert*-butylcatechol if not used immediately. The *picrate* has **m** 175-176°. [UV: Coleman & Fuoss *J Am Chem Soc* 77 5472 1955, DOI: 10.1021/ja01626a006; Overberger et al. *J Polymer Sci* 27 381 1958, DOI: 10.1002/pol.1958.1202711529; Petro & Smyth *J Am Chem Soc* 79 6142 1957, DOI: 10.1021/ja01580a010.] It

is used for alkylating SH groups in peptides [Anderson & Friedman *Can J Biochem* **49** 1042 1971, DOI: 10.1139/o71-152; Cavins & Friedman *Anal Biochem* **35** 489 1970, DOI:10.1016/0003-2697(70)90211-3]. [Beilstein **20** II 170, **20** III/IV 2887, **20**/6 V 213.]

Viologen (4,4'-dipyridyl dihydrochloride) [27926-72-3] $C_{10}H_{10}Cl_2N_2$, **M 229.1, m 278° (also reported m 302-306°, >300°, with sublimation), pK_1^{20} 3.17, pK_2^{20} 4.82.** Purify viologen by precipitation on adding excess of acetone to a concentrated solution of it in aqueous MeOH. It has also been recrystallised several times from MeOH or *iso*-propanol and dried at 70° under vacuum for 24 hours [Prasad et al. *J Am Chem Soc* **108** 5135 1986, DOI: 10.1021/ja00277a017], and recrystallised three times from MeOH/isopropanol [Stramel & Thomas *JCS Faraday Trans 2* **82** 799 1986, DOI: 10.1039/F29868200799; Michaelis & Hill *J Am Chem Soc* **55** 1481 1933, DOI: 10.1021/ja01331a027; Tilford et al. *J Am Chem Soc* **70** 4001 1948, DOI: 10.1021/ja01192a010]. [Beilstein **23** I 49.]

Visnagin (4-methoxy-7-methyl-5H-furo[3,2-g][1]benzopyran-5-one) [82-57-5] $C_{13}H_{10}O_4$, **M 230.2, m 140-142°, 142-145°.** Crystallise visnagin from water. It is soluble in $CHCl_3$ but slightly soluble in EtOH. [Aneja et al. *Tetrahedron* **3** 230 1958, DOI: 10.1016/0040-4020(58)80018-6; Beilstein **19** III/IV 2640.]

9H-Xanthene (dibenzopyran) [92-83-1] $C_{13}H_{10}O$, **M 182.2, m 100.5°, 101-102.5°, b 310-312°/760mm.** Crystallise dibenzopyran from *benzene, MeOH or EtOH. [Beilstein **17** II 72, **17** III/IV 614, **17**/2 V 252.]

Xanthine (2,6-dihydroxypurine, purine-2,6(1H,3H)dione) [69-89-6] $C_5H_4N_4O_2$, **M 152.1, pK_1 0.8 [protonation of imidazole 7(9)NH], pK_2 7.44 [monoanion 1(3)NH], pK_3 11.12 [dianion 1,3-N²⁻].** The *monohydrate* separates in a microcrystalline form on slow acidification with acetic acid of a solution of xanthine in dilute NaOH. It is also precipitated by addition of concentrated NH_3 to its solution in hot 2N HCl (charcoal). After washing with H_2O and EtOH, it is dehydrated by heating above 125°. Its solubility in H_2O is 1 in 14,000 parts at 16° and 1 in 1,500 parts of boiling H_2O , and separates as plates. It has no **m**, but the *perchlorate* has **m 262-264°** [Lister *Purines Part II, Fused Pyrimidines* Brown Ed, Wiley-Interscience pp 252-253 1971, ISBN 0-471-38205-1]. [Beilstein **26** H 447, **26** I 131, **26** II 260, **26** III/IV 2327.]

9-Xanthone (9-xanthenone) [90-47-1] $C_{13}H_8O_2$, **M 196.2, m 172-174°, 175.6-175.4°, b 349-350°/730mm.** Crystallise xanthone from EtOH (25ml/g) and dry at 100°. It has also been recrystallised from *n*-hexane (3x) and sublimed *in vacuo*. [Saltiel et al. *J Am Chem Soc* **108** 2674 1986, DOI: 10.1021/ja00270a028]. [Beilstein **17**/10 V 430.] **9-Xanthyrol (9-hydroxyxanthenone)** [90-46-0] $C_{13}H_{10}O_2$, **M 198.2, m 122-123°, 123-124°, 124-126°, 127-128°.** Crystallise xanthyrol from EtOH and dry at 40-50°. [Beilstein **17** III 1602, **17**/4 V 502.]

Xanthosine (2H₂O) [9-(β-D-ribose)purin-2,6(1H,3H)-dione] [5968-90-1] $C_{10}H_{12}N_4O_6$, **M 320.3, [α]_D²⁰ -53 (c 8, 0.3M NaOH), pK_1^{25} <2.5, pK_2^{25} 5.67, pK_3^{25} 12.85.** It crystallises from EtOH (anhydrous) or water (as dihydrate). [Howard et al. *J Chem Soc* 232 1949, DOI: 10.1039/JR9490000232; Beilstein **26** III/IV 2428.]

Xanthurenic acid (5,8-dihydroxyquinoline-2-carboxylic acid) [59-00-7] $C_{10}H_7NO_4$, **M 205.2, m 286°, 290-295°(dec), 297-298°(dec), $pK_{Est(1)} \sim 1.5$, $pK_{Est(2)} \sim 4.9$, $pK_{Est(3)} \sim 9.8$.** It is precipitated by the addition of 2N formic acid to its solution in hot 2M ammonia (charcoal). The solid is filtered off, dried in a vacuum at ~80° in the dark. Its UV (H_2O) has λ_{max} nm (ϵ M⁻¹cm⁻¹) at 243 (30,000) and 342 (6,500). The *methyl ester* has **m 262°** (from MeOH). It forms Cu^{2+} , Zn^{2+} , Fe^{2+} and Fe^{3+} salts. [Beilstein **22** III/IV 2513.]

Yohimbine (Rauwolfscine) [146-48-5] $C_{13}H_{10}O_2$, **M 354.5, 235-237°(dec), 278°(dec), [α]_D²⁰ +55.6 (c 2, EtOH), pK_1^{22} 3.0, pK_2^{22} 7.45.** Crystallise this *Rauwolfia* alkaloid ester from EtOH or aqueous EtOH (needles), and dry *in vacuo* to remove EtOH. Soluble in MeOH, $CHCl_3$, warm C_6H_6 , poorly soluble in Et_2O and H_2O . UV (MeOH): λ_{max} nm(log ϵ) at 226(4.56), 280(3.88) and 291(380). The *hydrochloride* [45-19-0] **m 302°(dec)** (EtOH plates) [α]_D²⁰ +105° (H_2O), is sol. in H_2O (83%) and EtOH (0.25%). [Wenkert et al. *J Am Chem Soc* **101** 5370 1979, DOI: 10.1021/ja00512a043; Beilstein **25** III/IV 1234.] It is a 5-HT_{1A} serotonin receptor agonist and α_2 -adrenoceptor antagonist. δ -Yohimbine is *ajmalicine* [483-04-5] and γ -yohimbine is *ajmaline* [4360-12-7].

MISCELLANEOUS As, B, P, Si, S, Se and Te COMPOUNDS.

This section contains miscellaneous organic compounds of As, B, P, Si, S, Se, Te, and *ammonium and metal salts of their acids, and salts of their bases*. See other sections and chapters for further entries of sulfur, phosphorous and silicon compounds; and for sulfur heterocyclic compounds see section on 'Heterocyclic Compounds' in this Chapter.

Acetonyltriphenylphosphonium chloride [1235-21-8] $C_{21}H_{20}ClOP$, M 354.8, m 237-238°, 243-245°, 244-246°(dec), 248-252°. Recrystallise it from $CHCl_3$ /* C_6H_6 /petroleum ether (b 60-80°) or by dissolving it in $CHCl_3$ and pouring it into dry Et_2O . It has UV($EtOH$) with λ_{max} nm(ϵ) at 255(3,600), 262(3,700), 268(4,000) and 275(3,100). The *iodide salt* crystallises from H_2O and has m 207-209°. [Ramirez & Dershowitz *J Org Chem* **22** 41 1957, DOI: 10.1021/jo01352a010.] It is an **IRRITANT** and is *hygroscopic*. When shaken with a 10% aqueous solution of Na_2CO_3 (8 hours) it gives *acetylmethylene triphenyl phosphorane* [1439-36-7] $C_{21}H_{19}OP$, M 318.4, which is recrystallised from $MeOH/H_2O$, and after drying at 70°/0.1mm has m 205-206° (207°). It has UV with λ_{max} nm(ϵ) at 268 (6600), 275 (6500) and 288 (5700), and its IR has ν_{max} at 1529(s), 1470(m), 1425(s), 1374(m), 1105(s) and 978(s) (cm^{-1}). [Ramirez & Dershowitz *J Org Chem* **22** 41, 44 1957, DOI: 10.1021/jo01352a010; *Beilstein* **16** H 761, **16** II 373.]

3R,4R,1'R-4-Acetoxy-3-[1-(tert-butylmethylsilyloxy)ethyl]-2-azetinone [76855-69-1] $C_{13}H_{25}NO_4Si$, M 287.4, m 107-108°, $[\alpha]_D^{20}$ +55 (c 0.5, toluene), $[\alpha]_D^{20}$ +53.7 (c 1.04, $CHCl_3$). Purify it by chromatography on silica gel (3 x 14cm) for 50g of ester using 20% $EtOAc$ in *n*-hexane. The eluate is evaporated, and the residue is recrystallised from hexane (white fluffy crystals). Avoid breathing it in and contact with eyes. [Leanza et al. *Tetrahedron* **39** 2505 1983, DOI: 10.1016/S0040-4020(01)92144-1.]

N-Acetyl-4-hydroxy-m-arsanilic acid (Acetarsol, Stovarsol, 3-acetamido-4-hydroxyphenylarsonic acid) [97-44-9] $C_8H_{10}AsO_5$, M 275.1, m 240-250°, pK_1 3.73, pK_2 7.9, pK_3 9.3. It crystallises from water in colourless prisms. It decomposes slowly on prolonged boiling in H_2O or dilute alkalis. The *N-propionyl derivative* recrystallises from H_2O with m 228-229°(dec). [Raiziss & Fisher *J Am Chem Soc* **48** 1323 1926, DOI: 10.1021/ja01416a030; Hewitt & King *J Chem Soc* 817 1926, DOI: 10.1039/JR9262900817; *Beilstein* **16** I 491, **16** II 521, **16** III 1129.] The drug is an anti-infective used in suppositories [Chen et al. *Int J STD AIDS* **10** 277 1999, DOI: 10.1258/0956462991913943].

Alizarin Red S (3,4-dihydroxy-9,10-dioxo-2-anthracene sulfonic acid, Na salt. H_2O , Alizarin sulfonate sodium salt, Alizarin carmine) [130-22-3] $C_{14}H_{22}NaO_7S$, M 342.3, decomposes on heating, CI 58005, pK_1^{25} <1, pK_2^{25} 5.49, pK_3^{25} 10.85 (11.01). Commercial samples contain large amounts of sodium and potassium chlorides and sulfates. It is purified by passing through a Sephadex G-10 column (size exclusion column), followed by elution with water, then 50% aqueous $EtOH$ [King & Pruden *Analyst (London)* **93** 601 1968, DOI: 10.1039/AN9689300601]. Finally dissolve it in $EtOH$ and precipitate it with Et_2O several times [Sacconi *J Phys Chem* **54** 829 1950, DOI: 10.1021/j150480a012; polarography: Furman & Stone *J Am Chem Soc* **70** 3055 1948, DOI: 10.1021/ja01189a064]. A 1% aqueous solution is used as an indicator (purple at pH 5.2 and yellow at pH 7.7), is a reagent for Al and Ca, and is used for microscopic staining [*Beilstein* **11** IV 683.]

Allyl trimethylsilane (2-propenyltrimethylsilane) [762-72-1] $C_6H_{14}Si$, M 114.3, b 82°/atm, 83.0-84.5°/atm, 84-88°/atm, 85.5-86.0°/atm, d_4^{20} 0.713, n_D^{20} 1.405. Fractionate it through an efficient column at atmospheric pressure. If impure, dissolve it in THF, shake it with H_2O (2x), dry (Na_2SO_4), filter and fractionate it. [Cudlin & Chvalovský *Coll Czech Chem Commun* **27** 1658 1962, DOI: org/10.1135/cccc19621658; *Beilstein* **4** IV 3927.]

2-Amino-4-sulfobenzoic acid [98-43-1] $C_7H_7NO_5S$, M 217.1, m (decomp on heating), d^{25} 1.709g/cm³, n_D^{20} 1.661, $pK_{Est(1)}$ < 0.0, $pK_{Est(2)}$ ~1.0, $pK_{Est(3)}$ ~4.2. It was prepared by dissolving toluene in the smallest possible volume of fuming H_2SO_4 (care) then diluted with cold H_2O , and neutralised with $CaCO_3$ then evaporated to a small volume. A solution of $KMnO_4$ is added, the mixture is diluted, heated until colourless, filtered, and saturated with HCl. $BaCl_2$ solution is added to precipitate the barium salt which is collected, dried and treated with the calculated amount of H_2SO_4 to precipitate $BaSO_4$, which is filtered off. Evaporation of the filtrate gave

2-nitro-4-sulfobenzoic acid [552-23-8] which is soluble in H₂O but crystallises from a small volume of it as short prisms of the *hydrate*, **m 130-131°**. It forms Ba, Cu, Ka, Ca and Mg salts readily. The *potassium 2-nitro-4-sulfobenzoate* [5344-48-9], prepared by neutralising the acid and evaporating to crystallise it, was reduced with excess of ammonium sulfide, evaporating, filtering and acidifying with HCl, whereby the *aminosulfobenzoic acid* separates and is recrystallised from hot H₂O. [Hart *Am Chem J* **2** 42–44 p 44 1880, DOI: 10.1021/ja02132a600; Love & Kormendy *J Org Chem* **27** 2177 1962, DOI: 10.1021/jo01053a068; Dorssen & Holleman *Recl Trav Chim Pays-Bas* **29** 368 1910, DOI: 10.1002/recl.19100291004; Holleman *Recl Trav Chim Pays-Bas* **24** 194 1905, DOI: 10.1002/recl.19050240602; Gless et al. US Pat 556560 15 Oct 1996 to Zeneca].

4-Amino-2-sulfobenzoic acid [527-76-4] has **m** (decomp on heating), **d**²⁵ **1.7g/cm³**, **n**_D²⁰ **1.662**, **pK_{Est(1)} < 0.0**, **pK_{Est(2)} ~1.0**, **pK_{Est(3)} ~4.2**. It was prepared by sulfonating *o*-nitrotoluene to form 2-nitrotoluene-4-sulfonic acid, followed by oxidation with KMnO₄ to 2-nitro-4-sulfobenzoic acid then reduction with ammonium sulfide. Crystallise the aminosulfobenzoic acid from H₂O (solubility is 0.3% at 25°). It forms a *monohydrate*, is insoluble in EtOH and Et₂O, and forms insoluble Ag and Ba salts. [Hart *Am Chem J* **2** 42–44 p 44 1880, DOI: 10.1021/ja02132a600; Love & Kormendy *J Org Chem* **27** 2177 1962, DOI: 10.1021/jo01053a068; Dorssen & Holleman *Recl Trav Chim Pays-Bas* **29** 368 1910, DOI: 10.1002/recl.19100291004; Holleman *Recl Trav Chim Pays-Bas* **24** 194 1905, DOI: 10.1002/recl.19050240602; Beilstein **14** H 877, **14** I 769, **14** IV 2834; Beilstein **19** I 356, **19** III/IV 1641 for 4-NO₂.]

Ammonium dodecylsulfate (ammonium laurylsulfate) [2235-54-3] **C₁₂H₂₉NO₄S**, **M 283.4**. Recrystallise it first from 90% EtOH and then twice from absolute EtOH, and finally dry it in a vacuum. It is an anionic high-foam surfactant detergent available also as a 30% solution in H₂O (d²⁵ 1.01g/l). [Beilstein **1** III 1786.]

Ammonium tetraphenylborate [14637-34-4] **C₂₄H₂₄BN**, **M 337.3**, **m ca 220°(dec)**. Dissolve it in aqueous Me₂CO and allow crystallisation to proceed slowly; otherwise very small crystals are formed. No trace of Me₂CO is left in the crystals after drying at 120° [Davies & Staveley *Trans Faraday Soc* **53** 19 1957, DOI: 10.1039/TF9575300019]. Also, the salt can be precipitated from a dilute AcOH solution of sodium tetraphenylborane in the presence of NH₄⁺ ions. After standing for 5 minutes, the precipitate is filtered off onto a sintered porcelain crucible, washed with very dilute AcOH and dried at room temperature for at least 24 hours [Wendlandt *Anal Chem* **28** 1001 1956, DOI: 10.1021/ac60114a021]. Alternatively, a solution of sodium tetraphenylborane (5% excess) in H₂O is added to NH₄Cl solution. After 5 minutes the precipitate is collected, washed several times with H₂O and recrystallised from aqueous Me₂CO. [Howick & Pflaum *Analyt Chim Acta* **19** 342 1958, DOI: 10.1016/S0003-2670(00)88173-8; Beilstein **16** IV 1625.] A precipitating agent for some inorganic and organometallic compounds (e.g. of Ni, Pd and Pt).

9-Anthraceneboronic acid [100622-34-2] **C₁₄H₁₁BO₂**, **M 222.0**, **m 214-216°, 203-250°**. Crystallise the boronic acid from dilute HCl (**m 180-184°**). The *disodium salt* has **m 209-213°**. [Beilstein **16** IV 1679.]

Anthraquinone Blue B (Acid Blue 45, 1,5-diamino-4,8-dihydroxy-9,10-anthraquinone-3,7-disulfonic acid di-Na salt) [2861-02-1] **C₁₄H₁₀N₂NaO₁₀S₂**, **M 474.3**, **m >300°**, **CI 63010**, **λ_{max} 595nm**, **pK_{Est(1)} ~<0**, **pK_{Est(2)} ~2**, **pK_{Est(3)} ~9**. Purify it by salting out an aqueous solution three times with sodium acetate, followed by repeated extraction with EtOH. Useful dye for Nylon. [McGrew & Schneider *J Am Chem Soc* **72** 2547 1950, DOI: 10.1021/ja01162a059; Beilstein **14** H 706, 725].

Anthraquinone Blue RXO [4403-89-8] **M 445.5**, **CI 1076**. Purify the dye by salting out an aqueous solution three times with sodium acetate, followed by repeated extraction with EtOH. Useful dye for Nylon. [McGrew & Schneider *J Am Chem Soc* **72** 2547 1950, DOI: 10.1021/ja01162a059]. [Beilstein **14** H 706, 725.]

Anthraquinone Green G [Acid Green 25, Alizarin Cyanine Green F, 1,4-bis-(4-methyl-2-sulfophenyl-1-amino)-9,10-anthraquinone di-Na salt] [4403-90-1 *Na salt*, 86923-71-9 *diNa salt*] **C₂₈H₂₂N₂NaO₈S₂**, **M 624.6**, **m 235-238°**, **CI 61570**, **λ_{max} 642nm**, **pK²⁵ >0**. Purify it by salting out three times from an aqueous solution with sodium acetate, followed by repeated extraction with EtOH [McGrew & Schneider *J Am Chem Soc* **72** 2547 1950, DOI: 10.1021/ja01162a059]. It is a green powder that is slightly soluble in Me₂CO, EtOH and pyridine. It is soluble in concentrated H₂SO₄ to give a blue solution that becomes turquoise in colour upon dilution. [UV: Allen et al. *J Org Chem* **7** 63 1942, DOI: 10.1021/jo01195a009; Beilstein **14** H 725.]

9,10-Anthraquinone-2,6-disulfonic acid (disodium salt) [853-68-9] $C_{14}H_6Na_2O_8S_2$, M 412.3, m >325°, $pK_{Est} \sim 0$ (for SO_3H). Crystallise it three times from water, in the dark [Moore et al. *JCS Faraday Trans 2* **82** 745 1986, DOI: 10.1039/F29868200745]. Its solubility in H_2O at ~25° is 1%. [Beilstein **11** IV 673.]

***o*-Arsanilic acid** [2045-00-3] $C_6H_8AsNO_3$, M 217.0, m 145-150°, 153°, 154-155°, pK_1^{22} 3.77 (AsO_3H_2), pK_2^{22} 8.66 (AsO_3H^-). Crystallise it from water or ethanol/ether. **POISONOUS**. [Beilstein **16** I 463.]

***p*-Arsanilic acid** [98-50-0] has m 232°, pK_1^{22} 4.05 (AsO_3H_2), pK_2^{22} 8.66 (AsO_3H^-). Crystallise it from water or ethanol/ether. **POISONOUS**. [Beilstein **16** I 466.]

Arsenazo I [Neothorin, Uranon, 3(2-arsonophenylazo)-4,5-dihydroxy-2,7-naphthalenedisulfonic acid tri Na salt] [520-10-5, 66019-20-3] $C_{16}H_{10}AsN_2Na_3O_{11}S_2$, M 614.3, m >300°, ϵ 2.6×10^4 at 500nm, pH 8.0, pK_1 0.6(0.8), pK_2 3.52, pK_3 2.97(AsO_3H_2), pK_4 8.20(AsO_3H^-), pK_5 9.98(OH), pK_6 15.0. A saturated aqueous solution of the free acid is slowly added to an equal volume of concentrated HCl. The orange precipitate is filtered off, washed with acetonitrile and dried for 1-2 hours at 110° [Fritz & Bradford *Anal Chem* **30** 1021 1958, DOI: 10.1021/ac60138a002]. It is then titrated with NaOH to form the di or tri Na salt as set out below. It has been used to titrate Ca and Mg (pH 10, violet/red-orange); and rare earths (pH 5.5-6.5, violet/red-orange), also Th^{4+} and U^{4+} (pH 1.7-3.0, violet/orange) and Y (pH 5.5-6.5, violet/red-orange) [Fritz et al. *Anal Chem* **30** 1111 1958, DOI: 10.1021/ac60138a032].

Arsenazo III [3,6-bis(2-arsonophenylazo)-4,5-dihydroxy-2,7-naphthalenedisulfonic acid di Na salt] [62337-00-2 Na salt, 1668-00-4 acid] $C_{22}H_{18}As_2N_4O_{14}S_2$, M 776.4 (acid), has m >320°, λ_{max} 350nm, pK_1 -2.7, pK_2 -2.7, pK_3 0.6, pK_4 0.8, pK_5 1.6, pK_6 3.4, pK_7 6.27, pK_8 9.05, pK_9 11.98, pK_{10} 15.1. Contaminants include monoazo derivatives, starting materials for synthesis and by-products. It is partially purified by precipitation of the dye from aqueous alkali on addition of HCl. More thorough purification is achieved by taking a 2g sample in 15-25ml of 5% aqueous NH_3 and filter. Add 10ml HCl (1:1) to the filtrate to precipitate the dye. Repeat the procedure and dissolve the solid dye (0.5g) in 7ml of a 1:1:1 mixture of *n*-propanol/880 NH_3 /water at 50°. After cooling, filter the solution and chromatograph the filtrate through a cellulose column using a 3:1:1 mixture of *n*-propanol/880 NH_3 /water as eluent. Collect the blue band and evaporate it to 10-15ml below 80°, then add 10ml of concentrated HCl to precipitate pure Arsenazo III. Wash it with EtOH and dry it in air [Borak et al. *Talanta* **17** 215 1970, DOI: 10.1016/0039-9140(70)80069-8]. The sodium salt is then obtained by dissolving it in the equivalent amount of dilute NaOH and freeze-drying. The purity of the dye can be checked by paper chromatography using M HCl as eluent. It is used for the spectrophotometric estimation of Th, U, Zr, rare earths, Cd and Zn [Michaylova & Yuroukova *Anal Chim Acta* **68** 73 1974, DOI: 10.1016/S0003-2670(01)85147-3.]

Benzaldehyde-2-sulfonic acid sodium salt [1008-72-6] $C_7H_5NaO_4S$, M 208.2, m decomposes on heating. It forms white prisms or plates by extracting with boiling EtOH, filtering, evaporating to dryness and recrystallising the Na salt from a small volume of H_2O . The ***N*-phenylhydrazone sodium salt** recrystallises from H_2O with m 174.5°. [Gnehm & Schüle *Justus Liebigs Ann Chem* **299** 347 1898, DOI: 10.1002/jlac.18982990212; Beilstein **11** IV 652.]

Benzeneselenenyl bromide (phenylselenenyl bromide) [34837-55-3] C_6H_5BrSe , M 236.0, m 58-62°, 60°, 62°, b 107-108°/15mm, 134°/35mm. Distil it in a vacuum, recrystallise it from petroleum ether, $CHCl_3$ (EtOH free), or Et_2O (cooling mixture) to give dark red or orange crystals. These sublime at 25°/0.001mm and are soluble in hexane, CH_2Cl_2 , Et_2O and THF. [Behaghel et al. *Chem Ber* **65** 812 1932, DOI: 10.1002/cber.19320650525; Pitteloud & Petrzilka *Helv Chim Acta* **62** 1319 1979, DOI: 10.1002/hlca.19790620445]. [Beilstein **6** III 1111.] **HIGHLY TOXIC**.

Benzeneselenenyl chloride (benzeneselenenyl chloride, phenylselenenyl chloride) [5707-04-0] C_6H_5ClSe , M 191.5, m 59-60°, 64-65°, b 92°/5mm, 120°/20mm. Purify it by distillation in a vacuum, and recrystallisation (orange needles) from hexane [Foster *J Am Chem Soc* **55** 822 1933, DOI: 10.1021/ja01329a063; Foster et al. *Recl Trav Chim Pays-Bas* **53** 405, 408 1934, DOI: 10.1002/recl.19340530504; Behaghel & Seibert *Chem Ber* **66** 708 1933, DOI: 10.1002/cber.19330660520]. [Beilstein **6** III 1110.] **HIGHLY TOXIC**.

Benzeneseleninic acid [6996-92-5] $\text{C}_6\text{H}_6\text{O}_2\text{Se}$, M 189.1, m 121-124°, 122-124°, b 114.4°/760mm, pK^{25} 4.70. Add 10% excess of 15M NH_3 to the solid acid and stir until the solid dissolves, filter, decolorise with charcoal (2x, Norite) and acidify by slow addition of 6M HCl, filter the solid off and wash it with H_2O . Dissolve the acid in the minimum volume of MeOH, and this solution is added dropwise to boiling H_2O until cloudiness appears. At this point add 25% more boiling H_2O , filter hot (decolorise if necessary) and cool rapidly, with scratching, to 0°. After 30 minutes the solid is filtered off and recrystallised as before but with very slow cooling. The colourless needles are filtered off and dried in a vacuum desiccator (CaCl_2) before the melting point is measured [McCullough & Gould *J Am Chem Soc* **71** 674 1949, DOI: 10.1021/ja01170a083]. The ***HNO*₃ complex** has m 112°. [Beilstein **11** H 422, **11** I 110, **11** III 716.] **TOXIC** solid.

Benzeneseleninic anhydride [17697-12-0] $\text{C}_{12}\text{H}_{10}\text{O}_3\text{Se}_2$, M 360.1, has m 124-126°, 164-165°, 165-170°, 170-173°. When the anhydride is recrystallised from $^*\text{C}_6\text{H}_6$ it has m 124-126°, but when this is heated at 140°/1 hour in a vacuum or at 90°/2 hours it has m 164-165° and gives a solid m 124-126° when then recrystallised from $^*\text{C}_6\text{H}_6$. Both depress the melting point of the acid PhSeO_2H . If the low melting anhydride is dissolved in $^*\text{C}_6\text{H}_6$ and seeded with the high melting anhydride, the high melting anhydride crystallises out. It readily absorbs H_2O to form the acid (PhSeO_2H , m 122-124°). Because of this, the commercial anhydride could contain up to 30% of the acid. It is best purified by converting to the ***HNO*₃ complex** (m 112°) and heating this *in vacuo* at 120°/72 hours to give the anhydride as a white powder m 164-165°. Alternatively, heat the anhydride *in vacuo* at 120°/72 hours until the IR shows no OH band. [Ayrey et al. *J Chem Soc* 2089 1962, DOI: 10.1039/JR9620002089; Barton et al. *JCS Perkin Trans 1* 567 1977, DOI: 10.1039/P19770000567; Beilstein **11** H 422, **11** I 110, **11** III 716, **11** IV 708.] **TOXIC** solid.

Benzeneselenol (phenylselenol, selenophenol) [645-96-5] $\text{C}_6\text{H}_6\text{Se}$, M 157.1, b 57-59°/8mm, 71-72°/18mm, 84-86°/25mm, d_4^{20} 1.480, n_D^{20} 1.616. Dissolve it in aqueous N NaOH, acidify this with concentrated HCl and extract with Et_2O , dry over CaCl_2 , filter, evaporate on a steam bath and distil the residue from a Claisen flask or through a short column collecting the middle fraction, and seal immediately in a glass vial, otherwise the colourless liquid becomes yellow. The alkali insoluble materials consist of ***diphenylselenide*** (b 167°/16mm) and ***diphenyldiselenide***, m 63° (from EtOH). **TOXIC**, use rubber gloves. It has a foul odour. [Foster *Org Synth Coll Vol* **3** 771 1955, DOI: 10.152227/orgsyn.024.0089; Beilstein **6** III 1104, 1110, **6** IV 1777.] The anion is a strong nucleophile [Sonoda in *Encyclopedia of Reagents for Organic Synthesis* (ed. L. Paquette) J. Wiley & Sons, NY 2005, DOI:10.1002/047084289X.rb018].

Benzeneselenonic acid (benzeneselenoic acid) [39254-48-3] $\text{C}_6\text{H}_6\text{O}_3\text{Se}$, M 205.1, m 64°, pK^{25} 4.79. It has been prepared by reaction of $^*\text{C}_6\text{H}_6$ with SeO_3 in liquid SO_2 , and isolated in good yield *via* its barium salt. Purify it by dissolving in H_2O and passing through a strong cation exchange resin (H^+ form). Evaporate the effluent under reduced pressure and dry the residue in a high vacuum to give colourless hygroscopic crystals [Dostál et al. *Z Chem* **6** 153 1966, DOI: 10.1002/zfch.19660060413; and IR: Dostál et al. *Chem Ber* **104** 2044 1971, DOI: 10.1002/cber.19711040704]. [Beilstein **11** H 422, **11** I 111.]

Benzenestibonic acid [535-46-6] $\text{C}_6\text{H}_7\text{O}_3\text{Sb}$, M 248.9, m >285°(dec). It crystallises from acetic acid (minute white crystals), or from EtOH/ CHCl_3 mixture on addition of water (large diamond-shaped glistening crystals). It is insoluble in H_2O , slightly soluble in EtOH, soluble in Me_2CO , warm CHCl_3 and amyl acetate. It has also been purified by dissolving it in aqueous Na_2CO_3 , filtering, then acidifying with dilute H_2SO_4 which precipitates the stibonic acid as a fine white powder. When moist it has a characteristic, not unpleasant, odour. [Schmidt *Justus Liebigs Ann Chem* **421** 174 1920, DOI: 10.1002/jlac.19204210204; May *J Chem Soc* **101** 1033 1912, DOI: 10.1039/CT9120101033.]

m-Benzenedisulfonic acid [98-48-6] $\text{C}_6\text{H}_6\text{O}_6\text{S}_2$, M 238.2, pK_{Est} <0. Free it from H_2SO_4 by conversion to the calcium or barium salts (using $\text{Ca}(\text{OH})_2$ or $\text{Ba}(\text{OH})_2$, and filtering). The calcium salt is then converted to the potassium salt, using K_2CO_3 . Both the potassium and the barium salts are recrystallised from H_2O , and the acid is regenerated by passing through the H^+ form of a strong cation exchange resin. The acid is recrystallised twice from conductivity water and dried over CaCl_2 at 25°. [Atkinson et al. *J Am Chem Soc* **83** 1570 1961, DOI: 10.1021/ja01468a008.] It has also been crystallised from Et_2O and dried in a vacuum oven. The ***S-benzylisothiuronium salt*** has m 214.3° (from EtOH/ H_2O). [Beilstein **11** IV 553.] It is best kept as the ***di-***

disodium salt, [831-59-4] **M 180.2**, $\text{C}_6\text{H}_4\text{O}_6\text{S}_2\text{Na}_2$, which decomposes on heating [Beilstein 11 H 199, 11 I 48, 11 II 213, 11 III 453.]

m-Benzenedisulfonyl chloride [585-47-7] **C₆H₄Cl₂O₂S₂, M 275.1**, has **m 59-60°, 63°**. Crystallise it from CHCl_3 (EtOH free, by passing through an alumina column) or C_6H_6 /petroleum ether and dry it in a 20mm vacuum. [Beilstein 11 IV 553.]

Benzene-1,2-dithiol (1,2-dimercaptobenzene) [17534-15-5] **C₆H₆S₂, M 142.2**, **m 24-25°, 27-28°, b 119-120°/17mm, pK_{Est(1)} ~6.1, pK_{Est(2)} ~9.5**. Likely impurities are the oxidation products, the disulfides which could be polymeric. Dissolve it in aqueous NaOH until the solution is alkaline. Extract with Et_2O and discard the extract. Acidify with cold HCl (diluted 1:1 by volume with H_2O) to Congo Red paper under N_2 and extract it three times with Et_2O . Dry the Et_2O with Na_2SO_4 , filter, evaporate and distil the residue under reduced pressure in an atmosphere of N_2 . The distillate solidifies on cooling. [UV: Dewar et al. *J Chem Soc* 3076 1958, DOI: 10.1039/JR9580003076; Adams & Ferretti *J Am Chem Soc* 81 4939 1959, DOI: 10.1021/ja01527a045; Ferretti *Org Synth Coll Vol* 5 419 1973, DOI: 10.15227/orgsyn.042.0054; Beilstein 6 IV 5651.]

Benzene-1,3-dithiol (1,3-dimercaptobenzene) [626-04-0] has **m 24-25°, 27-28°, b 62°/0.2mm, 128-129°/16mm, 244°/atm, 245°/atm, d₂₅ 1.236, pK_{Est(1)} ~6.0, pK_{Est(2)} ~8.0**. Purification is as for the preceding 1,2-isomer. [Beilstein 6 IV 5705.]

Benzenesulfinic acid [618-41-7] **C₆H₆O₂S, M 142.2**, **m 84°, pK²⁵ 2.16(2.74)**. The acid is purified by dissolving the Na salt in H_2O , acidifying to Congo Red paper with HCl and adding a concentrated solution of FeCl_3 whereby Fe sulfinate precipitates. Collect the salt, wash it with a little H_2O , drain, suspend it in H_2O and add a slight excess of 1.5M aqueous NaOH. The $\text{Fe}(\text{OH})_3$ precipitates, it is filtered off, the sulfinic acid in the aqueous solution is extracted with Et_2O , the extract is dried (Na_2SO_4) and evaporated to give colourless crystals of benzenesulfinic acid **m 84°** which are stored under N_2 in the dark, as it slowly oxidises in air to the sulfonic acid. [Beilstein 11 H 2, 26, 11 I 3, 11 II 3, 11 III 3, 11 IV 3.] **Sodium benzenesulfinate** [873-55-2] **C₆H₅O₂SNa, M 164.2**, prepared from the acid with one equivalent of NaOH has **m ~300°**. [Beilstein 11 IV 3.]

Benzenesulfonic acid [anhydrous 98-11-3; monohydrate 26158-00-9] **C₆H₆O₃S, M 158.2**, **m 43-44°, 50-55°(anhydrous), 65-66°, pK²⁵ -2.7, 0.70, (2.53?)**. Purify benzenesulfonic acid by dissolving it in a small volume of distilled H_2O and stirring with slightly less than the theoretical amount of BaCO_3 . When effervescence is complete and the solution is still acidic, filter off the insoluble **barium benzenesulfonate**. The salt is collected and dried to constant weight *in vacuo*, then suspended in H_2O and stirred with a little less than the equivalent (half mol.) of sulfuric acid. The insoluble BaSO_4 (containing a little barium benzenesulfonate) is filtered off and the filtrate containing the **free acid** is evaporated in a high vacuum. The oily residue will eventually crystallise when completely anhydrous. A 32% commercial acid is allowed to fractionally crystallise at room temperature over P_2O_5 in a vacuum desiccator giving fine colourless deliquescent plates **m 52.5°**. The anhydrous crystalline acid is deliquescent and should be stored over anhydrous Na_2SO_4 in the dark, and should be used in subdued sunlight as it darkens under sunlight. The main impurity is Fe which readily separates as the Fe salt in the early fractions [Taylor & Vincent *J Chem Soc* 3218 1952, DOI: 10.1039/JR9520003218]. The **S-benzylisothiuronium salt** has **m 148°** (from $\text{EtOH}/\text{H}_2\text{O}$). It is a strong acid, and an IRRITANT to the skin and eyes. [See Adams & Marvel *Org Synth Coll Vol* 1 84 1941, DOI: 10.15227/orgsyn.001.0021; Michael & Adair *Chem Ber* 10 583 1877, DOI: 10.1002/cber.187701001162; Beilstein 11 IV 27.]

Benzenesulfonyl chloride [98-09-9] **C₆H₅ClO₂S, M 176.6**, has **m 14.5°, b 120°/10mm, 177°/100mm, 251.2°/760mm(dec), d₄²⁰ 1.384**. Distil the sulfonyl chloride, preferably under a vacuum, then treat it with 3mole % each of toluene and AlCl_3 , and allow it to stand overnight. The sulfonyl chloride is distilled off at 1mm pressure and then carefully fractionally distilled at 10mm in an all-glass column. [Adams & Marvel *Org Synth Coll Vol* 1 84 1941, DOI: 10.15227/orgsyn.001.0021; Jensen & Brown *J Am Chem Soc* 80 4042 1958, DOI: 10.1021/ja01548a055; Beilstein 11 IV 49.] It is reactive and **TOXIC, handle with gloves**. With concentrated aqueous NH_3 it is converted into **benzenesulfonamide** [98-10-2] **C₆H₇NOS, M157.2**, with **m 153°** after recrystallisation from H_2O or aqueous EtOH. [Beilstein 11 IV 50.] **Benzenesulfonyl hydrazide (Porofor, BSH)** [98-09-9] **C₆H₈N₂O₂S, M 172.2**, has **m 101-103°, 103-104° (with evolution of N_2)**. The hydrazide separates out when benzenesulfonyl chloride is treated with aqueous hydrazine and cooling. It crystallises from H_2O or aqueous EtOH. [Beilstein 11 IV 94.]

Benzenesulfonic anhydride [512-35-6] $C_{12}H_{20}O_5S_2$, M 298.3, has m 88-91°. Crystallise the anhydride from Et_2O (m 88.5-91.5°), $CHCl_3$ or chlorobenzene (m 90-92°). Store it dry. [Field *J Am Chem Soc* 74 394 1952, DOI: 10.1021/ja01122a032; *Beilstein* 11 H 3, 11 I 11, 11 II 23, 11 III 50, 11 IV 50.]

Benzopurpurin 4B {3,3'-[(3,3'-dimethyl[1,1'-biphenyl]-4,4'-diyl)bis(azo)]bis[4-amino-1-naphthalene-sulfonic acid] di-Na salt, Direct red 2} [992-59-6] $C_{34}H_{26}N_6Na_2O_6S_2$, M 724.7, decomposes without melting, λ_{max} 500nm, CI 23500, $pK^{25} < 0$. It crystallises from H_2O , is soluble in $EtOH$, Me_2CO , $EtOCH_2CH_2OH$, insoluble in other organic solvents, and soluble in aqueous $NaOH$ or H_2SO_4 . It is a biological stain that is violet at pH 1.2 and red at pH 4.0 and is used for detecting Al, Mg, Hg, Au and U. Used also for dyeing rayon and cotton. [*Beilstein* 16 H 411, 16 II 224, 16 III 474, 16 IV 601.]

Benzotriazol-1-yloxytris(dimethylamino)phosphonium hexafluorophosphate (BOP, Castro's Reagent) [56602-33-6] $C_{12}H_{22}F_6N_6OP_2$, M 442.3, m > 130°(dec), 147-149°(dec). Dissolve it in CH_2Cl_2 , dry ($MgSO_4$), filter, concentrate it under a vacuum, then add dry Et_2O and filter off the first crop. Add CH_2Cl_2 to the filtrate and concentrate again to obtain a second crop. The solid is washed with dry Et_2O and dried in a vacuum. Also recrystallise it from dry Me_2CO/Et_2O and check the purity by NMR. Store it in the dark. [Castro et al. *Synthesis* 751 1976, DOI: 10.1055/s-002-1666; Nguyen et al *JCS Perkin Trans I* 1915 1987, DOI: 10.1039/P19870001915; Coste et al. *Tetrahedron Lett* 36 4253 1995, DOI: 10.1016/0040-4039(95)00736-V.]

S-Benzylisothiuronium chloride [538-28-3] $C_8H_{11}ClN_2S$, M 202.7, two forms, m 146-148° and 172-174°, $pK_{Est} \sim 9.8$ (free base). This useful compound which forms characteristic salts with carboxylic, sulfinic and sulfonic acids is prepared from benzyl chloride (126g) and thiourea (76g) in $EtOH$ (200ml), whereby a vigorous reaction occurs to form a homogeneous solution which is boiled for half an hour. On cooling, the chloride salt forms a crystalline magma which is filtered off and recrystallised from $EtOH$, or a mixture of equal volumes of concentrated HCl and H_2O . Crystallise the chloride from 0.2M HCl (2ml/g) or $EtOH$ and it dry in air. It is **dimorphic**. When the lower melting form is dissolved in $EtOH$ and seeded with crystals of the higher melting form, the higher melting form crystallises out. However, the same characteristic carboxylic, sulfinic and sulfonic isothiuronium salts are formed whichever crystals are used. The procedure for preparing the salts for characterisation of organic acids is as follows: To a strong solution of the Na or K salt of the organic acid in aqueous $EtOH$ or H_2O is added, rapidly with stirring, a slight excess of a 15% solution of S-benzylthiuronium chloride in hot $EtOH$. On cooling, the desired S-benzylthiuronium salt crystallises out of solution in a highly pure state. It can be recrystallised from $EtOH$. In a few cases the solution has to be evaporated in order to obtain crystals. This can happen with weak carboxylic acids where the salts readily hydrolyse. To avoid this, the solutions need to be as concentrated as possible, or better avoid the presence of H_2O in the solvent. This does not occur with strong acids that include the sulfinic and sulfonic acids. Careless manipulation of salts of very weak acids, e.g. repeated recrystallisation, leads to hydrolysis where the free base decomposes to form benzylmercaptan which results in a noxious odour. [Donleavy *J Am Chem Soc* 58 1004 1936, DOI: 10.1021/ja01297a048; *Beilstein* 6 III 1600.]

Benzyl mercaptan [100-53-8] C_7H_8S , M 124.2, b 70.5-70.7°/9.5mm, 194-195°/atm, d_4^{20} 1.058, n_D^{20} 1.5761, pK^{25} 9.43. Purify benzyl mercaptan via the **mercury salt** [see Kern *J Am Chem Soc* 75 1865 1953, DOI: 10.1021/ja01104a025], which crystallises from *benzene as needles (m 121°), and then dissolve it in $CHCl_3$. Pass H_2S gas through the solution to regenerate the mercaptan. The HgS that precipitates is filtered off and washed thoroughly with $CHCl_3$. The filtrate and washings are evaporated to remove $CHCl_3$; then the residue is fractionally distilled under reduced pressure. Store under N_2 as it oxidises in air to form **dibenzyl disulfide**. [Mackle & McClean, *Trans Faraday Soc* 58 895 1962, DOI: 10.1039/TF9625800895]. [*Beilstein* 6 IV 2632.]

Benzyl Orange [4-(4-benzylaminophenylazo)benzenesulfonic acid potassium salt] [589-02-6 K-salt, 36402-77-4 Na-salt] $C_{19}H_{16}N_3KO_3S$, M 405.5, $pK_{Est(1)} \sim < 0$, $pK_{Est(2)} \sim 3.8$. Crystallise it from H_2O .

Benzyltriphenylphosphonium chloride [1100-88-5] $C_{25}H_{22}ClP$, M 388.9, m 280° (sintering), 287-288°, 337°. Wash it with Et_2O and crystallise it from $EtOH$ (six-sided plates). It is **hygroscopic** and forms crystals with one molecule of H_2O . [Michaelis et al. *Justus Liebigs Ann Chem* 229 320 1885, DOI: 10.1002/jlac.18852290304; Kröhnke *Chem Ber* 83 295 1950, DOI: 10.1002/cber.19500830318; *Beilstein* 16 IV

994.]

Biphenyl-2-yl diphenyl phosphate [132-29-6] $\text{C}_{24}\text{H}_{19}\text{O}_4\text{P}$, M 402.4, b 280-290°/9mm, 494.4 \pm 24°/760mm, d₂₅²⁵ 1.184, n_D²⁵ 1.5925. Distil the ester in a vacuum, then percolate it through an alumina column. Pass the ester through a packed column maintained at 150° to remove residual traces of volatile materials by a counter-current stream of nitrogen at reduced pressure. [Dobry & Keller *J Phys Chem* **61** 1448 1957, DOI: 10.1021/j150556a052; *Beilstein* **6** III 3295, **6** IV 4585.]

Bis(*p*-tert-butylphenyl)phenyl phosphate [115-87-7] $\text{C}_{26}\text{H}_{31}\text{O}_4\text{P}$, M 438.5, b 281°/5.5mm, d₄²⁵ 1.108, n_D²⁵ 1.5412. Purify as for biphenyl-2-yl diphenyl phosphate (above). [*Beilstein* **6** III 1871, **6** IV 3310.]

Bis(2-chlorophenyl) phenyl phosphate [597-80-8] $\text{C}_{18}\text{H}_{13}\text{Cl}_2\text{O}_4\text{P}$, M 395.2, b 254°/4mm, n_D²⁵ 1.5767. Purify as for biphenyl-2-yl diphenyl phosphate (above). [Fox et al. *J Phys Chem* **59** 1097 1955, DOI: 10.1021/j150532a027; *Beilstein* **6** III 680, **6** IV 807.]

Bis(2-ethylhexyl) 2-ethylhexyl phosphonate [25103-23-5, 126-63-6, 1300-52-3] $\text{C}_{24}\text{H}_{51}\text{O}_3\text{P}$, M 418.6, b 479.1 \pm 14°/760mm, d 0.9 \pm 0.1g/cm³, n_D²⁵ 1.4473. Purify it by stirring a 0.4M solution in *benzene with an equal volume of 6M HCl at ca 60° for 8 hours. The *benzene layer is then shaken successively with equal volumes of water (twice), aqueous 5% Na₂CO₃ (three times), and water (eight times), followed by evaporation of the *benzene and distillation of the residue under reduced pressure at room temperature (using a rotating evacuated flask). It should be stored dry and in the dark [Peppard et al. *J Inorg Nucl Chem* **24** 1387 1962, DOI: 10.1016/0022-1902(62)80147-X]. Distil it in a vacuum, then percolate it through an alumina column before finally passing it through a packed column maintained at 150° where residual traces of volatile materials are removed by a counter-current stream of N₂ under reduced pressure [Dobry & Keller *J Phys Chem* **61** 1448 1957, DOI: 10.1021/j150556a052].

Bis(2-ethylhexyl) phosphoric acid ('diisooctyl' phosphate) [298-07-7, 27215-10-7] $\text{C}_{16}\text{H}_{35}\text{O}_4\text{P}$, M 322.4, m -60°, -50°, b 209°/10mm, 393°/760mm, d₄²⁰ 0.965, pK_{Est} ~1.7. Contaminants of commercial samples include the monoester, polyphosphates, pyrophosphate, 2-ethylhexanol and metal impurities. Dissolve the acid in *n*-hexane to give an 0.8M solution. Wash this with an equal volume of M HNO₃, then with saturated (NH₄)₂CO₃ solution, with 3M HNO₃, and twice with water [Petrov & Allen *Anal Chem* **33** 1303 1961, DOI: 10.1021/ac60178a004]. Similarly, the impure sodium salt, after scrubbing with petroleum ether, is acidified with HCl and the free organic acid is extracted into petroleum ether and purified as above [Peppard et al. *J Inorg Nucl Chem* **7** 231 1958, DOI: 10.1016/0022-1902(58)80075-5], or as described by Stewart & Crandall [*J Am Chem Soc* **73** 1377 1951, DOI: 10.1021/ja01147a533]. It can be purified *via* its copper salt [McDowell et al. *J Inorg Nucl Chem* **38** 2127 1976, DOI: 10.1016/0022-1902(76)80486-1]. [*Beilstein* **1** IV 1796.] It can be used as a solvent for the extraction of uranium and rare earth metals [Sato *Hydrometallurgy* **22** 121 1989, DOI: 10.1016/0304-386X(89)90045-5], and similarly it can be used for extracting Fe, but it must be in the reduced form Fe²⁺.

Bis-(4-fluoro-3-nitrophenyl) sulfone [312-30-1] $\text{C}_{12}\text{H}_6\text{F}_2\text{N}_2\text{O}_6\text{S}$, M 344.3, m 193-194°. Recrystallise the sulfone from Me₂CO and H₂O (5:1). It should give a yellow colour in aqueous base. [Zahn & Zuber *Chem Ber* **86** 172 1953, DOI: 10.1002/cber.19530860209; *Beilstein* **6** IV 172.]

2,4-Bis(methylthio)-1,3,2λ⁵,4λ⁵-dithiadiphosphetane-2,4-dithione (Davy's reagent methyl) [82737-61-9] $\text{C}_2\text{H}_6\text{P}_2\text{S}_6$, M 284.4, m 160°, b 382.5°/760mm, d 1.69g/cm³. It crystallises from *C₆H₆ in yellow plates or from hot trichlorobenzene. The low melting point reported in the literature (112° with gradual softening at 68-102°) has been attributed to the presence of elemental sulfur in the crystals. It has a **foul odour** and is a **suspected carcinogen**. [Yousif et al. *Tetrahedron* **40** 2663 1984, DOI: 10.1016/S0040-4020(01)96883-8; Scott et al. *J Org Chem* **22** 789 1957, DOI: 10.1021/jo01358a019.]

Bis-(phenylsulfonyl)imide (dibenzenedisulfimide) [2618-96-4] $\text{C}_{12}\text{H}_{11}\text{NO}_4\text{S}_2$, M 297.4, m 157-158°. Obtained by adding slowly a 1:1 solution of benzenesulfonyl chloride (2.2 mol) in Me₂CO to a 4% solution of NH₄Cl (1 mol) with stirring followed, at 0°, by addition of 10N aqueous NaOH until the pH rises to 8, and the

temperature is allowed to rise to 25°. Acetone was distilled off, and *diphenylsulfinimide sodium salt* separated [~90% yield, **m 327-329°** (also **314-316°** reported); and was recrystallised from H₂O in needles, and dried at 100°. It can similarly be prepared from benzenesulfonamide by reaction with benzenesulfonyl chloride in the presence of NaOH. The *free imide* is obtained on acidification, extraction into Et₂O, and finally crystallised as white crystals from Et₂O. It readily forms stable recrystallisable salts with a variety of amines, amino acids and esters, as well as a variety of heterocyclic compounds and alkaloids; and used as a method for purifying these compounds (see Tables 1 to 5 in Chapter 2). The *L-glutamic acid di-tert-butyl ester dibenzenesulfimide salt* [16874-06-9] has **m 139-141°**, $[\alpha]_{\text{D}}^{20} +12$ (c 2, MeOH); the *glycine tert-butyl ester dibenzenesulfimide salt* [77284-30-1] has **m 154-157°**; the *L-aspartic acid di-tert-butyl ester dibenzenesulfimide salt* [70534-48-4] has **m 150-155°**, $[\alpha]_{\text{D}}^{20} +6.6$ (c 2, pyridine); and the *L-proline tert-butyl ester dibenzenesulfimide salt* [83283-35-6] and are available commercially. *Bis-(4-chlorobenzenesulfonyl)imide sodium salt* crystallises as white needles from H₂O, which on drying at 100°, have **m 298°** (also **297-300°**, **>360°**, **dec**). The *free imide* is obtained on acidification, and crystallises from H₂O (**m 207-208°**) or from *C₆H₆ (**m 206-207°**). [Runge et al. *Chem Ber* **88** 533 1955, DOI: 10.1002/cber.19550880414; Runge et al. *Chem Ber* **86** 1571 1953, DOI: 10.1002/cber.19530861216; Runge & Pfeiffer *Chem Ber* **90** 1757 1957, DOI: 10.1002/cber.19570900910; Helferich & Grünert *Chem Ber* **73** 1131 1940, DOI: 10.1002/cber.19400731019; Helferich & Flechig *Chem Ber* **75** 532 1942, DOI: 10.1002/cber.19420750514; Beilstein **11** H 49, **11** IV 85, 122.] *Dibenzenesulfonimide* is soluble in alkaline solution, used in electroplating, and a primary brightner instead of saccharin with better leveling ability, and with lower consumption than saccharin. [See Tables 1 to 5 in Chapter 2.]

Bismuthiol I (2,5-dimercapto-1,3,4-thiadiazole) potassium salt [4628-94-8] C₂H₂N₂S₃, **M 226.4**, **m 274-276°(dec)**, **275-276°(dec)**, **~294°(dec)**, **pK_{Est(1)} ~4.1**. Usually contaminated with disulfide. Purify it by recrystallisation from EtOH (yellow needles). *Bismuthiol I* [1072-71-5] C₂K₂N₂S₃, **M 150.3**, is the *free acid* and has **m 165°(dec)** (**m 162°** and **168°** also reported), **pK_{Est(1)} ~4.5**. This reagent has been used for the spectroscopic detection of Bi, Cu, Pb and Sb, and forms complexes with Pd and Pt. [Busch *Chem Ber* **27** 2507 1894, DOI: 10.1002/cber.189402702251; Najumdar & Chakrabarty *Anal Chim Acta* **19** 372 1958, DOI: 10.1016/S0003-2670(00)88179-9; Mura et al. *Inorg Chim Acta* **97** 45 1985, DOI: 10.1016/S0020-1693(00)87988-8; Beilstein **27** H 677, **27** IV 7725.]

Bis-(2-nitrophenyl) disulfide [1155-00-6] C₁₂H₈N₂O₄S₂, **M 308.3**, **m 192-195°**, **195°**, **194-197°**, **198-199°**. Purify the disulfide by recrystallisation from glacial AcOH or from *C₆H₆ and the yellow needles are dried in an oven at 100° until the odour of the solvent is absent. It is sparingly soluble in EtOH and Me₂CO. It is used for the preparation of *orthanilic acid*, see below. [Bogert & Stull *Org Synth Coll Vol* **1** 220 1941, DOI: 10.15227/orgsyn.008.0064; Bauer & Cymerman *J Chem Soc* 3433 1949, DOI: 10.1039/JR9490003433; Beilstein **6** IV 1672.]

Bis[4-(1,1,3,3-tetramethylbutyl)phenyl]phosphate calcium salt (Selectophore) [40835-97-0] C₅₆H₈₄CaO₈P₂, **M 987.3**. The Ca diester salt is washed with H₂O (x3) and MeOH (x3) alternately and dried in a vacuum oven at 50°. If the Ca salt is contaminated with much Ca salt of the *monoester*, then it (10g) is converted to the *free acid* by adding 6N HCl (ca 10 volumes) and Et₂O (> 50 volumes) is added to it and stirred vigorously to form the free acids. When no white precipitate remains (ca 5 minutes), the Et₂O is separated, washed with H₂O (2x > 50 ml) and dried by filtering through a bed of anhydrous Na₂SO₄ (11 x 5 cm) which is then washed with Et₂O (2x > 50 ml). Evaporation gives an oil (TLC R_F 0.81 for diester and 0.50 for monoester). The oil is dissolved in *benzene (ca 25ml) and extracted with ethane-1,2-diol (25ml, 10x). After ten washings, a small sample of the *benzene layer is washed twice with H₂O to remove the diol and showed that it is pure *bis-[4-(1,1,3,3-tetramethylbutyl)-phenyl]phosphoric acid* by TLC, i.e. no monophosphate. To form the *Ca salt*, the oil is dissolved in MeOH and to it is added the equivalent amount of CaCl₂ together with aqueous NaOH to keep the pH >10. The resulting white precipitate is collected, washed alternately with 3 batches of H₂O and MeOH and dried in a vacuum oven at 50°. [Craggs et al. *J Inorg Nucl Chem* **40** 1483 1978, DOI: 10.1016/0022-1902(78)80455-2; Morton et al. *Anal Biochem* **157** 345 1986, DOI: 10.1016/0003-2697(86)90636-6.] It has been used as a component of a polyaniline Ca-selective conducting electrode membrane [Lindfors & Ivaska *Analyt Chim Acta* **437** 171 2001, DOI: 10.1016/S0003-2670(01)00996-5].

2,4-Bis(p-tolylthio)-1,3,2λ⁵,4λ⁵-dithiadiphosphetane-2,4-dithione (Heimgartner's reagent or Davy's rea-

gent *p*-tolyl) [114234-09-2] $C_{14}H_{14}P_2S_6$, **M 436.6, m 175-176°(dec)**. Recrystallise it from toluene (light yellow solid), wash it with Et_2O and dry *in vacuo*. It has IR (KBr) with ν_{max} at 3420w, 1597w, 1488w, 1387w, 1183w, 1020w, 941w, 811s and 684s cm^{-1} ; and MS has *m/e* at 218 (10%, $M^{+}/2$). [Wipf et al. *Helv Chim Acta* **70** 1001 1987, DOI: 10.1002/hlca.19870700412.]

***N,O*-Bis(trimethylsilyl)acetamide (BSA)** [10416-59-8] $C_8H_{21}NOSi_2$, **M 203.4, b 71-73°/35mm, d_4^{20} 0.836, n_D^{20} 1.4150**. Fractionate it through a spinning band column and collect liquid **b 71-73°/35mm**, and not higher because the main impurity MeCONHSiMe₃ distils at **b 105-107°/35mm**. It is used for derivatising alcohols and sugars [Klebe et al. *J Am Chem Soc* **88** 3390 1966, DOI: 10.1021/ja00966a038; see Blau, Karl; J. M. Halket *Handbook of Derivatives for Chromatography* (2nd ed.). John Wiley & Sons 1993, ISBN 0-471-92699-X; Matsuo et al. *Carbohydr Res* **241** 209 1993, DOI: 10.1016/0008-6215(93)80107-P; Johnson *Carbohydr Res* **237** 313 1992, DOI: 10.1016/S0008-6215(92)84254-P]. It is **FLAMMABLE** and **TOXIC**.

Bis(trimethylsilyl)acetylene (BTMSA) [14630-40-1] $C_8H_{18}Si_2$, **M 170.4, m 24-24.5°, 26°, b 134-136°/atm**. Dissolve it in petroleum ether and wash it with ice-cold dilute HCl. The petroleum ether extract is dried ($MgSO_4$), evaporated and fractionated at 760mm. Its solubility in H_2O is 31mg/L at ~25°. [Walton & Waugh *J Organomet Chem* **37** 45 1972, DOI: 10.1016/S0022-328X(00)89260-8; *Beilstein* **4** IV 3950.]

Bis(trimethylsilyl) selenide [4099-46-1] $C_6H_{18}SeSi_2$, **M 225.3, m -7°, b 31°/2mm, 45-46°/5.3mm, 176°/atm, d^{25} 0.9010g/cm³**. Purify by redistillation at high vacuum preferably in an inert atmosphere. Prepare it as follows: Under a dry inert (argon) atmosphere containing 1M Li(Et)₃BH in THF (105ml, 0.105mol) and cooled in an ice bath, is added selenium shot (3.95g, 0.0500mol). The mixture is slowly warmed to ~25°, with stirring during 2 hours, then cooled to ice-bath temperature and Me₃SiCl (12.6g, 0.117mol, see [75-77-4] below) is added all at once. The mixture is stirred for 2 hours at ~25°, the low boiling volatiles are removed at low vacuum, then the product is distilled at higher vacuum to give the colourless selenide [10.6g, 95%, **b 45-46°/5.3mm**, ¹H NMR (CDCl₃) one peak at δ 0.50]. Store at -35° in a drybox. [Detty & Seidler *J Org Chem* **47** 1354 1982, DOI: 10.1021/jo00346a041; see also Steigerwald et al. *J Am Chem Soc* **110** 3046 1988, DOI: 10.1021/ja00218a008.]

It is a colourless, air sensitive liquid with a strong odour, readily hydrolysed by moist air to give highly toxic H_2Se , so work under an efficient fume cupboard. Store under argon or N_2 at -20°, at which temperature it is a solid. It slowly deposits amorphous Se which is red in colour. Destroy the pot residue in an inert atmosphere after treating with cold MeOH. Any gases that escape should be absorbed in an aqueous alkaline solution. [Osawa & Sonoda 15 APR 2001 *e-EUROS Encyclopedia of reagents in organic synthesis*, DOI: 10.1002/047084289X.rb221.]

Bis(trimethylsilyl) sulfide (hexamethyldisilathiane) [3385-94-2] $C_6H_{18}SSi_2$, **M 178.5, b 65-67°/16mm, 162.5-163.5°/750mm, 164°/760mm, d_4^{20} 0.85, n_D^{20} 1.4598**. Dissolve it in petroleum ether (b ca 40°), remove the solvent and distil it. Redistil it under atmospheric pressure of dry N_2 . It is collected as a colourless liquid which solidifies to a white solid in Dry-ice. On standing for several days it turns yellow possibly due to liberation of sulfur. Store it below 4° under dry N_2 . It is prepared by adding Ag₂S (37g, 0.15mol) to Me₃SiI (40g, 0.2mol, see 16029-98-4 below) and heating under reflux for 14 hours when the temperature rises to 150°. Distil off the liquid from the solid and redistil to give the disilathiane (13g, 73%). The boiling time depended on the quality of the Ag₂S. [Eaborn *J Chem Soc* 3077 1950, DOI: 10.1039/JR9500003077; *Beilstein* **4** IV 4033; Fieser **8** 240, **15** 165.] Alternatively, sulfur (0.64g, 0.02mol) is added to 1M Li(Et)₃BH in THF (40ml, 0.04mol) with cooling in an ice bath. The mixture is then stirred for 0.5 hour at ~25° and Me₃SiCl (4.32g, 0.04mol, see 75-77-4 below) is added slowly as an exothermic reaction ensues, the mixture is stirred for 2 hours at ~25°. The low boiling volatiles are removed (at 30-40°/20mm), and the disilathiane (2.9g, 83%) is distilled off. Store it at -35° in a drybox. [Detty & Seidler *J Org Chem* **47** 1354 1982, DOI: 10.1021/jo00346a041.]

Bis(trimethylsilyl) telluride [4551-16-0] $C_6H_{18}TeSi_2$, **M 274.0, m 13.5°, b 74°/11mm**. Purify by fractional distillation (or bulb transfer) at reduced pressures under red light to minimise photochemical decomposition, also observed after 24 hours in dark storage at -20° under argon. It is prepared freshly from Te shot (3.56g, 0.0279mol) and 1M Li(Et)₃BH (58ml, 0.058mol) as described for the preceding Se compound. After 8 hours the mixture turned purple with a chalky white suspension. The flask is wrapped in foil, Me₃SiCl (7.00g, 0.0648mol, see 75-77-4 below) is added, stirred at ~25° for 6 hours, the volatiles are then removed (at 30-40°/20mm), and

then distilled to give the telluride (4.07g, 53%, **b** 49-51°/2.5mm, orange-brown liquid) which is collected in a foil-wrapped receiving flask to minimise exposure to light. Store it as a solid at -20° under Ar in the dark, or at -35° in a drybox also in the dark. [Detty & Seidler *J Org Chem* **47** 1354 1982, DOI: 10.1021/jo00346a041; for reactions with acyl halides to form C-C bonds see Severengiz et al. [*Angew Chem Int Ed* **24** 1041 1985, DOI: 10.1002/anie.198510411], and Severengiz & du Mont [*JCS Chem Commun* 820 1987, DOI: 10.1039/C39870000820].

The more stable *bis(tert-butyldimethylsilyl) telluride* [80594-86-1] $C_{12}H_{30}TeSi_2$, **M 358.1**, **m 46-49°**, **b 90-95°/5.5mm**, is prepared from Te shot (3.56g, 0.0279mol), 1M Li(Et)₃BH (58ml, 0.058mol), and *tert*-butylchlorodimethylsilane (9.74g, 0.0580mol, 18162-48-6 *below*) in a similar way. The product (7.77g, 79%) is isolated as a white solid after initial distillation as a colourless oil. It has ¹H NMR (CDCl₃) δ 1.00 (s, 9 H), 0.57 (s, 6 H). It is stable for several weeks at -20° under Ar in sealed flasks in the dark, but when stored at ~25° under red light it is noticeably decomposed after several days. [Detty & Seidler *J Org Chem* **47** 1354 1982, DOI: 10.1021/jo00346a041.]

9-Borabicyclo[3.3.1]nonane (9BBN) [*monomer* 280-64-8] [*dimer* 21205-91-4 or 70658-61-6] [*1:1 coordination compound with tetrahydrofuran* 76422-63-4] $C_8H_{15}B$, **M 122.0 (monomer), 244.0 (dimer)**, **m 141-143° (monomer), 150-152°, 154-155° (dimer)**, **b 195°/12mm**. It is available as the solid dimer or in tetrahydrofuran solution. The solid is relatively stable and can be purified by distillation in a vacuum (as dimer) and by recrystallisation from tetrahydrofuran (solubility at room temperature is 9.5%, 0.78M), filter off the solid under N₂, wash it with dry pentane and dry it *in vacuo* at *ca* 100°. The solid is a *dimer* (IR 1567cm⁻¹), stable in air (for *ca* 2 months), and can be heated for 24 hours at 200° in an inert atmosphere without loss of hydride activity. It is a *dimer* in tetrahydrofuran solution also (IR 1567cm⁻¹). It is sensitive to H₂O and air (O₂) in solution. Its concentration in solution can be determined by reaction with MeOH and measuring the volume of H₂ liberated, or it can be oxidised to *cis-cyclooctane-1,5-diol* (**m 73.5-74.5°**). [IR: Knights & Brown *J Am Chem Soc* **90** 5280 1968, DOI: 10.1021/ja01021a046; Brown et al. *J Am Chem Soc* **96** 7765 1974, DOI: 10.1021/ja00832a025; Brown et al. *J Org Chem* **41** 1778 1976, DOI: 10.1021/jo00872a025; Brown & Chen *J Org Chem* **46** 3978 1981, DOI: 10.1021/jo00333a009; Fieser **2** 31, **3** 24, **10** 48, **15** 43, **17**, 49.]

Borane pyridine complex [110-51-0] C_5H_5BN , **M 92.9**, **m 8-10°, 10-11°**, **b 86°/7mm, 100-101°/12mm**, **d₄²⁰ 0.785**. Dissolve it in Et₂O and wash it with H₂O in which it is insoluble. Evaporate the Et₂O and distil the residual oil to give better than 99.8% purity. Its vapour pressure is less than 0.1mm at room temperature. [Taylor et al. *J Am Chem Soc* **77** 1506 1955, DOI: 10.1021/ja01611a031; *Beilstein* **20** IV 2235.]

Borane triethylamine complex [1722-26-5] $C_6H_{15}N \cdot BH_3$, **M 115.0**, has **b 76°/4mm, 8°/7mm, 100-101°/12mm, d₄²⁰ 0.78**. Distil it in a vacuum using a 60cm glass helices-packed column. [Brown et al. *J Am Chem Soc* **64** 325 1942, DOI: 10.1021/ja01254a031; Ashby & Foster *J Am Chem Soc* **84** 3407 1962, DOI: 10.1021/ja00876a040; Matsumura & Tokura *Tetrahedron Lett* 4703 1968, DOI: 10.1016/S0040-4039(00)89911-6; *Beilstein* **4** IV 329.]

Borane trimethylamine complex [75-22-9] $C_3H_9N \cdot BH_3$, **M 73.0**, has **m 94-94.5°, 95°**, **b 171°/atm, 172°/760mm, d 0.81g/ml**. It is sublimed using equipment described in Burg and Schlesinger [*J Am Chem Soc* **59** 780 1937, DOI: 10.1021/ja01284a002]. Its vapour pressure is 86mm at 100°. It forms colourless hexagonal crystals varying from needles to short lumps, which are slightly soluble in H₂O (1.48% at 30°), EtOH (1%), hexane (0.74%), but very soluble in Et₂O, *C₆H₆ and AcOH. It is stable at 125°. [Burg & Schlesinger *J Am Chem Soc* **59** 780 1937, DOI: 10.1021/ja01284a002; Brown et al. *J Am Chem Soc* **64** 325 1942, DOI: 10.1021/ja01254a031; *Beilstein* **4** IV 140.]

2-Bromoallyltrimethylsilane [81790-10-5] $C_6H_{13}BrSi$, **M 193.2**, **b 64-66°/10mm, 82-85°/58-60mm, d₄²⁵ 1.13**. It is fractionally distilled through an efficient column. Store at 2-8°. It is **flammable**. [Trost & Chan *J Am Chem Soc* **104** 3733 1982, DOI: 10.1021/ja00377a038; Trost & Coppola *J Am Chem Soc* **104** 6879 1982, DOI: 10.1021/ja00388a112.] It has been used as a bifunctional molecular linchpin in a three component protocol using it [Smith III & Duffey *Synlett* 1363 2004, DOI: 10.1055/s-2004-825621].

4-Bromobenzenesulfonyl chloride [98-58-8] $C_6H_4BrClO_2S$, **M 255.5**, **m 73-75°, 74.3-75.1°, 75.4°, 75-76°, 77°**, **b 150.6°/13mm, 153°/15mm**. Wash the sulfonyl chloride with cold water, dry and recrystallise it from petroleum ether, or from ethyl ether cooled in powdered Dry-ice after the ether solution had been washed with

10% NaOH until colourless, then dry it with anhydrous Na₂SO₄. *Alternatively*, dissolve it in CHCl₃, wash it with H₂O, dry (Na₂SO₄), evaporate and recrystallise it from light petroleum (e.g. b 60-80°) or dry Et₂O. [Huntress & Carten *J Am Chem Soc* **62** 511 1940, DOI: 10.1021/ja01860a014.] Test for the SO₂Cl group by dissolving it in EtOH and boiling with NH₄CNS whereby a yellow amorphous precipitate forms on cooling. [Beilstein **11** IV 162.] On boiling the sulfonyl chloride (e.g. 0.5g) with aqueous concentrated NH₃ (5ml, sp.gr. 0.90) for 10min, cooling, adding cold H₂O (e.g. 10ml), collecting and washing thoroughly with H₂O, then drying it *in vacuo* followed by recrystallisation from aqueous EtOH provides pure **4-bromobenzenesulfonamide** [701-34-8] C₆H₆BrNO₂S, M 236.1, with m 161.4° (163-167° also reported). [Huntress & Carten *J Am Chem Soc* **62** 511 1940, DOI: 10.1021/ja01860a014; Beilstein **11** III 104.]

2-Bromo-1,3,2-benzodioxaborole [51901-85-0] C₆H₄BBrO₂, M 198.8, m 47°, 51-53°, b 76°/9mm. Keep at 20°/15mm for some time and then fractionally distil. [Gerrard et al. *J Chem Soc* 1529 1959, DOI: 10.1039/JR9590001529; Beilstein **6** IV 5612.]

(+)-3-Bromocamphor-8-sulfonic acid [5344-58-1] C₁₀H₁₅BrO₄S, M 311.2, m 195-196°(anhydrous), [α]_D²⁰ +88.3 (c 1, H₂O), pK ~0. Crystallise the acid from water. The *ammonium salt* has m 268-207°, [α]_D²⁰ +81.9 (c 2.2, H₂O). [Kauffman *J Prakt Chem* **33** 95 1966, DOI: 10.1002/prac.19660330512; Beilstein **11** H 319, **11** I 77, **11**, II 183, **11** III 595.]

IR(endo, anti)-3-Bromocamphor-8-sulfonic acid ammonium salt [55870-50-3, 14575-84-9] C₁₀H₁₅BrO₄S.NH₃ M 328.2, m 284-285°(dec), [α]_D²⁵ +84.8 (c 4, H₂O). Pass a hot aqueous solution of it through an alumina column to remove water-soluble coloured impurities which remain on the column when the ammonium salt is eluted with hot water. The salt is crystallised from water and is dried over CaCl₂ in a desiccator [Craddock & Jones *J Am Chem Soc* **84** 1098 1962, DOI: 10.1021/ja00866a006; Kauffmann *J Prakt Chem* **33** 295 1966, DOI: 10.1002/prac.19660330512]. [Beilstein **11** H 319, **11** I 77, **11**, II 183, **11** III 595.]

(+)-3-Bromocamphor-10-sulfonic acid hydrate [67999-30-8] C₁₀H₁₅BrO₄S.xH₂O M 329.2, has m 119-121°, [α]_D²⁰ +98.3 (c 1, H₂O), pK ~0. Crystallise the acid from water. [For 'Methods of optical resolution' see Boyle *Quart Rev Chem Soc* **25** 323 1971, DOI: 10.1039/QR9712500323; UV: Lowry & Owen *J Chem Soc* 606 1926, DOI: 10.1039/JR9262900606; Beilstein **11** II 181, **11** III 592.]

Bromocresol Green (3',3'',5',5''-tetrabromo-*m*-cresolsulfonephthalein) [76-60-8] C₂₁H₁₄Br₄O₅S, M 698.0, m 218-219°(dec), 225°(dec), pK²⁵ 4.51. Crystallise the dye from glacial acetic acid or dissolve it in aqueous 5% NaHCO₃ solution and precipitate it from the hot solution by dropwise addition of aqueous HCl. Repeat this until the UV/VIS-extinction does not increase at λ_{max} 423nm. It is an indicator: at pH 3.81 (yellow) and pH 5.4 (blue-green). [Beilstein **19/3** V 460.]

Bromocresol Purple (5',5''-dibromo-*o*-cresolsulfonephthalein) [115-40-2] C₂₁H₁₆Br₂O₅S, M 540.2, has m 241-242°(dec), pK₁ -2.15, pK₂ 6.3. Dissolve the dye in aqueous 5% NaHCO₃ solution and precipitate it from a hot solution by dropwise addition of aqueous HCl. Repeat this until the UV/VIS-extinction does not increase at λ_{max} 419nm. It can also be recrystallised from *benzene. It is an indicator: at pH 5.2 (yellow) and pH 6.8 (purple). [Beilstein **19/3** V 460.]

Bromophenol Blue (3,3',5,5'-tetrabromophenolsulfonephthalein, sultone form) [115-39-9] C₁₉H₁₀Br₄O₅S, M 670.0, has m 270-271°(dec), 273°(dec), 279°(dec), λ_{max} 422nm, pK²⁵ 3.62 (acidic). Crystallise the dye from *C₆H₆ or Me₂CO/AcOH, and dry it in air. It is an indicator: at pH 3.0 it is yellow and is purple at pH 4.6. For indicator use: Dissolve indicator (0.1g) in of 0.02M NaOH (7.45ml) and dilute with H₂O (750ml). [Beilstein **19/3** V 458.] **Bromophenol Blue solution (sodium salt, 0.04wt% in H₂O)** [62625-28-9] C₁₉H₉Br₄NaO₅S, M 691.9, λ_{max} 422nm, is available commercially. [Beilstein **19** 649.]

Bromopyrogallol Red (5,5'-dibromopyrogallolsulfonephthalein) [16574-43-9] C₁₉H₁₀Br₂O₈S, M 558.2 (576.2 for free sulfonic acid), has m 300°, λ_{max} 538nm (ε 54,500 H₂O pH 5.6-7.5), pK₁ 2.9, pK₂ 4.39, pK₃ 9.15, pK₄ 11.72. Crystallise the dye from 50% EtOH, or aqueous alkaline solution followed by acidification. It is a metal chromic indicator. [Suk *Coll Czech Chem Commun* **31** 3127 1966, DOI: org/10.1135/ccccc19663127; Beilstein **19/10** V 226].

Bromosulfalein (phenoltetrabromophthalein 3',3'-disulfonic acid disodium salt) [71-67-0] C₂₀H₈Br₄O₁₀S₂.2Na, M 838.0. Purify it by TLC on silica Gel G (Merck 250μ particle size) in two solvent

systems (BuOH/AcOH/H₂O 30:7.5:12.5 v/v, and BuOH/propionic acid/H₂O 30:20:7.5 v/v). When the solvent reaches a height of ~10cm, the plate is removed, dried in air and developed with NH₃ vapour giving blue-coloured spots. Also, the dye can be chromatographed on MN Silica Gel with *t*-BuOH/H₂O/*n*-BuOH (32:10:5 v/v) as eluent and visualised with a dilute KOH (or NaOH if the Na salt is required) spray. The product corresponding to bromosulfalein is scraped off and eluted with H₂O, filtered and evaporated to dryness in a vacuum. It is then dissolved in H₂O, filtered through Sephadex G-25 and evaporated to dryness. [UV and IR identification: Barbier & DeVeerd *J Pharm Sci* **57** 819 1968, DOI: 10.1002/jps.2600570521; NMR: Kato et al. *Chem Pharm Bull Jpn* **20** 581 1972, DOI: 10.1016/0003-2697(77)90512-7; McGuire et al. *Anal Biochem* **83** 75 1977, DOI: 10.1016/0003-2697(77)90512-7; *Beilstein* **18/9** V 461.]

Bromothymol Blue (3',3"-dibromothymolsulfonephthalein) [76-59-5] **C₂₇H₂₈Br₂O₅S**, **M 624.4**, has **m 201-203°**, **pK₁ -0.66**, **pK₂ 6.99**. Dissolve the dye in aqueous 5% NaHCO₃ solution and precipitate it from the hot solution by dropwise addition of aqueous HCl. Repeat this until the extinction at λ_{max} 420 nm does not increase. It is an indicator [prepared by dissolving it (0.1g) in 50% aqueous EtOH (100ml)]: aqueous solutions are yellow at pH 6.0, and blue at pH 7.6. [*Beilstein* **19/3** V 461.]

Bromotrimethylsilane (trimethylbromosilane, trimethylsilyl bromide) [2857-97-8] **C₃H₉BrSi**, **M 153.1**, **m -43.5° to -43.2°**, **b 40.5°/200mm**, **77.3°/735mm**, **79°/744mm**, **79.8-79.9°/754mm**, **d₄²⁰ 1.1805**, **n_D²⁰ 1.422**. Purify it by repeated fractional distillation and store it in sealed ampoules in the dark. [McCusker & Reilly *J Am Chem Soc* **75** 1583 1953, DOI: 10.1021/ja01103a019.] Also fractionate it through a 15-plate column (0.8 x 32cm packed with 1/16in single turn helices of Pt-Ir wire). [Gilliam et al. *J Am Chem Soc* **68** 1161 1946, DOI: 10.1021/ja01211a009; Pray et al. *J Am Chem Soc* **70** 433 1948, DOI: 10.1021/ja01181a528; *Beilstein* **4** IV 4008.]

But-3-enylboronic acid [379669-72-4] **C₄H₉BO₂**, **M 99.9**, **m 84-90°**, **pK_{Est} 8.8**. Recrystallise the acid from toluene and dry it *in vacuo*. [cf. Letsinger & Skoog *J Org Chem* **18** 895 1953, DOI: 10.1021/jo50013a019.]

Butylboronic acid (1-butanedihydroxyborane) [4426-47-5] **C₄H₁₁BO₂**, **M 101.9**, **m 90-92°**, **94-96°**, **pK_{Est} ~8.8**. Purify the acid by recrystallisation from *C₆H₆/petroleum ether and dry it *in vacuo*. [Corey & Cimprich *J Am Chem Soc* **116** 3151 1994, DOI: 10.1021/ja00086a066; Quallich et al. *J Am Chem Soc* **116** 8515 1994, DOI: 10.1021/ja00098a012; Seerden et al. *Tetrahedron Lett* **35** 4419 1994, DOI: 10.1016/S0040-4039(00)73373-9; *Beilstein* **4** IV 4383.]

(±)-sec-Butylboronic acid ([sec-butyl]-dihydroxyborane) [88496-88-2] **C₄H₁₁BO₂**, **M 101.9**, **m 86-89°**, **87-88°**, **pK_{Est} ~8.8**. Purify the acid by recrystallisation from *C₆H₆/petroleum ether and dry *in vacuo*. [McCusker et al. *J Am Chem Soc* **79** 5179 1957, DOI: 10.1021/ja01576a026; *Beilstein* **4** IV 4386.]

tert-Butyldicyclohexylphosphine (dicyclohexyl-tert-butylphosphine, Cy₂P^tBu) [93634-87-8] **C₁₆H₃₁P**, **M 254.4**, **m ~22-25°**, **d₄²⁵ 1.094**, **pK_{Est} ~8.7**. This phosphine was prepared by adding *t*-BuLi (18ml, 27.2mmol, 1.5M in pentane) dropwise to a solution of chloro-dicyclohexylphosphine (5.75g, 24.7mmol, [16523-54-9] see above) in THF (20ml) at -78° and the yellow suspension was allowed to warm to ~25° and stirred overnight. The mixture was evaporated to dryness, extracted with pentane (2 x 20ml), filtered through Celite, concentrated to 20ml, cooled to -78° and after 2 hours the **Cy₂P^tBu** crystallised in white crystals (5.6g, 89%) which were filtered off and washed with cold pentane under N₂ or argon. It is highly air sensitive, flammable and should be stored under N₂ or argon. It melts at room temperature. The ¹H NMR (CDCl₃) has peaks at δ: 1.90-1.05 (m, 22H, C₆H₁₁), 1.10 (d, 9H, C(CH₃)₃, ³J_{H-P} = 10.8Hz); the ¹³C NMR (CDCl₃) has peaks at δ: 33.11 (d, CMe₃, ¹J_{C-P} = 19.4Hz), 33.61 (d, C₁ C₆H₁₁, ¹J_{C-P} = 16.2Hz), 30.96, 27.82 (2d, C₂ C₆H₁₁, ²J_{C-P} = 9.7Hz), 30.33 (d, CH₃)₃, ²J_{C-P} = 13.2Hz), 27.69, 27.61 (2s, C₃ C₆H₁₁), 26.41 (s, C₄ C₆H₁₁); and ³¹P NMR (CDCl₃) has a peak at δ 28.58. [Jan et al. *J Organomet Chem* **606** 55 2000, DOI: 10.1016/S0022-328X(00)00287-4.]

tert-Butyldimethylsilyl chloride (TBDMSCl) [18162-48-6] **C₆H₁₅ClSi**, **M 150.7**, **m 87-89°**, **92.5°**, **b 125°/760mm**. Fractionally distil it at atmospheric pressure. [Sommer & Tyler *J Am Chem Soc* **76** 1030 1954, DOI: 10.1021/ja01633a032; Corey & Venkateswarlu *J Am Chem Soc* **94** 6190 1972, DOI: 10.1021/ja00772a043; *Beilstein* **4** IV 4076.] It is a useful derivatising reagent for analytical and preparative purposes [Lalonde & Chan

Synthesis 817 1985, DOI: 10.1055/s-1985-31361], and for selective cleavage of TBDMS-ethers [with *i*-Bu₂AlH: Corey & Jones *J Org Chem* **57** 1028 1992, DOI: 10.1021/jo00029a050; with H₂SiF₆ in MeCN: Pilcher et al. *J Org Chem* **57** 2492 1992, DOI: 10.1021/jo00034a057].

tert-Butyldimethylsilyl-N-methyltrifluoroacetamide (BSMTFA, MTBSTFA) [77377-52-7] C₉H₁₈F₃NOSi, **M 241.3, b 168-170°/760mm, 172-175°/atm, d**²⁵ **1.036g/cm³, n**_D²⁰ **1.401-1.403**. Obtained in 91% yield by reaction of *N*-methyl-2,2,2-trifluoroacetamide (1 equiv) in /MeCN (1:1 v/v) with NaH (1 equiv) followed by *tert*-BuMeSiCl (1.2 equiv) at 4°, and purified by distillation [Hwu & Chen *e-EROS Encyclopedia of Reagents for Organic Synthesis* 15 Apr 2001, DOI: 10.1002/047084289X.rb379m]. Moisture sensitive and best transferred via gas-tight syringe. This reagent successfully silylates *N,O*-diacylhydroxylamines in MeCN [Schraml et al. *Organometallics* **23** 2157 2004, DOI: 10.1021/om030684u], and used for the determination of nitrogen mustard hydrolysis products by silylation and GC/MS [Ohsawa & Seto *J Chromatogr A* **1122** 242 2006, DOI: 10.1016/j.chroma.2006.04.076].

tert-Butyldiphenylchlorosilane (TBDPSCI, tert-butylchlorodiphenylsilane) [58479-61-1] C₁₆H₁₉ClSi, **M 274.9, b 90°/0.015mm, 335°/760mm, d**₄²⁰ **1.057, n**_D²⁰ **1.568**. Purify it by repeated fractional distillation. It is soluble in DMF and pentane [Hanessian & Lavalee *Can J Chem* **53** 2975 1975, DOI: 10.1139/v75-419; Robl et al. *J Med Chem* **34** 2804 1991, DOI: 10.1021/jm00113a019]. [Beilstein **4** IV 4076 for *tert*-butylchlorodimethylsilane.]

***n*-Butylphenyl *n*-butylphosphonate** [36411-99-1] C₁₄H₂₃O₃P, **M 270.3**. Crystallise it three times from hexane as its compound with uranyl nitrate. See *tri-n-butyl phosphate* below.

***p*-tert-Butylphenyl diphenyl phosphate** [981-40-8] C₂₂H₂₃O₄P, **M 382.4, b 261°/6mm, n**²⁵ **1.5522**. Purify it by vacuum distillation, and percolation through an alumina column, followed by passage through a packed column maintained at 150° to remove residual traces of volatile materials in a counter-current stream of N₂ at reduced pressure [Dobry & Keller *J Phys Chem* **61** 1448 1957, DOI: 10.1021/j150556a052].

Cacodylic acid (dimethylarsinic acid) [75-60-5] C₂H₇O₂As, **M 138.0, m 195-196°, pK**₁²⁵ **1.57, pK**₂²⁵ **6.27** [Me₂As(O)OH]. Recrystallise it from warm EtOH (3ml/g) by cooling and filtering. Dry it in a vacuum desiccator over CaCl₂. It has also been recrystallised twice from propan-2-ol. [Kilpatrick *J Am Chem Soc* **71** 2607 1949, DOI: 10.1021/ja01176a003; Nichol *J Am Chem Soc* **72** 2367 1950, DOI: 10.1021/ja01162a006; Koller & Hawkrige *J Am Chem Soc* **107** 7412 1985, DOI: 10.1021/ja00311a032; Beilstein **4** H 610, **4** I 567, **4** II 993, **4** III 1818, **4** IV 3681.]

Cadion [1-(4-nitrophenyl)-3-(4-phenylazophenyl)-triazene] [5392-67-6] C₁₈H₁₄N₆O₂, **M 346.3, m 189°(dec)**. Commercial cadion is purified by recrystallisation from 95% EtOH and is dried *in vacuo*. It is stable in 0.2 N KOH (in 20% aqueous EtOH) at 25°. It is a sensitive reagent for Cd, and the Cd complex has λ_{max} (EtOH) 475nm. [Chavanne & Geronimi *Anal Chim Acta* **19** 377 1958, DOI: 10.1016/S0003-2670(00)88180-5; Beilstein **16** III 664.]

(1*R*)-(-)Camphor-10-sulfonic acid [35963-20-3] C₁₀H₁₆O₄S, **M 232.3, m 197.4-198°(dec), 197-198°, [α]**_D²⁰ **-20.7 (c 5.4, H₂O), pK**_{Est} **~ -1**. It forms prisms from AcOH or EtOAc, and is deliquescent in moist air. Store it in tightly stoppered bottles. The **(1*R*)-(-)-NH₄ salt** [82509-30-6] C₁₀H₁₉NO₄S, **M 249.3**, forms needles from H₂O with **m ~250°(dec)** and [α]_D²² -18.4, [α]_D¹⁶ -20.5 (c 5, H₂O). [Burgess & Lowry *J Chem Soc* **127** 271 1925, DOI: 10.1039/CT9252700271; Marsi et al. *J Am Chem Soc* **78** 3063 1956, DOI: 10.1021/ja01594a032]. The ***RS*-acid** [5872-08-2] recrystallises from AcOH. [60g of (±)-acid in 60ml of AcOH at 105° gave 40g of crystals has **m 202-203°**]. The ***1R*-(-) acid** has been obtained by the optical resolution of the *RS*-acid as follows: To a warm solution of the (±)-acid (50.0g, 0.215 mole) in H₂O (500 mL) is added *l*-brucine powder (83.6g, 0.215 mole) in small portions. After 24 hour at 25° the salt [56.8g, [α]_D²⁴ -16.1 (c 5.4, H₂O)]. By a process of fractional crystallisation the more soluble fairly pure **(-)-brucine (-)-camphorsulfonate** (12.5g, [α]_D²⁴ -29.3) can be obtained. The salt is then dissolved in warm H₂O (195 mL), treated with Ba(OH)₂·8H₂O (3.25g, 0.01 mole),

the precipitated mixture is kept at $\sim 100^\circ$ for 2 hours, then filtered to remove the insoluble brucine (7.09g, 90%). The filtrate is treated with 0.50 M H_2SO_4 , the precipitated Ba SO_4 is filtered off and the filtrate is evaporated to dryness *in vacuo*. The yellow-brown residue is extracted with hot anhydrous C_6H_6 (4 x 24 mL) and the extract, on refrigeration, gave colourless crystals (3.35g) of ***R*-sulfonic acid, m 197.4-198°(dec)**, $[\alpha]_{\text{D}}^{24} -18.52$ (c 5.4, H_2O); a further 0.37g can be recovered from the mother liquor. ***Silver (1R)-(-)-camphor-10-sulfonate*** is readily obtained by adding freshly prepared hydrated silver oxide [from AgNO_3 (22.2g, 0.129 mole) in H_2O (40 mL) basified with NaOH (6.0g, 0.15 mole), filtered and washed well with H_2O] to the sulfonic acid (28.0g, 0.12 mole) in H_2O (200 mL), stirred for several minutes, unreacted Ag_2O is filtered off, and the filtrate is evaporated to dryness which provided the ***silver salt*** (32.0g, 79%). The ***ethylamine salt***, $\text{C}_{10}\text{H}_{16}\text{O}_3\text{S} \cdot \text{C}_2\text{H}_7\text{N}$, has **m 160-161°** (from $\text{Et}_2\text{O}/\text{EtOH}$). [Marsi et al *J Am Chem Soc* **78** 3063 1956, DOI: 10.1021/ja01594a032; Bartlett & Knox *Org Synth* **45** 12 1965, DOI: 10.15227/orgsyn.045.0012; Beilstein **11** II 182, **11** III 584, **11** IV 642.]

(1S)-(+)-Camphor-10-sulfonic acid [3144-16-9] $\text{C}_{10}\text{H}_{16}\text{O}_4\text{S}$, **M 232.3**, **m 193°(dec)**, **197-198°**, $[\alpha]_{\text{D}}^{20} +27.5$ (c 10, H_2O), $[\alpha]_{\text{D}}^{20} +43.5$ (c 4.3, EtOH), **pK_{Est} ~ -1** . Crystallise the acid from ethyl acetate and dry it under vacuum (deliquescent). [Loudon *J Chem Soc* 823 1933, DOI: 10.1039/JR9330000819; Komppa *J Prakt Chem* **162** 19 1943, DOI: 10.1002/prac.19431620103; Beilstein **11** IV 642.] It can also be isolated from the less soluble ***(-)-brucine (+)-camphorsulfonate salt*** in the above optical resolution. See above for *RS*-isomer. The ***(1S)-(+)-NH₄ salt*** [82509-30-6] $\text{C}_{10}\text{H}_{19}\text{NO}_4\text{S}$, **M 249.3**, forms needles from H_2O with **m $\sim 250^\circ$ (dec)** and $[\alpha]_{\text{D}}^{22} +18.4$ (c 5, H_2O).

Camphor-10-sulfonyl chloride [*1S*-(+)- 21286-54-4, *1R*-(-)- 39262-22-1, (\pm)- 4552-50-5] $\text{C}_{10}\text{H}_{15}\text{ClO}_3\text{S}$, **M 250.7**, has **m 67-68°, 70°, 69-71°**, $[\alpha]_{\text{D}}^{20}$ (+) and (-) **32.2** (c 3, CHCl_3). If free from OH bands in the IR, then recrystallise it from Et_2O or petroleum ether; otherwise treat it with SOCl_2 at 50° for 30 minutes, evaporate, dry the residue over KOH in a vacuum and recrystallise it. The ***(\pm)-acid chloride*** [4552-50-5] has **m 85° (66-68°)** [Bartlett & Knox *Org Synth* **45** 14 1965, DOI: 10.15227/orgsyn.045.0014]. They are characterised as their ***amides*** (by reaction with $\text{NH}_4\text{OH}/0-10^\circ/1\text{hr}$ then $25^\circ/4\text{hr}$ and extraction into CH_2Cl_2 and evaporation) [*S* +, 60933-63-3; and *R* -, 72597-34-3], $\text{C}_{10}\text{H}_{17}\text{NO}_3\text{S}$, **M 231.3**, which form prisms from EtOH with **m 132°**, and $[\alpha]_{\text{D}}^{20}$ + and -22 and (c 1, MeOH). On repeated recrystallisation from EtOH the ***anilides*** have **m 120.5-121°**, and $[\alpha]_{\text{D}}^{25} +76$ or -76 (c 1, CHCl_3). [Read & Storey *J Chem Soc* 2761 1930, DOI: 10.1039/JR9300002761; Sutherland & Shriner *J Am Chem Soc* **58** 62 1936, DOI: 10.1021/ja01292a019; Halterman et al. *J Am Chem Soc* **109** 8105 1987, DOI: 10.1021/ja00260a037; Bartlett & Knox *Org Synth* **45** 55 1945, DOI: 10.15227/orgsyn.045.0055; Coll Vol **5** 196 1973, DOI: 10.15227/orgsyn.045.0014; Towson et al. *Org Synth Coll Vol* **8** 104 1993, DOI: 10.15227/orgsyn.069.0158; and Beilstein **11** IV 650.]

(4-Carbamylphenylarsylenedithio)diacetic acid (Arsenamide) [531-72-6] $\text{C}_{11}\text{H}_{12}\text{NO}_5\text{S}_2\text{As}$, **M 377.3**, **m 158-162°, 168-169°, pK 4.0**. Recrystallise it from H_2O (1.7g/25ml, containing a small amount of thiolacetic acid), MeOH or EtOH . It is insoluble in *iso*- PrOH . The ***disodium salt*** is obtained by titrating with NaOH to pH 7-8. [Maren *J Am Chem Soc* **68** 1864, 1946, DOI: 10.1021/ja01213a504; Gough & King *J Chem Soc* 669 1930, DOI: 10.1039/JR9300000669.] It has chemotherapeutic activity against *filaria* and *trichomonas*, and is used in the treatment of heart worm in dogs.

Catecholborane (1,3,2-Benzodioxaborole) [274-07-7] $\text{C}_6\text{H}_5\text{O}_2\text{B}$, **M 119.2**, **b 50°/50mm, 66°/80mm, 76-77°/100mm, 88°/165mm, d₄²⁰ 1.125, n_D²⁰ 1.507** (also available as a 1.0M solution in THF or toluene). It is a moisture-sensitive flammable liquid which is purified by distillation in a vacuum under a N_2 atmosphere and stored under N_2 at $0-4^\circ$. It liberates H_2 when added to H_2O or MeOH . A solution in THF, after 25 hours at 25° , has residual hydride of 95% (under N_2) and 80% (under air) [Brown & Gupta *J Am Chem Soc* **97** 5249 1975, DOI: 10.1021/ja00851a038].

Calconcarboxylic acid [3-hydroxy-4-(2-hydroxy-4-sulfo-1-naphthylazo)naphthalene-2-carboxylic acid] [3737-95-9] $\text{C}_{21}\text{H}_{14}\text{N}_2\text{O}_7\text{S}$, **M 428.4**, **m 300°, λ_{max} 560nm, pK₁ 1.2, pK₂ 3.8, pK₃ 9.26, pK₄ 13.14**. Purify it via its ***p-toluidinium salt***, viz: dissolve the dye in warm 20% aqueous MeOH , treat with an equivalent of *p*-toluidine and cool to precipitate the salt. Finally recrystallise the acid from hot water. [Itoh & Ueno *Analyst (London)* **95** 583 1970, DOI: 10.1039/AN9709500583.] It complexes with Ca^{2+} in presence of Mg^{2+} and other metal ions. It is thus used as an indicator for titrations of Ca^{2+} , which when EDTA is added, is unaffected by the presence of Mg^{2+} and other metal ions which are masked. [Patton & Reeder *Anal Chem* **28** 1026 1956,

DOI: 10.1021/ac60114a029; Prentoe & Prentoe *Analyst* **106** 227 1981, DOI: 10.1039/AN9810600227.]

Calmagite [1-(1-hydroxy-4-methyl-2-phenylazo)-2-hydroxynaphthalene-4-sulfonic acid] [3147-14-6] $C_{17}H_{14}N_2O_5S$, **M 358.4**, **m 300°**, **pK₁ 8.1**, **pK₂ 12.4**. A crude dye is extracted with anhydrous diethyl ether and forms red crystals from Me_2CO . It gives a red colour in H_2O at pH 7–9 and a blue colour at pH 9–11 which turns red on addition of Ca^{2+} or Mg^{2+} ions. [Lindstrom & Diehl *Anal Chem* **32** 1123 1960, DOI: 10.1021/ac60165a022]. Complexes with Ca, Mg and Th, and is used as an indicator for titrating Ca or Mg ions with EDTA.

Chloramine-T (*N*-chloro-*p*-toluenesulfonamide sodium salt) $3H_2O$ [7080-50-4] $C_7H_7ClNNaO_2S \cdot 3H_2O$, **M 281.7**, **m 168–170°(dec)**. Recrystallise it from hot water (2ml/g). Dry it in a desiccator over $CaCl_2$ where it loses water. Protect it from sunlight. It is used for the detection of bromate and halogens, and Co, Cr, Fe, Hg, Mn, Ni and Sb ions. [Campbell & Johnson *Chem Rev* **78** 65 1978, DOI: 10.1021/cr60311a005; Bremner *Synthetic Reagents* **6** 9 1985, Chattaway **87** 145 1905, DOI: 10.1039/CT9058700145; for selective oxidation of methionine see Trout *Anal Biochem* **93** 419 1979, DOI: 10.1016/S0003-2697(79)80173-6; Inglis *J Soc Chem Ind* **37** 288 1918, DOI: 10.1002/jctb.5000371401; *Beilstein* **11** H 107, **11** I 29, **11** II 62, **11** III 300, **2** IV 457.]

Chlorazol Sky Blue FF {Chicago Sky Blue 6B, Direct Blue 1, 6,6'-[(3,3'-dimethoxy[1,1'-biphenyl]-4,4'-diyl)bis(azo)bis(4-amino-5-hydroxy-1,3-naphthylenedisulfonic acid) tetra-Na salt [2610-05-1] $C_{34}H_{24}N_6Na_4O_{16}S_4$, **M 996.9**, **m >300°(dec)**. Free it from other electrolytes by adding aqueous sodium acetate to a boiling solution of the dye in distilled water. After standing, the salted-out dye is filtered on a Büchner funnel, the process being repeated several times. Finally, the precipitated dye is boiled several times with absolute EtOH to wash out any sodium acetate, then dried (as the sodium salt) at 105°. [McGregor et al. *Trans Faraday Soc* **58** 1045 1962, DOI: 10.1039/TF9625801045; *Beilstein* **16** I 346, **16** II 259.]

4-Chlorobenzenesulfonic acid [98-66-8] $C_6H_5ClO_3S$, **M 192.6**, **m 67°**, **b 149°/22mm**, **pK²⁵ < 0**. It is prepared by the sulfonation of chlorobenzene with Oleum and recrystallises from H_2O as a *hydrate*. The *anhydrous* acid is obtained on distillation, preferably under reduced pressure. [Cook & Cook *J Am Pharm Assoc* **38** 239 1949, DOI: 10.1002/jps.3030380503; *Beilstein* **11** H 63.]

4-Chlorobenzenesulfonyl chloride [98-60-2] $C_6H_4Cl_2O_2S$, **M 211.1**, has **m 53°**, **b 141°/15mm**. Crystallise it from ether in powdered Dry-ice, after the solution has been washed with 10% NaOH until colourless and dried (Na_2SO_4). Distil it *in vacuo* and store it in the absence of H_2O . **IRRITANT**. [*Beilstein* **11** IV 114.] **4-Chlorobenzenesulfonamide** [98-64-6] $C_6H_6ClNO_2S$, **M 191.6**, **m 144°**, is obtained by reaction of the sulfonyl chloride with aqueous 088 NH_3 and crystallised from H_2O . [*Beilstein* **11** IV 115.]

S-4-Chlorobenzylisothiuronium chloride [544-47-8] $C_8H_{10}ClN_2S$, **M 237.1**, **m 177–178°**, and **197°**, **201–203°**, **pK_{Est} ~9.6** (free base). Crystallise the salt from concentrated HCl by addition of water (1:1). Dry it in a vacuum over P_2O_5 . Also crystallise it from EtOH, wash the crystals with EtOH, then Et_2O to give the lower melting form **m 177–178°**. By evaporating the filtrate and washings to a quarter of the volume and adding an equal volume of Et_2O the higher melting form **m 201–203°** is obtained. [Harvey & Jensen *J Org Chem* **28** 470 1963, DOI: 10.1021/jo01037a047; *Beilstein* **6** III 1639, **6** IV 2778.] Its preparation and use as a reagent for characterising acids by forming salts which are best recrystallised from dioxane is described by Dewey and Sperry [*J Am Chem Soc* **61** 3251 1939, DOI: 10.1021/ja01267a005]. Both forms provide the same derivatives.

Chlorodicyclohexylphosphine [Cy_2PCl] [16523-54-9] $C_{12}H_{22}ClP$, **M 232.7**, **b 132–138°/3mm**, **165°/12mm**, **173–174°/17mm**, **182–183°/23mm**, **d₄²⁵ 1.054**, **n_D²⁰ 1.533**. Cy_2PCl can be obtained as a colourless oil from cyclohexyldichloro-phosphine and cyclohexylmagnesium chloride, also from dicyclohexyldiethylamino-phosphine and HCl in the presence of NH_4Cl in petroleum ether (b 70–90°) followed by fractional distillation. Alternatively, reaction of cyclohexylmagnesium chloride (from 12.6g of Mg and 62g of cyclohexyl chloride) in Et_2O (250ml) and PCl_3 (35g) in Et_2O (300ml) under N_2 followed by fractional distillation gives (22g, 37.2%) of Cy_2PCl . [Issleib & Seidel *Chem Ber* **92** 2681 1959, DOI: 10.1002/cber.19590921102; *Beilstein* **16** IV 968.]

Chlorodiphenylphosphine (diphenylphosphinous chloride) [1079-66-9] $C_{12}H_{10}ClP$, **M 220.6**, **m 15–16°**, **b 124–126°/0.6mm**, **174°/5mm**, **320°/atm**, **d₄²⁰ 1.229**, **n_D²⁰ 1.636**. This air-sensitive, pale yellow lachrymatory

liquid is purified by careful fractional distillation and discarding the lower boiling fraction which contains the main impurity PhPCl_2 (b 48-51°/0.7mm), and checking for impurities by NMR. [Weinberg *J Org Chem* **40** 3586 1975, DOI: 10.1021/jo00912a027; Horner et al. *Chem Ber* **94** 2122 1961, DOI: 10.1002/cber.19610940826.]

Chlorodi(o-tolyl)phosphine [36042-94-1] $\text{C}_{14}\text{H}_{14}\text{ClP}$, M 248.7, m 36-37°, b 120-122°/0.03mm, 146-147°/1.1mm. It is purified by fractional distillation in a vacuum (b 179-183°/7mm, 253-257°/15mm) and the distillate solidifies (m 36°, also reported is m 37°). [Weinberg *J Org Chem* **40** 3586 1975, DOI: 10.1021/jo00912a027; McEwen et al. *J Am Chem Soc* **100** 7304 1978, DOI: 10.1021/ja00491a030; *Beilstein* **16** H 769, **16** IV 970 for chlorodi(p-tolyl)phosphine.]

(Chloromethyl)dimethylvinylsilane [16709-86-7] $\text{C}_5\text{H}_{11}\text{ClSi}$, M 134.7, b 121-122°/760mm, 122-126°/atm, d_4^{25} 0.908, n_D^{20} 1.440. Distil the silane in a vacuum, but if it is suspect then dissolve it in Et_2O , shake it with saturated aqueous NH_4Cl , dry the Et_2O layer (anhydrous Na_2SO_4), filter, evaporate and fractionate in a vacuum. [Altamura et al. *J Org Chem* **60** 8403 1995, DOI: 10.1021/jo00131a015.]

Chloromethyl phenyl sulfide [7205-91-6] $\text{C}_7\text{H}_7\text{ClS}$, M 158.7, b 63°/0.1mm, 98°/12mm, 113-115°/20mm, d_4^{20} 1.184, n_D^{20} 1.5950. Dissolve the sulfide in CH_2Cl_2 or CCl_4 and dry it (CaCl_2), or pass it through a tube of CaCl_2 and distil it using a fractionating column. **Harmful vapours.** It gives the **sulfone** [7205-98-3] $\text{C}_7\text{H}_7\text{ClO}_2\text{S}$, M 190.7, which has b 130°/1mm and m 53° (from EtOH) on oxidation with permonophthalic acid. [*Beilstein* **6** IV 1507.] [Böhme et al. *Justus Liebigs Ann Chem* **563** 54 64 1949, DOI: 10.1002/jlac.19495630107.] [*Beilstein* **6** III 1002.] The **sulfoxide** [7205-94-9] $\text{C}_7\text{H}_7\text{ClOS}$, M 174.7, has b 109-111°/0.011mm, is a useful reagent for the synthesis of alkyl sulfoxides [Hojo et al. *Synthesis* 789 1977, DOI: 10.1055/s-1977-24581], and can be used as a thiol ester acyl anion equivalent [More & Wimple *J Org Chem* **43** 2713 1978, DOI: 10.1021/jo00407a040].

Chloromethylphosphonic acid dichloride [1983-26-2] $\text{CH}_2\text{Cl}_2\text{OP}$, M 167.4, b 50°/0.5mm, 52-53(59)°/2mm, 63-65°/3mm, 78-79°/10mm, 87-88°/15mm, 102-103°/30mm, d_4^{20} 1.638, n_D^{20} 1.4971. It is fractionally distilled using a short Claisen column and redistilled. The **aniline salt** has m 199-201°. The ^{31}P NMR has a single peak at -38 ± 2 ppm from 85% H_3PO_4 . [Kinnear & Perren *J Chem Soc* 3437 1952, DOI: 10.1039/JR9520003437; NMR: van Wazer et al. *J Am Chem Soc* **78** 5715 1956, McConnell et al. *J Org Chem* **22** 462 1957, DOI: 10.1021/jo01355a619; *Beilstein* **1** III 2593, **1** IV 3068.]

2-Chloro-2-oxo-1,3,2-dioxaphospholane [6609-64-9] $\text{C}_2\text{H}_4\text{ClO}_3\text{P}$, M 142.5, m 12°, 14°, 12-14°, b 89-91°/0.8mm, 88-89°/2mm, 303°/760mm, d_4^{20} 1.549, n_D^{20} 1.448. It should be distilled under high vacuum as some polymerisation occurs at atmospheric pressure. It has IR bands at 3012, 2933, 1477, 1366, 1325, 1040, 924 and 858 cm^{-1} . It is hydrolysed to $\text{HOCH}_2\text{CH}_2\text{OPO}_3\text{H}_2$ in 30 minutes in H_2O at 100° [IR: Cox & Westheimer *J Am Chem Soc* **80** 5441 1958, DOI: 10.1021/ja01553a031]. [*Beilstein* **1** IV 2419.]

Chlorophenol Red (3,3'-dichlorophenolsulfonephthalein) [4430-20-0] $\text{C}_{19}\text{H}_{12}\text{Cl}_2\text{O}_5\text{S}$, M 423.3, m dec on heating, λ_{max} 573nm, pK^{25} 5.96. Crystallise the dye from glacial acetic acid. It is an indicator which is yellow at pH 4.8 and violet at pH 6.7. [*Beilstein* **19/3** V 458.]

2-Chlorophenyl diphenyl phosphate [115-85-5] $\text{C}_{18}\text{H}_{14}\text{ClO}_4\text{P}$, M 360.7, b 236°/4mm, 413.9°/760mm, n_D^{25} 1.5707. Purify it by vacuum distillation, percolate it through a column of alumina, then pass it through a packed column maintained by a countercurrent stream of N_2 at reduced pressure [Dobry & Keller *J Phys Chem* **61** 1448 1957, DOI: 10.1021/j150556a052].

Chlorosulfonic (chlorosulfuric) acid [7790-94-5] HClO_3S , M 116.5, m -80°, b 60-64°/2-4mm, 74-75°/19mm, 151-152°/750mm, 151°/760mm, d_4^{20} 1.753, n_D^{25} 1.4929, pK^{25} -5.9 (aqueous H_2SO_4). Distil it in an all-glass apparatus, taking the fraction boiling at 156-158°/atm, or under reduced pressure. The colourless to pale yellow distillate has a pungent odour, fumes in air and reacts **EXPLOSIVELY** with water [Kaplan et al. *Inorg Synth* **4** 52 1953, DOI: 10.1002/9780470132357.ch17; Cremlyn *Chlorosulfonic acid: A Versatile Reagent*, Royal Society of Chemistry UK, 2002, 308 pp, ISBN 0854044981]. **LACHRYMATORY, CORROSIVE, attacks the eyes and skin, wear gloves and face shield, work in an efficient fume cupboard.**

Chlorotriphenylsilane (TPSCl, triphenylchlorosilane) [76-86-8] $C_{18}H_{15}ClSi$, *M* 294.9, *m* 90-92°, 91-93°, 91-96°, 94-95°, 97-99°, *b* 156°/1mm, 161°/0.6mm, 188-189°/3mm. Likely impurities are tetraphenylsilane, small amounts of hexaphenyldisiloxane and traces of triphenylsilanol. Purify it by distillation at low pressure, then crystallise it from EtOH-free $CHCl_3$, and from petroleum ether (*b* 30-60°) or hexane by cooling in a Dry-ice/acetone bath. It is moisture sensitive. [Allen & Modena *J Chem Soc* 3671 1957, DOI: 10.1039/JR9570003671; Curran et al. *J Am Chem Soc* 72 4471 1950, DOI: 10.1021/ja01166a038; Speier & Zimmerman *J Am Chem Soc* 77 6395 1955, DOI: 10.1021/ja01628a110; Thomas & Rochow *J Am Chem Soc* 79 1843 1957, DOI: 10.1021/ja01565a021; *Beilstein* 16 IV 1484.]

Chromeazurol S (Mordant Blue 29) [1667-99-8] $C_{23}H_{13}Cl_2Na_3O_9S$, *M* 605.3, λ_{max} 540nm, ϵ 7.80×10^4 (10M HCl), *CI* 43825, $pK_1^{25} < 0$, $pK_2^{25} 2.25$, $pK_3^{25} 4.88$, $pK_4^{25} 11.75$. The crude *phenolic triphenylmethanecarboxysulfonic acid tri-Na salt* (40g) is dissolved in water (250ml) and filtered. Then concentrated HCl (50ml) is added to the filtrate, with stirring. The precipitate is filtered off, washed with HCl (2M) and dried. It is redissolved in water (250ml), and precipitation is repeated twice more in a water bath at 70°. It is dried under vacuum over solid KOH (first), then P_2O_5 [Martynov et al. *Zh Analyt Khim* 32 519 1977]. It has also been purified by paper chromatography using *n*-butanol, acetic acid and water (7:3:1). First and second spots were extracted. It chelates Al and Be. It is used also for estimating fluoride. [*Beilstein* 11 IV 707.]

Congo Red (4B) {Cosmos Red, Cotton Red B, 3,3,'-[(1,1'-biphenyl)-4,4'-diylbis(azo)]bis[4-amino-1-naphthaleneulfonic acid] disodium salt} [573-58-0] $C_{32}H_{22}N_6Na_2O_6S_2$, *M* 696.7, *m* >360°, λ_{max} 488nm ($E_{1\%}^{1cm}$ 595) at pH 7.3, $pK_2^{28} 4.19$. Crystallise the dye from aqueous EtOH (1:3), and dry it in air. Its colour is yellow-red in H_2O and orange in EtOH. It is a useful biological stain. [*Beilstein* 6 I 342.]

Copper (I) thiophenolate [1192-40-1] C_6H_5CuS , *M* 172.7, *m* ca 280°, $pK_1^{25} 6.62$ (for PhS^-). The Cu salt can be extracted from a thimble (Soxhlet) with boiling MeOH. It is a green-brown powder that gives a yellow-green solution in pyridine. Wash it with EtOH and dry it in a vacuum. It can be precipitated from a pyridine solution by adding H_2O , collecting the precipitate, washing it with EtOH and drying in a vacuum. [Posner et al. *Synthesis* 662 1974, DOI: 10.1055/s-1974-23397; Krebs et al. *Chem Ber* 90 425 DOI: 10.1002/cber.19570900321; 1957, *Beilstein* 6 IV 1465.] It is a useful reagent for preparing Cu-*N*-heterocyclic carbenes [Cisnetti et al. *Tetrahedron Lett* 51 5226 2010, DOI: 10.1016/j.tetlet.2010.07.124].

***o*-Cresol Red [*o*-cresolphthalein, 3,4-benz-5,5-bis-(4-hydroxy-2-methylphenyl)-1(3*H*)-oxa-2-thiole 2,2-dioxide]** [1733-12-6] $C_{21}H_{18}O_5S$, *M* 382.4, *m* 290°(dec), λ_{max} 367nm and 570nm, $pK_1^{25} 1.26$, $pK_2^{25} 8.18$. Crystallise the reddish-brown powder from glacial acetic acid. Dry it in air. Alternatively, dissolve it in aqueous 5% $NaHCO_3$ solution and precipitate it from a hot solution by dropwise addition of aqueous HCl. Repeat the procedure until the UV maxima do not increase. It is an acid-base indicator volumetric analysis with two colour changes *viz.*: pH range 1.8–2.0 from Orange (acid) to Yellow, and at pH range 7.0–8.8 from Yellow to Violet (alkaline) (cf: pK values). [*Beilstein* 19 IV 1133.] The *sodium salt* [62625-29-0] $C_{21}H_{17}NaO_5S$, *M* 404.4, λ_{max} 425nm, has *m* 250°(dec).

Cupferron ammonium salt (*N*-nitroso-*N*-phenylhydroxylamine ammonium salt) [135-20-6] $C_6H_9N_3O_2$, *M* 155.2, *m* 150-155°(dec), 162.5-163.5°, 163-164°, $pK^{25} 4.16$ (free base). Recrystallise it twice from EtOH after treatment with Norite and finally once with EtOH. The crystals are washed with diethyl ether and air dried, then stored in the dark over solid ammonium carbonate. A standard solution (ca 0.05M prepared in air-free H_2O) is prepared daily from this material for analytical work and is essentially 100% pure. [Olsen & Elving *Anal Chem* 26 1747 1954, DOI: 10.1021/ac60095a017.] It can also be washed with Et_2O , dried and stored as stated. In a sealed, dark container it can be stored for at least 12 months without deterioration. The UV has λ_{max} at 260nm ($CHCl_3$). [Marvel *Org Synth Coll Vol* 1 177 1941, DOI: 10.15227/orgsyn.004.0019; Elving & Olson *J Am Chem Soc* 78 4206 1956, DOI: 10.1021/ja01598a006; *Beilstein* 16 IV 891.] Possible **CARCINOGEN**.

Cupric trifluoromethylsulfonate (copper II triflate) [34946-82-2] $C_2CuF_6O_6S_2$, *M* 361.7, *m* >300°, $pK^{25} < 3.0$ (for triflic acid). Dissolve it in MeCN, add dry Et_2O until cloudy and cool at -20° in a freezer. The light blue precipitate is collected and dried in a vacuum oven at 130°/20mm for 8 hours. It has λ_{max} at 737nm (ϵ 22.4 $M^{-1}cm^{-1}$) in AcOH. [Jenkins & Kochi *J Am Chem Soc* 94 843 1972, DOI: 10.1021/ja00758a024; Salomon &

Kochi *J Am Chem Soc* **95** 3300 1973, DOI: 10.1021/ja00791a038]. It was prepared from CuCO_3 in MeCN and $\text{CF}_3\text{CO}_2\text{H}$, and the **blue salt** has also been dried in a vessel at 0.1 Torr by heating with a Fischer burner [Andrist et al. *J Org Chem* **43** 3422 1978, DOI: 10.1021/jo00411a047]. It has been dried at 110-120°/5mm for 1 hour before use and forms a ***benzene complex** which should be handled in a dry box because it is air sensitive [Kobayashi et al. *Chem Pharm Bull Jpn* **28** 262 1980, DOI: org/10.1248/cpb.28.262; Salomon & Kochi *J Am Chem Soc* **95** 3300 1973, DOI: 10.1021/ja00791a038]. [Beilstein **3** IV 34.]

Cuprous (I) bromide dimethylsulfide complex [54678-23-8] $\text{C}_2\text{H}_6\text{BrCuS}$, **M 205.6, m ca 135°(dec)**. Purify it by recrystallisation in the presence of Me_2S . A solution of the complex (1.02g) in Me_2S (5ml) is slowly diluted with hexane (20ml), and the pure colourless prisms of the complex (0.96g) separate and are collected and dried, **m 124-129°(dec)**. The complex is insoluble in hexane, Et_2O , Me_2CO , CHCl_3 and CCl_4 . It dissolves in DMF and DMSO, but the solution becomes hot and green indicating decomposition. It dissolves in $^*\text{C}_6\text{H}_6$, Et_2O , MeOH and CHCl_3 if excess of Me_2S is added and a colourless solution is obtained. [House et al. *J Org Chem* **40** 1460 1975, DOI: 10.1021/jo00898a019.] Prior to use, the complex is dissolved in Me_2S and evaporated to dryness in the weighed reaction flask [Bourgain-Commerçon et al. *J Organomet Chem* **228** 321 1982, DOI: 10.1016/S0022-328X(00)84333-8]. If this reagent is to assist in methylations with MeMgBr (e.g. of fullerenes) it is always best to make a fresh preparation. CuBr (25g) is washed with MeOH (4 x 50ml) to remove coloured impurities and dried *in vacuo* for 1 hour. This white green-tinged CuBr is dissolved in Me_2S (60ml, redistilled b 38°, **use efficient fume cupboard**) and insoluble impurities are filtered off. Hexane (200ml) is added and the precipitated light-sensitive white crystalline complex is filtered off and washed with hexane (5 times) under suction and N_2 . It should be dried under a stream of dry N_2 , and should not be kept or stored under some pressure as it will lose the some of the disulfide ligand and its efficiency. [Wuts *Synth Commun* **2** 139 1981, DOI: 10.1080/00397918108064294; Matsuo et al. *Org Synth* **83** 80, 2006, DOI: 10.15227/orgsyn.083.0080.]

Cuprous iodide trimethylphosphite [34836-53-8] $\text{C}_3\text{H}_9\text{CuIO}_3\text{P}$, **M 314.5, has m 175-177°, 192-193°**. Cuprous iodide dissolves in a $^*\text{C}_6\text{H}_6$ solution containing trimethylphosphite to form the **complex**. The **complex** crystallises from $^*\text{C}_6\text{H}_6$ or petroleum ether. [Arbusoff *Chem Ber* **38** 1171 1905, DOI: 10.1002/cber.190503801212; Nishizawa *Bull Chem Soc Jpn* **34** 1170, 1177 1961, DOI: org/10.1246/bcsj.34.1170.] It is useful for preparing Cu-enolates [Ziegler & Fang *J Org Chem* **46** 825 1981, DOI: 10.1021/jo00317a042] and for the alkylation of trimethylsilylvinyl aluminates [Ziegler & Mikami *Tetrahedron Lett* **25** 131 1984, DOI: 10.1016/S0040-4039(00)99821-6].

Cyclohexyl mercaptan (cyclohexane thiol) [1569-69-3] $\text{C}_6\text{H}_{12}\text{S}$, **M 116.2, b 38-39°/12mm, 57°/23mm, 90°/100mm, 157°/763mm, d_4^{20} 0.949, n_D^{20} 1.493, pK_{Est} ~10.8**. Possible impurities are the sulfide and the disulfide. Purify the thiol by conversion to the Na salt by dissolving it in 10% aqueous NaOH, extract the sulfide and disulfide with Et_2O , and then acidify the aqueous solution (with cooling and under N_2) with HCl, extract with Et_2O , dry over MgSO_4 , evaporate and distil it in a vacuum (**b 41°/12mm**). The **sulfide** has **b 74°/0.2mm, $n_D^{18.5}$ 1.5162** and the **disulfide** has **b 110-112°/0.2mm, $n_D^{18.5}$ 1.5557**. The **Hg-mercaptide** has **m 77-78°** (needles from EtOH). [Naylor *J Chem Soc* 1532 1947, DOI: 10.1039/JR9470001532; Beilstein **6** H 8, **6** I 6, **6** II 14, **6** III 46, **6** IV 72.]

Decaborane (14) ($\text{B}_{10}\text{H}_{14}$) [17702-41-9] $\text{B}_{10}\text{H}_{14}$, **M 122.2, m 99.6-99.7°, 99.7-100°, b 100°/19mm, 213°/atm, d_4^{25} 0.94**. Purify decaborane by vacuum sublimation at 80°/0.1mm, followed by crystallisation from methylcyclohexane, CH_2Cl_2 , or dry olefin-free-*n*-pentane, the solvent being subsequently removed by storing the crystals in a vacuum desiccator containing CaCl_2 , and is stable at ~25° indefinitely. It is soluble in H_2O but is slowly decomposed to give H_2 . It is soluble in alkali, and on acidification it liberates H_2 . **TOXIC, excessive exposure causes headache, dizziness, muscle spasm and nausea**. [Greenwood in *Comprehensive Chemistry* (Ed Bailer et al.) Pergamon Press Vol **1** pp 818-837 1973.]

4,4-Diaminodiphenyl sulfide (4,4'-thioaniline) [139-65-1] $\text{C}_{12}\text{H}_{12}\text{S}$, **M 216.3, m 108-109°, pK^{20} 2.28 (1:1 EtOH/ H_2O)**. It separates as needles from EtOH and is a **possible mutagen**. The free base is used for the detection of NO_3^- ions. The **diacetate** crystallises from aqueous AcOH with **m 182°** and the **sulfoxide**, [119-59-5] **m 184°**, forms prisms from EtOH or H_2O . [Fuson & Melamed *J Org Chem* **13** 690 1948, DOI: 10.1021/

jo01163a012; *Beilstein* **13** III 1246, **13**, IV 1306.]

Di-*n*-amyl *n*-amylphosphonate [6418-56-0] $\text{C}_{15}\text{H}_{33}\text{O}_3\text{P}$, *M* 292.4, *b* 150-151°/2mm, n_{D}^{20} 1.4378. Purify it by three crystallisations of its *uranyl nitrate complex* from hexane (see *tributyl phosphate*). It extracts Zr^{2+} from NaCl solutions.

Dibenzyl disulfide [150-60-7] $\text{C}_{14}\text{H}_{14}\text{S}_2$, *M* 246.4, *m* 69°, 71-72°, 74-75°, *b* 142-148°/0.05-0.1mm, 210-216°/18mm. Crystallise the disulfide from EtOH (*m* 77°), petroleum ether or CS_2 (*m* 72°) or distil it. The AgNO_3 complex has *m* 103°. [*Beilstein* **6** H 465, **6** I 229, **6** II 437, **6** III 1635, **6** IV 2760.] It is a rubber antioxidant.

Dibenzyl sulfide [538-74-9] $\text{C}_{14}\text{H}_{14}\text{S}$, *M* 214.3, has *m* 44°, 47°, 48.5°, 50°. Crystallise the sulfide from EtOH/water (10:1), or repeatedly from Et_2O . It has also been purified by chromatography on Al_2O_3 (pentane as eluent), then recrystallised from EtOH [Kice & Bowers *J Am Chem Soc* **84** 2390 1962, DOI: 10.1021/ja00871a023]. Dry in a vacuum at 30° over P_2O_5 , fuse under N_2 and re-dry. [*Beilstein* **6** IV 2649.]

Di-*n*-butyl boron triflate (di-*n*-butylboryl trifluoromethanesulfonate) [60669-69-4] $\text{C}_9\text{H}_{18}\text{BF}_3\text{O}_3\text{S}$, *M* 274.1, *b* 37°/0.12mm, 60°/2mm, $\text{pK}^{25} < -3.0$ (for triflic acid). Distil it in a vacuum under argon and store it under argon. It should be used within 2 weeks of purchase or after redistillation. Use a short path distillation system. It has IR bands in CCl_4 with ν_{max} at 1405, 1380, 1320, 1200 and 1550cm^{-1} , and ^{13}C NMR (CDCl_3) with δ at 118.1, 25.1, 21.5 and 13.6. [Gage & Evans *Org Synth* **68** 83 1990, DOI: 10.15227/orgsyn.068.0083; Evans et al. *J Am Chem Soc* **103**, 3099 1981, DOI: 10.1021/ja00401a031.] **TOXIC.**

Di-*n*-butyl *n*-butylphosphonate [78-46-6] $\text{C}_8\text{H}_{18}\text{O}_3\text{P}$, *M* 250.3, *b* 150-151°/10mm, 160-162°/20mm, *b* 283°/atm, d_4^{25} 0.946, n_{D}^{25} 1.4302. Purify by three recrystallisations of its compound with *uranyl nitrate*, from hexane. For method, see *tributyl phosphate*. It has been used as an extraction solvent. [*Beilstein* **4** III 1782.]

Di-*n*-butyl cyclohexylphosphonate [1085-92-3] $\text{C}_{14}\text{H}_{25}\text{O}_3\text{P}$, *M* 272.3. The compound with *uranyl nitrate* is recrystallised three times from hexane. For method see *tributyl phosphate*.

Di-*tert*-butyl dichlorosilane (DTBCl₂) [18395-90-9] $\text{C}_8\text{H}_{18}\text{Cl}_2\text{Si}$, *M* 213.2, *m* -15°, *b* 190°/729mm, 195-197°/atm, d_4^{20} 1.01, n_{D}^{20} 1.457. Purify it by fractional distillation. It is a colourless liquid with a pleasant odour and does not fume in moist air, but does *not* titrate quantitatively with excess of dilute alkali. [Tyler et al. *J Am Chem Soc* **70** 2876 1948, DOI: 10.1021/ja01189a012.]

Di-*tert*-butylphosphine [819-19-2] $\text{C}_8\text{H}_{19}\text{P}$, *M* 146.2, *m* -1°, *b* 34-35°/2mm, 38-40°/13mm, 165°/760mm, d_4^{25} 0.951, $\text{pK}_{\text{Est}} \sim 8.7$. It is prepared by reaction of *tert*-butylmagnesium bromide with PCl_3 in the presence of LiAlH_4 , and purified by distillation in a vacuum. *Alternatively*, the intermediate *di-tert-butylchlorophosphine* [13716-10-4] is reduced separately and the product is purified by fractional distillation *in vacuo*. [Hoffmann & Schellenbeck *Chem Ber* **99** 1134 1966, DOI: 10.1002/cber.19660990408; Hoffmann & Schellenbeck *Chem Ber* **100** 692 1967, DOI: 10.1002/cber.19671000241; Crofts et al. *J Chem Soc C* 332 1970, DOI: 10.1039/J39700000332.] **Flammable and pyrophoric.**

Di-*tert*-butyl silyl bis(trifluoromethanesulfonate) [85272-31-7] $\text{C}_{10}\text{H}_{18}\text{F}_6\text{O}_6\text{S}_2\text{Si}$, *M* 440.5, *b* 73.5-74.5°/0.35mm, d_4^{20} 1.36, n_{D}^{20} 1.398, (see pK for triflic acid). Purify it by fractional distillation at high vacuum. It is a pale yellow liquid that should be stored under argon. It is less reactive than the diisopropyl analogue. The presence of the intermediate monochloro compound can be detected by ^1H NMR, (CHCl_3): *tert-Bu₂Si(OTf)₂* [δ 1.25s], but impurities have δ 1.12s for *tert-Bu₂Si(H)OTf* and δ 1.19s for *tert-Bu₂HSi(Cl)OTf*. [Corey & Hopkins *Tetrahedron Lett* **23** 4871 1982, DOI: 10.1016/S0040-4039(00)85735-4; Deslongchamps *Aldrichimica Acta* **17** 72 1984.] Used as a protecting group (Mukaiyama) reagent for e.g. 1,3-diols [Kimura et al. *Synlett* 2379 2006, DOI: 10.1055/s-002-5130; Hillaert & Van Calenberg *Org Lett* **7** 5769 2005, DOI: 10.1021/ol052335x]. **TOXIC.**

Dichloramine-T (*N,N*-dichloro-*p*-toluenesulfonamide) [473-34-7] $\text{C}_7\text{H}_7\text{Cl}_2\text{NO}_2\text{S}$, *M* 240.1, *m* 83°.

Crystallise it from petroleum ether (b 60-80°) or CHCl₃/petroleum ether. Dry it in air <55° and store in the dark. It is soluble in CHCl₃ (~1:1), *C₆H₆ (~1:1) and CCl₄ (~1:2.5). It is a *germicide* and *antibacterial* as it readily liberates chlorine, and it should not smell strongly of chlorine; otherwise it should be purified. *Alternatively*, dissolve ~15g of reagent in 75ml of hot acetic acid and precipitate it by addition of 37ml of N/10 bleaching powder solution (NaOCl), filter off, wash it with a dilute solution of the latter, dry it as above (**m 78-84°**) and recrystallise it. Excessive drying, 'Sharp drying', will decompose it. [Orton & Bradfield *J Chem Soc* 986 1927, DOI: 10.1039/JR9270000986; Krauss & Crede *J Am Chem Soc* 39 2720 1917, DOI: 10.1021/ja02257a024; Soper *J Chem Soc* 1899 1924, DOI: 10.1039/CT9242501899; see also *chloramine-T* (*monochloramine T Na salt*) in this Chapter.] [*Beilstein* 11 H 107, 11 I 27, 11 II 63, 11 III 301.]

2,6-Dichlorophenol-indophenol sodium salt (2H₂O) (DCPIP, DCIP) [620-45-1] C₁₂H₆Cl₂NNaO₂·xH₂O, **M 326.1**, ε 2.1 x 10⁴ at 600nm and pH 8, pK³⁰ 5.7 (oxidised form), pK₁³⁰ 7.0, pK₂³⁰ 10.1 (reduced form). Dissolve the green powder in 0.001M phosphate buffer, pH 7.5 (*alternatively*, about 2g of the dye is dissolved in 80ml of M HCl), and extracted into diethyl ether. The extract is washed with water, extracted with aqueous 2% NaHCO₃, and the *sodium salt* of the dye is precipitated by adding NaCl (30g/100ml of NaHCO₃ solution), then filtered off, washed with dilute NaCl solution and dried. It is *blue* in the oxidised form with λ_{max} at 605nm at pH ~8, but turns *red* in aqueous acidic solution. [Hiromi et al. *Anal Biochem* 101 421 1980, DOI: 10.1016/0003-2697(77)90512-7.] It has been used for the determination of vitamin C in fruits by capillary zone electrophoresis [Chiari et al. *J Chromatogr* 645 179 1993, DOI: 10.1016/0021-9673(93)80637-N]. The *acetate* [24857-20-3] **M 310.1** has **m 101-103°** (from Et₂O/petroleum ether) and **99.5-100.5°** (from Et₂O). [*Beilstein* 13 IV 1078-1079.]

Dicyclopentylphosphine [39864-68-1] C₁₀H₁₉P, **M 170.2**, b 76-78°/0.8mm, d₄²⁵ 0.933, pK_{Est} ~4.5. Purify dicyclopentylphosphine by distillation in a vacuum in a stream of N₂ or Ar as it is air sensitive and must be stored in an inert atmosphere. [cf. Neidergall & Langenfeld *Chem Ber* 95 64 1962, DOI: 10.1002/cber.19620950114; *Beilstein* 16 IV 947 for dicyclohexyl-phosphine.]

O,O-Diethyl-S-2-diethylaminoethyl phosphorothiolate [78-53-5] C₁₀H₂₄NO₃PS, **M 269.3**, m 98-99°, b 110°/0.2mm. Recrystallise it from isopropanol/diethyl ether. It is a very *toxic nerve agent* (similar to *Sarin* which is *iso-Pr-OP(=O)MeF*, [107-44-8] C₄H₁₀FO₂PS, **M 140.1**, m -57°, b 56°/16mm, 147°/atm), too dangerous to use in agriculture and is used in warfare. [Ailman & Magee in *Organo Phosphorus Compound* (Kosolapoff & Maier eds) Wiley Vol 7 pp 487-871 1976.] For the systematic behaviour of the *free base* and its salts see Metcalf et al. *J Economic Entomology* (ed. J.T. Trumble) pp 205-210 1957, DOI: <http://dx.doi.org/10.1093/jee/50.2.205>. **TAKE EXTREME CARE as with very toxic insecticides, use with protective clothing in an efficient fume cupboard.**

Diethylmethylsilane [760-32-7] C₅H₁₄Si, **M 102.3**, b 78.4°/760mm, 77.2-77.6°/atm, d₄²⁰ 0.71, n_D²⁰ 1.3984. Fractionally distil it through a ca 20-plate column, and the fraction boiling within a range of less than 0.5° is collected. It is a *flammable* and *irritant* liquid. [Price *J Am Chem Soc* 69 2600 1947, DOI: 10.1021/ja01203a010; *Beilstein* 4 III 1847, 4 IV 3894]

N,N-Diethyltrimethylsilylamine [996-50-9] C₇H₁₉NSi, **M 145.3**, m -10°, b 33°/26mm, 126.8-127.1°/738mm, 126.1-126.4°, 126.3°/760mm, d₄²⁰ 0.763, n_D²⁰ 1.411. Fractionate it through a 2ft vacuum-jacketed column containing Helipak packing with a reflux ratio of 10:1. [Sauer & Hasek *J Am Chem Soc* 68 241 1946, DOI: 10.1021/ja01206a028; Langer et al. *J Org Chem* 23 50 1958, DOI: 10.1021/jo01095a017; Rühlmann *J Prakt Chem* 9 315 1959, DOI: 10.1002/prac.19590090514; *Beilstein* 4 IV 4010.]

Diethyl trimethylsilyl phosphite [13716-45-5] C₇H₁₉O₃PSi, **M 210.3**, b 61°/10mm, 66°/15mm, d₄²⁰ 0.9476, n_D²⁰ 1.4113. Fractionate it under reduced pressure and has ³¹PMR: δ_p -128ppm relative to H₃PO₄. It is useful for preparing ketones from aldehydes *via* α-trimethylsilyloxy-phosphonates [Sekine et al. *Bull Chem Soc Jpn* 55 224 1982, DOI: org/10.1246/bcsj.55.224; Sekine et al. *J Org Chem* 46 2097 1981, DOI: 10.1021/jo00323a024; Evans et al. *J Am Chem Soc* 100 3467 1978, DOI: 10.1021/ja00479a031.]

Dihexadecyl phosphate (DHP) [2197-63-9] $C_{32}H_{67}O_4P$, **M 546.9**, **m 74-75°**, **75°**, **pK_{Est} ~1.2**. Recrystallise it from MeOH. Solutions of DHP were made in $CHCl_3$ or CH_2Cl_2 for making vesicles. It has been used for making surfactant vesicles in studies of the viscosity-dependent variations of the fluorescence yield ϕ_F and the polarity induced shift of the emission band maximum $\lambda_{F(max)}$ of a derivative of a [**p-(dialkylamino)benzylidene]malononitrile** [74677-08-0] fluorescence probe. [Luka *J Am Chem Soc* **106** 4386 1984, DOI: 10.1021/ja00328a016]. [Beilstein **1** IV 1880.] It forms vesicles with $Ru(bpy)_3^{2+}$ adsorbed on the outer and the inner surface of the vesicles whose absorption spectra are different [Tricot et al. *Aust J Chem* **38** 527 1985, DOI:10.1071/CH9850527].

1,2-Dihydroxybenzene-3,5-disulfonic acid, di-Na salt (TIRON) [149-45-1] $C_6H_4Na_2O_8S_2$, **M 332.2**, **ϵ 6.9 x 10⁴ at 260nm**, **pH 10.8**, **pK₁ and pK₂ <2 (for SO₃⁻)**, **pK₃ 7.7**, **pK₄ 12.6 (for OHs of disulfonate dianion)**. Recrystallise it from water [French & Adams *Analyst* **99** 551 1974, DOI: 10.1039/AN9749900551]. It is an indicator colour reagent for Fe, Mn, Ti and Mo ions, and complexes with Al, Cd, Co, Co, Fe (III), Mn, Pd, UO_2^{2+} , VO^{2+} and Zn. [Beilstein **11** IV 630.]

(±)-Diisooctyl phenylphosphonate (bis[2-ethylhexyl] phenylphosphonate) [49637-59-4] $C_{22}H_{39}O_3P$, **M 382.5**, **b 204-207°/4mm**, **d₄²⁰ 0.970**, **n_D²⁵ 1.4780**. Distil it in a vacuum, percolate it through a column of alumina, then pass it through a packed column maintained at 150° to remove residual traces of volatile materials in a countercurrent stream of N₂ under reduced pressure [Dobry & Keller *J Phys Chem* **61** 1448 1957, DOI: 10.1021/j150556a052]. [Beilstein **16** III 884.]

Diisopropyl chlorosilane (chlorodiisopropylsilane) [2227-29-4] $C_6H_{14}ClSi$, **M 150.7**, **b 59°/8mm**, **80°/10mm**, **200°/738mm**, **d₄²⁰ 0.9008**, **n_D²⁰ 1.4518**. Impurities can be readily detected by ¹H NMR spectroscopy. Purify it by fractional distillation [Gilman & Clark *J Am Chem Soc* **69** 1499 1947, DOI: 10.1021/ja01198a071; Allen et al. *J Chem Soc* 3668 1957, DOI: 10.1039/JR9570003668].

Dilongifolyl borane [77882-24-7] $C_{30}H_{51}B$, **M 422.6**, **m 169-172°**. Wash it with dry Et₂O and dry it in a vacuum under N₂. It has **m 160-161°** in a sealed evacuated capillary. It is sparingly soluble in pentane, tetrahydrofuran, carbon tetrachloride, dichloromethane, and chloroform, but the suspended material is capable of causing asymmetric hydroboration. Disappearance of solid indicates that the reaction has proceeded. [Jadhav & Brown *J Org Chem* **46** 2988 1981, DOI: 10.1021/jo00327a036; Herold et al. *Helv Chim Acta* **66** 744 1983, DOI: 10.1002/hlca.19830660304.]

4,4'-Dimethoxythiobenzophenone [958-80-5] $C_{15}H_{14}O_2S$, **M 258.3**, **m 120°**. Recrystallise the thioketone from a mixture of cyclohexane/dichloromethane (4:1) or EtOH (**m 119°**). [Bergmann & Wagenberg *Chem Ber* **63** 2585 1930, DOI: 10.1002/cber.19300630934; Beilstein **8** H 319, **8** II 355, **8** III 2658, **8** IV 2457.]

4-N,N'-Dimethylaminoazobenzene-4'-isothiocyanate {DABITC, 4-[(4-isocyanatophenyl)-azo]-N,N'-dimethylaniline} [7612-98-8] $C_{15}H_{14}N_4S$, **M 282.4**, **m 167° to 171°**, **170-171°**, **pK_{Est} ~2.5**. Crystallise DABITC by dissolving 1g in 150ml of boiling Me₂CO, filtering hot and allowing to cool at -20° overnight, collecting the solid and drying it in a vacuum. Solutions of it in pyridine should be used immediately, otherwise they decompose. It is soluble in dioxane (20mg/ml) to give a **red** solution. It is moisture sensitive and is used for microsequencing of peptides and proteins. [Chang *Methods Enzymol* **91** 79, 455 1983, DOI: 10.1016/S0076-6879(83)91012-1, DOI: 10.1016/S0076-6879(83)91043-1.] **IRRITANT**.

Dimethyldichlorosilane [75-78-5] $C_2H_6Cl_2Si$, **M 129.1**, **m -75.5°**, **b 68.5-68.7°/750mm**, **70.5°/760mm**, **d₄²⁰ 1.0885**, **n_D²⁰ 1.4108**. Other impurities are chlorinated silanes and methylsilanes. Fractionate it through a 3/8in diameter 7ft Stedman column rated at 100 theoretical plates at almost total reflux. See purification of MeSiCl₂ [75-54-7] Solutions in heptane, CH_3CCl_3 or, 1-chloronaphthalene are used for the silanisation of glassware and pipettes. [Sauer & Hadsell *J Am Chem Soc* **70** 3590 1948, DOI: 10.1021/ja01191a014; Beilstein **4** IV 4110.]

2,6-Dimethyl-1,10-phenanthrolinedisulfonic acid, di-Na salt (H₂O) (bathocuproine-disulfonic acid di-Na salt) [52698-84-7] $C_{26}H_{20}N_2Na_2O_6S_2 \cdot xH_2O$, **M 564.5**, **m ~300°**, **pK_{Est} ~0 (for free acid)**. Inorganic salts and some coloured species can be removed by dissolving the crude material in the minimum volume of water

and precipitating by adding EtOH. The purified reagent can be obtained by careful evaporation of the filtrate. Recrystallise it from EtOH and dry it in a vacuum at room temperature in the dark. The reagent is used for the analytical determination of Cu and Cu-protein complexes. [Lorenzo et al. *J Electroanal Chem* **356** 43 1993, DOI: 10.1016/0022-0728(93)80509-G; Watkins et al. *Microchem J* **16** 14 1971, DOI: 10.1016/0026-265X(71)90077-4; cf. Matsushita et al. *Clin Chim Acta* **216** 103 1993, DOI: 10.1016/0009-8981(93)90143-R.]

Dimethylphenylsilyl chloride (DMPSCI, chlorodimethylphenylsilane, phenyl dimethyl chlorosilane) [768-33-2] **M 170.7, b 79°/15mm, 85-87°/32mm, 196°/760mm, d_4^{20} 1.017, n_D^{20} 1.509.** Fractionate it through a 1.5 x 18inch column packed with stainless steel helices, or a spinning band column. [Daudt & Hyde *J Am Chem Soc* **74** 386 1952, DOI: 10.1021/ja01122a029; Lewis *J Am Chem Soc* **70** 1115 1948, DOI: 10.1021/ja01183a073; Eaborn *J Chem Soc* 494 1953, DOI: 10.1039/JR9530000494.] It is used for standardising MeLi or MeMgBr that form Me₃PhSi which is estimated by GC. [Maienthal et al. *J Am Chem Soc* **76** 6392 1954, DOI: 10.1021/ja01653a043; House & Respass *J Organomet Chem* **4** 95 1965, DOI: 10.1016/S0022-328X(00)82372-4; *Beilstein* **16** IV 1475.] **TOXIC and MOISTURE SENSITIVE.**

Dimethyl thiophosphonate (dimethyl hydrogen phosphonothiolate) [5930-72-3] **M 126.1, b 53-53.5°/16.5mm, 56-59°/9mm, 59°/16mm, d_4^{25} 1.1892, n_D^{20} 1.4768.** Fractionally distil the ester in a stream of dry N₂ or Ar at as high a vacuum as possible. Store in a sealed ampoule under argon as it has a foul odour. The IR (film) has ν_{\max} at ~800 cm⁻¹ (12.6 μ , P-S). [IR: McIvor et al. *Canad J Chem* **34** 1611 1956, DOI: 10.1139/v56-211; McIvor et al. *Canad J Chem* **36** 820 1958, DOI: 10.1139/v58-121; Tongcharoensirikul et al. *J Org Chem* **69** 2322 2004, DOI: 10.1021/jo035707t; *Beilstein* **1** IV 1258.]

2,4-Dinitrobenzenesulfonyl chloride (Kharasch reagent) [528-76-7] **C₆H₃ClN₂O₄S, M 234.6, m 94-95°, 96°.** Crystallise the yellow sulfonyl chloride from CCl₄. It is a useful protective group [Review: Kharasch et al. *Chem Rev* **39** 269 1946]. [Kharasch & Langford *Org Synth Coll Vol* **5** 474 1973, DOI: 10.15227/orgsyn.044.0047; *Beilstein* **6** II 316.]

2,4-Dinitrobenzenesulfonyl chloride [1656-44-6] **C₆H₃ClN₂O₆S, M 266.6, m 101° to 103°, 102°.** Crystallise the sulfonyl chloride from *benzene or *benzene/petroleum ether. Useful reagent for the Eschenmoser α,β -cleavage of α,β -epoxyketones which involves conformational control of halolactonisation [Corey & Sachdev *J Org Chem* **40** 579 1975, DOI: 10.1021/jo00893a008]. [*Beilstein* **11** H 78, **11** IV 214.]

Di-*n*-octyl phenylphosphonate (DOPP) [1754-47-8] **C₂₂H₃₉O₃P, M 382.5, b 207°/4mm, 219-220°/0.7mm, d_4^{20} 0.932, n_D^{25} 1.4780.** Purify it as described for diisooctyl phenylphosphonate and distil under high vacuum. [*Beilstein* **16** IV 1069.] It is used in ion-based potentiometric sensors for the flow-injection determination of promethazine hydrochloride in pharmaceutical formulations and human urine [Hassan et al. *Sensors (Basel)* **11** 1028 2011, DOI: 10.3390/s110101028].

(1,3-Dioxalan-2-ylmethyl)triphenylphosphonium bromide [52509-14-5] **C₂₂H₂₂BrO₂P, M 429.3, m 198-193°, 191.5-193°, 193-195°.** Wash the crystals with Et₂O, dry them in a vacuum and recrystallise them from CH₂Cl₂/dry Et₂O to give prisms with **m 172-174°**, which is raised to **191.5-193°** on drying at 56°/0.5mm. [Cresp et al. *JCS Perkin Trans 1* 37 1974, DOI: 10.1039/P19740000037.]

Diphenyl disulfide (phenyl disulfide) [882-33-7] **C₁₂H₁₀S₂, M 218.3, m 58° to 60°, 60.5°.** Crystallise the orange coloured disulfide from MeOH. [Alberti et al. *J Am Chem Soc* **108** 3024 1986]. Also crystallise it repeatedly from hot Et₂O, then dry it in a vacuum at 30° over P₂O₅, fuse it under N₂ and re-dry it; the whole procedure being repeated, with a final drying under a vacuum for 24 hours. Alternatively, recrystallise it from hexane/EtOH solution. [Burkey & Griller *J Am Chem Soc* **107** 246 1985, DOI: 10.1021/ja00287a044; *Beilstein* **6** H 323, **6** IV 1560.]

Diphenyldiselenide [1666-13-3] **C₁₂H₁₀Se₂, M 312.1, m 59-61°, 61°, 62-64°.** Crystallise it twice from hexane [Kice & Purkiss *J Org Chem* **52** 3448 1987, DOI: 10.1021/jo00391a054]. [*Beilstein* **6** IV 1781.]

Diphenyl hydrogen phosphate (diphenyl phosphate) [838-85-7] $C_{12}H_{11}O_4P$, M 250.2, m 62-70°, 66°, 67.5°, 69°, 66-70°, 99.5°, pK²⁰ 0.26. Crystallise it from $CHCl_3$ /petroleum ether. [Cherbuliez in *Organo Phosphorus Compounds* (Kosolapoff & Maier eds) Wiley Vol 6 pp 211-577 1973, *Beilstein* 6 IV 714.]

4,7-Diphenyl-1,10-phenanthroline-disulfonic acid, di-Na salt 3H₂O (bathophenanthroline-disulfonic acid di-Na salt) [52746-49-3] $C_{24}H_{14}N_2Na_2O_6S_2$, M 590.6, m 300°, pK_{Est} ~0 (for free acid). Dissolve the crude sample in the minimum volume of water and add EtOH to precipitate the contaminants. Carefully evaporate the filtrate to obtain pure material. It forms a **dark red complex** with Fe^{2+} with λ_{max} 535nm (ϵ 2.23 x 10⁴ mol⁻¹ cm⁻¹) [Imasaka et al. *Anal Chim Acta* 115 407 1980, DOI: 10.1016/S0003-2670(01)93187-3]. It is prepared by sulfonating **bathophenanthroline** {(4,7-diphenyl-1,10-phenanthroline) [1662-01-7] $C_{24}H_{16}N_2$, M 332.4, m 218-220°, pKa 4.67, green colour}, with $ClSO_3H$: to 100g of bathophenanthroline is added 0.5ml of Fe free $ClSO_3H$ and heated over a flame for 30 seconds (CARE). Cool and carefully add 10ml of pure distilled H₂O and warm on a water bath with stirring till all solid dissolves. A stock solution is made by diluting 3ml of this reagent to 100ml with 45% aqueous NaOAc, filter off the solid and store in a dark bottle. In this way it is stable for several months. [Landers & Zak *Am J Clinical Pathology* 29 590 1958, PMID: 13533371.]

Diphenylphosphinic acid [1707-03-5] $C_{12}H_{11}O_2P$, M 218.2, m 193-195°, 194-195°, pK²⁰ 1.72. Recrystallise it from 95% EtOH and dry it under vacuum at room temperature. Its solubility in 0.1N aqueous NaOH is 5g/100ml at ~25°. [see Kosolapoff *Organophosphorus Compounds* J Wiley, NY, 1950, Kosolapoff and Maier *Organic Phosphorus Compounds* Wiley-Interscience, NY, 1972-1976, *Beilstein* 16 IV 1036.]

Diphenyl phosphorazidate (DPPA, diphenyl phosphoryl azide, phosphoric acid diphenyl ester azide) [26386-88-9] $C_{12}H_{10}N_3O_3P$, M 275.2, b 134-136°/0.2mm, 157°/0.17mm, d₄²⁵ 1.277, n_D²⁰ 1.551. This azide, a **nitrene source**, is prepared by stirring a mixture of diphenyl phosphorochloridate (56.8g, 210mmol, freshly distilled at 165-168°/5mm, see [2524-64-3]), sodium azide (16.3g, 250mmol) and anhydrous Me₂CO (300ml, dried over K₂CO₃ and freshly distilled) in a dry atmosphere at 20-25° for 21 hours. The **lachrymatory** mixture is filtered in a fume cupboard, and the filtrate is concentrated in vacuum. The residue is purified by distillation through a short Vigreux column keeping the bath temperature below 200° to avoid decomposition of the **azidate**, and collecting the fraction with b 134-136°/0.2mm (49-52g, 84-89%). It is a colourless non-explosive oil which can be stored for long periods if it is protected from moisture and light. It has IR (neat) with ν_{max} at 3060 (w, C-H), 2170 (s, -N₃), 1590 (m), 1490 (s, arene C=C), 1270 (m, P=O) and 960 (s, P-O-aryl) cm⁻¹; and the ¹H NMR (CDCl₃) has δ at 7.0-7.3 (br s, aryl-H). [Shioiri & Yamada *Org Synth Coll Vol* 7 206 1990, DOI: 10.15227/orgsyn.062.0187.] It is a **useful activating agent** and a versatile reagent in organic synthesis (Cremlyn *Aust J Chem* 26 1591 1973, DOI:10.1071/CH9731591; see also references cited in *Org Synth Coll Vol* 7 206 1990, DOI: 10.15227/orgsyn.062.0187; above). It is commercially available on a 100-200 mesh, 1% cross-linked polydivinylbenzene support and bound (via *para* linkage) through one of its phenyl rings (**PS-DPPA**, 1.0-1.5mmol/g N₃ loading). **TOXIC**.

Diphenyl phosphoro chloridate (diphenyl phosphoryl chloride) [2524-64-3] $C_{12}H_{10}ClO_3P$, M 268.6, b 141°/1mm, 194°/13mm, 314-316°/272mm, d₄³⁰ 1.2960, n_D³⁵ 1.5490. Fractionally distil it in a good vacuum; better use a spinning band column. [Walsh *J Am Chem Soc* 81 3023 1959, DOI: 10.1021/ja01521a028; IR: Bellamy & Beecher *J Chem Soc* 475 1952, DOI: 10.1039/JR9520000475; *Beilstein* 6 IV 737.] Used for phosphorylating OH groups, e.g. of serines [Mora et al. *Tetrahedron Lett* 34 2461 1963, DOI: 10.1016/S0040-4039(00)60441-0] and of pyranoses [Sabesam & Neira *Carbohydr Res* 223 169 1992, DOI: 10.1016/0008-6215(92)80015-S]. **TOXIC, do not inhale vapours**.

Diphenylsilane [775-12-2] $C_{12}H_{12}Si$, M 184.1, b 75-76°/0.5mm, 95-95°/13mm, 113-114°/9mm, 124-126°/11mm, 134-135°/16mm, 235.9-238.5°/760mm, d₄²⁰ 1.0027, n_D²⁰ 1.5802. Dissolve it in Et₂O, mix slowly with ice-cold 10% AcOH. The Et₂O layer is then shaken with H₂O until the washings are neutral to litmus. Dry over Na₂SO₄, evaporate the Et₂O and distil the residual oil under reduced pressure using a Claisen flask with the take-off head modified into a short column. Ph₂SiH₂ boils at 257°/760mm, but it cannot be distilled at this temperature because exposure to **air leads to flashing**, decomposition and formation of silica. It is a colourless, odourless oil, miscible with organic solvents but not H₂O. A possible impurity is **Ph₃SiH** which has m 43-45°

and would be found in the residue. [West & Rochow *J Org Chem* **18** 303 1953, DOI: 10.1021/jo01131a012; Benkeser et al. *J Am Chem Soc* **74** 648 1952, DOI: 10.1021/ja01123a019; Gilman & Zuech *J Am Chem Soc* **81** 5925 1959, DOI: 10.1021/ja01531a021; *Beilstein* **16** IV 1366.]

Diphenylsilanediol [947-42-2] $C_{12}H_{12}O_2Si$, **M 216.1**, **m 148°(dec)**, **150°(dec)**. Recrystallise it from $CHCl_3$ /methyl ethyl ketone. The diol can act as an anticonvulsant like *phenytoin* (5,5-diphenylimidazolidine-2,4-dione, *Dilantin*) [57-41-0] $C_{15}H_{12}N_2O_2$, **M 252.3**, **m 293-295°**, Fawcett et al. *Can J Chem* **55** 3631 1977, DOI: 10.1139/v77-510]. [*Beilstein* **16** IV 1523.]

Diphenyl sulfide [139-66-2] $C_{12}H_{10}S$, **M 186.3**, **m -40°**, **b 145°/8mm**, **296°/atm**, d_4^{20} **1.114**, n_D^{20} **1.6327**. Wash the sulfide with aqueous 5% NaOH, then water. Dry it with $CaCl_2$, then with sodium. The sodium is filtered off, and the diphenyl sulfide is distilled under reduced pressure. [*Beilstein* **2** H 299, **6** IV 1488.]

Diphenyl sulfone [127-63-9] $C_{12}H_{10}SO_2$, **M 218.3**, has **m 123°, 125°, 129°**, **b 378-379°/atm**. Crystallise the sulfone from diethyl ether. Soluble in hot H_2O . It has been purified by zone melting. [*Beilstein* **6** H 300, **6** IV 1490.] Poisonous to insect eggs and spiders.

sym-Diphenylthiourea (thiocarbanilide) [102-08-9] $C_{13}H_{12}N_2S$, **M 228.3**, **m 152°, 153°, 154°, 155°**. Crystallise the thiourea from boiling EtOH by adding hot water and allowing to cool. [*Beilstein* **12** H 394, **12** IV 810.] It is a rubber vulcaniser.

Diphenyl p-tolyl phosphate [26444-49-5] $C_{19}H_{17}O_4P$, **M 340.3**, **m 18-29°**, **b 156-158°/0.002mm**, **235-255°/760mm**, d_4^{25} **1.5758**. Distil it in a vacuum, then percolate it through a column of alumina. Finally, pass it through a packed column maintained at 150° to remove traces of volatile impurities in a countercurrent stream of nitrogen under reduced pressure. [Dobry & Keller *J Phys Chem* **61** 1448 1947, DOI: 10.1021/j150556a052; *Beilstein* **6** IV 2130.]

Disodium 4,5(1,8)-dihydroxynaphthalene-2,7(3,6)-disulfonate 2H₂O (Chromotropic acid di-Na salt) [5808-22-0] $C_{10}H_6Na_2O_8S_2 \cdot 2H_2O$, **M 400.3**, **m >300°**, pK_1 **0.61(SO₃⁻)**, pK_2 **0.7(SO₃⁻)**, pK_3 **5.45(OH)**, pK_4 **15.5(OH)**. Recrystallise it from H_2O or H_2O by adding EtOH. It complexes with Ag, ClO_3^- , Cr, Hg, NO_2^- , NO_3^- and Ti. [*Beilstein* **11** H 72, **11** I 307, **11** II 174, **11** III 174, **11** IV 576.]

Disodium ethylenebis(dithiocarbamate) (Nabam) [142-59-6] $C_4H_6Na_2S_4$, **M 256.3**, **m 195-198°**, d^{20} **1.14**, $pK_{Est} \sim 3.0$. It crystallises (as *hexahydrate*) from aqueous ethanol. It is a fungicide, a skin irritant, a potential carcinogen and a hepatic microsomal monooxygenase inhibitor in rats [Périsquet & Derache *Toxicol Eur Res* **3** 285 1981, PMID: 7330871]. [*Beilstein* **4** III 149, **4** IV 234.]

Disodium-β-glycerophosphate [819-83-0 ($4H_2O$), 13408-09-8 ($5H_2O$)] $C_3H_7Na_2O_6P \cdot xH_2O$, **M 216.0** (anhydr), **m 96-98°, 102-104°**, pK_2^{25} **6.66** (free acid). Crystallise it from water (solubility at ~20° is 0.1g/ml) in which it forms various hydrates. [Attwood et al. *Biochem J* **253** 387 1988, DOI: 10.1042/bj253038; *Beilstein* **1** I 275, **1** IV 2766.] It is a tonic agent and used in the treatment of hyperphosphatemia.

Disodium naphthalene-1,5-disulfonate [1655-29-4] $C_{10}H_6Na_2O_6S_2$, **M 332.3**, $pK_{Est} \sim 0$. Recrystallise it from aqueous acetone. It darkens at high temperatures. [Okahata et al. *J Am Chem Soc* **108** 2863 1986, DOI: 10.1021/ja00271a013; *Beilstein* **11** IV 561.]

Disodium p-nitrophenylphosphate (6H₂O) (pNPP) [4264-83-9] $C_6H_4NNa_2O_6P \cdot 6H_2O$, **M 371.1**. Dissolve it in hot aqueous MeOH, filter and precipitate it by adding Me_2CO . Wash the solid with Me_2CO and repeat the purification. Aqueous MeOH and Et_2O can also be used as solvents. The white fibrous crystals contain less than 1% of free p-nitrophenol. It is a substrate for the determination of acid and alkaline phosphatases where it produces the soluble yellow end product (p-nitrophenol) that can be read off spectrophotometrically at 405nm. The pNPP enzymic reaction may be stopped with 3N NaOH, and its concentration is read at 405nm. [Assay: Axelrod *J Biol Chem* **167** 57 1947, <http://www.jbc.org/content/167/1/57>.] [*Beilstein* **6** IV 1327.]

Disodium phenylphosphate (2H₂O) [3279-54-7, 66778-08-3 ($2H_2O$)] $C_6H_5Na_2O_4P \cdot 2H_2O$, **M 254.1**, pK_1^{25}

1.46, pK_2^{25} 6.29 [for $\text{PhPO}(\text{OH})_2$]. Dissolve it in a minimum amount of methanol, filtering off any insoluble residue of inorganic phosphate, then precipitate it by adding an equal volume of Et_2O . Wash the solid with Et_2O and dry it in a vacuum [Tsuboi *Biochim Biophys Acta* **8** 173 1952, DOI: 10.1016/0006-3002(52)90027-9]. Store it in a sealed container at 2–8° as it is *hygroscopic*. Its solubility in H_2O at ~20° is 0.1g/ml. [Beilstein **6** IV 708.]

Dithizone (diphenylthiocarbazone) [60-10-6] $\text{C}_{13}\text{H}_{12}\text{N}_4\text{S}$, M 256.3, m 168°(dec), ratio of $\epsilon_{620\text{nm}}/\epsilon_{450\text{nm}}$ should be ≥ 1.65 , ϵ_{620} 3.4×10^4 (CHCl_3), pK_2 4.6. The crude dithizone is dissolved in CCl_4 to give a concentrated solution. This is filtered through a sintered glass funnel and shaken with 0.8M aqueous ammonia to extract *dithizonate ion*. The aqueous layer is washed with several portions of CCl_4 to remove undesirable materials. The aqueous layer is acidified with dilute H_2SO_4 to precipitate pure *dithizone*. This is dried in a vacuum. When only small amounts of dithizone are required, purification by paper chromatography is convenient. [Cooper & Hibbits *J Am Chem Soc* **75** 5084 1933, DOI: 10.1021/ja01116a054.] Instead of CCl_4 , CHCl_3 can be used, and the final extract, after washing with water, can be evaporated in air at 40–50° and dried in a desiccator. It complexes with Cd, Hg, Ni and Zn. [Beilstein **16** H 26, **16** IV 18.]

Di-*p*-tolyl phenylphosphonate [94548-75-1] $\text{C}_{20}\text{H}_{19}\text{O}_3\text{P}$, M 338.3, n_D^{25} 1.5758. Purify as described under diisooctyl phenylphosphonate.

Di-*p*-tolyl sulfone [599-66-6] $\text{C}_{14}\text{H}_{14}\text{O}_2\text{S}$, M 246.3, m 158–159°, 160°, 163.5°, b 405°/760mm. Crystallise the sulfone repeatedly from Et_2O or EtOH . It has been purified by zone melting. [Beilstein **6** H 419, **6** II 395, **6** III 1405, **6** IV 2174.]

1,3-Divinyl-1,1,3,3-tetramethyldisiloxane [2627-95-4] $\text{C}_8\text{H}_{18}\text{OSi}_2$, M 186.4, m -100°, -99°, -99.7°, b 128–129°/atm, 139°/760mm, 139–140°/760mm, d_4^{20} 0.811, n_D^{20} 1.4122. Dissolve it in Et_2O , wash it with H_2O , dry it over CaCl_2 , evaporate the solvent and distil the residue. It is an **IRRITANT**. [Kantor et al. *J Am Chem Soc* **77** 1685 1955, DOI: 10.1021/ja01611a090; Bažant & Matoušek *Coll Czech Chem Commun* **24** 3758 1959, DOI: org/10.1135/ccccc19593758; Beilstein **4** IV 4080.]

Eriochrome Black T [3-hydroxy-7-nitro-4-(1-hydroxy-2-naphthylazo)naphthalene-1-sulfonic acid Na salt] [1787-61-7] $\text{C}_{20}\text{H}_{12}\text{N}_3\text{NaO}_7\text{S}$, M 461.4, $A_{1\text{cm}}^{1\%}(\lambda_{\text{max}})$ 656(620nm) at pH 10, using the dimethylammonium salt, pK_2^{25} 5.81, pK_3^{25} 11.55. The sodium salt (200g) is converted to the *free acid* by stirring with 500ml of 1.5M HCl , and, after several minutes, the slurry is filtered on a sintered-glass funnel. The process is repeated and the material is air dried after washing with acid. It is then extracted with *benzene for 12 hours in a Soxhlet extractor, the *benzene solution is evaporated and the residue is air dried. A further desalting with 1.5M HCl (1L) is followed by crystallisation from dimethylformamide (in which it is very soluble) by forming a saturated solution at the boiling point, and allowing to cool slowly. The crystalline dimethylammonium salt so obtained is washed with *benzene and treated repeatedly with dilute HCl to give the insoluble *free acid* which, after air drying, is dissolved in alcohol, filtered and evaporated. The final material is air dried, then dried in a vacuum desiccator over $\text{Mg}(\text{ClO}_4)_2$. The purified acid is converted to the dimethylammonium salt with Me_2NH . [Diehl & Lindstrom *Anal Chem* **31** 414 1959, DOI: 10.1021/ac60147a026]. It is an indicator in the complexometry of alkaline earth metals. [Beilstein **16** IV 429.]

Eriochrome Blue Black B (Mordant Black, 3, 3-hydroxy-4-(1-hydroxy-2-naphthylazo)-naphthalene-1-sulfonic acid Na salt] [3564-14-5] $\text{C}_{20}\text{H}_{13}\text{N}_2\text{NaO}_5\text{S}$, M 416.4, has pK_2^{25} 7.0, pK_3^{25} 13.5. Free it from metallic impurities by three precipitations from aqueous solution by addition of HCl . The precipitated dye is dried at 60° under vacuum and converted into the Na salt with the calculated amount of alkali. It is an indicator in the complexometry of Al, Fe and Zr. [Beilstein **16** III 319.]

Eriochrome Blue Black R (Palatine Chrome Black 6BN, Mordant Black 17, 3-hydroxy-4-(2-hydroxy-1-naphthylazo)naphthalene-1-sulfonic acid Na salt] [2538-85-4] $\text{C}_{20}\text{H}_{13}\text{N}_2\text{NaO}_5\text{S}$, M 416.4, has pK_2^{25} 7.0, pK_3^{25} 13.5. Free it from metallic impurities by three precipitations from aqueous solution by addition of HCl . The precipitated dye is dried at 60° under vacuum and converted into the Na salt with the calculated amount of alkali. It is an indicator in the complexometry of Al, Fe and Zr. [Beilstein **16** H 297, **16** IV 428.]

Ethoxycarbonylmethylene triphenylphosphonium bromide [1530-45-6] $\text{C}_{22}\text{H}_{22}\text{BrO}_2\text{P}$, M 429.3, m 145–

150°, 155-155.5°, 158°(dec), pK_a 8.95-9.2 (deprotonation). Wash it with petroleum ether (b 40-50°) and recrystallise it from CHCl₃/Et₂O; and dry it in a high vacuum at 65°. Its solubility at ~20° in H₂O is 100g/L, in EtOH it is 350g/L, in CHCl₃ it is 550g/L, but it is insoluble in THF (<10g/L). [Isler et al. *Helv Chim Acta* **40** 1242 1957, DOI: 10.1002/hlca.19570400515; Wittig & Haag *Chem Ber* **88** 1654, 1664 1955, DOI: 10.1002/cber.19550881110.] [*Beilstein* **16** IV 980.] **IRRITANT.**

(Ethoxycarbonylmethylene)triphenylphosphorane [ethyl (triphenylphosphoranylidene)-acetate] [1099-45-2] **C₂₂H₂₁O₂P, M 348.4, has m 116-117°, 128-130°.** Crystallise it by dissolving it in AcOH and adding petroleum ether (b 40-50°) to give colourless plates. Its UV has λ_{\max} ($A_{1\text{mm}}^{1\%}$) at 222nm (865) and 268nm (116). Its solubility at ~20° in EtOH is 4.2g/10ml, in THF it is 1.3g/10ml, in CHCl₃ it is 2.9g/10ml, but it is insoluble in H₂O (<0.05g/10ml). [Isler et al. *Helv Chim Acta* **40** 1242 1957, DOI: 10.1002/hlca.19570400515; *Beilstein* **16** IV 977.]

Ethylarsonic acid [507-32-4] **C₂H₇AsO₃, M 154.0, m 94-95°, 99.5°, pK₁ 4.72 (As(OH)O⁻), pK₂ 8.00 [AsO₂²⁻].** Crystallise it from ethanol. Also it dissolves in excess EtOH, filter off the insoluble matter (salts?), evaporate the filtrate to dryness and recrystallise the residue from small volumes of EtOH or H₂O. [Quick & Adams *J Am Chem Soc* **44** 805 1922, DOI: 10.1021/ja01425a014; Banks et al. *J Am Chem Soc* **69** 927 1947, DOI: 10.1021/ja01196a057; *Beilstein* **4** H 614, **4** II 997, **4** III 1823, **4** IV 3682.]

Ethylene di(p-toluenesulfonate) (ethylene glycol ditosylate) [6315-52-2] **C₁₆H₁₈O₆S₂, M 370.44, m 124-127°, m 126-127°, m 128°.** It is prepared when ethylene glycol (0.5mole) and *p*-toluenesulfonyl chloride (1.0mole) are stirred in Me₂CO (100ml) in the presence of 25% aqueous NaOH (175ml, 44g, 1.1mole) at 2-10° for ~4 hours. Excess of ice-water is added, the solid is filtered off, washed with large volumes of H₂O, dried *in vacuo* and recrystallised from dry *C₆H₆ (absolute EtOH has also been used) to give the **di-tosylate (m 128°)** in 76% yield [Drahowzal & Klamann *Monatsh Chem* **82** 452 1951, DOI: 10.1007/BF00900841]. Alternatively, ethylene glycol (0.25mole), *p*-toluenesulfonyl chloride (0.52mole) in dry pyridine (80g, 1mole) are stirred gently at -10° to -4° for 2 hours then treated with excess of ice-water as above to give the **diester** in 87.3% yield [Drahowzal & Klamann *Monatsh Chem* **82** 460 1951, DOI: 10.1007/BF00900842]. The latter method is superior as there is no H₂O in the reaction medium. **Note** that attempts to prepare the **mono-p-tosylate ester** using equimolar amounts of glycol and sulfonyl chloride always produced the **diester** which crystallised as white plates (**m 126°**) from hot EtOH [Butler et al. *J Am Chem Soc* **57** 575 1935, DOI: 10.1021/ja01306a057]. [*Beilstein* **11** II 296, **11** III 225, **11** IV 290.]

S,S'-Ethylene di(p-toluenethiosulfonate) (ethylene dithiotosylate) [2225-23-2] **C₁₆H₁₈O₄S₄, M 402.5, m 75-76°.** This reagent is the lower homologue of **1,3-trimethylene di(thiotosylate)** [3866-79-3], and is made from potassium thiotosylate (45g, 0.20 mole, see below [28519-50-8]) and 1,2-dibromoethane (18.8g, 0.1 mole) in EtOH (200ml, containing 10-20mg of KI to activate the dibromide) by refluxing, with stirring in the dark under a N₂ atmosphere, for 8 hours. The solvent is evaporated *in vacuo*, and the white solid residue is washed, by decantation, with a mixture of EtOH (80ml) and H₂O (150ml), and then with H₂O (3 x 50ml), and recrystallised from EtOH (~150ml) to provide the crude thioester (28.7g) with m 72-75°. The **pure thioester** (24g, 60%), **m 75-76°**, is obtained as white crystals after three recrystallisations from EtOAc/EtOH; and its ¹H NMR (CDCl₃, TMS) has δ at 2.47 (s, 6H, 2CH₃), 3.31 (s, 4H, CH₂CH₂), 7.48 (d, *J* = 9Hz, 4H, Aromatic H) and 7.97 (d, *J* = 9Hz, 4H, Aromatic H). [Woodward et al. *Org Synth Coll Vol* **6** 1016 1988, DOI: 10.15227/orgsyn.054.0033.] Like the trimethylene homologue below it can form 1,3-dithiolanes with activated methylene groups and carbonyl compounds. [*Beilstein* **11** II 71.]

Ethynyl p-tolylsulfone (tosylacetylene) [13894-21-8] **C₉H₈O₂S, M 180.2, m 65-67°, 73-74°.** Recrystallise the sulfone from petroleum ether, *C₆H₆ or EtOH (**m 66°**), and dry it in a vacuum. [te et al. *J Am Chem Soc* **107** 686 1985, DOI: 10.1021/ja00289a024; *Beilstein* **6** III 1397, **6** IV 2160.]

Ethyl Orange (sodium 4,4'-diethylaminophenylazobenzenesulfonate) **C₁₆H₁₈NaN₃O₃S, [62758-12-7, 13545-67-0 free acid] M 355.4, pK_{Est} ~ 3.8.** Recrystallise it twice from water. It is an indicator with a visual transition interval from red to orange in the pH region 3.0—4.8 with λ_{\max} at 474nm. [*Beilstein* **16** IV 511.]

Ethyl trimethylsilylacetate (ETSA) [4071-88-9] $C_7H_{16}O_2Si$, M 160.3, b 75.5°/42mm, 157°/730mm, 156-158°/760mm, d_4^{20} 0.8762, n_D^{20} 1.4149. Purify it by distilling *ca* 10g of reagent through a 15cm, Vigreux column and then redistilling it through a 21cm glass helices-packed column [Hance & Hauser *J Am Chem Soc* **75** 994 1953, DOI: 10.1021/ja01100a511]. Alternatively, dissolve it in Et_2O , wash with H_2O , dilute Na_2CO_3 , dry over Na_2CO_3 , evaporate Et_2O , and distil it through a column of 15 theoretical plates. [Gold et al. *J Am Chem Soc* **70** 2874 1948, DOI: 10.1021/ja01100a511; *Beilstein* **4** IV 3974.]

Ethyl 3-(trimethylsilyl)propionate [17728-88-0] $C_8H_{18}O_2Si$, M 174.3, b 93°/40mm, 178°-180°/atm, d_4^{20} 0.8763, n_D^{20} 1.4198. Dissolve it in Et_2O , wash this with H_2O , dilute Na_2CO_3 , dry (Na_2SO_4), evaporate Et_2O and fractionally distil. It is a starting material for the synthesis of **2-(TMS-methyl)allylic alcohols** and related compounds [Vedejs et al. *J Org Chem* **47** 1534 1982, DOI: 10.1021/jo00347a034; Nishitani & Yamakawa *Tetrahedron Lett* **28** 655 1987, DOI: 10.1016/S0040-4039(00)95805-2]. [Sommer & Marans *J Am Chem Soc* **72** 1935 1950, DOI: 10.1021/ja01161a021; *Beilstein* **4** IV 3975.]

Ethyl triphenylphosphonium bromide [1530-32-1] $C_{20}H_{20}BrP$, M 371.3, m 203-205°. Recrystallise it from H_2O and dry it in a high vacuum at 100°. Its IR has bands at 1449, 1431 and 997 cm^{-1} . [Wittig & Wittenberg *Justus Liebigs Ann Chem* **606** 1 1957, DOI: 10.1002/jlac.19576060102; Bergmann & Dusza *J Org Chem* **23** 1245 1958, DOI: 10.1021/jo01103a002; *Beilstein* **16** IV 982.] It is a useful phase transfer catalyst, is *hygroscopic* and should be stored in a dry atmosphere.

Ethynyl trimethylsilane [1066-54-2] M 98.2, b 53°/atm, 52.5°/atm, d_4^{20} 0.71, n_D^{20} 1.3871. Distil it through an efficient column. The IR has bands at ν_{max} 2041 ($C\equiv C$) and 3289 ($\equiv C-H$) cm^{-1} . [Kröhnke & Gross *Chem Ber* **92** 22 1959, DOI: 10.1002/cber.19590920105; *Beilstein* **4** IV 3937.] A suitable substrate for Ni-catalysed cross-coupling with benzonitriles [Penney & Miller *Tetrahedron Lett* **45** 4989 2004, DOI: 10.1016/j.tetlet.2004.02.163], and used in the *microwave assisted* 1,3-dipolar cycloaddition reactions for preparing pyrazoles [Zrinski et al. *Heterocycles* **68** 1961 2006, DOI: 10.3987/COM-06-10803].

4-Fluorophenyl isothiocyanate [1544-68-9] C_7H_4FNS , M 153.2, m 24-26°, 26-27°, b 66°/2mm, 215°/atm, 228°/760mm, n_D^{20} 1.6116. A likely impurity is the symmetrical thiourea. Dissolve the isothiocyanate in dry $CHCl_3$, filter and distil the residue in a vacuum. It can also be steam distilled, the oily layer is separated, dried over $CaCl_2$ and distilled *in vacuo*. **Bis-(4-fluorophenyl)thiourea** has m 145° (from aqueous $EtOH$). [Browne & Dyson *J Chem Soc* 3285 1931, DOI: 10.1039/JR9310003285; Buu Hoi et al. *J Chem Soc* 1573 1955, DOI: 10.1039/JR9550001573; Dains et al. *Org Synth Coll Vol* **1** 447 1941, DOI: 10.15227/orgsyn.006.0072].

Fluorotrimethylsilane (trimethylsilyl fluoride, TMSF) [420-56-4] C_3H_9FSi , M 92.2, m -74°, b 16°/760mm, 16-18°/760mm, 19°/730mm, d_4^{20} 0.793. It is a **FLAMMABLE** gas that is purified by fractional distillation through a column at low temperature and with the exclusion of air. Preferably work in a vacuum line. [Booth & Suttle *J Am Chem Soc* **68** 2658 1946, DOI: 10.1021/ja01216a072; Reid & Wilkins *J Chem Soc* 4029 1955, DOI: 10.1039/JR9550004029; *Beilstein* **4** IV 4007.]

Hexaethyldisiloxane [924-49-0] $C_{12}H_{30}OSi_2$, M 246.5, b 114-115°/16mm, 235.5°/760mm, d_4^{20} 0.8443, n_D^{20} 1.4330. Distil in a vacuum, but it can be distilled at atmospheric pressure without decomposition. It is characterised by completely dissolving in concentrated H_2SO_4 . [Eaborn *J Chem Soc* 3077 1950, DOI: 10.1039/JR9500003077; *Beilstein* **4** IV 4055.] Used in the facile silylation of carboxylic acids to trimethylsilyl carboxylates in 74-97% yields [Matsumoto et al. *Chem Lett* 1475 1980, DOI: 10.1246/cl.1980.1475].

2,2,4,4,6,6-Hexamethylcyclotrisilazane [1009-93-4] $C_6H_{21}N_3Si_3$, M 219.6, m -10°, b 81-82°/19mm, 111-112°/85mm, 188°/756mm, d_4^{20} 0.9196, n_D^{20} 1.448. Purify it by fractional distillation at atmospheric pressure until the temperature reaches 200°. It is *moisture sensitive*. The residue in the flask can be distilled (b 225°/756mm) and is mostly octamethylcyclo-tetrasilazane which can be crystallised from petroleum ether (m

97°). [Brewer & Haber *J Am Chem Soc* **70** 3888 1948, DOI: 10.1021/ja01191a106; *Beilstein* **4** III 1887.]

2,2,4,4,6,6-Hexamethylcyclotrisiloxane [541-05-9] $C_6H_{18}O_3Si_3$, *M* 222.4, *m* 50-64°, 64.5°, 64-66°, *b* 131°/760mm, 134°/760mm. The cyclotrisiloxane is purified by stirring the molten compound with finely ground CaH_2 for 12 hours, filtered hot and sublimed under high vacuum. The sublimed *monomer* is then dissolved in dry *C_6H_6 , treated with PS^+Li^- (polystyrene lithium salt) for further purification. The *C_6H_6 solution is filtered, evaporated to dryness and the residue is sublimed under high vacuum again. A stock solution of the *siloxane* in *C_6H_6 can be used for further work [Bellas et al. *Macromolecules* **33** 6993 2000, DOI: 10.1021/ma000635i]. [*Beilstein* **4** IV 4123.]

Hexamethyldisilane [1450-14-2] $C_6H_{18}Si_2$, *M* 164.4, *m* 9-12°, 12-13°, 13.5°, *b* 113.1°/750mm, 112-114°/760mm, d_4^{20} 0.7272, n_D^{20} 1.4229. The most likely impurity is *trimethylchlorosilane* (cf. boiling point). Wash it with H_2O , cold concentrated H_2SO_4 , H_2O again, then aqueous $NaHCO_3$, dry over $CaSO_4$ and fractionate at atmospheric pressure. [Brown & Fowles *J Chem Soc* 2811 1958, DOI: 10.1039/JR9580002811.] A grossly impure sample (25% impurities) was purified by repeated spinning band distillation. This lowered the impurity level to 500 ppm. The main impurity was identified as *1-hydroxypentamethyldisilane*. The Si-Si bond is cleaved by strong nucleophiles and electrophiles. [Hiyama & Kuroboshi, 'Hexamethyldisilane' in Encyclopedia of Reagents for Organic Synthesis, 2001 John Wiley & Sons. DOI:10.1002/047084289X.rh015; *Beilstein* **4** IV 4277.] Used in low pressure chemical deposition of silicon carbide thin films [Chiu & Hsu *Thin Solid Films* **252** 13 1994, DOI: 10.1016/0040-6090(94)90818-4]. **IRRITANT and FLAMMABLE.**

Hexamethyldisilazane [HMDS, $(Me_3Si)_2NH$] [999-97-3] $C_6H_{19}NSi_2$, *M* 161.4, *m* -80°, -78°, *b* 125-125.6°/atm, 126°/760mm, d_4^{20} 0.7747, n_D^{20} 1.407. A possible impurity is Me_3SiCl . Wash it well with petroleum ether and fractionate it through a vacuum jacketed column packed with Helipac using a reflux ratio of 10:1. [Langer et al. *J Org Chem* **23** 50 1958, DOI: 10.1021/jo01095a017; Osthoff & Kantor 'Organosilazane Compounds' *Inorg Synth* **5** 5 1957, DOI: 10.1002/9780470132364.ch16. ISBN 978-0-470-13236-4; *Beilstein* **4** IV 4014.] It forms a *lithium salt*, $(Me_3Si)_2NLi$ [4039-32-1], which is soluble in THF and hexane; and ~ 1M solutions in these solvents are available commercially. **IRRITANT and FLAMMABLE.**

Hexamethyldisiloxane (HMDSO) [107-46-0] $C_6H_{19}OSi_2$, *M* 162.4, *m* -59°, *b* 99.4°/760mm, 100.4°/764mm, 101°/764mm, d_4^{20} 0.7633, n_D^{20} 1.3777. Fractionally distil through a column packed with glass helices with *ca* 15 theoretical plates. It is highly **flammable** and is an **irritant**. [Mills & McKenzie *J Am Chem Soc* **76** 2672 1954, DOI: 10.1021/ja01639a020; Csakvari et al. *J Organometal Chem* **107** 287 1976, DOI: 10.1016/S0022-328X(00)91519-5; *Beilstein* **4** IV 4018.] It is also a commercially available NMR reference standard, as a 25% solution in $^*C_6H_6-d_6$.

Hexamethylphosphoric triamide (HMPA) [680-31-9] $C_6H_{18}N_3OP$, *M* 179.2, *m* 7.2°, *b* 68-70°/1mm, 230-232°/740mm, 235°/760mm, d_4^{20} 1.024, n_D^{20} 1.460. The industrial synthesis is usually by treatment of $POCl_3$ with excess of dimethylamine in isopropyl ether. Impurities are water, dimethylamine and its hydrochloride. It is purified by refluxing over BaO or CaO at about 4mm pressure in an atmosphere of nitrogen for several hours, then distilled from sodium at the same pressure. The middle fraction (*b ca* 90°) is collected, refluxed over sodium under reduced pressure under nitrogen and distilled. It is kept in the dark under nitrogen, and stored in solid CO_2 . It can also be stored over 4A molecular sieves. **Alternatively**, it is distilled under vacuum from CaH_2 at 60° and is crystallised twice in a cold room at 0°, seeding the liquid with crystals obtained by cooling in liquid nitrogen. After about two-thirds are frozen, the remaining liquid is drained off [Fujinaga et al. *Pure Appl Chem* **44** 115 1975, DOI: 10.1351/pac197544010115]. For tests of purity see Fujinaga et al. in *Purification of Solvents*, Coetzee Ed., Pergamon Press, Oxford, 1982. For efficiency of desiccants in drying HMPA see Burfield and Smithers [*J Org Chem* **43** 3966 1978, DOI: 10.1021/jo00414a038; and Sammes et al. *JCS Perkin Trans 1* 281 1986, DOI: 10.1039/P19860000281]. [*Beilstein* **4** IV 284.] Useful aprotic solvent. **CARCINOGEN.**

Hexamethylphosphorous triamide [HMPT, tris(dimethylamino)phosphine] [1608-26-0] $C_6H_{18}N_3P$, *M* 163.2, *m* -44°, *b* 49-51°/12mm, 162-164°/atm, 176.2-178.9°/760mm, d_4^{20} 0.989, n_D^{20} 1.4636. It may contain more than 1% of phosphoric triamide. The yellow oil is first distilled at atmospheric pressure, then under

reduced pressure and stored under N₂. ¹H NMR (CDCl₃, 300MHz): δ 2.48 (d, *J* 9.1Hz, 18*H*) ppm; ³¹P NMR (CDCl₃, 162MHz, H₃PO₄, 40%): δ 122 ppm. It is air sensitive, **TOXIC**, and should not be inhaled. It is absorbed through the skin. [Mark *Org Synth Coll Vol* **5** 602 1973, DOI: 10.15227/orgsyn.046.0042; Harvey & Schneider in *e-EROS Encyclopedia of Reagents for Organic Synthesis* on line 22 APR 2013, DOI: 10.1002/047084289X.rh022.pub2; *Beilstein* **4** IV 274.]

Hydroquinone-2-sulfonic acid K salt [21799-87-1] C₆H₅KO₅S, M 228.3, m 250°(dec), pK_{Est(1)}~1, pK_{Est(2)}~8.5, pK_{Est(3)}~11. Recrystallise it from water or EtOH. [*Beilstein* **11** I 70, **11** II 170, **11** III 570.]

Hydroxynaphthol Blue tri-Na salt [1-(2-hydroxy-4-sulfo-1-naphthylazo)-2-naphthol-3,6-disulfonic acid tri-Na salt] [63451-35-4 for di-Na salt, 165660-27-5 for tri-Na salt] C₂₀H₁₁N₂Na₃O₁₁S₃, M 620.5, m dec on heating, pK_{Est} <0. The crude material is treated with hot EtOH to remove soluble impurities, then dissolve in 20% aqueous MeOH and chromatographed on a cellulose powder column with propanol/EtOH/water (5:5:4) as eluent. The upper of three zones are eluted to give the pure dye that is precipitated as the *monosodium salt trihydrate* by adding concentrated HCl to the concentrated eluate [Ito & Ueno *Analyst* **95** 583 1970, DOI: 10.1039/AN9709500583]. It can be converted to the *trisodium salt* by adding the calculated amount of alkali. The synthetic dye is a Ca indicator with λ_{max} at 650m, where in the pH range 12–13 the solution of the indicator is red-pink in the presence of Ca²⁺ and deep blue in the presence of excess of EDTA.

6-Hydroxy-2-naphthyl disulfide [6088-51-3] C₂₀H₁₄O₂S₂, M 350.5, m 221-222°, 226-227°, pK_{Est} ~9.0. It crystallises as leaflets from AcOH and is slightly soluble in EtOH, and AcOH, but is soluble in *C₆H₆ and in alkalies to give a yellow solution. [Zincke & Dereser *Chem Ber* **51** 352 1918, DOI: 10.1002/cber.19180510137.] The *acetyl derivative* has m 198-200° (from AcOH or dioxane/MeOH), and the *diacetyl derivative* has m 167-168° (from AcOH). A small amount of impure disulfide can be purified by dissolving it in a small volume of Me₂CO and adding a large volume of toluene, filtering rapidly and concentrating to one-third of its volume. The hot toluene solution is filtered rapidly from any tarry residue, and crystals separate on cooling. Recrystallisation from hot acetic acid gives crystals with m 220-223° [Barrett & Seligman *Science* **116** 323 1952, DOI: 10.1126/science.116.3013.323]. Store at -20°. [*Beilstein* **6** I 481.]

Hydroxy(tosyloxy)iodobenzene [phenyl(hydroxyl)tosyloxyiodine, hydroxy(4-methylbenzenesulfonato-O)-phenyliodine, Koser's reagent] [27126-76-7] C₁₃H₁₃IO₄S, M 392.2, m 131° to 137°, 134-136°, 135-138°, 134-136°, 136-138.5°. Possible impurities are tosic acid (removed by washing with Me₂CO) and acetic acid (removed by washing with Et₂O). It is purified by dissolving in the minimum volume of MeOH, adding Et₂O to cloud point and setting aside for the prisms to separate [Koser & Wettach *J Org Chem* **42** 1476 1977, DOI: 10.1021/jo00428a052; NMR: Koser et al. *J Org Chem* **41** 3609 1976, DOI: 10.1021/jo00884a028]. It has also been crystallised from CH₂Cl₂ (needles, m 140-142°) [Neiland & Karele *J Org Chem, USSR (Engl Transl)* **6** 889 1970]. Store in a refrigerator. It is a useful oxidant in organic synthesis [Kumar *Synlett* 2764 2007, DOI: 10.1055/s-2007-991080].

Indigocarmine (2[1,3-dihydro-3-oxo-5-sulfo-2*H*-indol-2-ylidene]-2,3-dihydro-3-oxo-1*H*-indole-5-sulfonic acid di-Na salt), **Acid Blue 74**, **Indigo-5,5'-disulfonic acid di-Na salt** [860-22-0] C₁₆H₈N₂Na₂O₈S₂, M 466.4, pK₁²⁰ 2.8, pK₂²⁰ 12.3. Its solubility in H₂O is 1g/100ml at 25°. It has been purified by dissolving in H₂O, filtering and adding EtOH to cause the salt to separate. Wash the solid with EtOH, Et₂O and dry *in vacuo*. Biological Stain. [Vörlander & Schubart *Chem Ber* **34** 1860 1901, DOI: 10.1002/cber.19010340290; UV: Smit et al. *Anal Chem* **27** 1159 1955, DOI: 10.1021/ac60103a035; Preisler et al. *J Am Chem Soc* **81** 1991 1959, DOI: 10.1021/ja01517a051; *Beilstein* **25** IV 1975.]

Iodomethyl trimethylsilane [4206-67-1] C₄H₁₁ISi, M 214.1, b 139.5°/744mm, d₄²⁰ 1.44, n_D²⁵ 1.4917. If slightly violet in colour, wash it with aqueous 1% sodium metabisulfite, H₂O, dry it (Na₂SO₄), and fractionally distil it at 760mm. [Whitmore & Sommer *J Am Chem Soc* **68** 481 1946, DOI: 10.1021/ja01207a036.]

Iodotrimethylsilane (trimethylsilyl iodide, TMSI) [16029-98-4] C₃H₉ISi, M 200.1, b 106.8°/742mm, 107.5°/760mm, d₄²⁰ 1.470. Add a little antimony powder and fractionate with this powder in the still. Stabilise

the distillate with 1% wt of Cu powder and store away from light. It is prepared by adding I_2 (124g, 0.49mol) in ~20g portions to trimethylphenylsilane (75g, 0.5ml, see 3385-94-2 below) containing aluminium iodide (1g) and refluxing after each addition until the I_2 is consumed. Further AlI_3 (1g) is added after ~60g of I_2 are used, and boiling continued (total 2 hours) when all I_2 is consumed. ***Me₃SiI*** (94g, 94%) is collected by fractionation as stated above. [Eaborn *J Chem Soc* 3077 1950, DOI: 10.1039/JR9500003077; review *A versatile organic reagent* see Olah & Narang *Tetrahedron* **38** 2225 1982, DOI: 10.1016/0040-4020(82)87002-6; *Beilstein* **4** IV 4009.] It has also been obtained from ***bis(trimethylsilyl) selenide*** (1.0g, 7.4mmol, 4099-46-1, *via* syringe) which is added to a solution of I_2 (1.14, 4.5mmol) in *m*-xylene (1ml) under argon causing a mild exothermic reaction. ***Me₃SiI*** (1.6g, 94%) is distilled onto a Cu wire (colourless liquid, b 106-107°/atm). The residue in the pot when washed with Et_2O and dried was pure Se (0.35g, 100%). Store at -35° in a drybox in the dark. [Detty & Seidler *J Org Chem* **47** 1354 1982, DOI: 10.1021/jo00346a041.]

Isopropyldimethyl chlorosilane [3634-56-8] $C_5H_{13}ClSi$, M 136.7, b 109.8-110.0°/738mm, d_4^{20} 0.88, n_D^{20} 1.4158. Probable impurity is Me_3SiCl (b 56.9°/783mm) which can be removed by efficient fractional distillation. [Sommer et al. *J Am Chem Soc* **76** 801 1954, DOI: 10.1021/ja01632a050; *Beilstein* **4** IV 4067.]

N*-Isopropylidene-*N'*-2-nitrobenzenesulfonyl hydrazine (IPNSBH, isopropylidenehydrazide *o*-nitrobenzenesulfonic acid)** [6655-27-2] $C_9H_{11}N_3O_4S$, M 257.3, m 132-135°, 139-140°. IPNSBH is more stable than ***NBSH (*vide infra*) from which it is readily prepared by dissolving it (1.14g, 1 equivalent) in Me_2CO (8ml) at 0°, and after 30 minutes the solvent is removed *in vacuo*, the residue is re-dissolved in Me_2CO (4ml), and added slowly into 150ml of hexanes. The fine powder is filtered off, rinsed with hexanes (2 x 5ml) and dried *in vacuo* to give ***IPNSBH*** as a white solid (1.20g, 89%; TLC has R_F 0.7, 66% $EtOAc$ in hexanes). Its FTIR (thin film) has ν_{max} at 1177s, 1347m, 1375s, 1551s, 3264s cm^{-1} ; its 1H NMR [400MHz, $CDCl_3$, 20°] has δ at 8.30-8.28 (m, 1H), 7.87-7.85 (m, 2H), 7.79-7.77 (m, 2H), 1.96 (s, 3H), 1.92 (s, 3H), and its ^{13}C NMR [500MHz, $CDCl_3$, 20°] has δ at 158.9, 134.2, 133.5, 132.9.2, 132.0, 125.4, 25.5, 17.2. ***IPNSBH*** promotes the same reactions as ***NBSH*** except that it is more stable. A solution of 0.02M ***IPNSBH*** in DMSO is undecomposed at 50° during 30 minutes, whereas ***NBSH*** is 60% decomposed, and at is stable at 75° or 100° for 30 minutes so it can be stored at room temperature for several months without deterioration; all the same store it under argon (as it is possibly flammable), and use gloves due to its **toxicity**. It is a reagent used for allylic transposition, debromination and reduction of alcohols in the presence of DEAD and Ph_3P . [Movassaghi et al. *Angew Chem Int Ed* **45** 5859 2006, DOI: 10.1002/anie.200602011; Movassaghi & Ahmad *J. Org. Chem.* **72** 1838 2007, DOI: 10.1021/jo062325p].

***N*-Lauroyl-*N*-methyltaurine sodium salt (sodium *N*-decanoyl-*N*-methyl-2-aminoethane sulfonate)** [4337-75-1] $C_{15}H_{30}NNaO_4S$, M 344.5, pK_{Est} ~1.5. It is prepared from methyldecanoate (at 180° under N_2) or decanoyl chloride and sodium *N*-methylethane sulfonate and purified by dissolving it in H_2O and precipitating by addition of Et_2O . It decomposes on heating. It is a useful surfactant detergent [Desseigne & Mathian *Mém Services Chim Etat Paris* **31** 359 1944, cf. *Chem Abstr* **41** 705 1947.]

Lawesson's Reagent [LR, 2,4-bis(4-methoxyphenyl)1,3,2,4-dithiadiphosphetane 2,4-disulfide, *p*-methoxyphenylthiophosphonic acid cyclic di(thioanhydride)] [19172-47-5] $C_{14}H_{14}O_2P_2S_4$, M 404.5, m 228-229.5° (sintering at 215°), 228-230°. The reagent has been washed with anisole or $C_6H_6^*$ and dried in a vacuum over paraffin wax and P_2O_5 , and heated above room temperature. It does not lose the solvents completely (possibly due to formation of clathrates). It loses anisole on heating at ~230°. Analytically pure reagent was obtained by recrystallisation from *o*-dichlorobenzene, washing the crystals with $C_6H_6^*$ and drying *in vacuo* at 150°. It is extremely **moisture sensitive**, liberating H_2S , and should be stored in sealed containers, and preferably weighed in a dry-box. Its molecular weight is consistent with the formula $(C_7H_7OPS_2)_2$ but slowly polymerises in solution at ~80-85°. **Polymerisation** is more rapid in polar than non-polar solvents. It is a very useful **thiating reagent** particularly for converting hydroxy to thiol substituents. [Nishio *JCS Perkin Trans I* 1113 1993, DOI: 10.1039/P19930001113; Nair et al. *Tetrahedron Lett* **47** 9329 2006, DOI: 10.1016/j.tetlet.2006.10.115; *Beilstein* **16** IV 1113.]

Lead diethyldithiocarbamate [17549-30-3] $C_{10}H_{20}N_2PbS_4$, M 503.7, pK_1^{25} 3.36 (for *N,N*-diethyldithio-

carbamate), decomposes on heating liberating toxic fumes. Wash it with H₂O and dry it at 60-70°, or dissolve it in the minimum volume of CHCl₃ and add the same volume of EtOH. Collect the solid that separates and dry it as before. *Alternatively*, recrystallise it by slow evaporation of a CHCl₃ solution at 70-80°. Filter the crystals, wash them with H₂O until all Pb²⁺ ions are eluted (check by adding chromate) and then dry it at 60-70° for at least 10 hours. [Lo et al. *Analyt Chem* **49** 1146 1977, DOI: 10.1021/ac50016a021.] It is **POISONOUS**.

Lissamine Green B {1-[bis-(4,4'-dimethylaminophenyl)methyl]-2-hydroxynaphthalene-3,6-disulfonic acid sodium salt, Acid Green 50, Wool Green S} [3087-16-9] C₂₇H₂₅N₂NaO₇S₂, M 576.6, m >200°(dec), CI 44090, λ_{max} 633nm. Crystallise it from EtOH/water (1:1, v/v). It is a food additive that can be used in mint sauce, desserts, sweets, ice creams and tinned peas, but has been prohibited in several countries. **Irritant**. [Beilstein **14** II 574.]

Lissapol C (mainly sodium salt of 9-octadecene-1-sulfate) [2425-51-6] C₁₈H₃₆NaO₄S, M 387.6. Reflux the salt with 95% EtOH, then filter to remove insoluble inorganic electrolytes. The alcoholic solution is then concentrated, and the residue is poured into dry acetone. The precipitate is filtered off, washed in acetone and dried under vacuum. It is an anionic surfactant detergent, and the effect of NaCl or non-ionic surfactants, e.g. polyethylene glycol (PEG), on its CMC (critical micellar concentration) has been investigated. [Biswas & Mukerji *J Phys Chem* **64** 1 1960, DOI: 10.1021/j100830a001].

Lissapol LS (mainly sodium salt of anisidine sulfate) [28903-20-0]. Reflux the salt with 95% EtOH, then filter to remove insoluble inorganic electrolytes. The alcoholic solution is then concentrated, and the residue is poured into dry acetone. The precipitate is filtered off, washed in acetone and dried under vacuum. It is an anionic surfactant detergent, and the effect of NaCl or non-ionic surfactants, e.g. polyethylene glycol (PEG), on its CMC (critical micellar concentration) has been investigated. [Biswas & Mukerji *J Phys Chem* **64** 1 1960, DOI: 10.1021/j100830a001].

Lithium dodecylsulfate [2044-56-6] C₁₂H₂₅LiO₄S, M 272.3. Recrystallise this detergent twice from absolute EtOH and dry it under vacuum. Critical Micellar Concentration (CMC) in H₂O is 8.77 x 10⁻³M (7-10mM at 20-25°). It is an anionic detergent that can be used instead of SDS for electrophoresis at low temperatures. [Mukerjee et al. *J Phys Chem* **71** 4166 1967, DOI: 10.1021/j100872a702; Mysels & Dulin *J Colloid Sci* **10** 461 1955, DOI: 10.1016/0095-8522(55)90064-7; Beilstein **1** IV 1847.]

Lithium trimethylsilanolate (trimethylsilanol Li salt) [2004-14-0] C₃H₉LiOSi, M 96.1, m 120°(dec in air). Wash it with Et₂O and petroleum ether. It sublimes at 180°/1mm as fine transparent needles. [Tatlock & Rochow *J Org Chem* **17** 1555 1952, DOI: 10.1021/jo50012a001; Beilstein **4** IV 3992.] Suspected **CARCINOGEN**.

Magnesium dodecylsulfate [3097-08-3] C₁₂H₂₅MgO₄S, M 555.1. Recrystallise it three times from EtOH and dry it in a vacuum. [Beilstein **1** I 1788, **1** IV 1849.]

Magnesium trifluoromethanesulfonate [60871-83-2] (CF₃SO₃)₂Mg, M 322.4, m >300°. Wash it with CH₂Cl₂ and dry it at 125°/2 hour and 3mmHg. An efficient catalyst for preparing dithioketals under mild conditions [Corey & Shimoji *Tetrahedron Lett* **24** 169 1983, DOI: 10.1016/S0040-4039(00)81357-X]. [Beilstein **3** IV 34.]

Manganous ethylenebis(dithiocarbamate) (Maneb) [12427-38-2] C₄H₆MnN₄S₄ (monomer), M 293.3, pK_{Est} ~ 3.0 (for -NCSSH). Crystallise this agriculture fungicide from EtOH. [Pease & Holt *J Agric Food Chem* **25** 561 1977, DOI: 10.1021/jf60211a043.] It is soluble in CHCl₃. It is a skin **irritant**. [Beilstein **4** III 149, **4** IV 234.]

1R,2S,5R-Menthyl phosphorochloridite (*l*-MenOPCl₂) [95456-31-8] C₁₀H₁₉Cl₂PO, M 257.1, b 110-112°/260Pa, (62°/150mm also reported). MenOPCl₂ is prepared by adding a solution of (*1R,2S,5R*)-(-)-*menthol* (78g, 0.50mol) in CH₂Cl₂ (50ml, or THF) dropwise to a solution of PCl₃ (137.5g, 1.0mol) in CH₂Cl₂ (50ml, or THF) during 30 minutes. The mixture is stirred for 1 hour under a N₂ atmosphere, the volatiles are

removed at $\sim 25^\circ/100\text{mm}$, and pure **MenOPCl₂** distilled at $\sim 62^\circ/150\text{mm}$ quantitatively. It has ^1H NMR (400MHz, C₆D₆, TMS) with δ at 0.64 (m, 1H, H-4), 0.77 (m, 10H, H-3, H-8, H-9, H-10), 1.15 (M, 3H, H-2, H-5, H-6), 1.44 (m, 2H, H-3, H-4), 1.98 (M, 1H, H-7), 2.30 (M, 1H, H-6), 4.40 (m, 1H, H-1); the $^{13}\text{C}\{^1\text{H}\}$ NMR (100MHz, C₆D₆, TMS) with δ at 16.1 (s, C-8), 21.2 (s, C-9), 22.0 (s, C-10), 22.8 (s, C-3), 25.1 (s, C-7), 31.4 (s, C-5), 33.8 (s, C-4), 43.3 (s, C-6), 48.7 (s, C-2), 83.5 (d, $^2J_{\text{PC}} = 9.8\text{Hz}$, C-1); and the ^{31}P NMR (121.4MHz, THF, external ref 85% H₃PO₄) with δ at 175.6 or (at 162MHz, C₆D₆) with δ at 176.1 (d, $^3J_{\text{PH}} = 13.5\text{Hz}$). This **resolving agent** (generally used successfully in solution without strict purification) is stable for weeks under inert atmosphere, but is very sensitive to moisture. [Brunel & Buono *J Org Chem* **58** 7313 1993, DOI: 10.1021/jo00077a072; Totland et al. *Macromolecules* **29** 6114 1996, DOI: 10.1021/ma960351r]. Note that Hey-Hawkins and co-workers found that a 5 molar excess of PCl₃ in a solvent-free system gave 98% of **menthyl phosphorodi-chloridite (l-MenOPCl₂)**, **b 110-112°/260Pa**; but when a 1:0.5 ratio of menthol to PCl₃ was used a 65% yield of **bis-menthyl phosphorochloridite [(l-MenO)₂PCl]**, **m 53-55°, b 130°/0.4Pa**, was obtained [Hey-Hawkins et al. *Eur J Inorg Chem* 2776 2009, DOI: 10.1002/ejic.200900304].

2-Mercaptopyridine N-oxide sodium salt (pyridinethione or pyrithione sodium salt) [3811-73-2] **C₅H₄NOS**, **M 149.1**, **m $\sim 250^\circ(\text{dec})$** , **pK₁ -1.95**, **pK₂ 4.65**. When recrystallised from water, it assayed as 98.7% pure based on AgNO₃ titration [Krivis et al. *Anal Chem* **35** 966 1963, DOI: 10.1021/ac60201a013; see also Oliveri-Vigh & Karageozian *Anal Chem* **48** 1001 1976, DOI: 10.1021/ac60371a021; and Barton & Crich *JCS Perkin Trans 1* 1603, 1613 1986, DOI: 10.1039/P19860001603; Reactions: Crich *Aldrichimica Acta* **20** 35 1987; derived free radicals: Ingold et al. *Tetrahedron Lett* **29** 917 1988, DOI: 10.1016/S0040-4039(00)82481-8]. [Beilstein **21/7** V 151.]

Metanilic acid (3-aminobenzenesulfonic acid) [121-47-1] **C₆H₇NO₃S**, **M 173.2**, **m $<300^\circ(\text{dec})$** , **301-306°(sesquhydrate, dec)**, **pK₁²⁵ <1** , **pK₂²⁵ 3.74**. Crystallise the acid from water (as the **hydrate**, solubility is $\sim 2.4\text{w/v\%}$ at 15°), under CO₂ in a semi-darkened room. The solution is **photosensitive**. Dry it over 90% H₂SO₄ in a vacuum desiccator. The solubility in H₂O of the **anhydrous** amino acid is $\sim 0.9\text{w/v\%}$ at 0° and $\sim 7\text{w/v\%}$ at $\sim 90^\circ$. The **sodium salt**, crystallises from H₂O, and decomposes above 300° . [Beilstein **14** H 688, **14** IV 2640.]

Metanil Yellow (3[{4-phenylamino}phenylazo]-benzenesulfonic acid) [587-98-4] **C₁₈H₁₄N₃NaO₃S**, **M 375.4**, **pK_{Est} <0** . It can be salted out from water three times with sodium acetate, then repeatedly extracted with EtOH [McGrew & Schneider *J Am Chem Soc* **72** 2547 1950, DOI: 10.1021/ja01162a059]. It is an indicator (2 drops of a 0.1% aqueous solution in 10ml H₂O) in the pH range of 1.5 (red) to 2.7 (yellow) with λ_{max} at 414nm. [Beilstein **16** II 168.]

(Methoxycarbonylmethyl)triphenylphosphorane [methyl (triphenylphosphoranylidene)-acetate] [2605-67-6] **M 334.4**, **m 162-163°, 168-172°, 169-171°**. Crystallise this **phosphorus ylide** by dissolving in it AcOH and adding petroleum ether (b 40-50°) to give colourless plates which are dried at $50^\circ/\text{water-pump vacuum}$. Its UV has λ_{max} ($A_{1\text{mm}}^{1\%}$) at 222nm (908) and 268nm (125) [Isler et al. *Helv Chim Acta* **40** 1242 1957, DOI: 10.1002/hlca.19570400515]. [Fieser **1** 112; Beilstein **16** IV 977.] It is a useful two-carbon homologation **Wittig reagent** for conversion of aldehydes to the respective α,β -unsaturated esters as in the synthesis of (-)-mellein and (+)-ramulosin [Islam et al. *Tetrahedron* **63** 1074 2007, DOI: 10.1016/j.tet.2006.11.068] and in the efficient preparation of pyrazoles *via* cascade reactions of *N*- and *P*- containing ylides with methyl diazoacetate/Et₃N [Tomilov et al. *Tetrahedron Lett* **48** 883 2007, DOI: 10.1016/j.tetlet.2006.11.133].

Methoxycarbonylmethyltriphenylphosphonium bromide [1779-58-4] **C₂₁H₂₀BrO₂P**, **M 415.3**, **m 163°, 165-170°(dec)**. Wash it with petroleum ether (b 40-50°), recrystallise it from CHCl₃/Et₂O and dry it at high vacuum at 65° . [Isler et al. *Helv Chim Acta* **40** 1242 1957, DOI: 10.1002/hlca.19570400515; Wittig & Haag *Chem Ber* **88** 1654, 1664 1955, DOI: 10.1002/cber.19550881110; Beilstein **16** IV 988.]

Methoxymethyl trimethylsilane (trimethylsilylmethyl methyl ether) [14704-14-4] **C₅H₁₄OSi**, **M 118.3**, **b 83°/740mm**, **d₄²⁵ 0.758**, **n_D²⁵ 1.3878**. It forms an azeotrope with MeOH (b 60°). If it contains MeOH (check IR for bands above 3000cm^{-1}), then wash with H₂O and fractionate. A possible impurity could be **chloromethyl trimethylsilane** (b $97^\circ/740\text{mm}$). [Speier *J Am Chem Soc* **70** 4142 1948, DOI: 10.1021/ja01192a049; Beilstein **4**

III 1844.] It is an **IRRITATING FLAMMABLE** liquid.

1-Methoxy-2-methyl-1-trimethylsiloxypropene (dimethyl ketene methyl trimethylsilyl acetal) [31469-15-5] $\text{C}_8\text{H}_{18}\text{O}_2\text{Si}$, **M 174.3**, **b 35-36°/15mm, 140-142°/atm**, d_4^{20} **0.858**, n_D^{20} **1.4150**. Add Et_2O , wash with cold H_2O , dry (Na_2SO_4), filter, evaporate Et_2O , and distil the oily residue in a vacuum. It is moisture sensitive. [Ainsworth et al. *J Organometal Chem* **46** 59 1972, DOI: 10.1016/S0022-328X(00)90475-3.]

trans-1-Methoxy-3-(trimethylsilyloxy)-1,3-butadiene (Danishefsky's diene) [54125-02-9] $\text{C}_8\text{H}_{16}\text{O}_2\text{Si}$, **M 172.3**, **b 68-69°/14mm, 70-72°/16mm**, d_4^{20} **0.885**, n_D^{20} **1.4540**. It may contain up to 1% of the precursor 4-methoxybut-4-ene-2-one. It is easily purified by distilling through a Vigreux column in a vacuum and taking the middle fraction. [Danishefsky & Kitihara *J Am Chem Soc* **96** 7807 1974, DOI: 10.1021/ja00832a031; Danishefsky *Acc Chem Res* **14** 400 1981, DOI: 10.1021/ar00072a006.] A useful electron rich functionalised Diels-Alder diene [Teyssot et al. *J Org Chem* **72** 2364 2007, DOI: 10.1021/jo062186b; Dai et al. *J Am Chem Soc* **129** 645 2007; DOI: 10.1021/ja065762u].

Methylarsonic acid [124-58-3] CH_3AsO_3 , **M 139.97**, **m 159.8°, 161°, b 393°/atm**, pK_1^{25} **4.58**, pK_2^{25} **7.82** [$\text{As}(\text{OH})_2$]. Recrystallise this herbicide from Me_2CO or absolute EtOH (white plates; its solubility is 28g/100ml EtOH). Inhalation causes coughing and soreness of throat — **TOXIC**. [Quick & Adams *J Am Chem Soc* **44** 805 1922, DOI: 10.1021/ja01425a014; *Beilstein* **4** H 613, **4** I 577, **4** II 996, **4** III 1822, **4** IV 3682.]

Methyl dichlorosilane (dichloro methylsilane) [75-54-7] $\text{CH}_3\text{Cl}_2\text{Si}$, **M 115.0**, **m -93°, -92.5°, b 41°/748mm, 40.9°/760mm, 40-45°/atm**, d_{27}^{27} **1.105**. Impurities are generally other chloromethyl silanes. Distil it through a conventional Stedman column of 20 theoretical plates or more. It should be protected from H_2O by storing over P_2O_5 . [Stock & Somieski *Chem Ber* **52** 695 1919, DOI: 10.1002/cber.19190520410; Sauer et al. *J Am Chem Soc* **68** 962 1946, DOI: 10.1021/ja01210a014; *Beilstein* **4** IV 4096.]

N-Methyl-N-nitroso-p-toluenesulfonamide (Diazald) [80-11-5] $\text{C}_8\text{H}_{10}\text{N}_2\text{O}_3\text{S}$, **M 214.2**, **m 58°, 61-62°. 62°**. Crystallise diazald from *benzene by addition of petroleum ether and store it in a refrigerator. It is soluble in most organic solvents including Et_2O , and liberates *diazomethane* on treatment with alkali. Store it in the cold. It remains active for many months if it is kept in a stoppered bottle in a cold room below 10°. [de Boer & Backer *Org Synth* **34** 96 1954, DOI: 10.15227/orgsyn.034.0096; *Org Synth Coll Vol* **4** 250 1963, DOI: 10.15227/orgsyn.036.0016; *Beilstein* **11** I 29.] Reagent for the preparation of *diazomethane*. It has also found use in the *nitrosation* of molybdenum and tungsten complexes [Legzdins et al. *Organometallics* **12** 1029 1993, DOI: 10.1021/om00028a017], and in *kinetic studies* of Diazald with *amines* [Garcia-Rio et al. *JCS Perkin 2* 29 1993, DOI: 10.1039/P29930000029] and *carbanions* [Leis et al. *JCS Perkin 2* 1233 1993, DOI: 10.1039/P29930001233].

Methyl Orange (sodium 4,4'-dimethylaminophenylazobenzenesulfonate, Orange III) [547-58-0] $\text{C}_{14}\text{H}_{14}\text{N}_3\text{NsO}_3\text{S}$, **M 327.3**, pK_1^{25} **3.56**, pK_2^{25} **6.49**. Recrystallise the orange-yellow powder twice from hot water (solubility at ~20° is 0.2w/w%), then wash it with a little EtOH followed by diethyl ether. It is an acid-base indicator (few drops of 0.1% aqueous solution): pH 3.1 (red) and pH 4.4 (yellow). [*Beilstein* **16** IV 510.]

Methylphenyl dichlorosilane (dichloro methyl phenylsilane) [149-74-6] $\text{C}_7\text{H}_8\text{Cl}_2\text{Si}$, **M 191.1**, **b 114-115°/50mm, 202-205°/atm, 205-206°/760mm**, d_4^{20} **1.173**, n_D^{20} **1.519**. Purify it by fractionation using an efficient column. It hydrolyses *ca* ten times more slowly than methyltrichlorosilane and *ca* sixty times more slowly than phenyltrichlorosilane. [Shaffer & Flanigen *J Phys Chem* **61** 1591 1957, DOI: 10.1021/j150558a004; *Beilstein* **16** III 1211, **16** IV 1517.]

Methylphosphonic acid [993-13-5] $\text{CH}_3\text{O}_3\text{P}$, **M 96.0**, **m 104-106°, 106°, 105-107°, 108°, 108.5°, pK_1^{25} 2.12, pK_2^{25} 7.29**. If it tests for Cl^- , then add H_2O and evaporate to dryness, repeat several times till free from Cl^- . The residue solidifies to a wax-like solid. *Alternatively*, dissolve the acid in the minimum volume of H_2O , add charcoal, warm, filter and evaporate to dryness in a vacuum over P_2O_5 . [Crofts & Kosolapoff *J Am Chem Soc* **75** 3379 1953, DOI: 10.1021/ja01110a024.] The *di-Na salt* is prepared from 24g of acid in 50ml of dry EtOH , and a solution of 23g Na dissolved in 400ml EtOH is added. A white precipitate is formed, but the mixture is

refluxed for 30 minutes to complete the reaction. Filter off the solid and recrystallise it from 50% EtOH. Dry the crystals in a vacuum desiccator. [Thompson *J Chem Soc* 3292 1952, DOI: 10.1039/JR9520003292; *Beilstein* 4 IV 3498.]

Methylphosphonic dichloride [676-97-1] $\text{CH}_3\text{Cl}_2\text{OP}$, M 132.9, m 33°, 33-37°, b 53-54°/10mm, 64-67°/20.5mm, 86°/44mm, 162°/760mm, d_4^{40} 1.4382. Fractionally redistil it until the purity as checked by hydrolysis and acidimetry for Cl^- is correct and the distillate should solidify on cooling. [Kinnear & Perren *J Chem Soc* 3437 1952, DOI: 10.1039/JR9520003437; Crofts & Kosolapoff *J Am Chem Soc* 75 3379 1952, DOI: 10.1021/ja01110a024; for IR see McIvor et al. *Can J Chem* 34 1611 1956, DOI: 10.1139/v56-211; Svara, Weferling & Hofmann *Phosphorus Compounds, Organic*, in *Ullmann's Encyclopedia of Industrial Chemistry*, Wiley-VCH, Weinheim, 2008; DOI: 10.1002/14356007.a19_545; L. Maier, *Organic Phosphorus Compounds*, 90.1 'A convenient one-step synthesis of alkyl- and arylphosphonyl dichlorides' in *Phosphorus, Sulfur, and Silicon and the Related Elements* 47 (3-4) 465-470 1990, DOI: 10.1080/10426509008038002; *Beilstein* 4 IV 3509.] **TOXIC**

Methyl Thymol Blue, sodium salt {MTB, 3,3'-bis[N,N'-di(carboxymethyl)amino]thymolsulfonephthalein sodium salt} [1945-77-3] $\text{C}_{37}\text{H}_{44}\text{N}_2\text{O}_{13}\text{S}$, M 844.8, ϵ 1.89×10^4 at 435nm, pH 5.5, pK_1^{25} 3.0, pK_2^{25} 3.3, pK_3^{25} 3.8, pK_4^{25} 7.4 (pK_4^{25} 7.2), pK_5^{25} 11.5 pK_6^{25} 13.4. The starting material for synthesis is **Thymol Blue**. Purify it as for **Xylenol Orange** [1611-35-4], see entry below in this Chapter. [Tereshin et al. *J Anal Chem USSR (Engl Trans)* 20 1138 1965, Körbl & Kakáč *Coll Czech Chem Commun* 23 889 1958, DOI: org/10.1135/cccc19580889; *Beilstein* 19/8 V 619.]

Methyl 4-toluenesulfonate [80-48-8] $\text{C}_8\text{H}_{10}\text{O}_3\text{S}$, M 186.2, m 25-28°, 28°, b 144.6-145.2°/5mm, 168-170°/13mm, d_4^{20} 1.23, n_D^{20} 1.5172. The ester is purified by distillation *in vacuo* and can be crystallised from petroleum ether or Et_2O /petroleum ether at low temperature. It is a powerful **methylating agent**, is **TOXIC** and is a **skin irritant**, so it is better to purify it by repeated distillation. [IR: Schreiber *Anal Chem* 21 1168 1949, DOI: 10.1021/ac60034a005; Buehler et al. *J Org Chem* 02 167 1937, DOI: 10.1021/jo01225a004; Roos et al. *Org Synth Coll Vol* 1 145 1941, DOI: 10.15227/orgsyn.009.0028; *Beilstein* 11 IV 247.]

Methyl trichlorosilane [75-79-6] $\text{CH}_3\text{Cl}_3\text{Si}$, M 149.5, b 13.7°/101mm, 64.3°/710.8mm, 65.5°/745mm, 66.1°/atm, d_4^{20} 1.263, n_D^{20} 1.4110. If not very pure, distil it before use. The purity is checked by ^{29}Si NMR (δ in MeCN is 13.14 ppm with respect to Me_4Si). Possible contaminants are other silanes which can be removed by fractional distillation through a Stedman column of >72 theoretical plates with total reflux and 0.35% take-off. This apparatus should be under N_2 gas at a rate of 12 bubbles/minute fed into the line using an Hg manometer to control the pressure. It is sensitive to H_2O . [Gilliam et al. *J Am Chem Soc* 73 4252 1951, DOI: 10.1021/ja01153a064; Olah et al. *J Org Chem* 48 3667 1983, DOI: 10.1021/jo00169a010; *Beilstein* 4 IV 4212.]

Methyl triethoxysilane [2031-67-6] $\text{C}_7\text{H}_{18}\text{O}_3\text{Si}$, M 178.31, b 142-144.5°/742mm, 141°/765mm, 141.5°/775mm, d_4^{20} 0.8911, n_D^{20} 1.3820. Purify by fractionating it repeatedly in a stream of N_2 through a 3' Heligrid packed Todd column. It is hydrolysed by H_2O and yields cyclic polysiloxanes on hydrolysis in the presence of acid in $^*\text{C}_6\text{H}_6$. [McBee et al. *J Am Chem Soc* 77 1292 1955, DOI: 10.1021/ja01610a066; Sprung & Guenther *J Am Chem Soc* 77 3990 1955, DOI: 10.1021/ja01620a013; *Beilstein* 4 IV 4204.]

Methyl trimethoxysilane [1185-55-3] $\text{C}_4\text{H}_{12}\text{O}_3\text{Si}$, M 136.2, has b 102°/760mm, 102-104°/atm, d_4^{20} 1.3687, n_D^{20} 1.3711. Likely impurities are **1,3-dimethyltetramethoxy disiloxane** (b 31°/1mm) and cyclic polysiloxanes, see previous entry methyl triethoxysilane. It hydrolyses in both aqueous acidic and basic solutions. [Tamborski & Post *J Org Chem* 17 1400 1952, DOI: 10.1021/jo50010a022; Seyferth & Rochow *J Org Chem* 20 250 1955, DOI: 10.1021/jo01120a016; *Beilstein* 4 IV 4203.]

N-Methyl-N-trimethylsilylacetamide [7449-74-3] $\text{C}_6\text{H}_{15}\text{NOSi}$, M 145.3, b 48-49°/11mm, 84°/13mm, 105-107°/35mm (solid at room temperature), 159-161°/atm, d_4^{20} 0.90, n_D^{20} 1.4379. A likely impurity is $\text{Et}_3\text{N.HCl}$ which can be detected by its odour. If it is completely soluble in $^*\text{C}_6\text{H}_6$, then redistil; otherwise dissolve it in this solvent, filter and evaporate first in a vacuum at 12mm, then fractionate; all operations should be carried out in a dry N_2 atmosphere. [Klebe et al. *J Am Chem Soc* 88 3390 1966, DOI: 10.1021/ja00966a038; Ried & Suarez-Rivero *Chem Ber* 96 1475 1963, DOI: 10.1002/cber.19630960544; *Beilstein* 4 IV 4011.]

Methyl trimethylsilylacetate [2916-76-9] $C_6H_{14}O_2Si$, M 146.3, b 38-39°/13mm, 65-68°/50mm, d_4^{20} 0.89. Dissolve it in Et_2O , shake with 1M HCl, wash with H_2O , aqueous saturated $NaHCO_3$, H_2O again, and dry it (a precipitate may be formed in the $NaHCO_3$ solution and should be drawn off and discarded). The solvent is distilled off, and the residue is fractionated through a good column. The IR ($CHCl_3$) has ν_{max} at $1728cm^{-1}$. [Fessenden & Fessenden *J Org Chem* **32** 3535 1967, DOI: 10.1021/jo01286a054; Matsuda et al. *J Org Chem* **45** 237 1980, DOI: 10.1021/jo01290a006; *Beilstein* **4** III 1855, **4** IV 3974 for Et ester.]

Methyl (±)-2-(trimethylsilyl)propionate [55453-09-3] $C_7H_{16}O_2Si$, M 160.3, m 32°, b 142.3 ±13°/760mm, 155-157°/atm, d_4^{20} 0.89. Dissolve it in Et_2O , wash it with aqueous $NaHCO_3$, H_2O , 0.1M HCl, H_2O again, dry ($MgSO_4$), evaporate and distil it. [Emde & Simchen *Synthesis* 867 1977, DOI: 10.1055/s-1977-24611; Oppolzer et al. *Tetrahedron* **39** 3695 1983, DOI: 10.1016/S0040-4020(01)88608-7; Crimmin et al. *JCS Perkin Trans 1* 541 1985, DOI: 10.1039/P19850000541.]

N-Methyl-N-trimethylsilyl trifluoroacetamide [24589-78-4] $C_6H_{12}F_3NOSi$, M 199.3, m 98-100°, b 78-79°/130mm, 131-132°/atm. Fractionate the amide through a 40mm Vigreux column. Usually it contains *ca* 1% of methyl trifluoroacetamide and 1% of other impurities which can be removed by gas chromatography or fractionating using a spinning band column. It is *moisture sensitive*, store appropriately. [Donike *J Chromatogr* **42** 103 1969, DOI: 10.1016/S0021-9673(01)80592-6; Donike *J Chromatogr* **103** 91 1975, DOI: 10.1016/S0021-9673(00)83805-4; *Beilstein* **4** IV 4011.]

Methyl triphenoxyphosphonium iodide (triphenylphosphite methiodide) [17579-99-6] $C_{19}H_{18}IO_3P$, M 452.2, m 114° (sealed tube), 146°. Gently heat the impure iodide with good grade Me_2CO . The saturated solution obtained is decanted rapidly from undissolved salt and treated with an equal volume of dry Et_2O . The iodide separates as flat needles which are collected by centrifugation, washed several times with dry Et_2O , and dried in a vacuum over P_2O_5 . For this recrystallisation, it is essential to minimise the time of contact with Me_2CO , and to work rapidly and with rigorous exclusion of moisture. If the crude material is to be used, it should be stored under dry Et_2O , and dried and weighed *in vacuo* immediately before use. [Landauer & Rydon *J Chem Soc* 2224 1953, DOI: 10.1039/JR9530002224; for NMR data see: Hudson et al. *JCS Perkin Trans 1* 982 1974, DOI: 10.1039/P19740000982; *Beilstein* **6** IV 704.]

Methyl triphenylphosphonium bromide [1779-49-3] $C_{19}H_{18}BrP$, M 357.2, m 229-230°(corr), 227-229°, 230-233°, 231-234°, 233°. If the solid is sticky, wash it with $*C_6H_6$ and dry it in a vacuum over P_2O_5 . [Marvel & Gall *J Org Chem* **24** 1494 1959, DOI: 10.1021/jo01092a026; Chem Ber **87** 1318 1954, Milas & Priesing *J Am Chem Soc* **79** 6295 1957, DOI: 10.1021/ja01580a049; Wittig & Schöllkopf *Org Synth* **40** 66 1960, DOI: 10.1002/cber.19540870919.] The *iodide*, on recrystallisation from H_2O , has m 187.5-188.5° [Mann et al. *J Chem Soc* 1130 1953, DOI: 10.1039/JR9530001130; Wittig & Geissler *Justus Liebigs Ann Chem* **580** 44 1953, DOI: 10.1002/jlac.19535800107]. [*Beilstein* **16** IV 981.]

Methyl vinyl dichlorosilane (dichloro methyl vinyl silane) [124-70-9] $C_3H_6Cl_2Si$, M 141.1, b 43-45.5°/11-11.5mm, 91°/742mm, 92.5°/743.2mm, 92.5-93°/atm, d_4^{20} 1.0917, n_D^{20} 1.444. Likely impurities are dichloromethyl-silane, butadienyl-dichloromethylsilane. Fractionate the silane through a column packed with metal filings (20 theoretical plates) at atmospheric or better under reduced pressure. [Shostakovskiy et al. *Izv Akad Nauk SSSR Ser Khim* **6** 1474 1957, DOI: 10.1007/BF01169751; Mironov & Petrov *Izv Akad Nauk SSSR Ser Khim* **7** 767 1958, DOI: 10.1007/BF00917295; *Beilstein* **4** III 1894, **4** IV 4184.]

Milling Red SWB {1-[4-[4-[4-toluenesulfonyloxy]phenylazo](3,3'-dimethyl-1,1'-biphenyl)-4'-azo]-2-hydroxynaphthalene-6,8-disulfonic acid di-Na salt, Acid Red 114} [6459-94-5] $C_{37}H_{28}N_4Na_2O_{10}S_3$, M 830.8, m dec >250°, CI 23635, λ_{max} ~514nm. Salt out three times with sodium acetate, then repeatedly extract it with $EtOH$ and dry the solid in air. [McGrew & Schneider *J Am Chem Soc* **72** 2547 1950, DOI: 10.1021/ja01162a059; *Beilstein* **16** II 140.] This textile Dye can also be used for staining epithelial intracellular bridges [Bredrick *Stain Technology* **39** 33 1964, DOI: 10.3109/10520296409061205]

Milling Yellow G {5-chloro-2-[3-methyl-4-[[4-(methylphenyl)sulfonyloxyphenyl]diazonyl]-5-oxo-4H-pyrazil-1-yl]benzenesulfonic acid Na-salt, Acid yellow 40} [6372-96-9, 51569-18-7, 88529-27-5] $C_{23}H_{18}ClN_4$

Na₂O₇S₂, M 582.0, CI 642, decomposes on heating. Salted out three times with sodium acetate, then repeatedly extracted with EtOH. [McGrew & Schneider *J Am Chem Soc* **72** 2547 1950, DOI: 10.1021/ja01162a059; *Beilstein* **16** II 125.] See also **Solochrome Violet R** [2092-55-9] below.

Naphthalene Scarlet Red 4R [1-(4-sulfonaphthalene-1-azo)-2-hydroxynaphthalene-6,8-disulfonic acid tri-Na salt, New Coccine, Ponceau 4R, Acid Red 18] [2611-82-7] **C₂₀H₁₁N₂Na₃O₁₀S₃, M 604.5, m>250°(dec), CI 16255, λ_{max} 350nm and 506nm.** Dissolve the dye in the minimum quantity of boiling water, filter and enough EtOH is added to precipitate *ca* 80% of the dye. This process is repeated until a solution of the dye in aqueous 20% pyridine has a constant extinction coefficient. [*Beilstein* **16** I 306.]

Naphthalene-1,4-disulfonic acid [92-41-1] **C₁₀H₈O₆S₂, M 288.2, pK_{Est} <0.** Crystallise the acid from concentrated HCl. The *disulfonamide* has **m 273°** (from EtOH). [*Beilstein* **11** II 119, **11** III 463.]

Naphthalene-1,5-disulfonic acid tetrahydrate (Armstrong's acid) [81-04-9 (anhydrous), 211366-30-2 (4 H₂O)] **C₁₀H₈O₆S₂, M 288.3, 360.4 (4 H₂O), has m 240-245° (anhydrous), d 1.704, pK_{Est} <0.** Crystallise the acid from concentrated HCl. It is soluble in EtOH and H₂O, but insoluble in Et₂O. [Lynch & Scanlan *Ind Eng Chem* **19** 1010 1927, DOI: 10.1021/ie50213a018; *Beilstein* **11** H 212.]

Naphthalene-1-sulfonic acid [85-47-2] **C₁₀H₈O₃S, M 208.2, m (hydrates) 97° to 79°, (2H₂O) 90°, (anhydrous) 139-140°, pK²⁰ -0.17.** Crystallise the acid from concentrated HCl and twice from H₂O. The *S-benzylisothiuronium salt* has **m 137°** (from aqueous EtOH). [*Beilstein* **11** H 155, **11** III 383, **11** IV 521.]

Naphthalene-1-sulfonyl chloride [85-46-1] **C₁₀H₇ClO₂S, M 226.7, has m 64-67°, 68°, b 147.5°/0.9mm, 194-195°/13mm.** If the IR indicates the presence of OH, then treat it with an equal weight of PCl₅ and heat it at *ca* 100° for 2 hours, cool and pour onto ice + H₂O, stir well and filter off the solid. Wash the solid with cold H₂O and dry the solid in a vacuum desiccator over P₂O₅ + solid KOH. Extract the solid with petroleum ether (b 40-60°), filter off any insoluble solid and cool. Collect the crystalline sulfonyl chloride and recrystallise it from dry Et₂O, petroleum ether or *C₆H₆/petroleum ether. If large quantities are available, then it can be distilled under high vacuum. [Fierz & Weissenbach *Helv Chim Acta* **3** 312 1920, DOI: 10.1002/hlca.19200030130.] The *sulfonamide* crystallises from EtOH (**m 150.5°**) or H₂O (**m 153°**). [*Beilstein* **11** H 175, **11** II 93, **11** IV 383.]

Naphthalene-2-sulfonic acid [120-18-3] **C₁₀H₈O₃S, M 208.2, has m 91° (anhydrous), 124-125° (anhydrous) pK_{Est} <1.** Crystallise the acid from concentrated HCl. The *S-benzylisothiuronium salt* has **m 192°** (from aqueous EtOH). [Berger **8** 432 1954, *Beilstein* **11** H 171, **11** IV 527.]

Naphthalene-2-sulfonyl chloride [93-11-8] **C₁₀H₇ClO₂S, M 226.7, has m 74-76°, 78°, 79°, 83°, b 147.7°/0.6mm, 201°/13mm.** Distil the chloride in a vacuum and/or recrystallise it (twice) from *benzene/petroleum ether (1:1 v/v). Purify it as the *1-sulfonyl chloride* above. [Fierz & Weissenbach *Helv Chim Acta* **3** 2312 1920, DOI: 10.1002/hlca.19200030130.] The *sulfonamide* [1576-47-2] **C₁₀H₉NO₂S, M 207.3, has m 217°** (from EtOH, **215° to 219°** also reported). [*Beilstein* **11** III 399, **11** IV 529.]

Naphthionic acid (4-aminonaphthalene-1-sulfonic acid, Piria's acid) [84-86-6] **C₁₀H₉NO₃S, M 223.3, m > 300°(dec), d₄²⁵ 1.673, pK²⁵ 2.68.** It crystallises from H₂O as needles of the *0.5 hydrate*. Its solubility in H₂O varies from 0.03w/v% at 0° to 0.22w/v% at 100°. Salt solutions fluoresce strongly blue. The *S-benzylisothiuronium salt* has **m 195°** (from aqueous EtOH). [*Beilstein* **14** IV 2793.]

Naphthol Yellow S (citronin A, Food Yellow 1, flavianic acid sodium salt, 8-hydroxy-5,7-dinitro-2-naphthalene sulfonic acid disodium salt) [846-70-8] **C₁₀H₄N₂Na₂O₈S, M 358.2, CI 10316, decomposes on heating.** This dye is a water-soluble greenish yellow powder. The *free sulfonic acid* can be recrystallised from dilute HCl (**m 150°**) or AcOH/EtOAc (**m 148-149.5°**). The disodium salt is then obtained by dissolving the acid in two equivalents of aqueous NaOH and evaporating to dryness, and drying the residue in a vacuum desiccator. The *sodium salt* can be recrystallised from the minimum volume of H₂O or from EtOH [Dermer & Dermer *J Am Chem Soc* **61** 3302 1939, DOI: 10.1021/ja01267a019]. It is a food colouring matter that has been banned by many countries [see https://www.sigmaaldrich.com/content/dam/sigma-aldrich/docs/Fluka/Brochure/1/food_color_additives.pdf]. [*Beilstein* **11** III 542.]

1,2-Naphthoquinone-4-sulfonic acid sodium salt (Folin's reagent, 3,4-dihydro-3,4-dioxo-1-naphthlene

sulfonic acid sodium salt) [521-24-4] $\text{C}_{10}\text{H}_5\text{NaO}_5\text{S}$, **M 260.2**, **m 289°(dec)**, **pK_{Est} <0**. It forms yellow crystals from aqueous EtOH and should be dried at 80° *in vacuo*. Its solubility in H_2O is 5%, and it is moderately soluble in Me_2CO , slightly soluble in EtOH but mostly insoluble in other organic solvents. It is a reagent for the detection and determination of amines, amino acids, peptides and proteins. Aqueous solutions of it fade in the presence of light, but addition of HCl slows down this process. [Folin (and Wu) *J Biol Chem* **51** 377 1922, <http://www.jbc.org/content/167/1/57>; Martin & Fieser *Org Synth Coll Vol* **3** 633 1955, DOI: 10.15227/orgsyn.021.0091; **estimation** of amino acid nitrogen in blood: Danielson *J Biol Chem* **101** 505 1933, <http://www.jbc.org/content/101/2/505>.] **UV estimation** of amines: Rosenblatt et al. *Anal Chem* **27** 1290 1955, DOI: 10.1021/ac60104a024]. [*Beilstein* **11** IV 668.]

2-Naphthylamine-1-sulfonic acid (Tobias acid) [81-16-3] $\text{C}_{10}\text{H}_9\text{NO}_3\text{S}$, **M 223.3**, **m >200°(dec)**, **pK₁²⁵ <1**, **pK₂²⁵ 2.35 (NH₂)**. Crystallise the acid under nitrogen from boiling water and dry it in a steam oven [Bryson *Trans Faraday Soc* **47** 522, 527 1951, DOI: 10.1039/TF9514700522]. [*Beilstein* **14** III 2240, **14** IV 2792.]

5-Naphthylamine-1-sulfonic acid (Laurent's acid) [84-89-9] $\text{C}_{10}\text{H}_9\text{NO}_3\text{S}$, **M 223.3**, has **m >200°(dec)**, **pK_{Est(1)} <1**, **pK₂²⁵ 3.69 (NH₂)**. Crystallise the acid under nitrogen from boiling water (solubility is 0.1w/v% at <10°) and dry it in a steam oven [Bryson *Trans Faraday Soc* **47** 522, 527 1951, DOI: 10.1039/TF9514700522]. [*Beilstein* **14** IV 2800.]

2-Naphthylamine-6-sulfonic acid (Broenner's acid) [93-00-5] $\text{C}_{10}\text{H}_9\text{NO}_3\text{S}$, **M 223.3**, has **m >200°(dec)**, **pK₂₅ 3.74**. Crystallise the acid from a large volume of hot water. The **diethylamine salt** has **m 190.5-192°** (from EtOH/*iso*BuOH), and the **S-benzylisothiuronium salt** has **m 330°** (from *n*BuOH). [*Beilstein* **14** H 760, **14** II 463, **14** III 2249, **14** IV 2804.]

1-Naphthyl thiourea (ANTU) [86-88-4] $\text{C}_{11}\text{H}_{10}\text{N}_2\text{S}$, **M 202.2**, **m 197.8°, 198°**. Crystallise ANTU from EtOH as colourless prisms. [*Beilstein* **12** III 2941, **12** IV 3086.] It is a rodenticide and fairly toxic.

2-Nitrobenzenesulfonyl chloride (NPS-Cl) [7669-54-7] $\text{C}_6\text{H}_4\text{ClNO}_2\text{S}$, **M 189.6**, **m 70° to 75°, 73-74.5°, 74.5-75°, 74-76°**. Recrystallise it from CCl_4 (2ml/g), filter off the solution at 5° (recovery 75%). It has also been recrystallised from petroleum ether (b 40-60°), dried rapidly at 50° and stored in a brown glass bottle, sealed well and stored away from moisture. [Hubacher *Org Synth Coll Vol* **2** 455 1943, DOI: 10.15227/orgsyn.015.0045; Ito et al. *Chem Pharm Bull Jpn* **26** 296 1978, *Beilstein* **6** I 157.]

2-Nitrobenzenesulfonyl chloride [1694-92-4] $\text{C}_6\text{H}_4\text{ClNO}_4\text{S}$, **M 221.6**, **m 63-67°, 68-69°, 69.5-69.7°, b 350.6°/760mm, d 1.606**. It is prepared by the oxidation of di-*o*-nitrophenyldisulfide with Cl_2 [Wertheim *Org Synth Coll Vol* **2** 471 1943, DOI: 10.15227/orgsyn.015.0055] or from *o*-nitrobenzenesulfonic acid and PCl_5 , then pouring into H_2O , washing the solid with H_2O , drying it *in vacuo* and recrystallising it from petrol, Et_2O /petrol, petroleum ether/ CCl_4 or petroleum ether. With ammonia it is converted to the **sulfonamide** [5455-59-4] **M 202.1**, **m 193°**, which forms needles from hot aqueous EtOH; and with phenol/pyridine or Na_2CO_3 , the **phenyl ester**, **m 57°**, is obtained which crystallises from EtOH. The **sulfonylazide**, **m 71-73°**, separates as needles from petroleum ether [Leffler & Tsuno *J Org Chem* **28** 902 1963, DOI: 10.1021/jo01039a004]. [*Beilstein* **11** H 67, **11** I 20, **11** II 32, **11** III 114, **11** IV 174.]

3-Nitrobenzenesulfonyl chloride [121-51-7] has **m 61-62°, 63.9-64.2°**. It can be prepared from *m*-nitrobenzenesulfonic acid and PCl_5 as for the isomer above and crystallised from petroleum ether/ CCl_4 or petroleum ether in needles. With ammonia it is converted to the **sulfonamide** [121-52-8] **M 202.1**, **m 167-168°**, which forms needles from hot aqueous EtOH and is a moderately strong base with **pK₂₀ 9.20**. With phenol/pyridine or Na_2CO_3 , the **phenyl ester**, **m 94°**, is obtained which crystallises from acetic acid. [Gilbert *Synthesis* 315 1977, DOI: 10.1055/s-1977-24376; *Beilstein* **11** H 69, **11** I 21, **11** II 33, **11** III 126, **11** IV 182.]

4-Nitrobenzenesulfonyl chloride (Nosyl chloride) [98-74-8] has **m 66-70°, 78-80°, 80°, b 143-144°/1.5mm, 180-181°/17mm**. It is prepared in the same way as for the *m*-isomer above from ***p*-nitro-benzenesulfonic acid**, and is purified by vacuum distillation and/or by recrystallisation from petroleum ether or Et_2O /petroleum ether. With ammonia it is converted to the **sulfonamide** [6325-93-5] **M 202.1**, **m 179-180°**, which forms prisms from hot aqueous EtOH and is a moderately strong base with **pK₂₀ 9.14**. [Prinsen et al. *Recl Trav Chim Pays-Bas* **84** 24 1965, DOI: 10.1002/recl.19650840104; *Beilstein* **11** H 72, **11** I 21, **11** II 34, **11** III 136, **11** IV 192.] It is highly efficient in solid phase peptide synthesis when using ***N*-nosyl- α -amino acids** [Leggio et al. *Tetrahedron* **63** 8164 2007, DOI: 10.1016/j.tet.2007.05.121.]

2-Nitrobenzenesulfonylhydrazine (NBSH) [606-26-8, 6655-77-2] $C_6H_7N_3O_4S$, **M 217.2, m 100-101°, 101° (dec)**. NBSH is unstable unless handled properly. If the material is suspect, it is preferable to prepare freshly thus: hydrazine monohydrate (12.1ml, 2.5 equivalents) is added to a solution of *o*-nitrobenzenesulfonyl chloride (22.2g, 1 equivalent) in dry THF (100ml) at -30° under argon whereby the solution became brown, white $N_2H_4 \cdot HCl$ deposited, and after 30 minutes TLC (2/1, EtOAc/hexanes) indicating that all the sulfonyl chloride was used up. EtOAc (200ml, at 23°) is added to the cold mixture and washed repeatedly with ice-cold 10% aqueous NaCl (5 x 150ml, contact time with each wash should not be <1 minute; the use of just distilled H_2O results in much lower yields). The organic layer is washed, dried (Na_2SO_4 at 0°), then added slowly to a stirred solution of hexanes (1.2L) at 23° during 5 minutes. The off-white NBSH that precipitates within 10 minutes is isolated by vacuum filtration, the solid is washed with hexanes (2 x 50ml, 23°) and dried at 23°/1.5mm for 14 hours to provide **pure NBSH** as an off-white powder (17.6g, 81%; TLC has R_F 0.19, EtOAc/hexanes 2/1). Store it at ~0-5° under argon, and use gloves due to its **toxicity**. Its IR (EtOAc) has ν_{max} at 1165, 1352, 1547, 3400—3100 cm^{-1} and its 1H NMR [300MHz, CD_3CN] has δ : 8.17-8.03 (1H, m), 7.91-7.78 (3H, m), 5.97 (1H, bs), 3.90 (2H, bs); and its ^{13}C NMR [75MHz, CD_3CN] has δ : 149.4, 135.5, 133.4, 133.2, 130.8 and 125.8. [Meyers et al. *J Org Chem* **62** 7507 1997, DOI: 10.1021/jo9710137; Dann & Davies *J Chem Soc* 1050 1929, DOI: 10.1039/JR9290001050; for use in the **Mitsunobu reaction** see: Hughes *Org React* **42** 335 1992, DOI: 10.1002/0471264180.or042.02.]

NBSH is a **versatile reagent** for synthesising allenes from propargyl alcohols [Meyers & Zheng *J Am Chem Soc* **118** 4492 1996], for reductive transposition of allylic alcohols [Myers & Zheng *Tetrahedron Lett* **37** 4841 1996, DOI: 10.1016/0040-4039(96)00965-3], and for deoxygenation of unhindered alcohols; each of which proceeds by **Mitsunobu displacement** of an alcohol with **NBSH** followed by *in situ* elimination of *o*-nitrobenzoylsulfinic acid to produce mono-alkyl diazene intermediates under mild reaction conditions, i.e. neutral pH and less than 23°, particularly useful with substrates that have sensitive substituents. [Mitsunobu *Synthesis* **1** 1981, DOI: 10.1055/s-1981-29317; Hughes *Org React* **42** 335 1992.] For the success of these reactions it is imperative that the reagent is as pure as possible. Solutions of NBSH in THF that are pale yellow are still suitable for these applications.

2-Nitrophenol-4-arsonic acid (Roxarsone, 4-hydroxy-3-nitrophenylarsonic acid) [121-19-7] $C_6H_6AsNO_6$, **M 263.0, m >300°, $pK_{Est(1)} \sim 4.4$ As(O)-(OH)-(O $^-$), $pK_{Est(2)} \sim 7.4$ (phenolic OH), $pK_{Est(3)} \sim 7.7$ As(O)-2(O $^-$).** It crystallises from water and is used for the spectroscopic detection of Zn. Used in poultry feed premixes. [Beilstein **16** II 468, **16** III 1073, **16** IV 1188.]

1-Nitroso-2-naphthol-3,6-disulfonic acid, di-Na salt, hydrate (Nitroso-R-salt) [525-05-3] $C_{10}H_5NNa_2O_8S_2$, **M 377.3, m >300°, $pK_{Est(1)} < 0$ (SO_3^-), pK_2 7.13 (OH).** Purify the salt by dissolution in aqueous alkali and precipitation by addition of HCl to give the **free acid** which is isolated, and dissolved in minimum volume of hot H_2O containing 2 equivalents of NaOH (solubility of di-Na salt at ~20° is 2.5w/w%), and cooled to give golden-yellow fan-shaped crystals. It complexes with Cd ($\log\beta_{11}$ 3.2, $\log\beta_{12}$ 5.6), Cu ($\log\beta_{11}$ 8.5, $\log\beta_{12}$ 14.6), La ($\log\beta_{11}$ 4.4, $\log\beta_{12}$ 7.8, $\log\beta_{13}$ 11.2), Ni ($\log\beta_{11}$ 6.9, $\log\beta_{12}$ 12.4), Pb ($\log\beta_{11}$ 4.6, $\log\beta_{12}$ 7.4), Pd ($\log K_{12}$ 8.8), Y ($\log\beta_{11}$ 4.5, $\log\beta_{12}$ 7.8, $\log\beta_{13}$ 11.3), is used for the **spectrophotometric determination** of Co, Fe, K, Ag, Ba, Cu (colour change pH 4-7, gold/green), Ni (colour change pH 3, yellow/green), and Pb. [Oka & Miyamoto *J Chem Soc Jpn* (Pure Chem Sectn) **76** 672 1955 (*Chem Abstr* 11882 1956I), *Beilstein* **11** IV 669.]

2-Nitroso-1-naphthol-4-sulfonic acid ($3H_2O$) [3682-32-4] $C_{10}H_7NO_5S$, **M 316.3, has m 140-145°(dec), 142-146°(dec), $pK_{Est} \sim 6.3$ (OH).** Crystallise the acid from dilute HCl solution. The crystals are dried over $CaCl_2$ in a vacuum desiccator. It has also been purified by dissolution in aqueous alkali and precipitation by addition of water. It is a reagent for cobalt. [Beilstein **11** H 331, **11** II 189, **11** III 621, **11** IV 668.]

2-Nitro-4-sulfobenzoic acid [552-23-8] $C_7H_5NO_7S$, **M 247.1, m 111°, d^{25}_4 1.8g/cm 3 , n_D^{20} 1.643, $pK_{Est} \sim 1.65$.** It is prepared by sulfonating 2-nitrotoluene to 2-nitro-4-toluenesulfonic acid which is oxidised with $KMnO_4$ [Hart *Am J* **1** 352 1879-1880]. **Alternatively**, boil under reflux a solution of 2-nitro-4-toluenesulfonic acid (2.0g, 8.4mmol) in NaOCl (20ml, 32.2mmol) for 1 hour, cool to room temperature, filter, acidify the filtrate with concentrated HCl (dropwise, CARE), filter off the solid and dry it *in vacuo* (92% yield) [Gless et al. to Zeneca Ltd, Pat US5565608 A, expired 19 Dec 2000]. Recrystallise the acid from dilute HCl (needles) and dry it *in vacuo*. It is stable in air but is *hygroscopic* in moist air. It is a standard in acid-base titrations. [Beilstein **11** III 685.]

Nuclear Fast Red (1-amino-2,4-dihydroxy-5,10-anthraquinone-3-sulfonic acid Na Salt) [6409-77-4] $\text{C}_{14}\text{H}_8\text{NNaO}_7\text{S}$, *M* 357.3, *m* >290°(dec), λ_{max} 518nm. A solution of 5g of the dye in 250ml of warm 50% EtOH is cooled to 15° for 36 hours, then filtered on a Büchner funnel, washed with EtOH until the washings are colourless, then with 100ml of Et₂O and dried over P₂O₅ *in vacuo*. It is a biological stain that is also used for the estimation of Ca. [Kingsley & Robnett *Anal Chem* **33** 552 1961, DOI: 10.1021/ac60172a020.]

***n*-Octadecylphosphonic acid (ODPA)** [4724-74-4] $\text{C}_{18}\text{H}_{38}\text{O}_3\text{P}$, *M* 334.5, *m* 95-98°, 98.5-99°, 98-100°, 100°, 100-101°, *b* ~463°/760mm (predicted), *d*₄²⁵ ~1.0g/ml, *n*_D²⁰ 1.444, *pK*_{Est} ~2.65. The acid is best prepared by reaction of sodium dibutylphosphite [$\text{NaPO}(\text{n-BuO})_2$] and *n*-octadecylbromide followed by hydrolysis of the dibutyl *n*-octadecylphosphonate formed. Thus, under N₂, Na (1.15g, 0.05 mole) is suspended in dry hexane, or heptane, (150ml) and refluxed gently while dibutylphosphite (9.7g, 0.05 mole) is added dropwise over 30 minutes. The boiling is continued until the Na completely dissolves (*ca* 4-5 hours) then *n*-octadecylbromide (0.05 mole) is added during 1 hour and gently refluxed for 5-6 hours while NaBr separated. The mixture is cooled, H₂O is added, the organic layer is collected, evaporated under reduced pressure (at room temperature when all the solvent is removed) and the residual oil (phosphonate ester) is distilled *in vacuo* (fractionating apparatus) to give **di-*n*-butyl *n*-octadecylphosphonate** (>80% yield, *b* 248-250°/2mm, *d*₄²⁵ 0.9037, *n*_D²⁵ 1.4499). The ester is hydrolysed by boiling under reflux with concentrated HCl (~75ml) overnight, then distilled from an oil bath whereby *n*-butanol and *n*-butylchloride and about half the volume of excess acid are distilled off, and cooled. The **phosphonic acid** that crystallises as crust is collected, dried and recrystallised from hexane or heptane (clusters of flat needles similar to naphthalene) and dried *in vacuo* over solid KOH and has *m* 98.5-99° (80-85% yield). [Kosolapoff *J Am Chem Soc* **67** 1180 1945, DOI: 10.1021/ja01223a045; see also Griffin & Wells *J Org Chem* **24** 2049 1959, DOI: 10.1021/jo01094a624.]

Alternatively, the phosphonic acid (100g) is dissolved in EtOH (150ml) by heating and stirring, and when clear, hexane (1L, total) is added slowly, set aside overnight then in a refrigerator for 6 hours and the crystals are collected by filtration. The crystals are stirred at 100° under vacuum to remove entrapped EtOH, toluene (300ml) is added slowly to the hot solid while stirring, heat is removed to give a white suspension, and the solid **ODPA** is collected at room temperature and dried to constant weight *in vacuo* (~24 hours, yield ~75g). {¹H} ³¹P NMR (162MHz, MeOH-*d*₄) δ 29.1. [Owen et al. *J Am Chem Soc* **130** 12279 2008, DOI:10.1021/ja804414f]. At 10⁻³ mol/L in Milli Q H₂O, ODPA molecules behave as free species as shown by ¹H NMR [Hauffman et al. *Langmuir*, **24** 13450 2008, DOI: 10.1021/la801978a].

ODPA is used for the preparation of thermal paper for receipts, tickets etc, capping nano materials (cf. e.g. CdTe quantum dots in *Nanotechnology* Chapter 7). **IRRITANT**, affects eyes, skin, harmful if ingested.

***n*-Octadecyl trichlorosilane (OTS)** [112-04-9] $\text{C}_{18}\text{H}_{37}\text{Cl}_3\text{Si}$, *M* 387.9, *m* 20°, 22°, *b* 160-162°/3mm, 185-199°/2-3mm, 223°/10mm, 280.3-283.2°/760mm, *d*₄³⁰ 0.98, *n*_D²⁰ 1.459. Purify it by fractional distillation at high vacuum. It is moisture sensitive. [Winstein & Seubold *J Am Chem Soc* **69** 2916-2917 1947, DOI: 10.1021/ja01203a513; Barry et al. *J Am Chem Soc* **69** 2916-2916 1947, DOI: 10.1021/ja01203a512; Beilstein **4** IV 4256.] It has been used for making SAMs (Self-Assembled Monolayers, see Chapter 7) [Kovaks (seminar, 7 May 2013), quantum.esu.edu; Kelkar et al. *Appl Surface Sci* **282** 291 2013, DOI: 10.1016/j.apsusc.2013.05.121].

Octamethyl cyclotetrasiloxane [556-67-2] $\text{C}_8\text{H}_{24}\text{O}_4\text{Si}_4$, *M* 296.6, *m* 17-19°, 17.58°, 18.5°, *b* 74°/20mm, 176.4°/760mm, *d*₄^{29.3} 0.9451, *n*_D³⁰ 1.3968. The solid exists in two forms, *m* 16.30° and 17.65°. Dry it over CaH₂ and distil it. Further fractionation can be effected by repeated partial freezing and discarding the liquid phase. [Patnode & Wilcock *J Am Chem Soc* **68** 358 1946, DOI: 10.1021/ja01207a004; Osthoff & Grubb *J Am Chem Soc* **76** 399 1954, DOI: 10.1021/ja01631a025; Hoffman *J Am Chem Soc* **75** 6313 1953, DOI: 10.1021/ja01120a515; Beilstein **4** IV 4125.]

Octamethyl trisiloxane [107-51-7] $\text{C}_8\text{H}_{24}\text{O}_2\text{Si}_3$, *M* 236.5, *m* -82°, -80°, *b* 151.7°/747mm, 153°/760mm. Distil it twice, the middle fraction from the first distillation is again distilled, and the middle fraction of the second distillation is used. [Patnode & Wilcock *J Am Chem Soc* **68** 358 1946, DOI: 10.1021/ja01207a004; Wolcock *J Am Chem Soc* **68** 691 1946, DOI: 10.1021/ja01208a050; Thompson *J Chem Soc* 1908 1953, DOI: 10.1039/JR9530001908; Beilstein **4** IV 4115.]

1-Octanethiol [111-88-6] $C_8H_{18}S$, M 146.3, m -49° , b $77-78^\circ/10\text{m}$, $197-199^\circ/\text{atm}$, $199.1^\circ/760\text{mm}$, $197-200^\circ/760\text{mm}$, d^{25}_4 0.843g/ml, n^{20}_D 1.452, [also available as a 5mM solution in EtOH, d^{25}_4 0.7.81g/ml], pKa 9.97. 1-Acetylthio-octane [$C_{10}H_{20}OS$, M 188.0, b $116-117^\circ/10\text{m}$, n^{20}_D 1.4641] prepared by reaction of thiolacetic acid and 1-octene (catalysed by a few milligrams of benzoyl peroxide) [Brown et al. *J Chem Soc* 2123 1951, DOI: 10.1039/JR9510002123], is hydrolysed with aqueous ethanolic KOH under N_2 to give **octane-1-thiol** in 85% yield, which is purified by distillation at $77-78^\circ/10\text{m}$, and is stored under N_2 . The **2,4-dinitrophenylthioether** has m $75-76^\circ$ after recrystallisation from light petroleum (b $60-80^\circ$), and the **mercuric salt** forms plates with m $71-72^\circ$ from EtOH. [Bateman et al. *J Chem Soc* 2838 1958, DOI: 10.1039/JR9580002838.] Alternatively, the thiol can be obtained by stirring a mixture of crude **S-octyl ethylxanthate** (0.1mol) [prepared by boiling under reflux a solution of potassium ethylxanthate (0.15mol, see [140-89-6] above) and *n*-octylchloride (0.1mol) in Me_2CO (200ml) for 8 hours}, the precipitated KCl is filtered off, the filtrate is evaporated off under reduced pressure, and the residue is washed with H_2O to remove unreacted K xanthate. The ester is recovered in 90-95% yield as an oil which is dried over anhydrous Na_2SO_4 . The crude ester in ethylenediamine (30ml) is heated at 30° for 3 hours under N_2 , poured into an ice-water solution of H_2SO_4 and extracted with $*C_6H_6$ (3 x 50ml). The extract is washed with 5% H_2SO_4 , dried (Na_2SO_4), filtered, the filtrate is evaporated and the oily thiol is distilled (b $71-72^\circ/10\text{mm}$) under a vacuum in a current of N_2 . [Mori & Nakamura *J Org Chem* 34 4170 1969, DOI: 10.1021/jo01264a095.] Store the foul smelling oil in a sealed container under N_2 or argon. Used to prepare functionalise gold nanoparticles, e.g. $Au-SCH_2(CH_2)_6CH_3$ [see MFCD09953506, PubChemID 24884475 in Chapter 7, 'Nanotechnology'], and for other applications in nanotechnology. **IRRITANT**, prevent contact with eyes and skin.

Octaphenyl cyclotetrasiloxane [546-56-5] $C_{48}H_{40}O_4Si_4$, M 793.2, m 200.5° , 201° , $201-202^\circ$, $203-204^\circ$, b $330-340^\circ/1\text{mm}$. Recrystallise it from AcOH, EtOAc, $*C_6H_6$ or $*C_6H_6/EtOH$. It forms two stable isomorphs and both forms, as well as the mixture, melt at $200-201^\circ$. There is a **metastable** form which melts at $187-189^\circ$. [Burkhard et al. *J Am Chem Soc* 67 2174 1945, DOI: 10.1021/ja01228a035; Hyde et al. *J Am Chem Soc* 69 488 1947, DOI: 10.1021/ja01195a003; *Beilstein* 16 IV 1530.]

Octyl trichlorosilane [5283-66-9] $C_8H_{17}Cl_3Si$, M 247.7, b $96.5^\circ/10\text{mm}$, $112^\circ/15\text{mm}$. $119^\circ/28\text{mm}$, $233^\circ/731\text{mm}$, $225-227^\circ/\text{atm}$, $229^\circ/760\text{mm}$, d^{20}_4 1.0744, n^{20}_D 1.4453. Purify the silane by repeated fractionation using a 15-20 theoretical plate glass column packed with glass helices. This can be done more efficiently using a spinning band column. Its purity can be checked by analysing for HCl (ca 0.5-1g of sample is dissolved in 25ml of MeOH, diluted with H_2O and the HCl formed by hydrolysis is titrated with standard alkali). It is moisture sensitive. [Whitmore *J Am Chem Soc* 68 475 1946, DOI: 10.1021/ja01207a035; El-Abbady & Anderson *J Am Chem Soc* 80 1737 1958, DOI: 10.1021/ja01540a058; *Beilstein* 4 III 1907, 4 IV 4253]

Orange I [tropaeolin 000 Nr1, 4-(4-hydroxy-1-naphthylazo)benzenesulfonic acid sodium salt] [523-44-4] $C_{16}H_{11}N_2NaO_4S$, M 350.3, m $>260^\circ(\text{dec})$. Purify the dye by dissolving it in the minimum volume of H_2O , adding, with stirring, a large excess of EtOH. The salt separates as orange needles. It is collected by centrifugation or filtration, washed with absolute EtOH (3x) and Et_2O (2x) in the same way, and dried in a vacuum desiccator over KOH. The **free acid** can be recrystallised from EtOH. [Slotta & Franke *Chem Ber* 64 86 1931, DOI: 10.1002/cber.19310640115; *Beilstein* 16 H 275, 16 II 117, 16 IV 410.] The purity can be checked by titration with titanium chloride [Klotz *J Am Chem Soc* 68 2299 1946, DOI: 10.1021/ja01215a051].

Orange II [tropaeolin 000 Nr2, CI Acid Orange 7, 4-(2-hydroxy-1-naphthylazo)benzenesulfonic acid sodium salt] [633-96-5] $C_{16}H_{11}N_2NaO_4S$, M 350.3, has m $164^\circ(\text{predicted})$. Purification is as for Orange I. Its solubility in H_2O is 40g/L at 25° . [Müller et al. *Helv Chim Acta* 35 2579 1952, DOI: 10.1002/hlca.19520350745.] Also purify it by extracting it with a small volume of cold water, then crystallising it by dissolving in boiling water, cooling to ca 80° , adding two volumes of EtOH and cooling. When cold, the precipitate is filtered off, washed with a little EtOH and dried in air. It can be salted out from aqueous solution with sodium acetate, then repeatedly extracted with EtOH. Meggy and Sims [*J Chem Soc* 2940 1956, DOI: 10.1039/JR9560002940], after crystallising the sodium salt twice from water, dissolved it in cold water (11ml/g) and added concentrated HCl to precipitate the **acid dye** which was separated by centrifugation, redissolved and again precipitated with acid. After washing the precipitate three times with 0.5M acid, it was dried over NaOH, recrystallised twice from absolute EtOH, washed with a little Et_2O , dried over NaOH and stored over concen-

trated H_2SO_4 in the dark. It can then be converted to the *pure salt* with the calculated amount of NaOH or Na_2CO_3 . [Beilstein 16 IV 408.] The purity, like Orange I above, can be checked by titration with titanium chloride [Klotz *J Am Chem Soc* 68 2299 1946, DOI: 10.1021/ja01215a051].

Orange G (Acid Orange 10, 1-phenylazo-2-naphthol-6,8-disulfonic acid di-Na salt) [1936-15-8] $\text{C}_{16}\text{H}_{10}\text{N}_2\text{Na}_2\text{O}_7\text{S}_2$, M 452.4, has $\text{pK}_{\text{Est}} \sim 9$. Recrystallise this dye from 75% EtOH, dry it for 3 hours at 110° and keep it in a vacuum desiccator over H_2SO_4 . The *free acid* crystallises from EtOH or concentrated HCl in deep red needles with a green reflex. A biological stain for connective tissue, pituitary acidophils, keratin, mucus, and in Flemming's triple stain on paraffin sections of onion root. [Oxidation-reduction potential of azo dyes: Conant & Pratt *J Am Chem Soc* 48 2468 1923, DOI: 10.1021/ja01420a039; Drew & Landquist *J Chem Soc* 292 1938, DOI: 10.1039/JR9380000292; Beilstein 16 H 301, 16 I 305, 16 II 141, 16 III 327.]

Orange RO {acid orange 8, 1,8-[bis(4-n-propyl-3-sulfophenyl-1-amino)]anthra-9,10-quinone di-Na salt} [5850-86-2] $\text{C}_{17}\text{H}_{13}\text{N}_2\text{NaO}_4\text{S}$, M 364.4, CI 15575, has λ_{max} 490nm. Salt it out three times with sodium acetate, then extract it repeatedly with EtOH and dry it *in vacuo*.

Orthanilic acid (2-aminobenzenesulfonic acid) [88-21-1] $\text{C}_6\text{H}_7\text{NO}_3\text{S}$, M 173.2, m $>300^\circ(\text{dec})$, pK^{25} 2.49. Crystallise orthanilic acid from aqueous solution, containing 20ml of concentrated HCl per L, then recrystallise it from distilled water, and dry it in a vacuum desiccator over Sicapent (m 315°). When an aqueous solution is chilled below 13.5° , the hydrated form of the acid is obtained. It is used for the determination of nitrite and nitrate. The *S-benzylisothiuronium salt* has m 137° (from H_2O). [Wertheim *Org Synth Coll Vol* 2 471 1943, DOI: 10.15227/orgsyn.015.0055; Beilstein 14 H 681, 14 I 714, 14 II 429, 14 III 1896, 14 IV 2638.] It promotes the formation of a reverse-turn in peptides, and induces a folded conformation when incorporated into peptides.

Pentafluorophenyl dimethylchlorosilane (Flophemesyl chloride) [20082-71-7] $\text{C}_6\text{H}_6\text{ClF}_5\text{Si}$, M 260.7, b $47^\circ/1\text{mm}$, 89-90°/10mm, d_{20}^{20} 1.4080-1.412, d_4^{30} 1.403, n_D^{20} 1.4460-1.4490. If it goes turbid on cooling due to separation of some LiCl, then dissolve it in Et_2O , filter and fractionate it in a vacuum. [Morgan & Poole *J Chromatogr A* 89 225 1974, DOI: 10.1016/S0021-9673(01)99397-5; Birkinshaw et al. *J Chromatogr A* 132 548 1977, DOI: 10.1016/S0021-9673(00)82921-0.] A sensitive derivatising reagent in electron capture gas chromatography for analysing steroids and aliphatic alcohols [Poole et al. *J Chromatogr A* 199 123 1980, DOI: 10.1016/S0021-9673(01)91366-4].

Phenol-2,4-disulfonic acid [96-77-5] $\text{C}_6\text{H}_6\text{O}_7\text{S}_2$, M 254.2, $\text{pK}_1 < 1$, $\text{pK}_2 < 1$, $\text{pK}_3 \sim 8.3$. Crystallise the acid from EtOH/diethyl ether. [Beilstein 11 H 250, 11 I 58, 11 II 139, 6 III 522.] **IRRITANT.**

Phenylarsonic acid (benzenearsonic acid) [98-05-5] $\text{C}_6\text{H}_7\text{AsO}_3$, M 202.0, m 155-158°(dec), 158-162°(dec), pK_1^{25} 3.65, pK_2^{25} 8.77. Crystallise it from H_2O (3ml/g) between 90° and 0° . Alternatively, dissolve 600g of the acid in 500ml of boiling H_2O , add 20g of Norite, filter hot, cool, filter off the crystals and dry them. On heating at ~ 154 - 160° it is converted to the *anhydride*. [Bullard & Dickery *Org Synth Coll Vol* 2 494 1943, DOI: 10.15227/orgsyn.015.0059; for the *4-nitro derivative* see Ruddy & Starkey *Org Synth Coll Vol* 3 665 1955, DOI: 10.15227/orgsyn.026.0060; Beilstein 16 H 868, 16 I 448, 16 II 457, 16 III 1057, 16 IV 1183.]

N-Phenyl-bis(trifluoromethanesulfonimide) [ditriflic phenylimide, N,N-bis(trifluoromethylsulfonyl) aniline] [37595-74-7] $\text{C}_8\text{H}_5\text{F}_6\text{NO}_4\text{S}_2$, M 357.3, m 93-94°, 100-102°, 99-105°. This ditriflic imide is a very *stable triflating agent*. If hydrolysis appears to have occurred, it would be better to prepare anew from purified aniline with 2 mols of redistilled trifluoroacetic anhydride in CH_2Cl_2 at -78° , with 2 mols of Et_3N in order to keep the aniline as the free base and to mop up the free acid formed. It is then evaporated *in vacuo* to give the crystalline reagent. [McMurry & Scott *Tetrahedron Lett* 24 979 1983, DOI: 10.1016/S0040-4039(00)81581-6; Crisp & Scott *Synthesis* 335 1985, DOI: 10.1055/s-1985-31204; Hendrickson & Bergeron *Tetrahedron Lett* 14 4607 1973, DOI: 10.1016/S0040-4039(01)87289-0; Mascarenas et al. *Tetrahedron* 47 3485 1991, DOI: 10.1016/S0040-4020(01)86410-3.]

1,4-Phenylene diisothiocyanate (bitoscanate) [4044-65-9] $\text{C}_6\text{H}_4\text{N}_2\text{S}_2$, M 192.3, m 129-131°, 130-131°, 132°, 129-133°. Purify bitoscanate by recrystallisation from AcOH, petroleum ether (b 40 - 60°), Me_2CO or

aqueous Me_2CO . [van der Kerk et al. *Recl Trav Chim Pays-Bas* **74** 1262 1955, DOI: 10.1002/recl.19550741007; Lieber & Slutkin *J Org Chem* **27** 2214 1962, DOI: 10.1021/jo01053a509; *Beilstein* **13** IV 174.] It is a CNS and gastrointestinal **toxin** in humans, harmful if swallowed.

Phenylboronic acid (benzeneboronic acid) [98-80-6] $\text{C}_6\text{H}_7\text{BO}_2$, **M 121.9**, **m $\sim 43^\circ$, 215-216 $^\circ$ (anhydride), 217-220 $^\circ$, 219 $^\circ$, b 265 $^\circ$ /atm, pK_a^{25} 8.83. It recrystallises from H_2O , but it can convert spontaneously to *benzeneboronic anhydride* or *phenylboroxide* on standing in dry air. A possible impurity is *dibenzeneborinic* acid which can be removed by washing with petroleum ether. Heating in an oven at 110 $^\circ$ /760mm for 1 hour converts it to the *anhydride* **m 214-216 $^\circ$** . Its solubility in H_2O is 1.1% at 0 $^\circ$ and 2.5% at 25 $^\circ$, and in EtOH it is 10% (w/v). [Gilman & Moore *J Am Chem Soc* **80** 3609 1958, DOI: 10.1021/ja01547a032.] If the acid is required, not the anhydride, the acid (from recrystallisation in H_2O) is dried in a slow stream of air saturated with H_2O . The *anhydride* is converted to the acid by recrystallisation from H_2O . The acid gradually dehydrates to the *anhydride* if left in air at room temperature with 30-40% relative humidity. The melting point is usually that of the anhydride because the acid dehydrates before it melts [Washburn et al. *Org Synth Coll Vol* **4** 68 1963, DOI: 10.15227/orgsyn.039.0003]. [Fieser **1** 833; *Beilstein* **16** IV 1654.] It is a useful component in Stille- [Chan & Mak *Tetrahedron* **50** 2003 1994, DOI: 10.1016/S0040-4020(01)85064-X; Gilbert & Wulff *J Am Chem Soc* **116** 7449 1994, DOI: 10.1021/ja00095a075] and Suzuki- [Chan & Mak *Tetrahedron* **50** 2003 1994, DOI: 10.1016/S0040-4020(01)85064-X; Shapiro & Gomez-Lor *J Org Chem* **59** 5524 1994, DOI: 10.1021/jo00098a006; Marck et al. *Tetrahedron Lett* **35** 3277 1994, DOI: 10.1016/S0040-4039(00)76884-5] cross coupling reactions, and in Suzuki and Suzuki-Miyaura bi-aryl cross-coupling reactions [Wu et al. *Tetrahedron Lett* **47** 9267 2006, DOI: 10.1016/j.tetlet.2006.10.127; Li et al. *Tetrahedron Lett* **47** 9239 2006, DOI: 10.1016/j.tetlet.2006.10.148].**

1,2-Phenylenephosphorochloridate (2-chloro-1,3,2-benzodioxaphosphole-2-oxide) [1499-17-8] $\text{C}_6\text{H}_4\text{ClO}_3\text{P}$, **M 190.5**, **m 52 $^\circ$, 58-59 $^\circ$, 59-61 $^\circ$, b 80-81 $^\circ$ /1-2mm, 118 $^\circ$ /10mm, 122 $^\circ$ /12mm, 125 $^\circ$ /16mm, 155 $^\circ$ /33mm. After distilling it in a vacuum, it sets to a colourless solid. It is soluble in petroleum ether, *benzene and slightly soluble in Et_2O . [Khawaja et al. *J Chem Soc (C)* 2092 1970, DOI: 10.1039/J39700002092; Anschütz & Broeker *Justus Liebigs Ann Chem* **454** 109 1927, DOI: 10.1002/jlac.19274540107; Fieser **1** 837; *Beilstein* **6** IV 5602.]**

Phenylisothiocyanate (phenyl mustard oil) [103-72-0] $\text{C}_7\text{H}_5\text{NS}$, **M 135.2**, **m -21 $^\circ$, b 95 $^\circ$ /12mm, 117.1 $^\circ$ /33mm, 221 $^\circ$ /760mm, d_4^{25} 1.1288, $\text{n}_\text{D}^{23.4}$ 1.64918. It is insoluble in H_2O , but soluble in Et_2O and EtOH. If impure (due to formation of thiourea), then steam distil it into a receiver containing 5-10ml of N H_2SO_4 . Separate the oil, dry over CaCl_2 and distil it under vacuum. [Dains et al. *Org Synth Coll Vol* **1** 447 1941, DOI: 10.15227/orgsyn.006.0072; *Beilstein* **12** IV 867.] Used for derivatisation.**

Phenyl methanesulfonate [16156-59-5] $\text{C}_7\text{H}_8\text{O}_3\text{S}$, **M 172.1**, **m 58-61 $^\circ$, 61-62 $^\circ$, 62-63 $^\circ$, b 279 $^\circ$ /atm, Crystallise the sulfonate from MeOH or from H_2O (**m 61.5 $^\circ$**). [*Beilstein* **6** H 176, **6** III 650, **6** IV 689.]**

Phenyl methanesulfonyl fluoride (PMSF) [329-98-6] $\text{C}_7\text{H}_7\text{FO}_2\text{S}$, **M 174.2**, **m 90-91 $^\circ$, 92-93 $^\circ$. Purify PMSF by recrystallisation from * C_6H_6 , petroleum ether or CHCl_3 /petroleum ether. [Davies & Dick *J Chem Soc* 483 1932, DOI: 10.1039/JR9320000483; cf. Tullock & Coffman *J Org Chem* **25** 2016 1960, DOI: 10.1021/jo01081a050.] It is a general **protease inhibitor** (specific for trypsin and chymotrypsin) and is a good substitute for diisopropylphosphorofluoridate [Fahrney & Gold *J Am Chem Soc* **85** 997 1963, DOI: 10.1021/ja00890a037]. [*Beilstein* **11** III 331.] It is used in the preparation of cell lysates as a protease inhibitor.**

Phenylphosphinic acid [benzenephosphinic acid, PhPH(O)(OH)] [1779-48-2] $\text{C}_6\text{H}_7\text{O}_2\text{P}$, **M 142.1**, **m 70 $^\circ$, 71 $^\circ$, 83-85 $^\circ$, 86 $^\circ$, pK_a^{25} 1.75. Crystallise it from H_2O (solubility is 7.7% at 25 $^\circ$). Also purify it by placing the solid in a flask covered with dry Et_2O , and allowed it to stand for 1 day with intermittent shaking. Et_2O is decanted off and the process repeated. After filtration, excess Et_2O is removed in a vacuum. It has also been recrystallised from * C_6H_6 . [Michaelis *Justus Liebigs Ann Chem* **181** 265 1876, DOI: 10.1002/jlac.18761810302; Banks & Skoog *Anal Chem* **29** 109 1957; for ^{31}P NMR of P compounds see Van Wazer et al. *J Am Chem Soc* **78** 5715 1956, DOI: 10.1021/ja01603a002; and for deuterium exchange see Reuben et al. *J Am Chem Soc* **85** 3093 1963, DOI: 10.1021/ja00903a009; *Beilstein* **16** IV 1033.]**

Phenylphosphonic acid [1571-33-1] $\text{C}_6\text{H}_7\text{O}_3\text{P}$, M 158.1, m 161°, 163°, 164.5-166°, $\text{pK}_{\text{a}}^{25}$ 7.43 (7.07). It is best to recrystallise it from H_2O by concentrating an aqueous solution to a small volume and allowing it to crystallise. Wash the crystals with ice cold H_2O and dry them in a vacuum desiccator over H_2SO_4 . [Lecher et al. *J Am Chem Soc* **76** 1045 1954, DOI: 10.1021/ja01633a035]. $\text{pK}_{\text{a}}^{25}$ values in H_2O are 7.07, and in 50% EtOH 8.26. [Jaffé et al. *J Am Chem Soc* **75** 2209 1953, DOI: 10.1021/ja01105a054; IR: Daasch & Smith *Anal Chem* **23** 853 1951, DOI: 10.1021/ac60054a008; *Beilstein* **16** IV 1068.]

Phenylphosphonic dichloride (P,P-dichlorophenyl phosphine oxide) [824-72-6] $\text{C}_6\text{H}_5\text{Cl}_2\text{OP}$, M 195.0, has m 3°, b 83-84°/1mm, 135-136°/23mm, d_4^{30} 1.977, n_D^{30} 1.5578. Fractionally distil it using a very efficient or a spinning band column. [Lecher et al. *J Am Chem Soc* **76** 1045 1954, DOI: 10.1021/ja01633a035; NMR: Muller et al. *J Am Chem Soc* **78** 3557 1956, DOI: 10.1021/ja01596a002; Van Wazer et al. *J Am Chem Soc* **78** 5715 1956, DOI: 10.1021/ja01603a002; IR: Daasch & Smith *Anal Chem* **23** 853 1951, DOI: 10.1021/ac60054a008; Fieser **1** 846; *Beilstein* **16** IV 1074.]

Phenylphosphonous acid [$\text{PhP}(\text{OH})_2$] [tautomer of phenylphosphinic acid – above] [121-70-0, 1779-48-2] $\text{C}_6\text{H}_7\text{O}_2\text{P}$, M 142.1, m 71°, $\text{pK}_{\text{a}}^{\text{Est}} < 0$, $\text{pK}_{\text{a}}^{17}$ 2.1. Crystallise it from hot H_2O or $^*\text{C}_6\text{H}_6$. [For deuterium exchange see Reuben et al. *J Am Chem Soc* **85** 3093 1963, DOI: 10.1021/ja00903a009; *Beilstein* **16** IV 1033.]

Phenylphosphonous dichloride (P,P-dichloro phenyl phosphine) [644-97-3] $\text{C}_6\text{H}_5\text{Cl}_2\text{P}$, M 179.0, has m -51°, b 68-70°/1mm, 224-226°/atm, d_4^{30} 1.9317, n_D^{35} 1.5962. Distil it in a vacuum by fractionating through a 20cm column packed with glass helices (better to use a spinning band column) [Buchner & Lockhart *J Am Chem Soc* **73** 755 1951, DOI: 10.1021/ja01146a076; NMR: Muller et al. *J Am Chem Soc* **78** 3557 1956, DOI: 10.1021/ja01596a002; IR: Daasch & Smith *Anal Chem* **23** 853 1951, DOI: 10.1021/ac60054a008]. It forms a yellow *Ni* complex: $\text{Ni}(\text{C}_6\text{H}_5\text{Cl}_2\text{P})_4$ (m 91-92°, from H_2O) [Quin *J Am Chem Soc* **79** 3681 1957, DOI: 10.1021/ja01571a020], and a yellow complex also with *molybdenum carbonyl*: $\text{Mo}(\text{CO})_3(\text{C}_6\text{H}_5\text{Cl}_2\text{P})_3$ (m 106-110° dec) [Abel et al. *J Chem Soc* 2323 1959, DOI: 10.1039/JR9590002323]. [*Beilstein* **16** IV 972.]

Phenyl phosphoryl dichloride [770-12-7] $\text{C}_6\text{H}_5\text{Cl}_2\text{O}_2\text{P}$, M 211.0, m -1°, b 103-104°/2mm, 110-111°/10mm, 130-134°/21mm, 241-243°/atm, d_4^{30} 1.4160, n_D^{30} 1.5216. Fractionally distil it under as high a vacuum as possible using an efficient fractionating column or a spinning band column. It should be redistilled if the IR is not very satisfactory [IR: Bellamy & Beecher *J Chem Soc* 475 1952, DOI: 10.1039/JR9520000475; Orloff et al. *J Am Chem Soc* **80** 727 1958, DOI: 10.1021/ja01536a052; for the diNa phenyl phosphate see Freeman & Colver *J Am Chem Soc* **60** 750 1938, DOI: 10.1021/ja01271a004]. [*Beilstein* **6** IV 737.] **HARMFUL VAPOURS.**

Phenylsilane (PhSiH_3) [694-53-1] $\text{C}_6\text{H}_8\text{Si}$, M 108.2, b 62°/100mm, 120°/atm, d_4^{25} 0.877, n_D^{20} 1.5125. It is best prepared by reduction of PhSiCl_3 with $\text{LiAlH}_4/\text{Et}_2\text{O}$: stir at room temperature overnight, pour onto crushed ice, extract with Et_2O , dry the extract (Drierite), filter, evaporate and distil the residual oil preferably in a vacuum [Benkeser et al. *J Am Chem Soc* **74** 648 1952, DOI: 10.1021/ja01123a019; Koch et al. *Recl Trav Chim Pays-Bas* **114** 206 1995, DOI: 10.1002/recl.19951140413; Finholt et al. *J Am Chem Soc* **69** 2692 1947, DOI: 10.1021/ja01203a041]. It is a **useful reagent** for deoxygenating phosphine oxides to phosphines, and *will reduced chiral phosphine oxides to chiral phosphines with retention of configuration* [Marsi *J Org Chem* **39** 265 1974, DOI: 10.1021/jo00916a041]. It is a reducing agent in radical reactions [Perez et al. *J Org Chem* **52** 5570 1967, DOI: 10.1021/jo00234a012], and deoxygenates primary and secondary alcohols *via* their xanthate or thionocarbonate esters (prepared from aryl-OCSCl and an inhibitor, e.g benzoyl peroxide) in 60-100 minutes in >87% yields [Barton et al. *Synlett* 435 1991, DOI: 10.1055/s-1991-20755]. [*Beilstein* **16** III 1198, **16** IV 1360.]

1-Phenylthiosemicarbazide [645-48-7] $\text{C}_7\text{H}_9\text{N}_3\text{S}$, M 167.2, m 200-201°(dec). Crystallise it from EtOH. [*Beilstein* **15** IV 183.]

4-Phenylthiosemicarbazide [5351-69-9] has m 138-140°, 140°. Crystallise it from EtOH. [*Beilstein* **12** IV 827.]

Phenylthio trimethylsilane (trimethyl phenylthio silane) [4551-15-9] $\text{C}_9\text{H}_{14}\text{SSi}$, M 182.4, b 72°/8mm, 95-99°/12mm, d_4^{25} 0.963, d_4^{30} 0.97, n_D^{20} 1.532. Purification is as for phenyl trimethylsilylmethyl sulfide.

1-Phenyl-2-thiourea [103-85-5] $\text{C}_7\text{H}_8\text{N}_2\text{S}$, M 152.1, m 145-150°, 154°. Crystallise the thiourea from water and dry it at 100° in air. [*Beilstein* **12** IV 804.]

Phenyl 4-toluenesulfonate [640-60-8] $\text{C}_{13}\text{H}_{12}\text{O}_3\text{S}$, M 248.3, m 94.5-95.5°, 96°, 95.8-97.8°. Crystallise the ester from MeOH or glacial acetic acid. [Beilstein 11 H 99, 11 II 47, 11 III 200, 6 IV 271.]

Phenyl trimethoxysilane (trimethoxysilyl benzene, PTMOS) [2996-92-1] $\text{C}_9\text{H}_{14}\text{O}_3\text{Si}$, M 198.3, m -25°, b 103°/20mm, 130.5-131°/45mm, 210-213°/atm, 233°/atm, d_4^{35} 1.022, n_D^{35} 1.4698. Fractionate it through an efficient column, but note that it forms an azeotrope with MeOH which is a likely impurity. [Kantor *J Am Chem Soc* 75 2712 1953, DOI: 10.1021/ja01107a048; Beilstein 16 IV 1556.] The hydrolysis and polymerisation of PTMOS produces vitreous solids [Jermouni et al. *J Mater Chem* 5 1203 1995, DOI: 10.1039/JM9950501203].

Phenyl trimethylsilylmethyl sulfide [(phenylthiomethyl)trimethylsilane] [17873-08-4] $\text{C}_{10}\text{H}_{16}\text{SSi}$, M 196.4, b 48°/0.04mm, 113-115°/12mm, 158.5°/52mm, d_4^{30} 0.9671, n_D^{30} 1.5380. If the sample is suspect, then add H_2O , wash it with 10% aqueous NaOH, H_2O again, dry (anhydrous CaCl_2) and fractionally distil it through a 2ft column packed with glass helices. [Cooper *J Am Chem Soc* 76 3713 1954, DOI: 10.1021/ja01643a035; Beilstein 6 IV 1505.]

Phosphine [7803-51-2] PH_3 , M 34.0, m -133°, b -87.5°/760mm, -87.7°/760mm, critical temperature 51.3°, d_4^{25} 1.5307, $\text{pK}^{25} \sim -14$ (extremely weak base). Phosphine is a gas with a very strong odour of fish and is **POISONOUS**. The gas is poorly soluble in H_2O (0.26ml/ml at 20°), and ignites spontaneously in air with a luminous glow. It has been prepared in various ways. A convenient preparation is to make aluminium phosphide by mixing 2 parts of Al powder and 1 part of red P on a piece of paper. By igniting the paper, the mixture becomes white-hot resulting in a spongy mass of aluminium phosphide [Hoffman *J Am Chem Soc* 43 1684 1921, DOI: 10.1021/ja01440a035; Bodoux *Bull Soc Chim Fr* 27 (3) 568 1902]. This phosphide reacts with cold H_2O to give a steady stream of PH_3 . [Use an efficient fume cupboard in these experiments.] Fortunately, the gas is available in metal cylinders, but all due precautions should still be taken. [Klement in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol 1 p 525-530 1963, Gokhale et al. *Inorg Synth* 9 56 1967, DOI: 10.1002/9780470132401.ch17; for basicity see Henderson & Streuli *J Am Chem Soc* 82 5791 1960, DOI: 10.1021/ja01507a008]. [For further detail see 'Inorganic Compounds' in Chapter 4.]

Polystyrenesulfonic acid sodium salt ($-\text{CH}_2\text{CH}(\text{C}_6\text{H}_4\text{SO}_3\text{Na})-$) [25704-18-1] M(average) ~70,000, ~200,000, ~1,000,000. Purify the polymeric anionic electrolyte by repeated precipitation of the sodium salt from an aqueous solution by MeOH, with subsequent conversion to the *free acid* by passage through an Amberlite IR-120 ion-exchange resin. [Kotin & Nagasawa *J Am Chem Soc* 83 1026 1961, DOI: 10.1021/ja01466a003] Recrystallise it from EtOH. Alternatively, purify it by passage through cation and anion exchange resins in series (Rexyn 101 cation exchange resin and Rexyn 203 anion exchange resin), then titrated it with NaOH to pH 7. The *sodium form* of polystyrenesulfonic acid is precipitated by addition of 2-propanol. Dry it in a vacuum oven at 80° for 24 hours, and finally increasing to 120° before to use. [Kowblansky & Ander *J Phys Chem* 80 297 1976, DOI: 10.1021/j100544a019.]

Pontacyl Carmine 2G (Acid Red 1, Amido Naphthol Red G, Azophloxine, 1-acetamido-8-hydroxy-7-phenylazonaphthalene-3,7-disulfonic acid di-Na salt) [3734-67-6] $\text{C}_{18}\text{H}_{13}\text{N}_3\text{Na}_2\text{O}_8\text{S}_2$, M 509.4, CI 18050, λ_{max} 506 and 532nm. Salt it out three times with sodium acetate, then repeatedly extract it with EtOH. See below. [McGrew & Schneider *J Am Chem Soc* 72 2547 1950, DOI: 10.1021/ja01162a059.]

Pontacyl Light Yellow GX [Acid Yellow 17, 1-(2,5-dichloro-4-sulfophenyl)-3-methyl-4-(4-sulfophenylazo)-5-hydroxypyrazole di-Na Salt] [6359-98-4] $\text{C}_{16}\text{H}_{10}\text{Cl}_2\text{N}_4\text{Na}_2\text{O}_7\text{S}_2$, M 551.3, CI 18965, has λ_{max} 400nm. Purify this yellow powder as for Pontacyl Carmine 2G above.

Potassium 4-acetylphenyltrifluoroborate [252726-24-2] $\text{C}_8\text{H}_7\text{BF}_3\text{KO}$, M 226.1, m >250°, 290°(dec). The salt is prepared by adding an aqueous solution of KHF_2 (41ml, 4.5M solution, 185mmol) to a solution of 4-acetylphenylboronic acid (10g, 61mmol) in MeOH (40ml) at ~25°, when a heavy precipitate deposits, but the suspension is stirred for 1 hour at 25° and the solid is filtered off, washed with MeOH and recrystallised from the minimum volume of Me_2CO to provide *p-AcC₆H₄-BF₃⁻ K⁺* (12, 87%), m >250°. Its ^1H NMR [$(\text{CD}_3)_3\text{CO}$, 500MHz] has δ at 7.74(d, $J = 7.8\text{Hz}$, 2H), 7.59 (d, $J = 7.8\text{Hz}$, 2H), 2.49 (s, 3H); the ^{13}C NMR [$(\text{CD}_3)_3\text{CO}$, 125MHz] has δ at 198.1, 134.2, 131.3 (d, $J = 1.5\text{Hz}$), 126.2, 26.4; the ^{19}F NMR (DMSO- d_6 , 470MHz) has δ at -140.3 (br s); the ^{11}B NMR (DMSO- d_6 , 64MHz) has δ at 3.27 (br s), and has the correct elemental analysis

for C and H. It has been used for inserting a **4-acetylphenyl group** into the position of a halogen or triflate group of various arenes and various heterocycles in Pd-catalysed **Suzuki-Miyaura cross-coupling** reactions [Molander & Biolatto *J Org Chem* **68** 4302 2003, DOI: 10.1021/jo0342368].

Potassium 3,5-bis(trifluoromethyl)phenyltrifluoroborate [166328-09-2] $\text{C}_8\text{H}_3\text{BF}_9\text{K}$, **M 320.0, m >250°**. This salt was prepared by the same procedure as the preceding salt in 89% yield and recrystallised from the minimum volume of Me_2CO . Its ^1H NMR ($\text{DMSO}-d_6$, 500MHz) has δ at 7.87(s, 2H), 7.71 (s, 1H), 2.49 (s, 3H); the ^{13}C NMR ($\text{DMSO}-d_6$, 125MHz) has δ at 131.2, 128.1 (q, $J = 31\text{Hz}$), 124.2 (q, $J = 272\text{Hz}$), 118.6 (s); the ^{19}F NMR ($\text{DMSO}-d_6$, 470MHz) has δ at -61.7, -141.6 (br d, $J = 71\text{Hz}$), and the ^{11}B NMR ($\text{DMSO}-d_6$, 64MHz) has δ at 2.57 (br s), and has the correct elemental analysis for C and H. It has been used for inserting a 3,5-bis(trifluoromethyl)phenyl group into the position of a halogen or triflate group of various arenes and various heterocycles in Pd-catalysed **Suzuki-Miyaura cross-coupling** reactions [Molander & Biolatto *J Org Chem* **68** 4302 2003, DOI: 10.1021/jo0342368; Vedejs et al. *J Org Chem* **60** 3020 1995, DOI: 10.1021/jo00115a016;]

Potassium 2,4-difluorophenyltrifluoroborate [871231-41-3] $\text{C}_6\text{H}_3\text{BF}_5\text{K}$, **M 220.0, m >300°**. This borate can be prepared by the same procedure as below and applied in similar reactions.

Potassium 2,6-difluorophenyltrifluoroborate [267006-25-7] has **m >300°**. This salt was prepared by the same procedure as the preceding bis(trifluoromethyl) salt in 90% yield except that the crystalline precipitate was washed with H_2O then Et_2O and dried at high vacuum in a Schlenk line, and had **m >250°**. Its ^1H NMR ($\text{DMSO}-d_6$, 500MHz) has δ at 6.66 (m, 2H), 7.10 (m, 1H); the ^{13}C NMR ($\text{DMSO}-d_6$, 125MHz) has δ at 168.7 (dd, $J = 242\text{Hz}$, 18H), 127.5 (t, $J = 11\text{Hz}$), 110.0 (dd, $J = 23\text{Hz}$, 8H), 118.6 (s); the ^{19}F NMR ($\text{DMSO}-d_6$, 470MHz) has δ at -103.7 (q, $J = 9.4\text{Hz}$), -132.6 (qt, $J = 43.9\text{Hz}$); and the ^{11}B NMR ($\text{DMSO}-d_6$, 64MHz) has δ at 2.17 (q, $J = 44\text{Hz}$), and has the correct elemental analysis for C and H. It has been used for inserting a **2,6-difluorophenyl group** into the position of a halogen or triflate group of various arenes and various heterocycles in Pd-catalysed **Suzuki-Miyaura cross-coupling** reactions [Molander & Biolatto *J Org Chem* **68** 4302 2003, DOI: 10.1021/jo0342368.]

Potassium 2,6-dimethylphenyltrifluoroborate [561328-67-4] $\text{C}_8\text{H}_9\text{BF}_3\text{KO}_2$, **M 212.1, has m >250°**. This salt was prepared by the same procedure as the preceding difluorophenyl salt in 88% yield, except that the crystalline precipitate was washed with H_2O then Et_2O and dried at high vacuum in a Schlenk line and had **m >250°**. Its ^1H NMR (acetone- d_6 , 500MHz) has δ at 6.83 (t, $J = 7.4\text{Hz}$, 1H), 6.75 (d, $J = 7.4\text{Hz}$, 2H), 2.40 (s, 6H); the ^{13}C NMR (acetone- d_6 , 125MHz) has δ at 142.5, 127.7, 125.9, 23.8; the ^{19}F NMR (acetone- d_6 , 470MHz) has δ at -132.5 (q, $J = 48\text{Hz}$); and the ^{11}B NMR (acetone- d_6 , 64MHz) has δ at 4.94 (br d, $J = 49\text{Hz}$), and has the correct elemental analysis for C and H. It has been used for inserting a **2,6-dimethylphenyl group** into the position of a halogen or triflate group of various arenes and various heterocycles in Pd-catalysed **Suzuki-Miyaura cross-coupling** reactions [Molander & Biolatto *J Org Chem* **68** 4302 2003, DOI: 10.1021/jo0342368; cf. Vedejs et al. *J Org Chem* **60** 3020 1995, DOI: 10.1021/jo00115a016]. **Potassium 4-tert-butylphenyltrifluoroborate** [423118-47-2] $\text{C}_{10}\text{H}_{13}\text{BF}_3\text{K}$, **M 240.1, m >230°**, can be similarly prepared from 4-tert-butylphenylboronic acid and used in Pd catalysed **Suzuki cross-coupling** reactions.

Potassium ethyl xanthate (potassium O-ethyl dithiocarbonate, potassium ethyl xanthogenate) [140-89-6] $\text{C}_3\text{H}_5\text{KOS}_2$, **M 160.3, m ~210°(dec), > 215°(dec), pK²⁵ 2.16 (for -S⁻)**. It forms white to pale yellow crystals, prepared by adding ethanolic KOH dropwise to an ethanolic solution of CS_2 to form the salt which can be coaxed further out of solution by adding Et_2O . Crystallise it from absolute EtOH, ligroin/ethanol or acetone by adding Et_2O . Wash it with ether, then dry it in a desiccator. Its solubility at 56° in EtOH is >1% and in Me_2CO is 8% (the Na salt has 44%), and very high in H_2O at 25°. Dry it *in vacuo*, if it contains H_2O , and store it in a tightly stoppered bottle away from light. [X-ray diffraction identification of xanthate derivatives of alcohols: Warren & Matthews *Anal Chem* **26** 1985 1954, DOI: 10.1021/ac60096a041; *Beilstein* **3** H 209, **3** I 84 **3** II 152, **3** III 336, **3** IV 402.]

Potassium 2-furantrifluoroborate [166328-14-9] $\text{C}_4\text{H}_3\text{BF}_3\text{KO}$, **M 174.0, m 200°(dec), >293-303°**. This salt has been prepared from furan (5.0ml, 68.7mmol, dried over 3Å Molecular Sieves) in dry THF (50ml) which was treated under N_2 with BuLi (42.0ml, 1.64M in pentane, 68.9mmol) and stirred at -5° for 3.5 hours to form **furyllithium**. The latter was treated with $\text{B}(\text{iso-PrO})_3$ and allowed to warm to ~25°, quenched with 10% aqueous HCl (~50ml), diluted with Et_2O (50ml) and the organic layer was extracted with 1N NaOH (2 X 50ml). The alk-

aline extracts were combined, acidified to pH 3 with 10% aqueous HCl, and the acid layer was then extracted with Et₂O (3 x 50ml). The combined Et₂O extracts were dried (Na₂SO₄) and evaporated to dryness. The residual 2-furanboronic acid was dissolved in MeOH (200ml) and H₂O (40ml) was treated with 3 equivalents of KHF₂ (16g, 206mmol), refluxed overnight and evaporated to dryness *in vacuo*. The residue was extracted with MeCN (2 x 30ml), filtered and the filtrate was evaporated to dryness, the residue was washed with Et₂O, dried and recrystallised from MeCN/EtOAc to give pure **2-furan-BF₃⁻ K⁺** (5.8g, 48%) as yellow crystals. Attempted TLC on Silica Gel with EtOAc caused hydrolysis of the salt to **2-furylboronic acid**. Its IR (KBr) had bands at ν_{\max} 1575 (C=C), 1005 (B-F), 970 (B-F) cm⁻¹, the ¹H NMR (CD₃CN, 200MHz) had δ at 7.44-7.33 (m, 1H), 6.25-6.17 (m, 1H), 6.17-6.10 (m, 1H); and the ¹¹B NMR (CD₃CN, 160MHz) has δ (from BF₃.OEt₂, 0 ppm) 1.8 (q, *J* 49Hz) and it had the correct elemental analysis for C and H. It is air stable and has been used successfully in Pd-catalysed **Suzuki-Miyaura cross-coupling** reactions [Vedejs et al. *J Org Chem* **60** 3020 1995, DOI: 10.1021/jo00115a016; Molander et al. *J Org Chem* **74** 973 2009, DOI: 10.1021/jo802590b].

Potassium isoamyl xanthate (potassium O-isoamyl dithiocarbonate) [928-70-1, 61792-26-5] **C₆H₁₁KOS₂, M 202.4, pK²⁵ 1.82 (pK⁰ 2.8 free acid)**. Crystallise it twice from acetone/diethyl ether. Dry it in a desiccator for two days and store it under refrigeration. Its solubility in Me₂CO is 2.3% at 56°. [See potassium ethyl xanthate above for storage. FOR X-ray diffraction identification of xanthate derivatives of alcohols see Warren & Matthews *Anal Chem* **26** 1985 1954, DOI: 10.1021/ac60096a041; *Beilstein* **3** III 340, **3** IV 404.]

Potassium 4-methoxyphenyltrifluoroborate [192863-36-8] **C₇H₇BF₃KO, M 214.0, m >300°**. The salt is prepared by mixing **4-methoxyphenylboronic acid** (0.65g, 4.3mmol) and KHF₂ (0.76g, 9.7mmol) in H₂O (1ml) and MeOH (1.8ml) at ~25° for 2 hours and the resulting yellow slurry is extracted into Me₂CO (10ml) and evaporated under a vacuum. The residue is dissolved consecutively in hot Me₂CO and THF, filtered and Et₂O is added to give a yellow solid which is filtered off, washed with Et₂O until free from the yellow colour, and the crystalline material is dried in a Schlenk vacuum line to give **p-MeOC₆H₄-BF₃⁻ K⁺** (0.83g, 92%). Its ¹H NMR (acetone-*d*₆, 500MHz) has δ at 7.38 (d, *J* = 8.2Hz, 2H), 7.69 (d, *J* = 8.2Hz, 2H), 3.70 (s, 3H); the ¹³C NMR (acetone-*d*₆, 125MHz) has δ at 158.9, 133.5, 112.8, 55.1; the ¹⁹F NMR (acetone-*d*₆, 470MHz) has δ (from .Cl₃, 0ppm) at -142.28 (br d, *J* = 66Hz); and the ¹¹B NMR (acetone-*d*₆, 64MHz) has δ (from BF₃.OEt₂, 0ppm) at 4.8 (br s, *J* = 17Hz), and it has the correct elemental analysis for C and H. It is air stable and has been used successfully in Pd-catalysed **Suzuki-Miyaura cross-coupling** reactions [Molander & Biolatto *Org Lett* **4** 1867 2002, DOI: 10.1021/ol025845p].

Potassium 3-methoxyphenyltrifluoroborate [438553-44-7] has **m >300°** and was prepared in the same manner from **3-methoxyphenylboronic acid** in 89% yield and had IR (KBr) with bands at ν_{\max} 1241 (? C=C), 987 (B-F) cm⁻¹; the ¹H NMR (acetone-*d*₆, N, 500MHz) had δ : 7.03 (m, 3H), 6.61 (m, 1H), 3.67 (s, 3H); the ¹³C NMR (acetone-*d*₆, 125MHz) had δ at 159.6, 128.2, 124.9, 117.6, 112.0 55.0; the ¹⁹F NMR (acetone-*d*₆, 470MHz) had δ (from CFCl₃, 0ppm) at -142.8 (br d, *J* = 66Hz); and the ¹¹B NMR (acetone-*d*₆, 64MHz) had δ (from BF₃.OEt₂, 0ppm) at 4.2 (br s, *J* 43Hz), and it had the correct elemental analysis for C and H [Molander & Biolatto *Org Lett* **4** 1867 2002, DOI: 10.1021/ol025845p; cf. Vedejs et al. *J Org Chem* **60** 3020 1995, DOI: 10.1021/jo00115a016].

Potassium 2-methoxyphenyltrifluoroborate [236388-46-8] has **m >300°** and can be prepared in the same manner from 2-methoxyphenylboronic acid and used successfully in Pd-catalysed **Suzuki-Miyaura cross-coupling** reactions.

Potassium methyltrifluoroborate [13862-28-7] **CH₃BF₃K, M 121.94, has m 168-183°, 183°**. This salt, which was used successfully in Pd-catalysed **Suzuki-Miyaura cross-coupling** is air stable, more robust and easier to handle than methylboronic acid. It is prepared by adding **trimethyl boroxine** (5.14g, 41mmol, 3 equivs, [823-96-1]) at room temperature to KHF₂ (19.0g, 243mmol, 6 equivs) in MeCN (200ml), and the mixture is cooled to 0° and stirred for 30 minutes. Then H₂O (4.5ml) is added and after stirring for 3 hours the solvent is evaporated and the resulting solid is dried thoroughly *in vacuo*. The residue is triturated with Me₂CO/MeOH (1:1, 100ml), filtered off, washed with the same mixture of solvents (100ml) and once with Me₂CO/MeOH (1:2, 100ml), and the remaining insoluble white solid is dried at high vacuum in a Schlenk line to give **MeBF₃⁻ K⁺** (12.0g, 80%) as a white powder **m 183°**. Its ¹H NMR (D₂O, 500MHz) has δ at -0.15 (s), the ¹³C NMR (D₂O, 125MHz) has δ at 1.40 to -1.14 (br s), ¹⁹F NMR (D₂O, 470MHz) has δ at -132.3 (q, *J* = 64Hz), and ¹¹B NMR (D₂O, 64MHz) has δ at 7.25 (q, *J* = 64Hz). and has the correct elemental analysis for C and H. It has been used for inserting a methyl group in the position of a halogen or triflate group of various arenes. [Molander et al. *J Org Chem* **68** 5534

2003, DOI: 10.1021/jo0343331.]

Potassium nonafluorobutane sulfonate [29420-49-3] $C_4H_9KO_3S$, **M 338.2**, **m >300°**. Wash it with H_2O and dry it *in vacuo*. When the K salt is distilled with 100% H_2SO_4 , it gives the *free acid* which can be re-distilled (**b 105°/22mm, 210-212°/760mm**) and then converted to the pure K salt. [Gramstad & Haszeldine *J Chem Soc* 2640 1957, DOI: 10.1039/JR9570002640; *Beilstein* 2 IV 818.] **IRRITANT**.

Potassium phenol-4-sulfonate (4-hydroxybenzene-1-sulfonic acid K salt) [30145-40-5] $C_6H_5KO_4S$, **M 212.3**, **m >300°**. Crystallise it several times from distilled water at 90°, after treatment with charcoal, and cooling to *ca* 10°. Dry it at 90-100°, *in vacuo*. [*Beilstein* 11 H 55, 11 I 242, 11 II 137, 11 III 498, 11 IV 582.]

Potassium phenyltrifluoroborate [11042-64-1, 153766-81-5] $C_6H_5BF_3K$, **M 184.0**, has **m 290°, 296°(dec), 296-301°**. The salt is obtained by adding excess of saturated aqueous KHF_2 (125ml, *ca* 4.5M solution, 563mmol) dropwise to a solution of *phenylboronic acid* (20g, 169mmol, [98-80-6]) in MeOH (50ml) with vigorous stirring. The precipitate is collected after 15 minutes, washed with cold MeOH and recrystallised from the minimum volume of MeCN and dried *in vacuo* to give $Ph-BF_3^- K^+$ (25.5g, 82%), **m 296°(dec)**. Its 1H NMR (CD_3CN , 200MHz) has δ at 7.44-7.41 (m, 2H), 7.22-7.05 (m, 3H); the ^{19}F NMR (CD_3CN , vs $CF_3C_6H_5$, 470MHz) has δ at -79 (1:1:1:1 q, $J = 57Hz$); and the ^{11}B NMR (CD_3CN , 160MHz) has δ : 4.1 (q, $J = 57Hz$), and it has the correct elemental analysis for C and H. [Vedejs et al. *J Org Chem* 60 3020 1995, DOI: 10.1021/jo00115a016; Thierig & Umland *Naturwissenschaften* 54 563 1967.] It is air stable and has been used successfully in Pd-catalysed **Suzuki-Miyaura cross-coupling** reactions [Vedejs et al. *J Org Chem* 60 3020 1995, DOI: 10.1021/jo00115a016; Molander & Biolatto *J Org Chem* 68 4302 2003, DOI: 10.1021/jo0342368].

Potassium 2-naphthalene-trifluoroborate [668984-08-5] $C_{10}H_7BF_3K$, **M 234.1**, **m >300°** can be prepared in the same manner from *2-naphthaleneboronic acid* and used successfully in Pd-catalysed **Suzuki-Miyaura cross-coupling** reactions.

Potassium pyridine-3-trifluoroborate [561328-69-6] $C_5H_4BF_3KN$, **M 185.0**, has **m 228-232°**. The 3-pyridine salt is prepared under N_2 from *3-bromopyridine* (3.46g, 21.9mmol) and *triisopropyl borate* (6.1ml, 26.4mmol) in a mixture of dry toluene (40ml) and dry THF (10ml) at -60° to which is added *n*-BuLi (39ml, 1.6M in hexane, 62mmol) dropwise, the mixture is stirred for 2 hours and allowed to warm to room temperature overnight. This mixture containing the *pyridine-3-boronic lithium salt* is cooled to 0° and an aqueous solution of KHF_2 (4.5M, 58mmol) is added dropwise and stirred for 4 hours then the mixture is evaporated to dryness. The residue is dissolved in MeOH, filtered, evaporated to dryness and the residual oil gave a solid (90%) under high vacuum. Its 1H NMR ($MeOD-d_3$, 500MHz) has δ at 8.57 (br s, 1H), 8.26 (br d, $J = 4.6Hz$, 1H), 7.90 (d, $J = 7.3$, 1H), 7.23 (dd, $J = 5.5, 6.6Hz$, 1H); the ^{13}C NMR ($DMSO-d_6$, 125MHz) has δ : 153.0, 145.7, 138.8, 122.0; the ^{19}F NMR ($DMSO-d_6$, 470MHz) has δ : -134.9 (s); and the ^{11}B NMR ($MeOD-d_3$, 64MHz) has δ : 4.0 (br s). It has been used for inserting a *2,6-dimethylphenyl group* into the position of a halogen or triflate group of various arenes and various heterocycles in Pd-catalysed **Suzuki-Miyaura cross-coupling** reactions [Molander & Biolatto *J Org Chem* 68 4302 2003, DOI: 10.1021/jo0342368; Molander et al. *Org Lett* 12 5783 2010, DOI: 10.1021/ol102717x.]

Potassium tetraphenylborate [3244-41-5] $C_{24}H_{20}BK$, **M 358.3**, has **m >300°**. Precipitate it from a solution of KCl acidified with dilute HCl, then crystallise it twice from acetone, wash it thoroughly with water and dry it at 110° [For the amperometric titration-method for determination of K see Findeis & de Vries *Anal Chem* 28 1899 1956, DOI: 10.1021/ac60120a026]. It has also been recrystallised from conductivity water. [*Beilstein* 16 IV 1625.]

Potassium 3-thiophenyltrifluoroborate [192863-37-9] $C_4H_3BF_3KS$, **M 190.0**, has **m >260°(dec)**. This salt, which was used successfully in Pd-catalysed **Suzuki-Miyaura cross-coupling** is air stable, more robust and easier to handle than 3-thienylboronic acid. *3-Thiophenylboronic acid* (4.97g, 26.2mmol) and KHF_2 (5.14g, 65.8mmol) in a Nalgene (polyethylene) bottle (100ml) are stirred vigorously with MeOH (7.5ml) and H_2O (14ml) for 2 hours, and the amber solid formed is set aside at 4° for 2 hours. The solid was collected washed with the minimum of cold MeOH and dissolved in hot Me_2CO filtered, and the filtrate was cooled to 25° and Et_2O was added in portions with stirring, until the supernatant showed no cloudiness. The mixture was set aside at 4° for 1 hour until crystallisation was complete. The crystals were collected washed with a little cold Et_2O and dried *in vacuo* to give $C_4H_3S-BF_3^- K^+$ (4.68g, 94%). Its 1H NMR [$(CD_3)_3CO$, 500MHz] has δ at 7.20 (s, 1H), 7.14 (m, 2H); the ^{13}C NMR [$(CD_3)_3CO$, 125MHz] has δ at 131.8, 125.2, 122.3; the ^{19}F NMR [$(CD_3)_3CO$, 470MHz] has δ at -139.5 (d, $J = 75Hz$), and it has the correct elemental analysis for C and H. It has been used

for inserting a *thiophenyl group* into the position of a halogen or triflate group of various arenes and various heterocycles. [Molander & Biolatto *J Org Chem* **68** 4302 2003, DOI: 10.1021/jo0342368;]. **Potassium 5-methyl-2-thiophenyltrifluoroborate** [871231-40-2] $C_5H_5BF_3KS$, *M* 204.1, *m* >220°, >250° (*dec also reported*) can be similarly prepared from *5-methylthiophenyl-2-boronic acid* and used in Pd catalysed **Suzuki cross-coupling** reactions.

Potassium trimethylsilanolate (trimethylsilanol K salt) [10519-96-7] C_3H_9KOSi , *M* 128.3, has *m* 131-135° (*cubic form*), 135-138°, d_4^{25} 1.11, 125°*dec* (*orthorhombic form*). Recrystallise it from H_2O and dry it at 100°/1-2mm. [Hyde et al. *J Am Chem Soc* **75** 5615 1953, DOI: 10.1021/ja01118a042; prepn and hydrolysis, IR: Tatlock & Rochow *J Org Chem* **17** 1555 1952, DOI: 10.1021/jo50012a001; *Beilstein* **4** IV 3992.]

Propargyl triphenyl phosphonium bromide [2091-46-5] $C_{21}H_{18}BrP$, *M* 381.4, *m* 179°. It recrystallises from 2-propanol as white plates. It also crystallises from EtOH with *m* 156-158°. Its IR has ν_{max} at 1440, 1110 cm^{-1} (P-C str). [Elter & Oediger *Justus Liebigs Ann Chem* **682** 62 1965, DOI: 10.1002/jlac.19656820106; Schweizer et al. *J Org Chem* **42** 200 1977, DOI: 10.1021/jo00422a003; X-ray crystallography: Steiner *Acta Cryst* **C52** 2263 1996, DOI: 10.1107/S0108270196004490].

Propenyloxy trimethylsilane [1833-53-0] $C_6H_{14}OSi$, *M* 130.3, *b* 93-95°/atm, d_4^{20} 0.786, n_D^{20} 1.395. Purify it by fractional distillation using a very efficient column at atmospheric pressure. It usually contains 5% of hexamethyl-disiloxane that boils at 99-101°, but is generally non-reactive and need not be removed. [Hauser & Hance *J Am Chem Soc* **74** 5091 1952, DOI: 10.1021/ja01140a029.] It has been distilled under N_2 through a 15cm column packed with glass helices. Fraction *b* 99-104° is further purified by gas chromatography through a Carbowax column (Autoprep A 700) at a column temperature of 87°, and has a retention time of ~9.5 minutes. [Krüger & Rochow *J Organomet Chem* **1** 476 1964, DOI: 10.1016/S0022-328X(00)94483-8.]

1-Propenyltrimethylsilane (cis and trans mixture) [17680-01-2] $C_6H_{14}Si$, *M* 114.3, *b* 85-88°, n_D^{20} 1.4121. Dissolve ~20g in THF (200ml), shake it with H_2O (2x 300 ml), dry (Na_2SO_4) and fractionate. This is a mixture of *cis* and *trans* isomers which can be separated by gas chromatography on an $AgNO_3$ column [for preparation: see Seyferth & Vaughan *J Organomet Chem* **1** 138 1963, DOI: 10.1016/S0022-328X(00)87444-6] at 25° with He as carrier gas at 9 psi. The *cis-isomer* has n_D^{25} 1.4105, and the *trans-isomer* has n_D^{25} 1.4062. [Seyferth et al. *Pure Appl Chem* **13** 159 1966, DOI: 10.1351/pac196613010159.]

1-Propyl-3-(p-chlorobenzenesulfonyl) urea [94-20-2] $C_{10}H_{13}ClN_2O_3S$, *M* 260.7, *m* 126-128°, 127-129°. Crystallise the urea from aqueous EtOH. It is soluble in $CHCl_3$ and Me_2CO , slightly soluble in Et_2O and $*C_6H_6$. [*Beilstein* **11** IV 119.]

Propylphosphonic acid (1-propanephosphonic acid) [4672-38-2] $C_3H_9O_3P$, *M* 124.1, *m* 71°, 73°, 73.8°, pK_1^{25} 2.49, pK_2^{25} 8.18 (H_2O). The phosphonic acid is purified by recrystallisation from hexane, heptane or $*C_6H_6$ to give long colourless needles, and is dried *in vacuo* over KOH. It is best prepared from di-*n*-butylphosphite (50mmol), which is converted into its sodium salt in dry hexane (150ml, by stirring under reflux until Na has dissolved, ~3-5 hours), treated with an equivalent of *n*-propylbromide and refluxed gently for 5-6 hours. After cooling, the mixture is washed thoroughly with H_2O , the organic layer is dried by distillation under a vacuum, and the residual *dibutyl n-propylphosphonate* is refluxed with 50-70ml of concentrated HCl overnight, and distilled from an oil bath to remove, BuCl and BuOH until *ca* 30ml are left, then carefully evaporated *in vacuo* and the residual solid is recrystallised from $*C_6H_6$ and/or hexane to give *n-propylphosphonic acid* (~80% yield) *m* 72.5-74.5°. **Diethyl propanephosphonate** [18812-51-6] $C_7H_{17}O_3P$, *M* 180.2 has *b* 88-89°/9mm, 94-96°/12mm, d_4^{25} 1.010, n_D^{20} 1.4172, and its ^{13}C NMR (25MHz, $MeCO$ d_6 , TMS) has δ_C at 61.8 (ester CH_3 , $^3J_{P,C-3}$ = 5.9 Hz), 61.3 (ester CH_2 , $^2J_{P,C-2}$ = 6.2 Hz), 28.2 (P- CH_2 , $^1J_{P,C-1}$ = 140.4 Hz), 16.7 ($-CH_2-CH_2-CH_3$, $^2J_{P,C-2}$ = 5.2 Hz) and 15.4 (propane CH_3 , $^3J_{P,C-3}$ = 16.2 Hz) ppm [Ernst *Org Mag Res* **9** 35 1977, DOI: 10.1002/mrc.1270090108]. [Kosolapoff *J Am Chem Soc* **67** 1180 1945, DOI: 10.1021/ja01223a045; *Beilstein* **4** H 596.]

Propylphosphonic anhydride (2,4,6-tripropyl-1,3,5,2,4,6-trioxatriphosphorinane-2,4,6-trioxide, T3P) [68957-94-8] $C_9H_{21}O_6P_3$, *M* 318.2 (cyclic trimer), has *b* 200°/0.3mm 200-350°/0.01-50mm. This reagent is prepared by heating *n*-propylphosphonic acid with acetic anhydride at 70-100°, and the polymeric phosphonic acid anhydride intermediate gives the trimeric cyclic anhydride on distilling at 200-350°/0.01-50mm. The an-

hydride is immediately made into 50% w/w solutions in DMF, CH_2Cl_2 , EtOAc or BuOAc, which are also commercially available. It was originally used as a **peptide coupling reagent**, but has found many applications which require removal of the elements of water from organic molecules. It allows the synthesis of a variety of heterocyclic compounds, acylation reactions which involves C-C coupling; in the presence of DMSO, alcohols can be oxidised to ketones; and with $\text{NH}_2\text{OH}\cdot\text{HCl}$ and Et_3N in MeCN, TBDMSCl is a selective reagent for hydroxyamidation of carboxylic acids. [Ariel & Llanes *Synlett* 1328 2007, DOI: 10.1055/s-2007-980339; Wehner et al. PCT Int. Appl. WO 2005014604 2005, *Chem Abstr* **142** 198208 2005, Wissmann & Kleiner *Angew Chem Int Ed* **19** 133 1980, DOI: 10.1002/anie.198001331; Escher & Bünning *Angew Chem Int Ed* **25** 277 1986, DOI: 10.1002/anie.19860277.]

1-Pyrenesulfonic acid [26651-23-0, 654055-00-2 hydrate] $\text{C}_{16}\text{H}_{10}\text{O}_3\text{S}\cdot x\text{H}_2\text{O}$, **M 202.2, m 125-129° (hydrate), >350°, $\text{pK}_{\text{Est}} < 0$** . Crystallise the sulfonic acid from EtOH/ H_2O . The **sulfonyl chloride** has **m 120°(dec)**. [Vollmann et al. *Justus Liebigs Ann Chem* **531** 1 32 1937, DOI: 10.1002/jlac.19375310102; and Tietze & Bayer *Justus Liebigs Ann Chem* **540** 189 1939, DOI: 10.1002/jlac.19395400113; *Beilstein* **11** H 198, **11** III 448.] The pyrene is a fluorogenic group.

1,3,6,8-Pyrenetetrasulfonic acid [6528-53-6] $\text{C}_{16}\text{H}_{10}\text{O}_{12}\text{S}_4$, **M 522.5**, has **m >400°, $\text{pK}_{\text{Est}} < 0$** . Crystallise the tetrasulfonic acid from water. The **tetra-Na salt** [59572-10-0] **M 614.5**, crystallises also from H_2O . [Tietze & Bayer *Justus Liebigs Ann Chem* **540** 189 1939, DOI: 10.1002/jlac.19395400113; *Beilstein* **11** III 486.]

3-(2-Pyridyl)-5,6-diphenyl-1,2,4-triazine-*p,p'*-disulfonic acid, monosodium salt (H_2O) [Ferrozine] [63451-29-6] $\text{C}_{20}\text{H}_{13}\text{N}_4\text{NaO}_6\text{S}_2$, **M 510.5, m >350°(dec)**. Purify it by recrystallisation from water or by dissolving it in the minimum volume of water, followed by addition of EtOH to precipitate the pure salt. It is light sensitive, complexes with Fe and used as a spectrophotometric reagent for Fe [Stokey *Anal Chem* **42** 779 1970, DOI: 10.1021/ac60289a016],

Pyrocatechol Violet (tetraphenolictriphenylmethanesulfonic acid Na salt) [115-41-3] $\text{C}_{19}\text{H}_{14}\text{O}_7\text{S}$, **M 386.4, ϵ 1.4×10^4 at 445nm in acetate buffer pH 5.2-5.4, $\text{pK}_{\text{Est}(1)} > 0$ (SO_3H), $\text{pK}_{\text{Est}(2)} \sim 9.4$, $\text{pK}_{\text{Est}(3)} \sim 13$** . It is recrystallised from glacial acetic acid. It is very **hygroscopic** and is a colour indicator standard for metal complex titrations: *viz* metal (pH, colour change, medium): Cd (10, blue/red-purple, NH_3 -buff); Cu (5-6.3, blue/yellow, AcOH-buff; 6-7, blue/yellow-green, pyridine; 9.3, blue/purple, NH_3 -buff); Fe^{2+} (3-6, blue/yellow, AcOH-buff); Ga (3.8, blue/yellow, AcOH-buff), In (5-6, blue/yellow, AcOH-buff); Mg (10, blue/red-purple, NH_3 -buff); Mn (9.3, blue/red-purple, NH_3 -buff. hydroxylamine); Ni (8-9.3, blue/red-violet, NH_3 -buff); Pb (5.5, blue/yellow, urotropin); Th (2.5-3.5, red/yellow, HNO_3 -soln); and Zn (10, blue/red-violet, NH_3 -buff). [Bishop *Indicators* pp 127-128, Oxford Pergamon Press 1972, Library of Congress Catalog Card No 78-171464, ISBN: 0080166172, 9780080166179; Ryba et al. *Coll Czech Chem Commun* **21** 349 1956, DOI: org/10.1135/cccc19560349; Cifka et al. *Coll Czech Chem Commun* **21** 1418 1956, DOI: org/10.1135/cccc19561418; Ryba et al. *Coll Czech Chem Commun* **23** 71 1958, DOI: org/10.1135/cccc19580071; Šír and R. Přibil *Coll Czech Chem Commun* **21** 866 1956, DOI: org/10.1135/cccc19560866; *Beilstein* **19/3** V 703.]

Pyrogallol Red (tetraphenolic xanthyliumphenylsulfonate) [32638-88-3] $\text{C}_{19}\text{H}_{12}\text{O}_8\text{S}$, **M 418.4, m >300°(dec), ϵ 4.3×10^4 at 542nm, pH 7.9-8.6, pK_1 2.71, pK_2 6.60, pK_3 10.41, pK_4 12.16 (5% aqueous EtOH)**. It is recrystallised from aqueous alkaline solution (Na_2CO_3 or NaOH) by precipitation on acidification. Filter the dye off and dry it in a vacuum. [Suk *Coll Czech Chem Commun* **31** 3127 1966, DOI: 10.1135/cccc19663127; *Beilstein* **19** H 407, **19** II 417, **19** III/IV 599, **19/10** V 226.]

Rose Bengal [Acid Red 94, 4,5,6,7-tetrachloro-2',4',5',7'-tetraiodofluorescein di-Na or di-K salt] [*di-Na salt* 632-69-9] $\text{C}_{20}\text{H}_2\text{Cl}_4\text{I}_4\text{Na}_2\text{O}_5$, **M 1017.6 (di-Na salt) [*di-K salt* 11121-48-5] M 1049.8 (di-K salt)**. This biological stain can be purified by chromatography on silica TLC using a 35:65 mix of EtOH/acetone as eluent. It is a biological stain for dead (not healthy) cells and mucous strands; and used for staining corneas and assessment of ocular surface damage [Feenstra et al. *Ophthalmology* **99** 605 1992, DOI: 10.1016/S0161-6420(92)31947-5; Doughty *Cont Lens Anterior Eye* **36(6)** 272 2013, DOI: 10.1016/j.clae.2013.07.008]. [*Beilstein* **19** II 261, **19** III/IV 2926.]

Selenopyronine (3,6-dimethylaminoselenaxanthene HCl) [85051-91-8] $C_{17}H_{19}N_2Se$. HCl, M 366.8, λ_{max} 571nm (ϵ 81,000). Purify by recrystallising it as the *hydrochloride* from hydrochloric acid. Alternatively recrystallisation from 10% HBF_4 provides the *borotetrafluoroborate salt* [84835-12-1]. [Fanghaenel et al. *J Phys Chem* **91** 3700 1987, DOI: 10.1021/j100297a048]. [*Beilstein* **18** II 434.]

Selenourea [630-10-4] CH_4N_2Se . M 123.0, m 200°(slow heating), 202-205°, 205-207°(dec), 214-215°(dec), **Kofler 235°(dec)**. Recrystallise it from the least volume of H_2O using Norite (preferably under N_2) to form colourless needles which are dried over P_2O_5 . It is air and light sensitive. It slowly turns moderately dark on storage even below 0°. Authors state that the solid can be kept in a refrigerator for extended periods with no apparent change other than moderate darkening. [For reactions with ketones and I_2 see King & Hlavacek *J Am Chem Soc* **73** 1864 1951, DOI: 10.1021/ja01148a529; synthesis: Bacher *Recl Trav Chim Pays-Bas* **62** 580 1943, DOI: 10.1002/recl.19430620903; synthesis: Hope *Acta Chem Scand* **18** 1800 1964, DOI: 10.3891/acta.chem.scand.18-1800]. The *Se-methyl iodide* provides yellow crystals from EtOH/Et₂O with m 187-188°(dec). *Methyl isoselenourea sulfate* (from 185g of selenourea and Me_2SO_4/N_2) is dissolved in 1L of hot H_2O (4-5 drops of 18M H_2SO_4 are added), and the metallic Se which cleaved off in the initial reaction is removed by filtering through a steam jacketed funnel, and the filtrate is cooled, when the *sulfate* (194g) crystallises out. A further amount (42g, total yield 85%, m 205-207°) can be obtained by reducing the filtrate to 100ml and adding an equal volume of EtOH. $MeSeH$ is **not** liberated on treatment with NaOH, but a yellow oil separates [Dunbar & Painter *J Am Chem Soc* **69** 1833 1947, DOI: 10.1021/ja01199a079]. The *N,N*-dimethyl *derivative* crystallises from H_2O or EtOH as colourless needles which slowly turn pink, then grey on standing, and although slightly soluble in *benzene it can be recrystallised from it and has m 167-170°(dec) [Zingaro et al. *J Org Chem* **18** 292 1953, DOI: 10.1021/jo01131a010; IR: Jensen & Nielsen *Acta Chem Scand* **20** 597 1966, DOI: 10.3891/acta.chem.scand.20-0597; *Beilstein* **3** IV 435.]

Silicon tetraacetate [562-90-3] $C_8H_{12}O_8Si$, M 264.3, m 110-111°, 111-115°, b 148°/5-6mm, pK_1^{25} 9.7, pK_2^{25} 11.9 (for H_4SiO_4 free acid). It can be crystallised from mixtures of CCl_4 and petroleum ether or Et₂O, or from acetic anhydride and then dried in a vacuum desiccator over KOH. Ac_2O adheres to the crystals and is removed first by drying at room temperature, then at 100° for several hours. It is soluble in Me_2CO , is very *hygroscopic* and effervesces with H_2O . It decomposes at 160-170°. Store at 0° to 6°. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 701 1963, *Beilstein* **2** H 171.]

Silver diethyldithiocarbamate [1470-61-7] $C_5H_{10}AgNS_2$, M 256.1, m 172°, 172-175°, 174°, 176-78°, pK_1^{25} 3.36 (for *N,N*-diethyldithio-carbamate). Purify it by recrystallisation from pyridine or CO_2 . Store it in a desiccator in a cool and dark place. [*Beilstein* **4** III 224, **4** IV 391.]

Silver tosylate [16836-95-6] $C_7H_7AgO_3S$, M 279.1. The anhydrous salt is obtained by recrystallisation from H_2O . Store it in the dark. [Claesson & Wallin *Chem Ber* **12** 1848 1879, DOI: 10.1002/cber.187901202170; *Beilstein* **11** H 97, 99.] Used for converting halides to tosylates [Kornblum et al. *J Am Chem Soc* **81** 4113 1959, DOI: 10.1021/ja01524a080; Hoffmann *J Chem Soc* 6748 1965, DOI: 10.1039/JR9650006748].

Silver trifluoromethanesulfonate (Silver triflate) [2923-28-6] $CAgF_3O_3S$, M 256.9, has m 286°, 356°, 356-357°. Recrystallise it twice from hot CCl_4 [Alo et al. *JCS Perkin Trans 1* 805 1986, DOI: 10.1039/P19860000805]. Store it in the dark. It is a useful halogen abstraction agent. [*Beilstein* **3** IV 34.]

Sodium *n*-alkylsulfates. Recrystallise these salts from EtOH/ Me_2CO [Hashimoto & Thomas *J Am Chem Soc* **107** 4655 1985, DOI: 10.1021/ja00302a010].

Sodium amylpenicillin [575-47-3] $C_{14}H_{21}N_2NaO_4S$, M 350.4, 336.4, m 188°(dec, anhydrous) [α]_D²³ +319° (c 1, H_2O). The *monohydrate* crystallises from moist acetone or moist ethyl acetate. Dry it in a vacuum. [Wintersteiner in 'The Chemistry of Penicillin', Clarke, Johnson and Robinson eds, Princeton University Press, p 470 1949.]

Sodium 9,10-anthraquinone-1,5-disulfonate (H_2O) [853-35-0] $C_{14}H_8Na_2O_8S_2$, M 412.3, m >300°. Separate it from insoluble impurities by continuous extraction with water. Recrystallise it twice from hot water and dry it under vacuum. [*Beilstein* **11** II 195, **11** III 634.]

Sodium 9,10-anthraquinone-1-sulfonate (H₂O) (Golden salt) [107439-61-2, 128-56-3 (Anhydr)] C₁₄H₇NaO₅S, M 328.3, 310.3 (Anhydr), m >300°, pK_{Est} ~<0 (SO₃H). Recrystallise it from hot water (4ml/g) after treatment with active charcoal, or from water by addition of EtOH. Dry it under vacuum over CaCl₂, or in an oven at 70°. Store it in the dark. [Beilstein 11 II 192, 11 III 626, 11 IV 670.]

Sodium 9,10-anthraquinone-2-sulfonate (H₂O) (9,10-anthraquinone-2-sulfonic acid [Na salt, H₂O]) [131-08-8, 152377-35-1 (Free acid)] C₁₄H₇NaO₅S, M 328.3, C₁₄H₈O₅S, M 287.3, has m >300°, pK_{Est} ~<0 (SO₃H). It crystallises from H₂O or MeOH (charcoal). It is useful for the determination of DNA as an *in situ* fluorescence photochemical probe [Li et al. *Analyt Chim Acta* 340 291 1997, DOI: 10.1016/S0003-2670(96)00446-1], and in *time-resolved fluorescence* studies of excited states in porphyrin-quinonesulfonic acid reversed micelles [Costa and Bookfield *JCS Faraday Trans 2*, 82 991 1986, DOI: 10.1039/F29868200991]. [Beilstein 11 IV 671.]

Sodium benzenesulfinate (benzenesulfinic acid Na salt) [873-55-2] C₆H₅NaO₂S, M 164.2, m >300°, pK₂₅ 2.16 (2.74, for PhSO₂H). Dissolve it in the minimum volume of O₂ free H₂O (prepared by bubbling N₂ through for 2 hours) and adding O₂ free EtOH (prepared as for H₂O), set aside at 4° overnight under N₂, filter, wash with EtOH, then Et₂O and dry *in vacuo*. The Na salt is relatively stable to air oxidation, but is best kept under N₂ in the dark. Also recrystallise it from EtOH and dry it at 120° for 4 hours in a vacuum. [Kornblum & Wade *J Org Chem* 52 5301 1987, DOI: 10.1021/jo00233a001; Beilstein 11 H 6, 11 IV 3.] Useful for the preparation of vinyl sulfones by reaction with vicinal dibromides [Guan et al. *Synthesis* 1465 2007, DOI: 10.1055/s-2007-966039], and aryl sulfones by Cu(II) acetate-catalysed cross-coupling with boronic acids [Huang & Batey *Tetrahedron* 63 7667 2007, DOI: 10.1016/j.tet.2007.05.029].

Sodium benzenesulfonate [515-42-4] C₆H₅NaO₃S, M 180.2, has pK₁²⁵ 0.70 (2.55) (for PhSO₃H₂). Crystallise it from EtOH or aqueous 70-100% MeOH, and dry it under a vacuum at 80-100°. [Fieser 1 1044; Beilstein 11 H 28, 11 I 10, 11 II 18, 11 III 33, 11 IV 27.]

Sodium bis(trimethylsilyl)amide (hexamethyl disilazane sodium salt) [1070-89-9] C₆H₁₈NaSi, M 183.4, m 165-167°(sintering at 140°), 171-175°, b 170°/2mm. It can be sublimed at 170°/2mm (bath temperature 220-250°) onto a cold finger, and can be recrystallised from *C₆H₆ (its solubility is: 10g in 100ml at 60°). It is soluble in THF, *C₆H₆ (see above) toluene, slightly soluble in Et₂O and is decomposed by H₂O. [Wannagat & Niederprüm *Chem Ber* 94 1540 1961, DOI: 10.1002/cber.19610940618.] It is available commercially under N₂ in Sure/Seal bottles in tetrahydrofuran (various concentrations) and at ~0.6M in toluene. [Fieser 1 1046; Beilstein 4 IV 4014.] It is a strong *deprotonating agent*, of esters and ketones to generate enolates and can be titrated in THF solution using *4-phenylbenzylidenbenzylamine* as indicator [Watson, B. T & Lebel, H. 'Sodium bis(trimethylsilyl)amide' in *Encyclopedia of Reagents for Organic Synthesis* (Ed: L. Paquette) 2004, J. Wiley & Sons, NY, DOI: 10.1002/047084289X.rs071m.pub2].

Sodium 4-bromobenzenesulfonate [5015-75-8, 79326-93-5 (Free acid monohydrate)] C₆H₄BrNaO₃S, M 258.7, m >300°. Crystallise it from MeOH, EtOH or distilled water. The *free acid monohydrate* has m 90-93° (M 255.1). [Beilstein 11 H 570, 11 I 14, 11 II 30, 11 III 97.]

Sodium cacodylate (3H₂O) [124-65-2, 6131-99-3] C₂H₁₂AsNaO₅, M 214.0, has m 60°, 77-80°, pK₂₅ 6.25. Recrystallise it from aqueous EtOH. Solubility in H₂O at 20° is 0.5M. [Beilstein 4 H 612, 4 I 576, 4 IV 1818, see cacodylic acid above.]

Sodium 4-chlorobenzenesulfonate [5138-90-9] C₆H₄ClNaO₃S, M 214.6, has pK_{Est} <0 (for SO₃H). Crystallise it twice from MeOH and dry it under vacuum. [Beilstein 11 IV 107.]

Sodium 3-chloro-5-methylbenzenesulfonate [5138-92-1] C₇H₆ClNaO₃S, M 228.7, has pK_{Est} <0 (for SO₃H). Crystallise it twice from MeOH and dry it under vacuum. [Beilstein 2 I 22.]

Sodium p-cymenesulfonate [77060-21-0] C₁₀H₁₃NaO₃S, M 236.3. Dissolve the salt in water, filter and evaporate to dryness. Recrystallise it twice from absolute EtOH and dry it at 110°.

Sodium 1-decanesulfonate [13419-61-9] C₁₀H₂₁NaO₃S, M 244.33, has m >300°. Recrystallise it from absolute EtOH and dry it over silica gel. [Beilstein 4 IV 62.]

Sodium n-decylsulfate [142-87-0] C₁₀H₂₁NaO₄S, M 260.3. Rigorously purify it by continuous Et₂O extraction of a 1% aqueous solution for two weeks. [Beilstein 1 IV 1818.] It has *detergent* activity, and is *toxic* to aquatic organisms.

Sodium dibenzylthiocarbamate [55310-46-8] $C_{15}H_{14}NNaS_2$, M 295.4 (anhydr), has m 230°(dec), 235°(dec), pK^{20} 3.13 (for monobenzyl-dithiocarbamic acid). The *free acid*, when recrystallised twice from dry Et_2O , has m 80-82°. The *Na salt* is reprecipitated from aqueous EtOH or EtOH by addition of Et_2O or Me_2CO [Linder et al. *Anal Chem* 50 896 1978, DOI: 10.1021/ac50029a019]. The *NH₄ salt* has m 130-133°, the *Cu salt* (yellow crystals) has m 284-286°, and the *Ti salt* has m 64-70°. [Beilstein 12 IV 2275.]

Sodium 2,5-dichlorobenzenesulfonate [5138-93-2] $C_6H_3Cl_2NaO_3S$, M 249.0, has $pK_{Est} < 0$ (for SO_3H). Crystallise it from MeOH, and dry it under vacuum. [Beilstein 11 III 93.]

Sodium diethyldithiocarbamate (3H₂O) [20624-25-3, 148-18-5] $C_5H_{10}NNaS_2$, M 225.3, 171.3 (anhydr), has m 94-96°(anhydrous), 95°, 98°, d 1.1 g/cm³, pK^{20} 3.65 (diethyldithiocarbamic acid). Recrystallise it from a small volume of H_2O , or dissolve it in the minimum volume of H_2O , add cold Me_2CO , collect it and dry it in air (cf. the dimethyl analogue [128-04-1] below). [Beilstein 4 III 224, 4 IV 390.] It complexes with softer metals [Cotton et al. *Advanced Inorganic Chemistry* (6th ed.), 1999, Wiley-Interscience NY, ISBN 0-471-19957-5], is a spin trap used with Fe(II) to detect NO in brain, liver and kidney [Vanin et al. *Methods Enzymology* 359 27 2002, DOI: 10.1016/s0076-6879(02)59169-2] and has fungicidal activity.

Sodium di(ethylhexyl)sulfosuccinate (Aerosol-OT, sodium docusate) [577-11-7] $C_{20}H_{37}NaO_7S$, M 444.6, has m 173-179°(dec). Dissolve it in MeOH and the inorganic salts which precipitate are filtered off. Water is added and the solution is extracted several times with hexane. The residue is evaporated to one-fifth its original volume, *benzene is added and azeotropic distillation is continued until no water remains. The solvent is evaporated. The white residual solid is crushed and dried *in vacuo* over P_2O_5 for 48 hours [El Seoud & Fendler *JCS Faraday Trans 1* 71 452 1975, DOI: 10.1039/F19757100452]. [Beilstein 4 IV 114.] It solubilises major myelin trans membrane proteolipids, and forms reverse micelles in hydrocarbon solvents [Saitoh et al. *J Chromatogr A* 1042 185 2004, DOI:10.1016/j.chroma.2004.04.010]. Used for making microemulsions for the simultaneous determination of natural and synthetic estrogens [Tripodi et al. *Electrophoresis* 27 4431 2006, DOI: 10.1002/elps.200600280]. It has laxative properties.

Sodium 2,2'-dihydroxy-1-naphthaleneazobenzene-5'-sulfonate See Solochrome Violet R [2092-55-9] below.

Sodium 2,4-dihydroxyphenylazobenzene-4'-sulfonate (Resorcinol yellow, Tropaeolin O, Acid Orange 6) [547-57-9] $C_{12}H_9N_2NaO_5S$, M 316.3. Recrystallise it from absolute EtOH, and it has high solubility in H_2O giving a yellow-red colour. It has λ_{max} at 490nm and CI (colour index) 14270; and is an indicator in the alkaline range imparting a yellow colour at pH ~11 and a brown-orange colour at pH ~12.6. Plasma is stained by it. [Beilstein 16 H 275, 16 IV 410.]

Sodium p-(p-dimethylaminobenzeneazo)-benzenesulfonate [23398-40-5] $C_{14}H_{14}N_3NaO_3S$, M 327.3. Recrystallise it from water and dry it *in vacuo*. [Beilstein 16 H 331, 16 I 317, 16 II 169, 16 III 370, 16 IV 510.]

Sodium 2,4-dimethylbenzenesulfonate [827-21-4] $C_8H_9NaO_3S$, M 208.2. Crystallise it from MeOH and dry it under vacuum. [Beilstein 11 III 339.]

Sodium 2,5-dimethylbenzenesulfonate (sodium p-xylenesulfonate) [827-19-0] $C_8H_9NaO_3S$, M 208.2. Dissolve it in distilled water, filter it, then evaporate it to dryness. Recrystallise it twice from absolute EtOH or MeOH and dry it at 110° under vacuum. [Beilstein 11 III 93, 11 IV 502.]

Sodium dimethyldithiocarbamate hydrate [128-04-1 (hydrate), 207233-95-2 (anhydrous)] $C_3H_6NNaS_2$, M 143.2 (anhydr), m 106-108°, 120-122°, pK^{25} 3.36 (diethyldithiocarbamic acid). Crystallise it from a small volume of H_2O , or dissolve it in the minimum volume of H_2O and add cold Me_2CO , collect it and dry it in air. The solubility in Me_2CO is ~12.5g/100ml. The *dihydrate* loses H_2O on heating at 115° to give the *hemi-hydrate* that decomposes on further heating [Kulka *Can J Chem* 34 1093 1956, DOI: 10.1139/v56-142]. [Beilstein 4 IV 233.] Fungicide and bactericide.

Sodium N,N-dimethylsulfanilate [2244-40-8] $C_8H_{10}NNaO_3S$, M 223.2, has m >300°. It crystallises from water. [See Beilstein 4 IV 270.]

Sodium 1-dodecanesulfonate [2386-53-0] $C_{12}H_{25}NaO_3S$, M 272.4, has m >300°. Recrystallise it twice from EtOH and dry it in an oven at 105° for 2 hours. It picks up moisture to form the 3.5 hydrate. Its *hydrate* crystallises in several phases. Forms micelles. [Tartar & Wright *J Am Chem Soc* 61 539 1939, DOI: 10.1021/ja01872a001; Wright & Tartar *J Am Chem Soc* 61 544 1939, DOI: 10.1021/ja01872a002; for preparation see Reed & Tartar *J Am Chem Soc* 57 571 1935, DOI: 10.1021/ja01306a055; Beilstein 4 III 27, 4 IV 64.]

Sodium 4-dodecylbenzenesulfonate [25155-30-0] $C_{18}H_{29}NaO_3S$, M 348.5. It crystallises from propan-2-ol or H_2O . [Gray et al. *J Org Chem* 20 511 1955, DOI: 10.1021/jo01122a014; Beilstein 11 IV 514.] This anionic detergent is a SKIN IRRITANT.

Sodium dodecylsulfate (SDS, sodium laurylsulfate) [151-21-3] $C_{12}H_{25}NaO_4S$, M 288.4, m 204-207°. Purify this *anionic detergent* by Soxhlet extraction with petroleum ether for 24 hours, followed by dissolution in acetone/MeOH/H₂O 90:5:5(v/v) and recrystallisation [For reversed micelles see Politi et al. *J Phys Chem* **89** 2345 1985, DOI: 10.1021/j100257a040]. It has been purified by two recrystallisations from absolute EtOH, aqueous 95% EtOH, MeOH, isopropanol or a 1:1 mixture of EtOH/isopropanol to remove dodecanol, and dried under vacuum. [Ramesh & Labes *J Am Chem Soc* **109** 3228 1987, DOI: 10.1021/ja00245a008]. SDS has also been purified by repeatedly foaming whereby a 0.15% aqueous solution is made to foam and the foam is discarded, then the H₂O is removed *in vacuo* and the residue is diluted to the required concentrations [Cockbain & McMullen *Trans Faraday Soc* **47** 322 1951, DOI: 10.1039/TF9514700322], or by liquid-liquid extraction [see Harrold *J Colloid Sci* **15** 280 1960, DOI: 10.1016/0095-8522(60)90029-5]. Dry it over silica gel. For DNA work it should be dissolved in excess MeOH passed through an activated charcoal column and evaporated until it crystallises out. It has also been purified by dissolving in hot 95% EtOH (14ml/g), filtering and cooling, then drying in a vacuum desiccator. *Alternatively*, it is crystallised from H₂O, vacuum dried, washed with anhydrous Et₂O and dried *in vacuo* again. These operations are repeated five times [for micellar effects see Maritato et al. *J Phys Chem* **89** 1341 1985, DOI: 10.1021/j100254a007; Lennox and McClelland *J Am Chem Soc* **108** 3771 1986, DOI: 10.1021/ja00273a035; Dressik et al. *J Am Chem Soc* **108** 7567 1986, DOI: 10.1021/ja00284a021]. [Beilstein **1** IV 1847.] It is very useful in electrophoresis (PAGE) and chromatography for denaturing proteins for the determination of their **sub-unit** molecular weights [Weber & Osborne *J Biol Chem* **244** 4406 1969, <http://www.jbc.org/content/244/16/4406>; Shapiro et al. *Biochem Biophys Res Commun* **28** 815 1967, DOI: 10.1016/0006-291X(67)90391-9].

Sodium ethylmercurithiosalicylate (Thimerosal) [54-64-8] $C_9H_9HgNaO_2S$, M 404.8, m ~230°. Recrystallise this antibacterial/antifungal salt from EtOH/Et₂O. It is soluble in H₂O (1g/ml) and EtOH (0.13g/ml). **HIGHLY TOXIC**. [Trikojus *Nature* **158** 472 1946, DOI: 10.1038/158472a0; Beilstein **10** III 213.] **Sodium ethylsulfate** [546-74-7] $C_2H_5NaO_4S$, M 148.1. Recrystallise it three times from MeOH/Et₂O and dry it in a vacuum. It is also available commercially as a solution in MeOH (1.0g/ml). [Beilstein **1** H 326, **1** I 164, **1** III 1317, **1** IV 1325.]

Sodium formaldehyde sulfoxylate dihydrate (sodium hydroxymethylsulfinate, Rongalite) [149-44-0] CH_3NaO_3S , M 118.1(anhydr), m 63-64° (dihydrate). It crystallises from H₂O (solubility is 0.05g/ml at ~20°) as the *dihydrate* and decomposes at higher temperatures. Store it in a closed container in a cool place. It is insoluble in EtOH and Et₂O and is a good reducing agent. [For X-ray structure see Truter *J Chem Soc* 3064 1955, DOI: 10.1039/JR9550003064.] **Note** that this compound {HOCH₂SO₂Na} should **not be confused** with formaldehyde sodium bisulfite adduct {HOCH₂SO₃Na} from which it is prepared by reduction with Zn. [Beilstein **1** IV 3052.]

Sodium hexadecylsulfate (cetyl sodium sulfate) [1120-01-0] $C_{16}H_{33}NaO_4S$, M 344.5, m 187-192°. Recrystallise it from absolute EtOH or MeOH and dry it in vacuum [Abu-Hamdiyyah & Rahman *J Phys Chem* **91** 1530 198, DOI: 10.1021/j100290a0487]. It is a low risk **irritant** of eyes, skin and respiratory tract, but handle with care.

Sodium 2-hydroxy-4-methoxybenzophenone-5-sulfonate [6628-37-1] $C_{14}H_{11}NaO_6S$, M 330.3. Crystallise it from MeOH and dry it under vacuum.

Sodium p-hydroxyphenylazobenzene-p'-sulfonate [2623-36-1] $C_{12}H_9N_2NaO_4S$, M 299.2. Recrystallise it from 95% EtOH. **Sodium metanilate** [1126-34-7] $C_6H_6NNaO_3S$, M 195.2, has m >300°. It crystallises from hot water. [Beilstein **14** H 688.]

Sodium methanethiolate [sodium methylmercaptide] [5188-07-8] CH_3NaS , M 70.1, has pK²⁵ 10.33 (MeS⁻). Dissolve the salt (10g) in EtOH (10ml) and add Et₂O (100ml). Cool and collect the precipitate, wash it with Et₂O and dry it in a vacuum. It is a white powder that is very soluble in EtOH and H₂O. [Chakraborty *e-EROS (Encyclopaedia of Reagents for Organic Synthesis)* October pp 1-5 2014, DOI: 10.1002/047084289X.rn01716; Billmann & Jensen *Bull Soc Chim Fr* **3** 2318 1936, Beilstein **1** III 1212.] C-21 thioethers of 16-prednisolone carboxylates were easily prepared from the corresponding mesylate esters by reaction with sodium thiolates (with 1.5 equivs of RSN₃/35°/1h) in 60-75% yields [Khan & Lee *Synth Commun* **409** 2007, DOI: 10.1080/00397910601038954].

Sodium 3-methyl-1-butanefulfonate [5343-41-9] $\text{C}_5\text{H}_{11}\text{NaO}_3\text{S}$, M 174.2. It crystallises from 90% MeOH.

Sodium 1-naphthalenesulfonate [130-14-3] $\text{C}_{10}\text{H}_7\text{NaO}_3\text{S}$, M 230.2. Recrystallise it from water or aqueous acetone [Okadata et al. *J Am Chem Soc* **108** 2863 1986]. [Beilstein **11** IV 521.]

Sodium 2-naphthalenesulfonate [532-02-5] $\text{C}_{10}\text{H}_7\text{NaO}_3\text{S}$, M 230.2. It crystallises from hot 10% aqueous NaOH or water and is dried in a steam oven. [Beilstein **11** IV 521.]

Sodium 2-naphthylamine-5,7-disulfonate (Amido-G-acid) [79004-97-0] $\text{C}_{10}\text{H}_7\text{NNa}_2\text{O}_6\text{S}_2 \cdot 6\text{H}_2\text{O}$, M 455.4. Crystallise it from water (charcoal, solubility at 20° is 72g/100ml) and dry it in a steam oven. Keep away from light. [Beilstein **14** H 784, **14** II 473, **14** IV 2811.]

Sodium 1-octanesulfonate H_2O [5324-84-5] $\text{C}_8\text{H}_{17}\text{NNaO}_3\text{S}$, M 216.3. Recrystallise it from absolute EtOH. [Beilstein **4** IV 58.] Used as an **ion-pairing reagent** for HPLC of peptides and proteins.

Sodium phenol-4-sulfonate ($2\text{H}_2\text{O}$) (4-hydroxybenzenesulfonic acid Na salt) [825-90-1, 10580-19-2 H_2O] $\text{C}_6\text{H}_5\text{NaO}_4\text{S}$, M 232.2, has m 300°. It crystallises from hot water (1g/1ml) by cooling to 0°, or from MeOH, and is dried in vacuum. [Beilstein **11** II 134.]

Sodium piperazine- N,N' -bis(2-ethanesulfonate) H_2O (PIPES-Na salt) [76836-02-7] $\text{C}_8\text{H}_{16}\text{N}_2\text{Na}_2\text{O}_6\text{S}_2$, M 364.3. It crystallises from water and EtOH. [Beilstein **23/2** V 380.] Useful in buffers.

Sodium isopropyl xanthate (sodium *O*-isopropylthiocarbonate) [140-93-2] $\text{C}_4\text{H}_7\text{NaOS}_2$, M 158.1, $\text{pK}^{25}_{\text{a}}$ 2.16 (for $-\text{S}^-$). It crystallises from ligroin/ethanol.

Sodium sulfanilate (sodium *p*-aminobenzenesulfonic acid) [515-74-2] $\text{C}_6\text{H}_6\text{NNaO}_3\text{S}$, M 195.2. It crystallises from water. [Beilstein **14** IV 2655.]

Sodium taurocholate [2-(3 α ,7 α ,12 α -trihydroxy-24-oxo-5- β -cholan-24-ylamino)ethanesulfonic acid sodium salt monohydrate] [145-42-6; 312693-83-7; 345909-26-4 ($x \text{ H}_2\text{O}$)] $\text{C}_{26}\text{H}_{44}\text{NNaO}_7\text{S}$, M 555.7 (monohydrate), m 168°dec (hydrate), $[\alpha]_{\text{D}}^{20} +23.9$ (c 2.5, H_2O), pK of acid is 1.4. The non-sulfated bile salt has been synthesised from *ethyl cholate* (m 162-163°, crystallised from EtOAc/petroleum ether b 30-60° 2:8) *via* the *hydrazide* (m 210°, sintering at 202°), which was diazotised to the *azide* (NaNO_2/HCl at 0-2°) and condensed with taurine in aqueous N NaOH at 8-14°/45 minutes. The acidified product was converted to *Na taurocholate* which was prepared and purified by precipitation with saturated aqueous NaCl and Et_2O (84% recovery; note that crystallisation does not occur unless enough H_2O is present) [Cortese *J Am Chem Soc* **59** 2532 1937, DOI: 10.1021/ja01291a014]. It was also purified by recrystallisation from aqueous EtOH/ Et_2O , or by gel chromatography using Sephadex LH-20. It is a useful **anionic detergent** for solubilising proteins and bilirubin [Woslewitz & Schroebler *Experientia* **35** 717 1979, PMID: 38133]. It has a CMC of 3-11 mM at 20-25°, with an average micellar weight of 2100. It is hydrolysed by mineral acids to cholic acid and taurine. [Tanaka *Z physiol Chem* **220** 39 1933, Beilstein **10** III 1655, **10** IV 2078.]

Sodium tetradecylsulfate (sodium meristyl sulfate) [1191-50-0] $\text{C}_{14}\text{H}_{30}\text{NaO}_4\text{S}$, M 316.4. It recrystallises from absolute EtOH [Abu Hamdiyyah & Rahman *J Phys Chem* **91** 1531 1987, DOI: 10.1021/j100290a048]. It is **hygroscopic**. [Beilstein **1** H 716, **1** IV 1866.] Used as a cosmetic ingredient.

Sodium tetrakis-(4-fluorophenyl)borate hydrate (Cesibor) [207683-22-5] $\text{C}_{24}\text{H}_{20}\text{BF}_4\text{NaO}_2$, M 450.2. This gravimetric reagent for Cs is purified by passing a solution (10g in 100ml H_2O) through a column of Dowex 50Wx4 (Na form) and eluting with dilute NaCl. Extract the 250-275ml eluate with Et_2O (3 x 50ml), add dry xylene (200ml), evaporate the Et_2O off *in vacuo*, immerse the xylene in a bath at 50° and the salt crystallises out. It is **hygroscopic**. [Moore et al. *Anal Chim Acta* **35** 1 1966, DOI: 10.1016/S0003-2670(01)81620-2 ; Tsubouci et al. *Anal Chem* **57** 783 1985, DOI: 10.1021/ac00280a051.]

Sodium tetraphenylborate [tetraphenyl boron Na] [143-66-8] $\text{C}_{24}\text{H}_{20}\text{BNa}$, M 342.2. Dissolve the **borate salt** in dry MeOH and add dry Et_2O . Collect the solid and dry it in a vacuum at 80°/2mm for 4 hours. It can also be extracted (Soxhlet) using CHCl_3 , and it crystallises from CHCl_3 as snow-white needles. It is freely soluble in H_2O , Me_2CO but insoluble in petroleum ether and Et_2O . An aqueous solution has pH ~ 5 and can be stored for days at 25° or lower, and for 5 days at 45° without deterioration. Its solubility in polar solvents increases with decrease in temperature [Wittig & Raff *Justus Liebigs Ann Chem* **573** 195 1951, DOI: 10.1002/jlac.19515730118]. The **salt** can also be recrystallised from acetone/hexane or CHCl_3 , or from Et_2O /cyclohexane (3:2) by warming the solution to precipitate the compound. Dry it in a vacuum at 80°. It dissolves in Me_2CO at 50-60° to give a clear solution. After standing at this temperature for 10 minutes the mix-

ture is filtered rapidly through a pre-heated Büchner funnel, cooled and the crystals are collected and dried in a vacuum desiccator at room temperature for 3 days [Abraham et al. *JCS Faraday Trans 1* **80** 489 1984, DOI: 10.1039/F19848000489]. If the product gives a turbid aqueous solution, the turbidity can be removed by treating with freshly prepared alumina gel and filtering. [Beilstein **16** IV 1624.]

Sodium thioglycolate (mercaptoacetic acid Na salt) [367-51-1] $\text{C}_2\text{H}_3\text{NaO}_2\text{S}$, **M 114.1**, **m >300°**. It crystallises from 60% EtOH (charcoal). It is *hygroscopic*. In concentrated aqueous solutions (~70%) it forms 1-2% of *thioglycolides* at ca percent per month which hydrolyse to the monomer on treatment of acid or alkali. Store at -20°. [Beilstein **3** IV 600.]

Sodium 4-toluenesulfinate [824-79-3, 7257-26-3 hydrate] $\text{C}_7\text{H}_7\text{NaO}_2\text{S}$, **M 178.2**, **m >290°**, **pK²⁵ 2.80 (1.99)(for -SO₂⁻)**. Recrystallise the salt from water (to constant UV spectrum) and dry it under vacuum, or extract it with hot *benzene, then dissolve it in EtOH/H₂O and heat with decolorising charcoal. The solution is filtered and cooled to give crystals of the *dihydrate*. Store in a cool dry place. [Beilstein **11** I 718, **11** IV 9]

Sodium 4-toluenesulfonate [657-84-1] $\text{C}_7\text{H}_7\text{NaO}_3\text{S}$, **M 194.2**, has **pK²⁵ -1.34 (for -SO₃⁻)**. Dissolve it in distilled water, filter it to remove insoluble impurities and evaporate it to dryness. Then recrystallise it from MeOH or EtOH, and dry it at 110°. Its solubility in EtOH is not high (maximum 2.5%), so that Soxhlet extraction with EtOH may be preferable. **Sodium p-toluenesulfonate** has also been crystallised from Et₂O and dried under a vacuum at 50°. [Beilstein **11** I 4, **11** II 6, **11** IV 241, cf. Gibson et al. *J Chem Soc* 874 1923, DOI: 10.1039/CT9232300874.]

Sodium 2,2',4-trihydroxyazobenzene-5'-sulfonate [3564-26-9] $\text{C}_{12}\text{H}_9\text{N}_2\text{NaO}_6\text{S}$, **M 300.3**, **pK₁ 0.10**, **pK₂ 6.14**, **pK₃ 7.72**, **pK₄ > 13**. Purify the dye by precipitating the *free acid* from aqueous solution using concentrated HCl, then wash it and extract it with EtOH in a Soxhlet extractor. Evaporation of the EtOH leaves the purified *acid* which is converted to the sodium salt with an equivalent of NaOH or Na₂CO₃. The characteristic constants (four pK_a values) and the UV spectra of the various species have been determined and the dye can be used for the determination of Zr [pK's and UV: Fletcher *Analyt Chem* **32** 1822 1960, DOI: 10.1021/ac50153a036; complexing with Zr: Fletcher *Analyt Chem* **32** 1827 1960, DOI: 10.1021/ac50153a037].

Sodium 2,4,6-trimethylbenzenesulfonate [6148-75-0] $\text{C}_9\text{H}_{11}\text{NaO}_3\text{S}$, **M 222.1**, has **m >300°**. Crystallise it twice from MeOH and dry it under vacuum. [Beilstein **11** III 345.]

Sodium trimethylsilanolate (sodium trimethylsilanol) [18027-10-6] $\text{C}_3\text{H}_9\text{NaSi}$, **M 112.2**, has **m 230°(dec)**. It is very soluble in Et₂O and *C₆H₆ but moderately soluble in petroleum ether. It is purified by sublimation at 130-150° in a high vacuum. [Hyde et al. *J Am Chem Soc* **75** 5615 1953, DOI: 10.1021/ja01118a042; Tatlock & Rochow *J Org Chem* **17** 1555 1952, DOI: 10.1021/jo50012a001; Beilstein **4** III 1856.]

Sodium 3,5-xylenesulfonate [30587-85-0] $\text{C}_8\text{H}_9\text{NaO}_3\text{S}$, **M 208.2**. Dissolve it in distilled water, filter, then evaporate it to dryness and recrystallise it twice from absolute EtOH and then dry it at 110°. [Beilstein **11** H 126, **11** I 34.]

Solochrome Violet R [4-hydroxy-3-(2-hydroxynaphthyl-1-ylazo)benzenesulfonic acid sodium salt, Java Chrome Violet B] [2092-55-9] $\text{C}_{16}\text{H}_{11}\text{N}_2\text{NaO}_5\text{S}$, **M 366.3**, **CI 15670**, **λ_{max} 501nm**, **pK₂²⁵ 7.22 (OH)**, **pK₃²⁵ 13.39 (OH)**. Convert the *acid* to the *monosodium* salt by precipitation with NaOAc/AcOH buffer of pH 4, then purify by precipitating the *free acid* from aqueous solution with concentrated HCl, wash and extract it with EtOH in a Soxhlet extractor. The acid precipitates on evaporating the EtOH and is reconverted to the sodium salt as described for *Chlorazole Sky Blue FF*. Dry it at 110°. It is *hygroscopic*. [Coates & Rigg *Trans Faraday Soc* **57** 1088 1961, DOI: 10.1039/TF9615701088; Beilstein **16** II 127.] It coordinates with metal ions, e.g. Cu²⁺ and Mg²⁺.

Strontium thiosalicylate (5H₂O) [15123-90-7] $\text{C}_7\text{H}_5\text{O}_2\text{SSr}$, **M 330.7**. It crystallises from hot water (0.5g/ml) by cooling to 0°. [Beilstein **10** IV 272.]

Sulfaguanidine (4-aminobenzenesulfonylguanidine) [57-67-0] $\text{C}_7\text{H}_{10}\text{N}_4\text{O}_2\text{S}$, **M 214.2**, **m 189-190°**, **b 488.4°/760mm**, **pK₁ 0.48**, **pK₂ 2.75**. Crystallise the antibacterial from hot water (7ml/g). [Beilstein **14** III 1970, **14** IV 2668.]

Sulfanilic acid (4-aminobenzenesulfonic acid) [121-57-3] $C_6H_7NO_3S$, M 173.2, m 288°(dec), $pK_1^{25} < 1$, pK_2^{25} 3.23. Crystallise the acid (as *dihydrate*) from boiling water. Dry it at 105° for 2-3 hours, then over 90% H_2SO_4 in a vacuum desiccator. The *S-benzylisothiuronium salt* has m 187° (from aqueous EtOH). [Beilstein 14 IV 2655.]

***o*-Sulfobenzoic acid (H_2O)** [123333-68-6 (H_2O), 632-25-7] $C_7H_6O_5S$, M 202.2, m 68-69°, 68-72° (hydrate?), 141°, $pK_{Est(1)} < 1$, $pK_{Est(2)} \sim 3.1$ (CO_2H). Crystallise the acid from water. The *S-benzylisothiuronium salt* has m 205.5-206.5° (from aqueous EtOH). [Beilstein 1 II 215, 1 III 658.] The *mono-ammonium salt* [6939-89-5] M 219.5, crystallises from water.

5-Sulfosalicylic acid [5965-83-3] $C_7H_6O_6S$, M 218.2, m 108-110°, 113°, 120°, $pK_1^{25} < 0$, pK_2^{25} 2.67, pK_3^{25} 11.67. Crystallise the acid from H_2O . Alternatively, it is converted to the *mono-sodium salt* which is crystallised from H_2O and washed with a little H_2O , EtOH and then Et_2O . The *acid* is recovered by acidifying. The *S-4-chlorobenzylisothiuronium salt* has m 181° (from dioxane). [Beilstein 11 H 411, 11 II 232, 11 III 704.] It precipitates proteins in affected urine; also used for integral colour anodising.

Sulfur trioxide pyridine complex [26412-87-3] $C_5H_5NO_3S$, M 159.2, m 155-165°, ~160°, 175°. Wash the solid with a little CCl_4 , then H_2O to remove traces of pyridine sulfate, and dry it over P_2O_5 . [Baumgarten *Chem Ber* 59 1166 1926, DOI: 10.1002/cber.19260590615; Beilstein 20/5 V 184]. It is useful for the sulfation of alcohols, sulfonations, deoxygenations and other reductions or oxidations involving DMSO [Olah et al. *Synthesis* 59 1979, DOI: 10.1055/s-1979-28557; Olah et al. *Synthesis* 984 1979, DOI: 10.1055/s-1979-28902; for conversion of α -amino and peptide aldehydes from respective alcohols without racemisation see Hamada & Shiori *Chem Pharm Bull Jpn* 30 1921 1982, DOI: org/10.1248/cpb.30.1921].

Tetrabutylammonium borohydride [33725-74-5] $C_{16}H_{40}BN$, M 257.3, m 123-128°, 128-129°. Purify it by recrystallisation from EtOAc followed by careful drying under vacuum at 50-60°. Samples purified in this way showed no signs of loss of *active H* after storage at room temperature for more than 1 year. Nevertheless samples should be stored at ca 6° in tightly stoppered bottles if they are to be kept for long periods (~1 year), with little loss of activity. It is soluble in CH_2Cl_2 and $CHCl_3$, moderately soluble in $*C_6H_6$, and poorly soluble in Et_2O and H_2O . [Raber & Guida *J Org Chem* 41 690 1976, DOI: 10.1021/jo00866a022; Brändström et al. *Tetrahedron Lett* 13 3173 1972, DOI: 10.1016/S0040-4039(01)93995-4.]

Tetrabutylammonium tetrafluoroborate [429-42-5] $C_{16}H_{36}BF_4N$, M 329.3, has m 161.8°, 161-163°, pK^{25} -4.9 (for HBF_4). Recrystallise it from H_2O , aqueous EtOH or from EtOAc by cooling in Dry-ice. Also recrystallise it from ethyl acetate/pentane or dry acetonitrile. Dry it at 80° under vacuum. [Detty & Jones *J Am Chem Soc* 109 5666 1987, DOI: 10.1021/ja00253a017; Hartley & Faulkner *J Am Chem Soc* 107 3436 1985, DOI: 10.1021/ja00298a007.] The *acetate* has m 118±2° (from BuCl), the *bromide* has m 118° (from EtOAc) and the *nitrate* has m 120° (from $*C_6H_6$). [Witschonke & Kraus *J Am Chem Soc* 69 2472 1947, DOI: 10.1021/ja01202a067; Wheeler & Sandstedt *J Am Chem Soc* 77 2024 1955, DOI: 10.1021/ja01612a102; Beilstein 4 IV 558.]

Tetraethoxysilane (tetraethyl orthosilicate) [78-10-4] $C_8H_{20}O_4Si$, M 208.3, m -77°, b 165-166°/atm, 169°/760mm, d_4^{20} 0.933, n_D^{25} 1.382. Fractionate it through an 80cm Podbielniak type column with a heated jacket and partial take-off head. It is soluble in EtOH, and is slowly decomposed by H_2O . It is *flammable*, and irritates the eyes and mucous membranes. [Sumrell & Ham *J Am Chem Soc* 78 5573 1956, DOI: 10.1021/ja01602a030; Bradley et al. *J Chem Soc* 5020 1952, DOI: 10.1039/JR9520005020; Beilstein 1 IV 1360.]

Tetraethylammonium hexafluorophosphate [429-07-2] $C_8H_{20}F_6NP$, M 275.2, m >300°, 331°(dec), $pK_1^{25} \sim 0.5$, pK_2^{25} 5.12 (for fluorophosphoric acid H_2PO_3F). Dissolve the salt (0.8g) in hot H_2O (3.3ml) and cool to crystallise. Yield of prisms is 0.5g. Its solubility in H_2O is 8.1g/L at 19° [Lange & Müller *Chem Ber* 63 1058 1930, DOI: 10.1002/cber.19300630510]. [Beilstein 4 III 199.] Used in electrolytes for electrochemical synthesis [Ruhlmann & Giraudeau *Chem Commun* 2007 1996, DOI: 10.1039/CC9960002007; Webster et al. *J*

Phys Chem **100** 10288 1996, DOI: 10.1021/jp9601173].

Tetraethylammonium tetrafluoroborate [429-06-1] $\text{C}_8\text{H}_{20}\text{BF}_4\text{N}$, **M 217.1**, **m 235°**, **356-367°**, **275-277°**, **289-291°**. **pK²⁵ -4.9 (for HBF₄)**. Dissolve the salt in hot MeOH, filter and add Et₂O. It is soluble in ethylene chloride [Thompson & Kraus *J Am Chem Soc* **69** 1016 1947, DOI: 10.1021/ja01197a012; Wheeler & Sandstedt *J Am Chem Soc* **77** 2025 1955, DOI: 10.1021/ja01612a103]. It has also been recrystallised three times from a mixture of ethyl acetate/hexane (5:1) or MeOH/petroleum ether, then kept at 95° for 48 hours under vacuum [Hartley & Faulkner *J Am Chem Soc* **107** 3436 1985, DOI: 10.1021/ja00298a007; Huang et al. *Anal Chem* **58** 2889 1986, DOI: 10.1021/ac00126a070]. It is used as a supporting electrolyte. [*Beilstein* **4** IV 333.]

Tetraethylsilane [631-36-7] $\text{C}_8\text{H}_{20}\text{Si}$, **M 144.3**, **m -82.5°**, **-82°**, **b 153.8°/760mm**, **d₄³⁰ 0.77**, **n_D³⁰ 1.427**. Fractionate it through a 3ft vacuum-jacketed column packed with 1/4" stainless steel saddles. The material is finally percolated through a 2ft column packed with alumina and maintained in an inert atmosphere. [Staveley et al. *J Chem Soc* 1992 1954, DOI: 10.1039/JR9540001992; Altshuller & Rosenblum *J Am Chem Soc* **77** 272 1955, DOI: 10.1021/ja01607a007; *Beilstein* **4** H 625, **4** IV 3895.]

1.1.3.3-Tetraisopropyldisiloxane [18043-71-5] $\text{C}_{12}\text{H}_{30}\text{OSi}_2$, **M 246.5**, **b 129-130°/6mm**, **d₄³⁰ 0.89**, **n_D³⁰ 1.47**. Fractionate it under reduced pressure in a N₂ atmosphere. [Gilman & Clark *J Am Chem Soc* **69** 1499 1947, DOI: 10.1021/ja01198a071.]

Tetrakis(hydroxymethyl)phosphonium chloride (THPC) [124-64-1] $\text{C}_4\text{H}_{12}\text{ClO}_4\text{P}$, **M 190.6**, **m 151°** (available also as an 80% solution in H₂O, d 1.341g/ml at 25° with ²⁰_D 1.512). THPC is prepared in an efficient fume cupboard by placing a mixture of 40% formaldehyde (90ml) and concentrated aqueous HCl (40ml, sp. gr. 1.2) into a 100ml 3-bulb Ladenburg flask (filling about half of the neck of the flask), the latter is provided with a stopper through which is passed a gas delivery tube that reaches the bottom. The flask is inclined to an almost horizontal position, when PH₃ (from a cylinder) is passed to displace the air, and the solution is heated to ~ 80° [*note* that at lower temperatures the reaction is slower, and at higher temperatures the vapour pressure of the solution decreases gas absorption]. The bulbs of the flask help to hold back the gas which results in a more complete reaction. The complete reaction may take up to many hours. Some H₂ and PH₃ which escape from the side neck of the flask should be vented through the flue of the fume cupboard. The mixture is then evaporated with stirring on a steam bath or *in vacuo* (~ 50° to 60°) until a white granular solid is obtained (50g, ~ 88%). Its purity is good enough for most purposes. However, it can be purified to analytical purity by recrystallising 1g of THPC from 50ml of AcOH. The low melting flat needles obtained which contain AcOH of crystallisation are converted to the pure substance, **m 151°**, by blowing dry air at 100° over the crystals, or by drying them at 100° *in vacuo* to give a constant weight. It can also be recrystallised from absolute EtOH. [For a Kg scale laboratory preparation see Reeves et al. *J Am Chem Soc* **77** 3923 1955, DOI: 10.1021/ja01619a074.] It is very *deliquescent* and should be kept in the presence of a desiccant. It is readily soluble in MeOH, slightly soluble in CHCl₃, but insoluble in Et₂O. It is quite stable, its aqueous solutions can be boiled without decomposition, and it is unaffected by dilute acids. The ¹H NMR (60MHz, D₂O, DSS) has δ for 4CH₂ at 4.77 (s, *J*_{PCH} = 1.7Hz and *J*_{13CH} = ~153Hz), and the ³¹P NMR (40MHz, D₂O, external 85% H₃PO₄) has δ at -25.8. With the stoichiometric amount of alkali, the *free tris(hydroxymethyl)phosphine* (see [2767-80-8] above) is obtained as an oil together with formaldehyde. However with excess of alkali on THPC, H₂ evolution occurs with the liberation of HCHO and formation of *tris(hydroxymethyl)phosphine oxide* (see [1067-12-5] below) **m 54-55°**. The reaction with alkali becomes more complicated when the mixture is heated, and *bis(hydroxymethyl)phosphinic acid* is formed. An 80% w/v aqueous solution of THPC with d₄²⁰ 1.33 is available commercially. Also a 70-75% aqueous solution of *bis[tetrakis(hydroxymethyl)phosphonium] sulfate* [55566-30-8] **M 406.3** is commercially available. [Hoffman *J Am Chem Soc* **43** 1684 1921, DOI: 10.1021/ja01440a035; Hoffman *J Am Chem Soc* **52** 2995 1930, DOI: 10.1021/ja01370a065; *Beilstein* **1** IV 3062.]

Tetramethoxysilane (tetramethyl orthosilicate) [681-84-5] $\text{C}_4\text{H}_{12}\text{O}_4\text{Si}$, **M 152.2**, **m 4-5°**, **b 122°/760mm**, **d₄²⁰ 1.023**, **n_D²⁰ 1.3688**. Purification is as for tetraethoxysilane. It has a vapour pressure of 2.5mm at 0°. [Sakurai in *Encyclopedia of Reagents for Organic Synthesis* 2001 John Wiley & Sons, NY, DOI: 10.1002/047084289X.rs012; IR: Sternbach & MacDiarmid *J Am Chem Soc* **81** 5109 1959, DOI: 10.1021/ja01528a023; *Beilstein* **1** IV 1266.]

Tetramethylammonium borohydride (TMAB) [16883-45-7] $C_4H_{16}BN$, **M 89.0**. Recrystallisation of the borohydride from H_2O three times yields *ca* 94% pure compound. Dry in high vacuum at 100° for 3 hours. The solubility in H_2O is 48% (20°), 61% (40°), in EtOH it is 0.5% (25°), and in MeCN it is 0.4% (25°). It decomposes slowly in a vacuum at 150° , but rapidly at 250° . The rate of hydrolysis of $Me_4N.BH_4$ (5.8M) in H_2O at 40° is constant over a period of 100 hours at 0.04% of original wt/hour. The rate decreases to 0.02%/hour in the presence of Me_4NOH (5% of the wt of $Me_4N.BH_4$). [Banus et al. *J Am Chem Soc* **74** 2346 1952, DOI: 10.1021/ja01129a048; *Beilstein* **4** IV 148.]

Tetramethylammonium hexafluorophosphate [558-32-7] $C_4H_{12}F_6NP$, **M 219.1**, has **m** $>300^\circ$, **d**₄²⁵ **1.617**, **pK**₁²⁵ **~ 0.5**, **pK**₂²⁵ **5.12** (for fluorophosphoric acid H_2PO_3F). The salt (0.63g) is recrystallised from boiling H_2O (76ml), yielding pure (0.45) $Me_4N.PF_6$ after drying at 100° . It is a good supporting electrolyte. [Lange & Müller *Chem Ber* **63** 1058 1930, DOI: 10.1002/cber.19300630510; *Beilstein* **4** III 110.]

Tetramethylammonium tetrafluoborate [661-36-9] $C_4H_{12}N^+BF_4^-$, **M 161.0**, has **m** **414° , 415° , 418°** . It is prepared by the reaction of tetramethylammonium halide or hydroxide with 40% fluoboric acid (or monohydroxy or methoxy fluoboric acids), in aqueous solution. After concentrating the solutions, adding Et_2O and shaking, the white salts are collected and dried under vacuum. It can be purified by dissolving in the minimum volume of H_2O (or MeOH) shaking with Et_2O , collecting the solid and drying in a vacuum until free of OH bands in the IR spectra. Whichever way it is prepared, the tetrafluoborate (not the trifluohydroxyborate) salt is formed as clearly shown by the absence of OH bands near $3000cm^{-1}$ in the IR spectra. [Moss & Sharp *J Inorg Nucl Chem* **13** 328 1960, DOI: 10.1016/0022-1902(60)80315-6; Wheeler et al. *J Am Chem Soc* **76** 6323 1954, DOI: 10.1021/ja01653a024; Wheeler & Nuttle *J Am Chem Soc* **76** 6322 1954, DOI: 10.1021/ja01653a023; Wheeler & Sandsted *J Am Chem Soc* **77** 2025 1955, DOI: 10.1021/ja01612a103.] [*Beilstein* **4** II 559, **4** IV 148.]

Tetramethylammonium triacetoxyborohydride [109704-53-2] $C_{10}H_{22}BNO_6$, **M 263.1**, has **m** **$93-98^\circ$, $96.5-98^\circ$** . If impure, wash it with freshly distilled Et_2O and dry it overnight in a vacuum to give a free flowing powder. Check 1H NMR, and if still suspect prepare it freshly from Me_4NBH_4 and AcOH in C_6H_6 and store it away from moisture [Banus et al. *J Am Chem Soc* **74** 2346 1952, DOI: 10.1021/ja01129a048; Evans & Chapman *Tetrahedron Lett* **27** 5939 1986, DOI: 10.1016/S0040-4039(00)85367-8; Iwamoto et al. *Tetrahedron Lett* **44** 7239 2003, DOI: 10.1016/j.tetlet.2003.08.009; Shangguan et al. *Org Lett* **9** 1093 2007, DOI: 10.1021/ol063143k]. It is an **IRRITANT** and **MOISTURE SENSITIVE**.

Tetramethylammonium triphenylborofluoride [437-11-6] $C_4H_{12}N^+C_{18}H_{15}BF_4^-$, **M 392.2**. Crystallise it from acetone or acetone/ethanol. [Gordon *Annual Review of Physical Chemistry* **1** 59 1950, DOI: 10.1146/annurev.pc.01.100150.000423.]

2,4,6,8-Tetramethylcyclotetrasiloxane (TMCTS) [2370-88-9] $C_4H_{16}O_4Si_4$, **M 240.4**, **m** **-69°** , **b** **$134^\circ/750mm$, $134.5-134.9^\circ/755mm$** , **d**₄²⁰ **0.99**, **n**_D²⁰ **1.3872**. It is purified by repeated redistillation, and fractions with the required 1H NMR data are collected. [Sokolov *J Gen Chem USSR (Engl Transl)* **29** 262 1959, Sauer et al. *J Am Chem Soc* **68** 962 1946, DOI: 10.1021/ja01210a014]. [*Beilstein* **4** IV 4099.] A useful component of photochemically formed SiO_x monolayers on oxide (TiO_2) semiconductors [Tada & Tanaka *Thin Solid Films* **281** 404 1996, DOI: 10.1016/0040-6090(96)08692-0].

1,1,3,3-Tetramethyldisiloxane [3277-26-7] $C_4H_{14}OSi_2$, **M 134.3**, **m** **-78°** , **b** **$70.5-71^\circ/731mm$, $71-72^\circ/atm$** , **d**₄³⁰ **0.75**, **n**_D²⁵ **11.367**. Possible impurity is 1,1,5,5-tetramethyl-3-trimethylsiloxytrisiloxane **b** $154-155^\circ/733mm$. Fractionate it, collect fractions boiling below 80° and re-fractionate it. Its purity can be analysed by alkaline hydrolysis and measuring the volume of H_2 liberated followed by gravimetric estimation of silica in the hydrolysate. It is unchanged when stored in glass containers in the absence of moisture for 2-3 weeks. Small amounts of H_2 are liberated on long storage. *Care should be taken when opening a container due to developed pressure*. [Speier et al. *J Am Chem Soc* **79** 974 1958, DOI: 10.1021/ja01561a054; Emeléus & Smythe *J Chem Soc* 609 1958, DOI: 10.1039/JR9580000609; IR: Kriegsmann *Z Anorg Chem* **299** 78 1959, DOI: 10.1002/zaac.19592990110; *Beilstein* **4** IV 3991.]

N,N,N',N'-Tetramethylphosphonic diamide (methylphosphonic bis-dimethylamide) [2511-17-3] $C_5H_{15}N_2OP$, **M 150.2**, **b** **$60.5^\circ/0.6mm$, $138^\circ/32mm$, $230-230^\circ/atm$** , **d**₄³⁰ **1.0157**, **n**_D³⁰ **1.4539**. Dissolve it in heptane or ethylbenzene, shake this with 30% aqueous NaOH, stir for 1 hour, separate the organic layer and fractionate. [Kosolapoff & Payne *J Org Chem* **21** 413 1956, DOI: 10.1021/jo01110a009.] Its IR (film) has ν_{max} at 1480, 1460, 1300, 1184, 1065 and $988-970cm^{-1}$ [Harvey & Mayhood *Can J Chem* **33** 1552 1955,

DOI: 10.1139/v55-190].

Tetramethylsilane (TMS) [75-76-3] $C_4H_{12}Si$, M 88.2, m -99° , b $26-28^\circ/atm$, $26.3^\circ/760mm$, d_4^{20} 0.639, n_D^{20} 1.359. Distil it from concentrated H_2SO_4 (after shaking with it) or $LiAlH_4$, through a 5ft vacuum-jacketed column packed with glass helices into an ice-cooled condenser, then percolate it through silica gel to remove traces of halide. [For preparation on a 250g scale see Whitmore & Sommer *J Am Chem Soc* **68** 481 1946, DOI: 10.1021/ja01207a036; *Beilstein* **4** IV 3875.]

2,4,6,8-Tetramethyl tetravinyl cyclotetrasiloxane [2554-06-5] $C_{12}H_{24}O_4Si_4$, M 344.7, m -44° , -43.5° , b $111-112^\circ/10mm$, $145-146^\circ/13mm$, $224-224.5^\circ/758mm$, d_4^{25} 0.997, n_D^{20} 1.434. A 7ml sample can be distilled in a small Vigreux column at atmospheric pressure without polymerisation or decomposition. It is soluble in cyclohexane. [Kantor et al. *J Am Chem Soc* **77** 1685 1955, DOI: 10.1021/ja01611a090; *Beilstein* **4** IV 4184.]

Tetraphenylarsonium (V) chloride hydrate [507-28-8] $C_{24}H_{20}AsCl$, M 418.8(anhydr), m $258-260^\circ$, $261-263^\circ$. A neutralised aqueous solution is evaporated to dryness. The residue is extracted into absolute EtOH, evaporated to a small volume and precipitated by addition of absolute Et_2O . It is again dissolved in a small volume of absolute EtOH or ethyl acetate and re-precipitated with Et_2O . Alternatively, it is purified by adding concentrated HCl to precipitate the *chloride dihydrate*. Redissolve in water, neutralise with Na_2CO_3 and evaporate to dryness. The residue is extracted with $CHCl_3$ and finally crystallised from CH_2Cl_2 or EtOH by adding Et_2O . If the aqueous layer is somewhat turbid treat it with Celite and filter it through filter paper. It can be dehydrated before use in a vacuum. The *tetrafluoroborate* salt has m $293-295^\circ$ (needles from MeCN), and the *picrate* salt has m $203-204^\circ$ (from EtOH). [Blicke & Monroe *J Am Chem Soc* **57** 720 1935, DOI: 10.1021/ja01307a038; Duke & Brown *J Am Chem Soc* **76** 1443 1954, DOI: 10.1021/ja01634a091; Popov & Humphrey *J Am Chem Soc* **81** 2043 1959, DOI: 10.1021/ja01518a003; Singhal & Raj *Synth Inorg Met-org Chem* **23** 1011 1993, DOI: 10.1080/15533179308016878; *Beilstein* **16** III 1006, **16** IV 1170.] It is a useful reagent for Cd, Hg, Zn, as well as ClO_4 , and IO_4 among other ions. **POISONOUS**.

Tetraphenylarsonium iodide [7422-32-4] $C_{24}H_{20}I$, M 510.2. It crystallises from MeOH. [Blicke & Monroe *J Am Chem Soc* **57** 720 1935, DOI: 10.1021/ja01307a038; Chatt & Mann *J Chem Soc* 1192 1940, DOI: 10.1039/JR9400001192.] [For crystal structure see Dean et al. *Acta Cryst Section C* **59** 484 2006, DOI: 10.1107/S0108270103022650]. **POISONOUS**.

Tetraphenylarsonium perchlorate [3084-10-4] $C_{24}H_{20}ClO_4$, M 482.8, has pK^{25} -2.4 to -3.1 (for $HClO_4$). It crystallises from MeOH. [Horner & Haufe *Chem Ber* **101** 2903 1968, DOI: 10.1002/cber.19681010841.] **POISONOUS** and possibly explosive.

Tetraphenylbiphosphine (Ph_2P-PPh_2) [1101-41-3] $C_{24}H_{20}P_2$, M 370.4, m 120.5° (evacuated tube), $120-122^\circ$ (sealed tube), b $258-260^\circ/1mm$. $pK_{Est} > 0.0$. This useful precursor of diphenylphosphine compounds is made by heating a mixture of diphenylphosphine (6.0g, 32.5mmol, 829-85-6) and chlorodiphenylphosphine (7.1g, 32.5mmol, 1709-66-9) in petroleum ether (100ml, b $90-100^\circ$, freshly distilled over Na) under N_2 at reflux for 3.5 hours during which time all the HCl has evolved. The white crystalline biphosphine that separates (~80%) on cooling is collected, washed with ligroin, dried *in vacuo* and recrystallised or distilled at high vacuum, preferably in a N_2 atmosphere. It is soluble in C_6H_6 , toluene, CCl_4 , and pyridine (yellow solution), and slightly soluble in Et_2O , EtOH and ligroin. The determined molecular weight was 391.6, 374.9 (method of Rast). When it is heated for 3 hours at $250-300^\circ$ under N_2 , phosphorus is liberated. The brown residue is boiled for 1 hour with dilute aqueous NaOH and 3% of H_2O_2 , filtered, cooled, the solid that is collected gives *Ph₃PO* (m 153° and mixed m with authentic Ph_3PO) after crystallisation from petroleum ether (b $90-100^\circ$). *Diphenylphosphonic acid* [m $191-192^\circ$ and mixed melting point with authentic $Ph_2P(O)OH$] can be isolated from the mother liquors, making a total yield of products almost quantitative. Reaction of a suspension of the biphosphine (3.3g) in CCl_4 under N_2 with Br (1ml) in CCl_4 is decolourised as it becomes clear. The mixture is then distilled and the oily residue is redistilled to give *bromodiphenylphosphine* (b $146.5-148^\circ/2.5mm$) as a colourless oil in almost quantitative yield. Bubbling dry air through a suspension of the biphosphine (3.3g) in dry toluene (30ml) at $\sim 0^\circ$ for 3 hours causes the mixture to become yellow in colour, and after dilution with toluene (30ml) and boiling for a short period it becomes clear. On cooling, and recrystallising the solid that separates from fresh toluene provides the white *dioxide $Ph_2P(O)-P(O)Ph_2$* (m 167° , evacuated tube, 1.9g, 53%), with a molecular weight of 418.5, 409.8 (measured by the method of Rast; the required value is 402.4). [Kuchen

& Buchwald *Chem Ber* **91** 2871 1958, DOI: 10.1002/cber.19580911246; *Beilstein* **16** H 2871.]

Sodium dipenylphosphine (Ph_2PNa) is readily prepared by adding Na to a solution of the diphosphine in dry Et_2O or THF (exothermic!), whereby the colour becomes yellow and the orange Na salt separates. It is sensitive to moisture (giving Ph_2PH) and to CO_2 (giving Ph_2PCOOH); and the necessary precautions have to be taken. It is a useful reagent for preparing a variety of Ph_2P-R derivatives and it is best to prepare it freshly when required. [Kuchen & Buchwald *Chem Ber* **92** 227 1959, DOI: 10.1002/cber.19590920126.]

Tetraphenylphosphonium chloride [2001-45-8] $C_{24}H_{20}ClP$, **M 374.9, m 273-275°**. Crystallise the chloride from acetone and dry at 70° under vacuum. It can also be recrystallised from a mixture of 1:1 or 1:2 dichloromethane/petroleum ether, the solvents having been dried over anhydrous K_2CO_3 . The purified salt is dried at room temperature under a vacuum for 3 days, and at 170° for a further 3 days. It also crystallises from isoPrOH/ Et_2O or EtOH/ Et_2O . **Extremely hygroscopic**. [Wittig & Geissler *Justus Liebigs Ann Chem* **580** 44, 50 1953, DOI: 10.1002/jlac.19535800107; Willard et al. *J Am Chem Soc* **70** 737 1948, DOI: 10.1021/ja01182a088; *Beilstein* **16** III 851, **16** IV 984.] [For crystal structure of the iodide see Dean et al. *Acta Cryst Section C* **59** m484 2006, DOI: 10.1107/S0108270103022650].

Tetraphenylsilane [1048-08-4] $C_{24}H_{20}Si$, **M 336.4, m 231-233°, 234-235°, 236.5°, 237°, 236-238°, b 228°/3mm, 479.6°/760mm**. It crystallises from *benzene as clear colourless bladed needles. It decomposes at ~360°/~760mm on attempted distillation. [George et al. *J Am Chem Soc* **77** 6647 1955, DOI: 10.1021/ja01629a079; Polis *Chem Ber* **98** 1540 1885, DOI: 10.1002/cber.188501801334; Drew & Landquist *J Chem Soc* 1480 1935, DOI: 10.1039/JR9350001480; *Beilstein* **16** I 525, **16** II 606, **16** III 1199, **16** IV 1372.]

Tetrasodium pyrene-1,3,6,8-tetrasulfonate [59572-10-0] $C_{16}H_6Na_4O_{12}S_4$, **M 610.5**. Recrystallise this salt from aqueous acetone [Okahata et al. *J Am Chem Soc* **108** 2863 1986, DOI: 10.1021/ja00271a013].

Thexyl dimethyl silyl chloride (TDSCl, dimethyl-[2,3-dimethyl-2-butyl] chlorosilane) [67373-56-2] $C_8H_{19}ClSi$, **M 178.8, b 55-56°/10mm, 158-159°/720mm, d_4^{20} 0.970, n_D^{20} 1.428**. Purify this chlorosilane by fractional distillation, and store it in small aliquots in sealed ampoules. It is very sensitive to moisture and is estimated by dissolving an aliquot in excess of 0.1M NaOH and titrating with 0.1M HCl using methyl red as indicator [Szabó et al. *Helv Chim Acta* **67** 2128 1984, DOI: 10.1002/hlca.19840670813].

N-(Thexyl dimethylsilyl)dimethylamine (N-[2,3-dimethyl-2-butyl]dimethylsilyl dimethyl-amine) [81484-86-8] $C_{10}H_{25}NSi$, **M 187.4, b 156-160°/720mm**. Dissolve the amine in hexane, filter, evaporate and distil. It is a colourless oil which is extremely sensitive to moisture. It is best to store small quantities in sealed ampoules after distillation. For estimation of purity, crush an ampoule in excess 0.1N HCl and titrate the excess acid with 0.1M NaOH using methyl red as indicator. [Szabó et al. *Helv Chim Acta* **67** 2128 1984 DOI: 10.1002/hlca.19840670813.]

Thioacetanilide [637-53-6] C_8H_9NS , **M 151.2, m 75-76°, 76-79°, b 225.1°/atm, d 1.159, pK_{Est} ~13.1**. Crystallise thioacetanilide from H_2O and dry it *in vacuo*. [*Beilstein* **12** I 193, **12** II 142, **12** III 464, **12** IV 378.]

Thiobenzanilide [636-04-4] $C_{13}H_{11}NS$, **M 213.2, m 101.5-102°, pK_{Est} ~12.6**. Crystallise thiobenzanilide from MeOH at Dry-ice temperature.

Thio-Michler's Ketone (TMK, [4,4'-bis(dimethylamino)thiobenzophenone] [1226-46-6] $C_{17}H_{20}N_2S$, **M 284.4, m 102-106°, λ_{max} 457 nm (ϵ 2.92 x 10⁴ in 30% aqueous *n*-propanol)**. Purify the thioketone by recrystallisation from hot EtOH or by trituration with a small volume of $CHCl_3$, followed by filtration and washing with hot EtOH [Tarbell & Wystrach *J Phys Chem* **68** 2110 1946, DOI: 10.1021/ja01214a514]. It also recrystallises from $CHCl_3$ / MeOH. [*Beilstein* **14** H 101, **14** I 395, **14** II 60.]

1-Thionaphthol [529-36-2] $C_{10}H_8S$, **M 160.2, b 106°/1.5mm, 208.5°/200mm, d_4^{20} 1.161, n_D^{20} 1.6802, pK^{25} 6.34**. It is steam volatile and is purified by distillation in the absence of O_2 , as it oxidises to the disulfide. It is soluble in Et_2O and EtOH but very slightly soluble in H_2O and dilute alkalis. The *S-ethyl derivative*,

[17539-31-0] **M** 188.2, has **b** 175-176°/25mm, and **d**^o 1.120. [Beilstein 6 III 2943, 6 IV 4241.] It may cause eye, skin, respiratory and digestive tract irritation.

2-Thionaphthol [91-60-1] has **m** 79-81°, **81.8-82.4°**, **82°**, **b** 153.5°/15mm, **286°/760mm**, **pK**²⁵ **6.47**. It is steam volatile. It has to be distilled under argon or N₂, as it oxidises to the disulfide, and crystallises from EtOH. It is very soluble in Et₂O and petroleum ether, but slightly soluble in H₂O. The *S*-methyl derivative has **m** 104-105° (from *C₆H₆/petroleum ether), and the *S*-ethyl derivative [32551-87-4] **M** 188.2, has **m** 16° and **b** 175-170.5°/15mm. The *S*-acetate [831-23-2] has **m** 53.5° and **b** 191°/15mm, and the diethylamine salt forms yellow needles **m** 107° from dioxane. [Beilstein 6 H 657, 6 I 316, 6 II 610, 6 III 3006, 6 IV 4312.]

Thiophenol (benzenethiol) [108-98-5] **C**₆**H**₆**S**, **M** 110.2, **m** -14.9°, -15°, **b** 46.4°/10mm, **168.0°/760mm**, **d**₄²⁰ **1.073**, **n**_D²⁰ **1.5897**, **pK**²⁵ **6.62**. Dry thiophenol with CaCl₂ or CaSO₄, and distil it at 10mm pressure or at 100mm (**b** 103.5°) in a stream of nitrogen. The **2,4-dinitrophenyl thioether** has **m** 121° (from EtOH), and the **2,4-dinitrophenyl sulfone** has **m** 161° (from EtOH). [Beilstein 6 IV 1463.]

Thiosalicylic (2-mercaptobenzoic) acid [147-93-3] **C**₇**H**₆**O**₂**S**, **M** 154.2, **m** 164-165°, 167°, **165-168°**, **d** 1.49, **pK**₁²⁵ **3.54**, **pK**₂²⁵ **8.80**. Crystallise the thio acid from hot EtOH (4ml/g), after adding hot distilled water (8ml/g) and boiling with charcoal. The hot solution is filtered, cooled, the solid is collected and dried *in vacuo* (P₂O₅). Crystallise it from AcOH and sublime *in vacuo*. [Beilstein 10 IV 272.]

o-Toluenesulfonamide [88-19-7] **C**₇**H**₉**NO**₂**S**, **M** 171.2, has **m** 155.5°. Crystallise the amide from hot H₂O (**m** 153°), then from EtOH or Et₂O/petroleum ether. The *N*-o-toluenesulfonylphthalimide has **m** 182° (from EtOH). [Evans & Dehn *J Am Chem Soc* 51 3651 1929, DOI: 10.1021/ja01387a027; Beilstein 11 H 86, 11 I 23, 11 II 39, 11 III 167, 11 IV 229.]

p-Toluenesulfonic acid [6192-52-5] **C**₇**H**₈**O**₃**S**, **M** 190.2, **m** 38° (anhydrous), **m** 105-107° (monohydrate), **pK**²⁵ **1.55**. Purify the acid by precipitation from a saturated solution at 0° by introducing HCl gas. It can also be crystallised from concentrated HCl, then crystallised from dilute HCl (charcoal) to remove benzenesulfonic acid. It has been crystallised from EtOH/water. Dry it in a vacuum desiccator over solid KOH and CaCl₂. *p*-Toluenesulfonic acid can be dehydrated by azeotropic distillation with *benzene or by heating at 100° for 4 hours under water-pump vacuum. The *anhydrous acid* can be crystallised from *benzene, CHCl₃, ethyl acetate, anhydrous MeOH, or from acetone by adding a large excess of *benzene. It can also be dried under vacuum at 50°. The *S*-benzylisothiuronium salt has **m** 182° (from aqueous EtOH). [Beilstein 11 IV 241.]

p-Toluenesulfonyl chloride (tosyl chloride) [98-59-9] **C**₇**H**₇**ClO**₂**S**, **M** 190.7, has **m** 66-69°, **67.5-68.5°**, **69°**, **b** 138-139°/9mm, **146°/15mm**, **167°/36mm**. Material that has been standing for a long time contains tosic acid and HCl and has **m** *ca* 65-68°. It is purified by dissolving (10g) in the minimum volume of CHCl₃ (*ca* 25ml) filtered, and diluting with five volumes (i.e. 125ml) of petroleum ether (**b** 30-60°) to precipitate impurities. The solution is filtered, clarified with charcoal and concentrated to 40ml by evaporation. Further evaporation to a very small volume gives 7g of white crystals which are analytically pure, **m** 67.5-68.5°. (The insoluble material is largely tosic acid and has **m** 101-104°.) [Pelletier *Chemistry and Industry* 1034 1953.] It also crystallises from toluene/petroleum ether in the cold, from petroleum ether (**b** 40-60°) or *benzene. Its solution in diethyl ether has been washed with aqueous 10% NaOH until colourless, then dried (Na₂SO₄) and crystallised by cooling in powdered Dry-ice. It has also been purified by dissolving in *benzene, washing with aqueous 5% NaOH, then dried with K₂CO₃ or MgSO₄, and distilled under reduced pressure and can be sublimed at high vacuum [Ebel *Chem Ber* 60 2079 1927, DOI: 10.1002/cber.19270600909]. [Beilstein 11 IV 375.] It is best prepared by the chlorosulfonation of toluene [Linder & Rodefild *Ullmann's Encyclopedia of Industrial Chemistry*, Wiley-VCH, DOI: 10.1002/14356007.a03_507; for reactions see Whittaker et al. e-EROS (*Encyclopedia of Reagents for Organic Synthesis*) 15 September 2006, DOI: 10.1002/047084289X.rt136.pub2]. **p-Toluenesulfonamide** [70-55-3] has **m** 137-137.5°, **138°**. Crystallise the amide from hot water, then from EtOH or Et₂O/petroleum ether. [Beilstein 11 H 104, 11 IV 376.]

p-Toluenesulfonic acid hydrazide (tosylhydrazide) [1576-35-8] **C**₇**H**₁₀**N**₂**O**₂**S**, **M** 186.2, **m** 108-110°, **109-110°**. Dissolve the hydrazide in hot MeOH (~1g/4ml), filter through Celite and precipitate the material by adding 2-2.5 volumes of distilled H₂O. Dry it in air or in a vacuum. It solubility in H₂O is 5g/L at 15°. [Friedman et al. *Org Synth Coll Vol* 5 1055 1973, DOI: 10.1522/orgsyn.040.0093; Beilstein 11 II 66.]

***p*-Toluenethiol (*p*-thiocresol)** [106-45-6] C_7H_8S , **M 124.2, m 43.5-44°, pK²⁵ 6.82**. Crystallise the thiol from petroleum ether (b 40-70°). The **2,4-dinitrophenyl thioether** has **m 103°** (from EtOH), and the **2,4-dinitrophenyl sulfone** has **m 190°** (from EtOH). [Beilstein 6 IV 2153.]

***p*-Toluenethiosulfonic acid potassium salt (potassium *p*-toluenethiosulfonate)** [28519-50-8] $C_7H_7KO_2S_2$, **M 226.4, m 227-229°**. When a solution of KOH (64.9g, 86.5%, 1.0 mole) in H₂O (28ml) is cooled in an ice bath while it is saturated with H₂S (in an efficient **fume cupboard**) and excess of H₂S is flushed out with N₂ (important to remove excess of H₂S), a fresh solution of **KHS** is obtained. This is diluted with H₂O (117ml) and stirred, under N₂ at 55-60°, while ground *p*-toluenesulfonyl chloride (95.3g, 0.5mole, freed from any acid by dissolving in *C₆H₆, washing with 5% aqueous NaOH, drying with Na₂SO₄, evaporating and distilling at 146°/15mm) at such a rate as not to allow the temperature to rise above 60°. The deep yellow colour of the solution disappears after 90g is added and the chloride ceases to dissolve. The mixture is filtered rapidly with suction through a warmed funnel, and the filtrate is set aside at 0-5° for many hours. The solid is filtered off, dissolved in hot 80% EtOH (200ml), filtered to remove traces of S, cooled for many hours at 0-5°, and the white crystals of **potassium *p*-toluenethiosulfonate** (48-55g, 42-49%) are collected and air dried. [Woodward et al. *Org Synth Coll Vol* 6 1016 1988, DOI: 10.15227/orgsyn.054.0033; Beilstein 11 IV 482.]

2-*p*-Toluidinylnaphthalene-6-sulfonic acid (2,6-TNS)[7724-15-4] $C_7H_5NO_3S$, **M 313.9, decomp on heating, pK_{Est} ~ 0**. Crystallise the acid twice from 2% aqueous KOH and dry it under high vacuum for 4 hours at room temperature. It also crystallises from H₂O. It is tested for purity by TLC on silica gel with isopropanol as solvent. The free acid is obtained by acidifying a saturated aqueous solution. The **sodium salt** [53313-85-2] $C_7H_4NNaO_3S$, **M 335.4**, crystallises from H₂O and decomposes on heating. [Beilstein 14 H 762.]

***p*-Tolyl disulfide** [103-19-5] $C_{14}H_{14}S_2$, **M 246.4, m 43-46°, 45°, 45-46°, 47.5°, 168°/45mm**. Purify it by chromatography on alumina using hexane as eluent, then crystallise it from MeOH, and/or distil it in a vacuum. [Kice & Bowers *J Am Chem Soc* 84 2384 1962, DOI: 10.1021/ja00871a022; Beilstein 6 H 245, 6 I 212, 6 II 400, 6 III 1432, 6 IV 3206.]

***p*-Tolylsulfonylmethyl isocyanide (tosylmethyl isocyanide, TOSMIC)** [36635-61-7] $C_9H_9NO_2S$, **M 195.2, m 109-113°, 112-115°, 114-115°(dec), 116-117°(dec)**. Use an efficient fume cupboard. Purify TOSMIC by dissolving (50g) in CH₂Cl₂ (150ml) and passing it through a column (40x3cm) containing neutral alumina (100g) in CH₂Cl₂ and eluting with CH₂Cl₂. A nearly colourless solution (700ml) is collected, evaporated *in vacuo* and the residue (42-47g) of **TOSMIC (m 113-114° dec)** is recrystallised once from MeOH (**m 116-117°dec**). [Hoogenboom et al. *Org Synth* 57 102 1977, DOI: 10.15227/orgsyn.057.0102; van Leusen et al. *Tetrahedron Lett* 13 2367 1972, DOI: 10.1016/S0040-4039(01)85304-1.] It also crystallises from EtOH (charcoal) [Saito & Itano, *JCS Perkin Trans 1* 1 1986, DOI: 10.1039/P19860000001].

If the reagent had deteriorated considerably it can be synthesised in two steps. **Firstly: *N*-(*p*-tolylsulfonylmethyl)formamide** is prepared by adding a mixture of aqueous 34-37% formalin (378g, 350ml, ~4.4moles), H₂O (750ml), excess of formamide (680g, 600ml, 15.5moles) and formic acid (244g, 200ml, 5.3moles) to sodium *p*-toluenesulfinate (267g, 1.5moles), and stirring at 90° to form a clear solution; and then heating at 90-95° is continued for 2 hours (prolonged heating diminishes the yield). The mixture is cooled (ice-salt bath) with continuous stirring, and set aside at -20° overnight after seeding. The white solid is collected, washed by stirring with ice-H₂O (3 x 250ml) and drained well. To remove occluded H₂O, the solid is dissolved in CH₂Cl₂, H₂O is removed (separating funnel), the organic layer is dried (MgSO₄), filtered, evaporated, and the residue is dried (over P₂O₅ at 70°) *in vacuo*, to give crude product (134-150g, 42-47%, **m 106-110°**) which is used in the following step. Pure ***N*-(*p*-tolylsulfonylmethyl)formamide, m 108-110°**, can be obtained by recrystallisation from *C₆H₆ or 95% aqueous EtOH.

In the **second step**, with exclusion of moisture, POCl₃ (84g, 50ml, 0.55mole) in 1,2-dimethoxyethane (60ml) is added dropwise to a stirred solution of the crude preceding **formamide** (107g, 0.50mole) in 1,2-dimethoxyethane (250ml), anhydrous Et₂O (100ml) and Et₃N (255g, 350ml, 2.52moles) cooled to -5° (ice-salt bath), at such a rate as to keep the temperature between -5° and 0° (requires ~1 hour). During the reaction, the formamide dissolves, white Et₃NH⁺ salts separate, and at the end of the reaction the suspension turns brown in

colour. If the mixture is not brownish in colour then more POCl_3 needs to be added. Finally the mixture is stirred for 30 minutes at 0° and ice- H_2O (1.5L) is added to give a clear brown solution before a fine brown crystalline solid separates. Stir the mixture for 30 minutes at 0° , collect the solid, wash it with cold H_2O (250ml), dissolve the solid in warm C_6H_6 (400ml at $40\text{--}60^\circ$), separate the H_2O , dry the organic layer (MgSO_4), filter, add charcoal (2g), heat at 60° for 5 minutes, filter, and add petroleum ether (1L, b $40\text{--}60^\circ$) with swirling. After 30 minutes, the solid is filtered off, dried *in vacuo* to give **crude TOSMIC** (74–82g, 76–84%) **m 111–114°** (dec), which can be used for most purposes. Analytically pure product can be obtained by a *third* purification through neutral Al_2O_3 as stated above. It has IR (Nujol) with ν_{max} at 2150 ($\text{N}=\text{C}$), 1320 and 1155 (SO_2) cm^{-1} , and the ^1H NMR (CDCl_3) has δ at 7.7 (q, 4H, C_6H_4), 4.6 (s, 2H, CH_2) and 2.5 (s, 3H, CH_3). [Hooogenboom et al. *Org Synth Coll Vol* **6** 987 1988, DOI: 10.15227/orgsyn.057.0102.] It is a versatile reagent, provides a formaldehyde anion or dianion equivalent, is used for the synthesis of various keto compounds and under reducing conditions amino and methylamino compounds have been prepared from the same intermediates, and it has been used for the preparation of several classes of heterocyclic compounds [van Leusen *Lect Heterocycl Chem* **5** S111 1980, see also a supplementary issue of Vol **17** of *J Heterocycl Chem* 1980]. **TOXIC, handle with care.**

Tribenzyl chlorosilane [18740-59-5] $\text{C}_{21}\text{H}_{21}\text{ClSi}$, **M 336.9**, **m 139–142°, 141–142°, b 300–360°/100mm**. It is recrystallised three times from petroleum ether (in slender colourless needles, **m 141°**). It is sparingly soluble in cold petroleum ether but is soluble in Et_2O . It does not fume in moist air but is decomposed by H_2O to give **tribenzyl silanol m 106°** (from petroleum ether). [Robinson & Kipping *J Chem Soc* **93** 439 1908, DOI: 10.1039/CT9089300439; Jenkins & Post *J Org Chem* **15** 556 1950, DOI: 10.1021/jo01149a018; *Beilstein* **16** H 906, **16 IV** 1498.]

Tribenzyl phosphine [7650-89-7] $\text{C}_{21}\text{H}_{21}\text{P}$, **M 304.4**, **m 96–101°, b 203–210°/0.5mm, $\text{pK}_{\text{Est}} \sim 8.8$** . Dissolve it in Et_2O , dry it over Na_2SO_4 , evaporate and distil it in an inert atmosphere. The distillate solidifies on cooling and is sublimed at $140^\circ/0.001\text{mm}$. This has **m 92–95°(evacuated capillary)**. When air is bubbled through an Et_2O solution, it is oxidised to **tribenzyl phosphine oxide**, [4538-55-0], **M 320.4**, **m 209–212°** (evacuated capillary) (it crystallises from Me_2CO). [Hinton & Mann *J Chem Soc* 2835 1959, DOI: 10.1039/JR9590002835; *Beilstein* **16** H 771, **16 IV** 961.]

Tri-*n*-butyl borate [688-74-4] $\text{C}_{12}\text{H}_{27}\text{BO}_3$, **M 230.2**, **m -70° , b $110^\circ/11\text{mm}$, $136^\circ/30\text{mm}$, 232.4° , d_4^{20} 0.857, n_D^{20} 1.4092**. The chief impurities are *n*-butyl alcohol and boric acid (from hydrolysis). It must be handled in a dry-box and can readily be purified by fractional distillation, under reduced pressure. [O'Brien *Aust J Chem* **10** 91 1957, DOI: 10.1071/CH9570091; Gerrard & Lappert *J Chem Soc* 2545, 2547 1951, DOI: 10.1039/JR9510002545; *Beilstein* **1** IV 1544.]

Tri-*n*-butyl chlorosilane [995-45-9] $\text{C}_{12}\text{H}_{27}\text{ClSi}$, **M 234.9**, **b $93\text{--}94^\circ/4.5\text{mm}$, $134\text{--}139^\circ/16\text{mm}$, $250\text{--}252^\circ/\text{atm}$, $142\text{--}144^\circ/29\text{mm}$, d_4^{20} 0.88, n_D^{20} 1.447**. Fractionally distil this silane, and store it in small aliquots in sealed ampoules. [Noller & Post *J Am Chem Soc* **74** 1361 1952, DOI: 10.1021/ja01125a600; Gilman et al *J Org Chem* **24** 219 1959, DOI: 10.1021/jo01084a018; *Beilstein* **4** IV 4072.]

Tri-*n*-butyl phosphate (butyl phosphate) [126-73-8] $\text{C}_{12}\text{H}_{27}\text{O}_4\text{P}$, **M 266.3**, **m -80° , b $47^\circ/0.45\text{mm}$, $98^\circ/0.1\text{mm}$, $121\text{--}124^\circ/3\text{mm}$, $136\text{--}137^\circ/5.5\text{mm}$, $166\text{--}167^\circ/17\text{mm}$, $177\text{--}178^\circ/27\text{mm}$, $289^\circ/760\text{mm}$ (some dec), d_4^{20} 0.980, n_D^{20} 1.44249**. The main contaminants in commercial samples are organic pyrophosphates, mono- and di- butyl phosphates and butanol. It is purified by washing successively with 0.2M HNO_3 (three times), 0.2M NaOH (three times) and water (three times), then fractionally distilled under vacuum. [Yoshida *J Inorg Nucl Chem* **24** 1257 1962, DOI: 10.1016/0022-1902(62)80202-4.] It has also been purified *via* its **uranyl nitrate addition compound**, obtained by saturating the crude phosphate with uranyl nitrate. This compound is crystallised three times from *n*-hexane by cooling to -40° , and then decomposed by washing with Na_2CO_3 and water. Hexane is removed by steam distillation; the water is then evaporated under reduced pressure, and the residue is distilled under reduced pressure. [Siddall & Dukes *J Am Chem Soc* **81** 790 1959, DOI: 10.1021/ja01513a007.] **Alternatively**, wash it with water, then with 1% NaOH or 5% Na_2CO_3 for several hours, then finally with water. Dry it under reduced pressure and fractionate it carefully under vacuum. It is a stable colourless oil, sparingly soluble in H_2O (1ml dissolves in 165ml of H_2O), but freely miscible in organic

solvents. [Kuivila & Masterton *J Am Chem Soc* **74** 4953 1952, DOI: 10.1021/ja01139a512; Cox & Westheimer *J Am Chem Soc* **80** 5441 1958, DOI: 10.1021/ja01553a031; ³¹P NMR: Van Wazer *J Am Chem Soc* **78** 5715 1956, DOI: 10.1021/ja01139a512; Fertig et al. *J Chem Soc* 1488 1957, DOI: 10.1039/JR9570001488; *Beilstein* **1** IV 1531.]

Tri-*n*-butyl phosphine (TBP) [998-40-3] $C_{12}H_{27}P$, **M 202.3**, **b 109-110°/10mm**, **115-116°/12mm**, **149.5°/50mm**, **240.4-242.2°/atm**, d_4^{20} **0.822**, n_D^{20} **1.4463**, pK_{Est} **~7.6**. Fractionally distil TBP under reduced pressure in an inert atmosphere (N₂) through an 8inch gauze-packed column (**b 110-111°/10mm**), and redistil it in a vacuum, then seal it in thin glass ampoules. It is easily oxidised by air to **tri-*n*-butylphosphine oxide**, [814-29-9], **m 64-49°**, **b 150°/1.5mm**, **293-296°/745mm**. It has a characteristic odour, it is soluble in EtOH, Et₂O, and *C₆H₆, but is insoluble in H₂O; and is less easily oxidised by air than the lower molecular weight phosphines. It forms **complexes**, e.g. with CS₂ (**1:1**) **m 65.5°** (from EtOH). [Davies & Jones *J Chem Soc* 33 1929, DOI: 10.1039/JR9290000033; Chernick & Skinner *J Chem Soc* 1401 1956, DOI: 10.1039/JR9560001401; *Beilstein* **4** IV 3436.] Unlike with Ph₃P, (MeO)₃P, (Me₂N)₃P or Me₃P, when (*n*-Bu)₃P is complexed with DEAD, best results are achieved in the **Mitsunobu lactonisation** of the final stage in the synthesis of the strained lactone **(-)-Echinospurin** (XK-213) [Smith et al. *J Am Chem Soc* **114** 2567 1992, DOI: 10.1021/ja00033a033; Rehnberg & Magnusson *J Org Chem* **55** 5467 1990, DOI: 10.1021/jo00307a017].

Tri-*tert*-butylphosphine [13716-12-6] $C_{12}H_{27}P$, **M 202.3**, has **m ~30-35°**, **62.3°**, **b 103-103°/13mm**, d_4^{20} **0.834**, pK^{25} **11.4**. It is prepared from *tert*-butylmagnesium chloride and PCl₃ in Et₂O and finally purified by vacuum fractionation and distillation. [Ger Pat, IG Farbenind DRP 730 638 1939, DRP *Org Chem* **3** 1144.] A 1.0M solution in toluene is commercially available (d^{25} **0.861 g/l**). [*Beilstein* **4** III 1771.]

It is used with Pd₂(dba)₃ and Bu₃SnF in the Pd-catalysed regiocontrolled α -arylation of trimethylsilyl enol ethers with aryl halides [Iwama & Rawal *Org Lett* **8** 5725 2006, DOI: 10.1021/ol062093g], is a useful ligand for Pd-catalysed **Suzuki-type coupling** of arylboronic acids with phenyliodonium ylides of hydroxyquinones [Kazantzi et al. *Synlett* 2597 2006, DOI: 10.1055/s-2006-950447], and is a good ligand for Ni(acac)₂ in catalysing the **cross-coupling** of aryl- and heteroaryl- halides with aryl Grignard reagents [Böhm et al. *Angew Chem Int Ed* **39** 1602 2000, DOI: 10.1002/(SICI)1521-3773(20000502)39:9<1602>.

Tri-*tert*-butylphosphine oxide [6866-70-2] $C_{12}H_{27}PO$, **M 218.3**, has **m 64-69°**, **70°**, **77°**. The oxide has been prepared by treating *t*-Bu₃P (1.7g, 85mmol) in Et₂O (20ml) containing a catalytic amount of KI (0.1g) with 30% of H₂O₂ until the colour of iodine disappears. The mixture is diluted with H₂O, basified with aqueous NaOH, the ethereal layer is evaporated, and the **phosphine oxide** (1.5g, 81%, **m 77°**) is obtained by sublimation at 50°/0.1mm. It is a crystalline **hygroscopic** solid that is soluble in most organic solvents, and is stable to 250°. Its IR (Nujol) has ν_{max} at 1160 (P=O, s) and 815 (P-C, br s) cm⁻¹; the ¹H NMR (C₆D₆, TMS) has δ_H at 0.59 (d, ³J_{P-H} = 12.1Hz, C-Me₃); the ¹³C NMR (C₆D₆, TMS) has δ_C at 29.08 (s, CH₃), 38.95 (d, ¹J_{P-C} = 51.27, C), and its ³¹P NMR (C₆D₆, external H₃PO₄) has δ at 60.86. [Schmidbauer & Blaschke *Z Naturforsch* **33B** 1556 1978, Rankin et al. *JCS Dalton Trans* 827 1985, DOI: 10.1039/DT9850000827.]

Tri-*n*-butyl phosphite [102-85-2] $C_{12}H_{27}PO_3$, **M 250.3**, **b 114-115°/5mm**, **122°/12mm**, **130°/17mm**, **137°/26mm**, d_4^{20} **0.926**, n_D^{20} **1.4924**. Fractionate the phosphite through an efficient column. It is stable in air but is slowly hydrolysed by H₂O. [Gerrard *J Chem Soc* 1464 1940, DOI: 10.1039/JR9400001464; Fertig et al. *J Chem Soc* 1488 1957, DOI: 10.1039/JR9570001488; for complexing with peroxymolybdc acids see Fields *J Am Chem Soc* **80** 2358 1958, DOI: 10.1039/JR9570001488; DOI: 10.1021/ja01543a004; for preparations and bond refractions see Gillis et al. *J Am Chem Soc* **80** 2999 1958, DOI: 10.1021/ja01545a025; *Beilstein* **1** IV 1527.]

B-Trichloroborazine (B-trichloroborazole) [933-18-6, 26445-82-9] $H_3B_3Cl_3N_3$, **M 183.8**, **m 83.9-84.5°(sealed evacuated cappillary)**, **87°**, **b 88-92°/21mm**, d_4^{25} **1.58**. Purify the borazine by distillation from mineral oil. It sublimes at 70°/1mm. [Brown & Laubengayer *J Am Chem Soc* **77** 3699 1955, DOI: 10.1021/ja01619a007; Emeléus & Videla *J Chem Soc* 1306 1959, DOI: 10.1039/JR9590001306.] It is extremely sensitive to moisture and reacts with H₂O exothermically to give boric acid and NH₄Cl. Store it in sealed tubes. It is soluble in *C₆H₆, cyclohexane, CS₂, CHCl₃, CCl₄, and C₆H₅Cl without decomposition; reacts vigorously MeOH or EtOH liberating HCl, but is insoluble in pyridine and PhNO₂. [*Beilstein* **4** III 174, **1** IV 305.]

Trichloromethyl trimethylsilane (trimethylsilyl trichloromethane) [5936-98-1] $\text{C}_4\text{H}_9\text{Cl}_3\text{Si}$, M 191.6, m 130-132°, b 146-156°/749mm, 159.1 \pm 35°/760mm. This silane distils at atmospheric pressure without decomposition and readily sublimes at 70°/10mm. It has one peak in the ^1H NMR spectrum (CD_2Cl_2) with δ at 0.38. [Speier *J Am Chem Soc* **73** 824 1951, DOI: 10.1021/ja01146a099; Hergott & Simchen *Synthesis* 626 1980, DOI: 10.1055/s-1980-29144; *Beilstein* **4** IV 3892.]

Trichlorosilane (SiHCl_3 , silicochloroform) [10025-78-2] HCl_3Si , M 135.4, m -126°, b 31.8°/atm, 32-34°/atm, 36.5°/atm, d_4^{25} 1.342, n_D^{20} 1.4020. This volatile trichlorosilane can be obtained from silane and HCl in the presence of AlCl_3 , and is purified by fractional distillation at atmospheric pressure in the strict absence of moisture. However, SiHCl_3 is best prepared by placing finely powdered silicon (purified by boiling with HCl and dilute HF) ground with *ca* 10% of CuCl_2 in a pyrex glass tube (fitted with an adapter and condenser which should extend into the middle of the distilling flask) and heating (carefully at first) in a furnace while a slow stream of absolutely dry HCl gas (from a gas cylinder, or generated from NaCl and concentrated H_2SO_4) is passed over the mixture with a furnace temperature of 300°. The receiver is cooled by an acetone/Dry-ice mixture, and the crude product is distilled from the receiver. HCl comes off first then SiHCl_3 distils over at 36.5° in 50% yield. With careful fractionation, the forerun SiH_2Cl_2 , can be recovered. At 300°, AlCl_3 disproportionates SiHCl_3 to SiH_2Cl_2 and SiCl_4 . [Note that by using an 4:1 mixture of H_2/HCl the yield of SiH_2Cl_2 , [4109-96-0] M 101.0, m -122°, b 8.5°/atm, d^{122° 1.22, can be greatly improved; and SiH_3Cl , which has m -118°, b -30.5°/atm, d^{113° 1.15. SiHCl_3 is a water clear, very volatile liquid which fumes in moist air as it is hydrolysed by H_2O , but is remarkably inert towards metals even the alkali metals such as Na. It is a good **reducing agent**, is soluble in most organic solvents, e.g. C_6H_6 , CHCl_3 , and CS_2 , but not in protic solvents in which it may decompose. It is **TOXIC, should not be inhaled**, and should be handled under an efficient fume hood. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 691-692 1963, Fritz *Z Anorg Allg Chem* **280** 134 1955, DOI: 10.1002/zaac.19552800111.]

Triethoxysilane [998-30-1] $\text{C}_6\text{H}_{16}\text{O}_3\text{Si}$, M 164.3, m -170°, b 131.2-131.8°/atm, 131.5°/760mm, 134-135°/atm, d_4^{20} 0.98753, n_D^{20} 1.4377. Fractionate it using a column packed with glass helices of *ca* 15 theoretical plates in an inert atmosphere. Store it in aliquots in sealed ampoules because it is sensitive to moisture. [Spauschus et al. *J Am Chem Soc* **72** 1377 1950, DOI: 10.1021/ja01159a082; MacKenzie et al. *J Am Chem Soc* **72** 2032 1950, DOI: 10.1021/ja01161a042; Havill et al. *J Org Chem* **13** 280 1948, DOI: 10.1021/jo01160a017.] A useful reagent for attaching groups onto silica surfaces [Corriu et al. 'Triethoxysilane' in *e-EROS Encyclopedia of Reagents for Organic Synthesis* 2001 John Wiley & Sons, NY, DOI: 10.1002/047084289X.rt215.pub3; *Beilstein* **1** IV 1359.]

Triethylborane [97-94-9] $\text{C}_6\text{H}_{15}\text{B}$, M 98.0, m -92.5°, b 94-97°, 94-95°, n_D^{20} 1.378, d_4^{20} 0.678. This borane distils at 56-57°/220mm. It can also be purified *via* its **ammonia addition complex** which is distilled in a high vacuum, decomposed with dry HCl, and the Et_3B is distilled out. It is commercially available as a 15% solution in hexane or as 1M solution in hexane. [Brown *J Am Chem Soc* **67** 374 1945, DOI: 10.1021/ja01219a007; Bamford et al. *J Chem Soc* 468 1946, DOI: 10.1039/JR9460000468; Lin *J Organomet Chem* **317** 277 1986, DOI: 10.1016/0022-328X(86)80538-1; *Beilstein* **4** III 1957, **4** IV 4359.]

Triethyl borate (boric acid triethyl ester, boron triethoxide) [150-46-9] $\text{C}_6\text{H}_{15}\text{BO}_3$, M 146.0, m -84.5°, -84.8°, b 44.5°/45mm, 118°/760mm, n_D^{20} 1.378, d_4^{20} 0.864. Dry the ester over sodium, then distil it. Also fractionate it through a gauze packed column. [Charnley et al. *J Chem Soc* 2288 1952, DOI: 10.1039/JR9520002288; for tributyl borate see Johnson & Tompkins *Org Synth Coll Vol* **2** 106 1943, DOI: 10.15227/orgsyn.013.0016; *Beilstein* **1** III 1339, **1** IV 1365.]

Triethyl phosphate [78-40-0] $\text{C}_6\text{H}_{15}\text{O}_4\text{P}$, M 182.2, m -56.4°, b 40-42°/0.25-0.3mm, 98-98.5°/8-10mm, 90°/10mm, 130°/55mm, 204°/680mm, 215-216°/760mm, d_4^{25} 1.608, n_D^{20} 1.4053. Dry the phosphate by refluxing it with solid BaO and then fractionally distil it under reduced pressure. It is kept over Na and distilled. Store it in the receiver protected from light and moisture. *Alternatively*, it is dried over Na_2SO_4 and distilled under reduced pressure. The middle fraction is stirred for several weeks over anhydrous Na_2SO_4 and again fractionated under reduced pressure until the specific conductance reaches a constant low value of κ^{25} 1.19×10^8 , κ^{40} 1.68×10^8 , and κ^{55} $2.89 \times 10^8 \text{ ohm}^{-1} \text{ cm}^{-1}$. It has also been fractionated carefully under reduced pressure.

ure through a glass helices-packed column. It is soluble in EtOH, Et₂O and H₂O (dec). [Estok & Wendlandt *J Am Chem Soc* **77** 4767 1955, DOI: 10.1021/ja01623a021; Hoffmann et al. *J Am Chem Soc* **78** 6413 1956, DOI: 10.1021/ja01605a033; (P NMR) Muller et al. *J Am Chem Soc* **78** 3557 1956, DOI: 10.1021/ja01596a002; French et al. *J Chem Soc* 3582 1959, DOI: 10.1039/JR9590003582; IR: Bellamy & Beecher *J Chem Soc* 475 1952, DOI: 10.1039/JR9520000475; McIvor et al. *Can J Chem* **36** 820 1958, DOI: 10.1139/v58-121; Kosolapoff *Organophosphorus Compounds*, Wiley p 258 1950, *Beilstein* **1** IV 1339.]

Triethylphosphine [554-70-1] C₆H₁₅P, M 118.2, b 100°/7mm, 127-128°/744mm, d₄¹⁵ 0.812, n_D¹⁸ 1.457, pK²⁵ 8.69 (also available as a 1.0M solution in THF). All operations should be carried out in an efficient fume cupboard because it is flammable, toxic and has a foul odour. This liquid has the odour of hyacinth flowers. Purify the phosphine by fractional distillation at atmospheric pressure in a stream of dry N₂, as it is oxidised by air to the oxide. In 300% excess of CS₂ it forms *Et*₃PCS₂ (m 118-120° crystallising from MeOH) that decomposes in CCl₄ to give *Et*₃PS as a white solid m 94° when recrystallised from EtOH. [Screttas & Isbell *J Org Chem* **27** 2573 1962, DOI: 10.1021/jo01054a064; Henderson & Streuli *J Am Chem Soc* **82** 5791 1960, DOI: 10.1021/ja01507a008; pK: Henderson & Streuli *J Am Chem Soc* **82** 5791 1960, DOI: 10.1021/ja01507a008; see also trimethylphosphine.] Store it in sealed vials under N₂.

Alternatively, dissolve it in Et₂O and shake it with a solution of AgI and KI to form the insoluble complex. Filter off the complex, dry it over P₂O₅ and the Et₃P is regenerated by heating the silver iodide complex in a tube attached to a vacuum system. It has the odour of hyacinths (see above). [Hewitt & Holliday *J Chem Soc* 530 1953, DOI: 10.1039/JR9530000530; Screttas & Isbell *J Org Chem* **27** 2573 1962, DOI: 10.1021/jo01054a064; Kosolapoff *Organophosphorus Compounds*, Wiley p 31 1950, *Beilstein* **4** IV 3431.] **Triethylphosphine oxide** [597-50-2] C₆H₁₅OP, M 134.2, has m 48-50°, 49°, 48-52°, b 84-85°/3mm, and is readily obtained by oxidation of triethylphosphine.

Triethyl phosphite [122-52-1] C₆H₁₅O₃P, M 166.2, b 48-49°/11mm, b 52°/12mm, 57.5°/19mm, 155°/atm, 157.9°/757mm, d₄²⁰ 0.9687, n_D²⁰ 1.4135. Treat the ester with Na (to remove water and any dialkyl phosphonate), then decant and distil it under reduced pressure, with protection against moisture; or distil it in a vacuum through an efficient Vigreux column or a column packed with Penn State 0.16 x 0.16inch protruded nickel packing and a variable volume take-off head. [Ford-Moore & Perry *Org Synth Coll Vol* **4** 955 1963, DOI: 10.15227/orgsyn.031.0111; Kosolapoff *Organophosphorus Compounds*, Wiley p 203 1950, *Beilstein* **1** IV 1333.]

Triethyl phosphonoacetate (triethyl carboxymethyl phosphonate) [867-13-0] C₈H₁₇O₅P, M 224.2, b 83-84°/0.5mm, 103°/1.2mm, 143-144°/11mm, 260-262°/atm, d₄²⁰ 1.1215, n_D²⁰ 1.4310. Purify the phosphonoacetate by fractional distillation, preferably *in vacuo*. The ³¹P NMR has a P resonance at 19.5 relative to orthophosphate. [Kosolapoff *J Am Chem Soc* **68** 1103 1946, DOI: 10.1021/ja01210a058; Kosolapoff & Powell *J Am Chem Soc* **72** 4198 1950, DOI: 10.1021/ja01165a104; Speziale & Freeman *J Org Chem* **23** 1883 1958, DOI: 10.1021/jo01106a015; *Beilstein* **4** IV 3613.] It is deprotonated with MeONa to form a **phosphonate ylide** which undergoes a **HWE (Homer-Wadsworth-Emmons) reaction** with a carbonyl groups to generate an *E*-alkene with very high regioselectivity [Wadsworth & Emmons *J Am Chem Soc* **83** 1733 1961, DOI: 10.1021/ja01468a042; Wadsworth & Emmons *Org Synth Coll Vol* **5** 547 1973; DOI: 10.15227/orgsyn.045.0044].

Triethyl phosphonoformate (ethyl diethoxyphosphinylformate) [1474-78-8] C₇H₁₅O₅P, M 210.2, has b 70-72°/0.1mm, 122.5-123°/8mm, 130-131°/10mm, 138.2°/12.5mm, d₄²⁰ 1.22, n_D²⁰ 1.423. Dissolve it in Et₂O, shake this with H₂O (to remove any trace of NaCl impurity), dry (Na₂SO₄), evaporate and distil it using an efficient fractionating column. [Nylén *Chem Ber* **57** 1023 1924, DOI: 10.1002/cber.19240570625; Reetz et al. *J Am Chem Soc* **77** 3813 1955, DOI: 10.1021/ja01619a040; Monson *Advanced Organic Synthesis* Academic Press p 89 1972, *Beilstein* **3** II 103.]

Triethyl 2-phosphonopropionate (ethyl 2-(diethoxyphosphinyl)propionate) [3699-66-9] C₉H₁₉O₅P, M 238.2, has b 76-77°/0.2mm, 143-144°/12torr, 137-138.5°/17mm, d₄²⁰ 1.096, n_D²⁰ 1.432. Purify the ester by fractional distillation with high reflux ratio, preferably using a spinning band column. [Kosolapoff & Powell *J Am Chem Soc* **72** 4198 1950, DOI: 10.1021/ja01165a104; Kresze et al. *Justus Liebigs Ann Chem* **756** 112 1972, DOI: 10.1002/jlac.19727560111; *Beilstein* **4** IV 3617.] See HWE reaction above.

Triethylsilane [617-86-7] $C_6H_{15}SiH$, M 116.3, b 105-107°/atm, 107-108°/atm, d_4^{20} 0.734, n_D^{20} 1.414. Reflux triethylsilane over molecular sieves, then distil it. It is passed through neutral alumina before use [Randolph & Wrighton *J Am Chem Soc* **108** 3366 1986, DOI: 10.1021/ja00272a035]. Useful reducing agent in stereochemically controlled reduction of 2-chromanols [Li et al. *Org Lett* **8** 4711 2006, DOI: 10.1021/ja00272a035]. [Fieser & Fieser's *Reagents for Org Synth* **1** 1218 1967, *Beilstein* **4** IV 3895.]

Triethylsilyloxy-1,4-pentadiene (1,4-pentadien-3-yloxy-trimethylsilane) [62418-65-9] $C_{11}H_{22}OSi$, M 198.4, b 72-74°/12mm, 140°/atm, d_4^{20} 0.842, n_D^{20} 1.439. Dissolve the diene in pentane, wash this with H_2O , dry (Na_2SO_4), evaporate, and distil it under vacuum. R_F values on Kieselgel 60 are 0.15 (pentane) and 0.60 ($*C_6H_6$). [IR, NMR, MS: Oppolzer et al. *Helv Chim Acta* **64** 2002 1981, DOI: 10.1002/hlca.19810640705.]

Tri-*n*-hexylborane [1188-92-7] $C_{18}H_{39}B$, M 266.3, b 127°/1.5mm, b 185-188°/30mm. Treat the borane with hex-1-ene and 10% anhydrous Et_2O for 6 hours at gentle reflux under N_2 , then distil it in a vacuum through an 18inch glass helices-packed column under N_2 taking the fraction b 130°/2.1mm to 137°/1.5mm. The distillate may still contain some *di-n-hexylborane* [Brown & Subba Rao *J Am Chem Soc* **81** 6423 1959, DOI: 10.1021/ja01533a023; Mirviss *J Am Chem Soc* **83** 3051 1961, DOI: 10.1021/ja01475a020]. [*Beilstein* **4** IV 4362.]

Triisomyl phosphate (TAP) [919-62-0] $C_{15}H_{33}O_4P$, M 308.4, b 143°/3mm. Purify the ester by repeated crystallisation of its addition compound with *uranyl nitrate* from hexane. Decompose the complex, and distil the ester at high vacuum. [Siddall *J Am Chem Soc* **81** 4176 1959, DOI: 10.1021/ja01525a015.] [see *tributyl phosphate* and Cherbuliez in *Organo Phosphorus Compounds* (Kosolapoff & Maier eds) Wiley Vol **6** pp 211-577 1973.] It is a useful liquid extractant for reactor fuel reprocessing [Syed Hadi & Shulka *J Radioanal Nucl Chem* **258** 563 2003, DOI: 10.1023/B:JRNC.0000011753.65662.64].

Triisobutyl phosphate [126-71-6] $C_{12}H_{27}O_4P$, M 266.3, has b 119-129°/8-12mm, 192°/760mm, d_4^{20} 0.962, n_D^{20} 1.421. Purify the phosphate by repeated crystallisation of its addition compound with *uranyl nitrate* from hexane. (see *tributyl phosphate*.) [Siddall *J Am Chem Soc* **81** 4176 1959, DOI: 10.1021/ja01525a015; see Cherbuliez in *Organo Phosphorus Compounds* (Kosolapoff & Maier eds) Wiley Vol **6** pp 211-577 1973.] This phosphate is a strong solvent for liquefying concrete, for textile auxiliaries and paper coating compounds. See preceding liquid extractant.

Triisooctyl thiophosphate [30108-39-5] $C_{24}H_{51}O_3PS$, M 450.7. Purify the ester by passing its solution in CCl_4 through a column of activated alumina. [See Ailman & Magean in *Organo Phosphorus Compound* (Kosolapoff & Maier eds) Wiley Vol **7** pp 487-465 1973, and tri-*p*-tolyl phosphate below.] Used to extract Ag.

Tri-isopropyl borate [(iso-PrO)₃B, boric acid tri-isopropyl ester, Boron iso-propoxide] [5419-55-6] $C_9H_{21}BO_3$, M 188.1, m -59°, b 75°/76mm, 90°/120mm, 139-141°/760mm, d_4^{25} 0.815, n_D^{20} 1.376. The borate ester is prepared in 85% yield from *iso*-propanol (250ml) and $NaBH_4$ (6.84g) followed by dropwise addition of AcOH (0.18g) over a period of 11 minutes, and then refluxing for 4 hours (fume cupboard and a 'Dry-Ice' condenser as 16.15L of H_2 are released, CARE due to its flammability). Then fractionate the ester through a Widmer column. [Brown et al. *J Am Chem Soc* **78** 3613 1956, DOI: 10.1021/ja01596a015.] The ester is a good reagent for borylation, e.g. *ortho*-borylating 1-substituted naphthalenes, by reacting with 1-halo- or cyano-naphthalenes (after treatment with LiTMP) to form 2-boryl esters which are later used in Pd-catalysed cross-coupling reactions [Lysén et al. *Synthesis* 3478 2006, DOI: 10.1055/s-2006-950239]. [*Beilstein* **1** H 363, **1** II 382, **1** III 1468, **1** IV 1488.]

Triisopropyl phosphite [116-17-6] $C_9H_{21}O_3P$, M 208.2, b 58-59°/7mm, 63-64°/11mm, 197-199°/760mm, d_4^{25} 0.844, n_D^{25} 1.4082. Distil the phosphite from sodium, under vacuum, through a column packed with glass helices. (This removes any dialkyl phosphonate.) [Ford-Moore & Williams *J Chem Soc* 1465 1947, DOI: 10.1039/JR9470001465; Arbuzoff *Chem Ber* **38** 1171 1905, DOI: 10.1002/cber.190503801212; see Verkade & Coskren in *Organo Phosphorus Compound* (Kosolapoff & Maier eds) Wiley Vol **2** pp 1-187 1972, *Beilstein* **1** IV 1476.]

Trimesitylphosphine [23897-15-6] $C_{27}H_{33}P$, M 388.5, m 185-188°, 205-206°, $pK_{Est} \sim 8.0$. It recrystallises from EtOH [Boere et al. *J Am Chem Soc* **109** 7781 1987, DOI: 10.1021/ja00259a029]. The *P*-methyl iodide has m 269° (yellow powder from EtOH or H₂O). [Beilstein **16** H 774.]

Trimethallyl phosphate [14019-81-9] $C_{12}H_{21}O_4P$, M 260.3, b 134.5-140°/5mm, n_D^{25} 1.4454. Purify it as for triisooamyl phosphate. [Cherbuliez in *Organo Phosphorus Compounds* (Kosolapoff & Maier eds) Wiley Vol **6** pp 211-577 1973.]

Trimethoxysilane [2487-90-3] $C_3H_{10}O_3Si$, M 122.2, m -114.8°, 81.1°/760mm, 84°/atm, d_4^{20} 0.957, n_D^{20} 1.359. Likely impurities are Si(OMe)₄ and H₂Si(OMe)₂. Efficient fractionation is essential for removing these impurities [IR: Sternbach & MacDiarmid *J Am Chem Soc* **81** 5109 1959, DOI: 10.1021/ja01528a023; Helferich & Hausen *Chem Ber* **57** 795 1924, DOI: 10.1002/cber.19240570516; Beilstein **1** IV 1266.]

Trimethyl borate (methylborate, trimethoxyboron) [121-43-7] $C_3H_9BO_3$, M 103.9, m -34°, b 67-68°/742mm, 68-69°/atm, d_4^{20} 0.928, n_D^{20} 1.3610. Carefully fractionate the borate through a gauze-packed column. Re-distil and collect it in weighed glass vials and seal them. Keep it away from moisture. It undergoes alkyl exchange with alcohols and forms azeotropes, e.g. with MeOH: the azeotrope consists of 70% (MeO)₃B and 30% MeOH with b 52-54°/760mm, d 0.87. [Charnley et al. *J Chem Soc* 2288 1952, DOI: 10.1039/JR9520002288; Gerrard & Lappert *Chemistry and Industry* 53 1952, Schlesinger et al. *J Am Chem Soc* **75** 213 1953, DOI: 10.1021/ja01097a056.] It was also dried with Na and then distilled. [Beilstein **1** IV 1269.]

Trimethyl boroxine (Me₃B₃O₃) [823-96-1] $C_3H_9B_3O_3$, M 125.5, m -38°, b 80°/742mm, 79.3°/755mm, d_4^{20} 0.902, n_D^{20} 1.362. Possible impurity is methylboronic acid. If present, then add a few drops of concentrated H₂SO₄ and distil it immediately, then fractionate it through an efficient column. [McCusker et al. *J Am Chem Soc* **79** 5179 1957, DOI: 10.1021/ja01576a026; IR: Goubeau & Keller *Z Anorg Allgem Chem* **272** 303 1953, DOI: 10.1002/zaac.19532720510; Beilstein **4** IV 4378.] Useful in the preparation of CBS (Corey-Bakshi-Shibata) catalysts for asymmetric reductions [Gilmore & Jones *Tetrahedron Asym* **14** 2115 2003, DOI: 10.1016/S0957-4166(03)00401-4].

1,3-S,S'-Trimethylene di(*p*-toluenethiosulfonate) [1,3-di(*p*-tosylthio)propane, 1,3-trimethylene di(thio-tosylate)] [3866-79-3] $C_{17}H_{20}O_4S_4$, M 416.6, m 64-67°. This reagent is prepared from potassium thiosulfonate (40g, 0.18 mole, see above [28519-50-8]) and 1,3-dibromopropane (20g, 0.1 mole) which are added to a solution of 95% EtOH (150ml) containing KI (10-20mg, to activate the dibromide), and the stirred mixture is boiled under reflux in the dark under N₂ for 8 hours. After cooling to ~25°, the mixture is diluted with an equal volume of cold H₂O and shaken. After the mixture settles, the supernatant is removed by decantation, the pale yellow residual oil is washed with cold H₂O (3 x 200ml), once with cold 95% EtOH (100ml) and once with cold absolute EtOH (100ml) also by decantation. The oil is then dissolved in Me₂CO (10ml), diluted with hot absolute EtOH (80ml), and stirred under N₂ at 0°. If some oil separates, it should be re-dissolved by adding enough Me₂CO (say ~5ml). The solution is seeded with crystals (a small portion of the oil stored at -30° for a few days crystallises and issued a seed*), stirred under N₂ at 0° for 1 hour then for several hours at -30°. The microcrystals are collected (20.2g) and have m 63.5-65.0°. This can be used for reactions; but if purer material is required then it should be recrystallised three times from EtOH (180ml, 9 parts) to provide white needles (17.2g, 41%) with m 66-67°. The *thioester* always separates as an oil when solutions are cooled to room temperature.

*If difficulties are experienced to crystallise the ester then seed crystals could be obtained by chromatography through a column of Woelm neutral alumina (Activity Grade I) and eluting with *C₆H₆. The centre fractions giving solid with m 67° are combined and recrystallised from EtOH (9parts) to give the pure *trimethylene dithiotosylate* with m 67.5°. On a Durapak-Carbowax 400/Poracil C column (Waters Associates, 3ft x 0.125in) with CHCl₃ as eluent it exhibits a single peak. It has IR (CHCl₃) with ν_{max} at 810 (m), 1015 (w), 1075 (s), 1140 (s), 1180 (w), 1300 (m), 1325 (s), 1410 (w), 1440 (w), 1490 (w), 1590 (w), 2930 (w), and 3030 (w) cm⁻¹; and the ¹H NMR (CDCl₃, TMS) has δ at 1.98 (quintet, J = 7Hz, 2H, CH₂CH₂CH₂), 2.43 (s, 6H, 2CH₃), 2.97 (t, J = 7Hz, 4H, CH₂CH₂CH₂), 7.30 (d, J = 9Hz, 4H, Arom) and 7.75 (d, J = 9Hz, 4H, Arom). [Woodward et al. *Org Synth Coll Vol* **6** 1016 1988, DOI: 10.1522/orgsyn.054.0033.]

This reagent is useful for the synthesis of 1,3-dithianes from carbonyl compounds and activated methylene

groups. Thus a ketone (0.163mmol), reagent (0.218mmol) and dry NaOH (0.907mmol) in EtOH (6ml) are refluxed for 6 hours, dilute with CH_2Cl_2 , wash with aqueous NaHCO_3 , evaporate the extract and purify the residue through a silica gel column by eluting with 1:2 EtOAc/hexane containing 2% of *N*-methylpyrrolidine to give the desired **1,3-dithiane** [McMurry et al. *J Org Chem* **49** 3803 1984, DOI: 10.1021/jo00194a026; see also Saksena & Ganguly *Tetrahedron Lett* **22** 5227 1981, DOI: 10.1016/S0040-4039(01)92466-9]. It also forms 1.3-dithianes, e.g. **2,2-(trimethylenedithio)cyclohexanone**, from the respective enamine, i.e. 1-pyrrolidinecyclohex-1-ene [Woodward et al. *Org Synth* **54** 39 1974, DOI: 10.15227/orgsyn.054.0039].

Trimethylphenylsilane (phenyltrimethylsilane) [768-32-1] $\text{C}_9\text{H}_{14}\text{Si}$, **M 150.3**, **b 67.3°/20mm**, **98-99°/80mm**, **170.6°/738mm**, d_4^{25} **0.8646**, n_D^{20} **1.491**. Fractionally distil the silane at atmospheric or reduced pressure (Podbielniak column and estimate it by GC with a column packed with Silicone Fluid No 710 on Chromosorb P support. [Gilman et al. *J Org Chem* **18** 1743 1953, DOI: 10.1021/jo50018a019; Maienthal et al. *J Am Chem Soc* **76** 6392 1954, DOI: 10.1021/ja01653a043; House & Respass *J Organomet Chem* **4** 95 1965, DOI: 10.1016/S0022-328X(00)82372-4; Roberts et al. *J Am Chem Soc* **71** 2923 1949, DOI: 10.1021/ja01176a10; Freiser et al. *J Am Chem Soc* **75** 2821 1953, DOI: 10.1021/ja01108a009; Beilstein **16** I 525, **16** II 605, **16** III 1198, **16** IV 1361.]

Trimethyl phosphate [512-56-1] $\text{C}_3\text{H}_9\text{O}_4\text{P}$, **M 140.1**, **m -46°**, **b 77°/12mm**, **94°/22mm**, **110°/60mm**, **197.2°/atm**, **196-197°/760mm**, d_4^{20} **1.0213**, n_D^{20} **1.3961**. Purify the phosphate by fractionation through an efficient column at high reflux ratio. It is quite soluble in H_2O ; the solubility is 1:1 at 25°. [Becher *J Am Chem Soc* **74** 2923 1952, DOI: 10.1021/ja01131a502; IR: Bergmann et al. *J Chem Soc* 847 1952, DOI: 10.1039/JR9520000847; McIvor et al. *Can J Chem* **36** 820 1958, DOI: 10.1139/v58-121; Kosolapoff *Organophosphorus Compounds*, Wiley p 258 1950, and Cherbuliez in *Organo Phosphorus Compounds* (Kosolapoff & Maier eds) Wiley Vol **6** pp 211-577 1973, Beilstein **1** IV 1259.]

Trimethylphosphine [594-09-2] $\text{C}_3\text{H}_9\text{P}$, **M 76.1**, **m -86°**, **b 38-39°/atm**, **38-40°/760mm**, pK^{25} **8.65**, (also available as a 1.0M solution in THF or toluene). All operations should be carried out in an efficient fume cupboard because it is flammable, toxic and has a foul odor. Distil trimethylphosphine at atmospheric pressure in a stream of dry N_2 (apparatus should be held together with springs to avoid loss of gas from increased pressure in the system) and the distillate run into a solution of AgI in aqueous KI whereby the **silver complex** $[\text{Me}_3\text{P}(\text{AgI})_4]$ separates steadily. Filter off the complex, wash it with saturated aqueous KI solution, then H_2O and dry it in a vacuum desiccator over P_2O_5 . The dry complex is heated in a flask (in a stream of dry N_2) in an oil bath at 140°, when pure Me_3P distils off (bath temperature can be raised up to 260°). The vapour pressure of Me_3P at 20° is 466mm and the **b** is **37.8°** [Thomas et al. *Inorg Synth* **9** 59 1967, DOI: 10.1002/9780470132401.ch18]. Alternatively, freshly distilled Me_3P (6g) is shaken with a solution of AgI (13.2g, 1.1mol) in saturated aqueous KI solution (50ml) for 2 hours. A white solid, not wetted with H_2O , separates rapidly. It is collected, washed with the KI solution, H_2O , and dried [Mann et al. *J Chem Soc* 1828 1937, DOI: 10.1039/JR9370001828]. The **silver complex** is stable if kept dry in the dark, in which state it can be kept indefinitely. Me_3P can be generated from the complex when required. Store it under N_2 in a sealed container. It has been distilled in a vacuum line at -78° *in vacuo* and condensed at -96° [IR and NMR: Crosbie & Sheldrick *J Inorg Nucl Chem* **31** 3684 1969, DOI: 10.1016/0022-1902(69)80363-5]. The pK^{22} by NMR was 8.80 [Silver & Lutz *J Am Chem Soc* **83** 786 1961, DOI: 10.1021/ja01465a008; pK^{25} 8.65: Henderson & Streuli *J Am Chem Soc* **82** 5791 1960, DOI: 10.1021/ja01507a008]. This **electron-rich phosphine ligand** when complexed with various metals makes efficient catalysts for numerous cross-coupling and other reactions. The $[\text{Me}_2\text{P}(\text{AgI})_4]$ complex [12389-34-3] is a flammable solid which has **m 140-142°**. It is decomposed by heating gently in one arm of an inverted U tube. The other arm is kept in a freezing mixture. The complex dissociates, and pure Me_3P collects in the cold arm and is used at once. It should not be allowed to come in contact with air [for AsMe_3 see Mann & Wells *J Chem Soc* 702 1938, DOI: 10.1039/JR9380000702]. The **CS_2 complex** has **m 119°** (crystallising from 95% EtOH) and decomposes in CCl_4 to give Me_3PS **m 154°** (from EtOH) [Soretas & Isbell *J Org Chem* **27** 2573 1962, DOI: 10.1021/jo01054a064].

Trimethylphosphine HCl is unstable and volatilises at 75°/0.4mm (120°/14mm). [Brown *J Am Chem Soc* **67** 503 1945, DOI: 10.1021/ja01219a514; IR: Wagstaffe & Thompson *Trans Faraday Soc* **40** 41 1944, DOI: 10.1039/TF9444000041; Kosolapoff *Organo-phosphorus Compounds*, Wiley p 31 1950, Beilstein **4** IV 3429.]

Trimethyl phosphite [121-45-9] $\text{C}_3\text{H}_9\text{O}_3\text{P}$, M 124.1, m -78° , b $22^\circ/23\text{mm}$, $86\text{--}86.5^\circ/351\text{mm}$, $111\text{--}112^\circ/760\text{mm}$, $111^\circ/\text{atm}$, d_4^{20} 1.0495, n_D^{20} 1.408. Treat the phosphite with Na (to remove water and any dialkyl phosphonate), then decant and distil it with protection against moisture. It has also been treated with sodium wire for 24 hours, then distilled in an inert atmosphere onto activated molecular sieves [Connor et al. *JCS Dalton Trans* 511 1986, DOI: 10.1039/DT9860000511]. It can be fractionally distilled using a spinning band column at high reflux ratio. It is a colourless liquid which is slowly hydrolysed by H_2O . [Gillis et al. *J Am Chem Soc* **80** 2999 1958, DOI: 10.1021/ja01545a025; ^{31}P NMR: Callis et al. *J Am Chem Soc* **79** 2719 1957, DOI: 10.1021/ja01568a015; Kosolapoff *Organophosphorus Compounds*, Wiley p 203 1950, Fieser **1** 1233; *Beilstein* **1** IV 1256.]

Trimethylsilyl acetamide [13435-12-6] $\text{C}_5\text{H}_{13}\text{NOSi}$, M 131.3, m $38\text{--}43^\circ$, $46\text{--}49^\circ$, $52\text{--}54^\circ$, b $84^\circ/18\text{mm}$, $185\text{--}186^\circ/\text{atm}$. Distil the amide repeatedly in an inert atmosphere with all operations to be performed in an anhydrous atmosphere. In the presence of moisture, **trimethylsilanol** (b $31\text{--}34^\circ/26\text{mm}$) is formed and is a likely impurity (check by NMR). [Birkofer et al. *Chem Ber* **96** 1473 1963, DOI: 10.1002/cber.1963096054; for reactions of the dianion see Kuzma et al. *J Org Chem* **49** 2015 1984, DOI: 10.1021/jo00185a038; Fieser **1** 1235; *Beilstein* **4** IV 4011.]

Trimethylsilyl acetonitrile (TMSAN) [18293-53-3] $\text{C}_5\text{H}_{11}\text{NSi}$, M 113.2, has b $49\text{--}51^\circ/10\text{mm}$, $65\text{--}70^\circ/20\text{mm}$, d_4^{20} 0.8729, n_D^{20} 1.4420. Check if NMR and IR spectra show impurities; if present dissolve it in $^*\text{C}_6\text{H}_6$ (10 volumes), wash it with buffer (AcOH/AcONa pH ca 7) several times, dry (CaCl_2) it, filter, evaporate and distil it. Its IR (CCl_4) has ν_{max} at $2215 (\text{CN}) \text{ cm}^{-1}$, and ^1H NMR (CCl_4) with δ at 0.23 (s, 9H, SiMe_3), and 1.53 (s, 2H, CH_2CN). [Matsuda et al. *JCS Perkin Trans 1* 26 1979, DOI: 10.1039/P19790000026; *Beilstein* **4** IV 3974.] This nitrile cyanomethylates various carbonyl compounds under Lewis base-catalysis [Kawano et al. *Chem Lett* **34** 1508 2005, DOI: org/10.1246/cl.2005.1508].

Trimethylsilyl azide [4648-54-8] $\text{C}_3\text{H}_9\text{N}_3\text{Si}$, M 115.2, has b $52\text{--}53^\circ/175\text{mm}$, $92\text{--}95^\circ/\text{atm}$, $95\text{--}99^\circ/\text{atm}$, d_4^{20} 0.878, n_D^{20} 1.441. Distil the azide through a Vigreux column in a N_2 atmosphere maintaining the oil bath temperature thermostat at $135\text{--}140^\circ$. Check the purity by ^1H NMR [CHCl_3 , δ : single peak at 13cps from Me_4Si]. Likely impurities are siloxane hydrolysis products. The azide is thermally stable even at 200° when it decomposes slowly without explosive violence. All the same, it is advisable to carry out the distillation behind a thick safety screen in a fumehood because unforeseen **EXPLOSIVE** azides may be formed on long standing. [Birkofer & Wegner *Org Synth Coll Vol* **6** 1030 1988, DOI: 10.15227/orgsyn.050.0107.]

Trimethylsilyl chloride (TMCS trimethyl chlorosilane, chlorotrimethylsilane) [75-77-4] $\text{C}_3\text{H}_9\text{ClSi}$, M 108.6, has m -40° , b $56\text{--}57^\circ/\text{atm}$, $58^\circ/760\text{mm}$, d_4^{20} 0.86, n_D^{20} 1.388. Likely impurities are other chlorinated methylsilanes and tetrachlorosilane (b 57.6°), some of which can form azeotropes. To avoid the latter, very efficient fractional distillation is required. It has been fractionated through a 12 plate glass helices-packed column with only the heart-cut material being used. It has also been fractionated through a 90cm with 19mm diameter Stedman column. Purify it by redistilling from CaH_2 before use. [Sauer et al. *J Am Chem Soc* **70**, 4254 1948, DOI: 10.1021/ja01192a511; Langer et al. *J Org Chem* **23** 50 1958, DOI: 10.1021/jo01095a017; Fieser **1** 1232; *Beilstein* **4** IV 4007.] **FLAMMABLE and CORROSIVE.**

Trimethylsilyl chloroacetate [18293-71-5] $\text{C}_5\text{H}_{11}\text{ClO}_2\text{Si}$, M 166.7, has m -20° , b $57\text{--}58^\circ/14\text{mm}$, $70\text{--}71^\circ/30\text{mm}$, $159^\circ/760\text{mm}$, d_4^{20} 1.057, n_D^{20} 1.4231. Purify the chloroacetate by repeated fractionation, and taking the fractions with clean NMR spectra. [Anderson *J Am Chem Soc* **74** 2371 1952, DOI: 10.1021/ja01129a502; *Beilstein* **4** IV 4004.] It has been used for the synthesis of substituted acetic acids [Akita et al. *J Organomet Chem* **348** 91 1988, DOI: 10.1016/0022-328X(88)80342-5].

Trimethylsilyl cyanide [7677-24-9] $\text{C}_4\text{H}_9\text{NSi}$, M 99.2, has m $8\text{--}11^\circ$, $10.5\text{--}11.5^\circ$, $11\text{--}12^\circ$, $12\text{--}12.5^\circ$, b $54\text{--}55^\circ/87\text{mm}$, $67\text{--}71^\circ/168\text{mm}$, $114\text{--}117^\circ/760\text{mm}$, $118\text{--}119^\circ/760\text{mm}$, d_4^{20} 0.79, n_D^{20} 1.43916. The material should have only one sharp signal in the ^1H NMR (in CCl_4 with CHCl_3 as internal standard) with δ at 0.4, and the IR with ν_{max} at 2210cm^{-1} ($\text{C}\equiv\text{N}$) [McBride & Beachall *J Am Chem Soc* **74** 5247 1952, DOI: 10.1021/ja01141a003; Prober *J Am Chem Soc* **77** 3224 1955, DOI: 10.1021/ja01617a023]; otherwise purify it by fractionating through an 18 x 1/4inch column. [Evers et al. *J Am Chem Soc* **81** 4493 1959, DOI: 10.1021/ja01526a013.] It has also been carefully distilled using a 60cm vacuum jacketed column. If the volume of sample is small, the cyanide can be chased (in the distillation) with xylene that had been previously distilled over P_2O_5 . It is **HIGHLY TOXIC** and **FLAMMABLE**. [Evans et al. *J Org Chem* **39** 914 1974, DOI: 10.1021/jo00921a012; *Beilstein* **4** IV 3893.]

Trimethylsilyldiazomethane (diazomethyl)trimethylsilane) [18107-18-1] $\text{C}_4\text{H}_{10}\text{N}_2\text{Si}$, M 114.2, distils

between 0°/100mm and 40°/15mm. This diazo-silane is non-explosive, and non-mutagenic, and the careful precautions used in preparing diazomethane are not strictly necessary here. It is a good safe and **stable substitute for diazomethane** and excess of reagent can be evaporated off. It is available commercially as 2.0M solutions in Et₂O or hexane. Solutions in hexane can be stored for periods of more than 6 months at 0° in the absence of light without noticeable decomposition. Care should be taken, however, when opening a container, as it may become pressurised by N₂ which may be released, and should be done at ~0°. Several methods were used for preparing this diazo-silane and some start from trimethylsilylmethylmagnesium chloride, e.g. by reaction with diazomethane (7-74%, Lappert et al. *J Chem Soc (A)* 2954 1970, DOI: 10.1039/J19700002954; Martin *Synth Commun* **13** 809 1983, DOI: 10.1080/00397918308063714), or with TsN₃ (~17%, Barton & Hoekman *React Inorg. Met-Org Chem* **9** 297 1979, DOI: 10.1080/00945717908069745], but the transfer of the azo group is most practical, high yielding and used on a large scale when diphenyl phosphorazidate is used, and is described here.

Trimethylsilylmethylmagnesium chloride [13170-43-9] is prepared in an inert atmosphere (e.g. argon) under strictly anhydrous conditions from Mg turning (10.7g, 0.44g-atom), anhydrous Et₂O (40ml) and 1,2-dibromoethane (0.1ml to initiate reaction) which are stirred at ~25° for 15 minutes, then 10ml of a mixture of trimethylsilylmethyl chloride (45.4g, 370mmol, freshly distilled at 97°/atm, see [2344-80-1]) in anhydrous Et₂O (100ml) is added with stirring all at once to start the reaction followed by the rest in a dropwise manner at such a rate as to maintain gently reflux during the addition (~2 hours). After the exothermic reaction subsides, reflux and stirring are continued for 1 hour then the mixture is cooled to ~25°, and this Grignard reagent is used in the next step. Under strictly anhydrous conditions and in a maintained argon atmosphere, a solution of diphenyl phosphorazidate (91.2g, 330mmol, freshly distilled at 134-136°/0.2mm see [26386-88-9]) in anhydrous Et₂O (350ml) is cooled and stirred in an ice-NaCl bath until the temperature is -10°, and the preceding Grignard reagent is added dropwise (via a funnel or a cannula) at such a rate as to maintain the inner temperature of the solution below 0° (addition requires ~ 1.5 hours, forming a large amount of white precipitate when two-thirds of the reagent is added). After addition is complete, the ice-salt bath is replaced by an ice bath while stirring for 2 hours, then stirring is continued for 14-16 hours. The mixture is again cooled in an ice-salt bath until the inner temperature drops to -15°, and cold H₂O (35ml) is added dropwise while keeping the inner temperature below 0° (requiring ~ 1 hour), and the resulting yellow silyldiazomethane mixture is stirred for a further 0.5 hours. The mixture is filtered, the white solid is washed with Et₂O (3 x 100ml), the combined filtrate and washings are washed with cold H₂O (2 x 100ml), dried (Na₂SO₄), and the filtrate is placed in a 1L flask equipped with a Teflon-coated magnetic stirrer bar and a 30cm Vigreux column (1.5cm diameter). The mixture is slowly concentrated to *ca* 200ml by distillation at ~760mm with a bath temperature below 45° during ~6 hours (the colour of the distilled Et₂O is yellow due to some co-distillation of the diazo-silane. This rate of distillation is important as a faster rate results in decreased yields of the diazo-silane. The concentration time can be reduced to ~ 4 hours if a 30cm Widmer column is used. The remaining deep-yellow solution is then distilled, in the same equipment, by reducing the pressure to 100mm at 0° (bath temperature) and then at 15mm and 40° (bath temperature) while the distillate is collected in a receiver in a Dry-ice/Me₂CO bath until distillation is complete. The distillate is dried (MgSO₄), filtered and diluted with redistilled hexane (100ml). This solution is then concentrated again by distillation through the 30cm Vigreux column until the temperature of the vapour reaches 68° (with final oil bath temperature at 87°) which requires ~3 hours. About 80-110ml of the residual yellow hexane solution contains 220-230mmol of **trimethylsilyldiazomethane** (67-70% based on the phosphorazidate). Its IR (hexane) has ν_{\max} at 2075, 1260 and 885 cm⁻¹, the ¹H NMR (100MHz, hexane, CHCl₃ as internal standard) has δ at 0.16 (s, 9H, -Si(CH₃)₃) and 2.58 (s, 1H, -CHN₂), and the ¹H NMR (100MHz, *C₆H₆) has δ at -0.03 (s, 9H, -Si(CH₃)₃) and 2.23 (s, 1H, -CHN₂).

The concentration of **trimethylsilyldiazo-methane** in hexane can be determined by adding 91mg (0.5mmol) of dibenzyl (see [103-29-7]) in 1ml of the hexane solution containing it, and measuring the ¹H NMR. The concentration of the azo-silane (*x* mmol/ml) is calculated from the formula: $x = 2a/b$, where *a* is the integral value (mm) of the methine proton (δ : 2.58) of **trimethylsilyldiazomethane**, and *b* is the integral value (mm) of the methylene protons (δ : 2.99) of dibenzyl. [Shioiri et al. *Org Synth Coll Vol* **8** 612 1993, DOI: 10.15227/orgsyn.068.0001].

It has been used for stereo and/or regio selective dipolar cycloadditions, e.g. with chiral acrylamides it produces Δ^2 -pyrazoline carboxylic acids (azaprolines) after protio-desilylation [Mish et al. *J Am Chem Soc* **119** 8379 1997, DOI: 10.1021/ja971708p], or with diethyl *trans*-glutaconate where the pyrazoline ester is oxidised to the respective pyrazole [Di & Rein *Tetrahedron Lett* **45** 4703 2004, DOI: 10.1016/j.tetlet.2004.04.097]. It also acts

as a carbene source in stereoselective [1 + 4] annulation reactions with silylvinylketenes to form a variety of cyclopent-2-enones in good to excellent yields [Moser et al. *J Org Chem* **71** 6542 2006, DOI: 10.1021/jo060994x]. It is commercially available in Et₂O (2.0M, d 0.733g/ml at 25°) and in hexanes (2.0M, d 0.733g/ml at 25°)

2-Trimethylsilyl-1,3-dithiane [13411-42-2] C₇H₁₆S₂Si, M 192.2, b 54.5°/0.17mm, 100°/8mm, d₄²⁰ 1.04, n_D²⁰ 1.533. Fractionally distil the dithiane through an efficient column and collect the fractions that have the correct NMR and IR spectra. Its ¹H NMR (CCl₄) has τ at 6.36 (SiMe₃), 9.87 (SCHS) and dithiane H at 7 and 8 (ratio 1:9:4:2) from Me₄Si; the UV has λ_{max} at 244nm (ε 711), sh 227nm (ε 800). [Corey et al. *J Am Chem Soc* **89** 434 1967, DOI: 10.1021/ja00978a048; for use as an acyl anion equivalent see Smith & Baldi *J Amer Chem Soc* **119** 6925 1997, DOI: 10.1021/ja00978a048; Smith et al. *J Am Chem Soc* **125** 14435 2003, DOI: 10.1021/ja0376238].

2-(Trimethylsilyl)ethanesulfonyl chloride (SES-Cl) [106018-85-3] C₅H₁₃ClO₂SSi, M 200.8, b 60°/0.1mm, 146.8°/760mm, d₄²⁵ 1.059, n_D²⁵ 1.4444. Check IR; if the bands at ~3200 (OH) cm⁻¹ are strong, then much of the SES-Cl had hydrolysed, and it should be treated with POCl₃ (with cooling) and stirred at ~25° for about 1 hour, poured into ice cold H₂O, extracted with CH₂Cl₂, washed with NaHCO₃, dried (Na₂SO₄), evaporated, and it distils as a yellow oil in a vacuum. This procedure is used for converting the Na salt [18143-30-1] to SES-Cl. It reacts with amines to form amides, e.g. SES-NRR', which on heating with CsF (i.e. F⁻ ions) in DMF at 95° provide the amine (NHRR'), SO₂ and CH₂=CH₂ [Weinreb et al. *Tetrahedron Lett* **27** 2099 1986, DOI: 10.1016/S0040-4039(00)84458-5]. [Ribière et al. *Chem Rev* **106** 2249 2006, DOI: 10.1021/cr0300587]

Trimethylsilyl ethanol [2916-68-9] C₅H₁₄OSi, M 118.3, b 53-55°/11mm, 71-72°/10mm, 75°/41mm, 95°/100mm, 211-213°/760mm, d₄²⁵ 0.8254, n_D²⁵ 1.4220. If the NMR spectrum is not clean, then dissolve the alcohol in Et₂O, wash it with aqueous NH₄Cl solution, dry (Na₂SO₄), evaporate and distil it. The **3,4-dinitrobenzoyl derivative** has m 66° (from EtOH). [NMR: Speier et al. *J Am Chem Soc* **79** 974 1957, DOI: 10.1021/ja01561a054; *Z Naturforsch* **14b** 137 1959, *Beilstein* **4** IV 3951.] Protecting reagent for phosphate [Wada & Sekine *Tetrahedron Lett* **35** 757 1994, DOI: 10.1021/ol061461d; Sawabe et al. *Tetrahedron Lett* **33** 7685 1992, DOI: 10.1016/0040-4039(93)88016-C; Chao et al. *J Org Chem* **59** 6687 1994, DOI: 10.1021/jo00101a029] and carbonyl groups [Kita et al. *JCS Perkin Trans I* 2639 1993, DOI: 10.1039/P19930002639]; also used to prepare *N*-protected amines via alcoholysis of the respective isocyanates [Kobayashi & Motoyama *Synlett* 2670 2006, DOI: 10.1055/s-2006-950435; Crawley & Funk *Org Lett* **8** 3995 2006, DOI: 10.1021/ol061461d].

2-(Trimethylsilyl)ethoxymethyl chloride (SEMCl) [76513-69-4] C₆H₁₅ClOSi, M 166.7, b 57-59°/8mm, 170-172°/atm, d₄²⁰ 0.942, n_D²⁰ 1.4350. Dissolve SEMCl in pentane, dry it (MgSO₄), evaporate and distil the residual oil in a vacuum. Stabilise it with 10 ppm of diisopropylamine. Store it under N₂ in a sealed container in a refrigerator. [Lipshutz & Pegram *Tetrahedron Lett* **21** 3343 1980, DOI: 10.1016/S0040-4039(00)78684-9.] SEM is a good protecting group for alcohols forming SEM-ethers which are stable over a wide pH range but can be selectively cleaved with fluoride ions under mild aprotic conditions [Wada et al. *Tetrahedron Lett* **36** 1683 1995, DOI: 10.1016/0040-4039(95)00130-5].

2-(Trimethylsilyl)ethoxymethyltriphenylphosphonium chloride (SEM-triphenylphosphonium chloride) [82495-75-8] C₂₄H₃₀ClOPSi, M 429.0, m 140-142°, 145-149°. Wash the solid with AcOH and recrystallise it from CH₂Cl₂/EtOAc. Dry it in a vacuum desiccator. *Hygroscopic*. The ¹H NMR (CDCl₃) has δ at -0.2 (s, Me₃Si), 0.8 (t, 8Hz, CH₂Si), 3.83 (t, 8Hz, OCH₂), 5.77 (d, J_{PH} = 4Hz, P⁺-CH₂O) and 7.70 (m, aromatic H). [Schönauer & Zbiral *Justus Liebigs Ann Chem* 1031 1983, DOI: 10.1002/jlac.198319830615.]

Trimethylsilylethyl phenylsulfone (phenyl-2-trimethylsilylethylsulfone) [73476-18-3] C₁₁H₁₈O₂SSi, M 242.4, m 50-53°, 52°. Dissolve the sulfone in Et₂O, wash it with saturated HCO₃ followed by saturated NaCl, H₂O and dried (MgSO₄). Filtration followed by evaporation leaves residual crystals with m 52°. [Hsiao & Shechter *Tetrahedron Lett* **23** 1963 1982, DOI: 10.1016/S0040-4039(00)87234-2; Hsiao & Shechter *J Org Chem* **53** 2688 1985, DOI: 10.1021/jo00247a006.]

Trimethylsilyl isocyanate [1118-02-1] $\text{C}_4\text{H}_9\text{NOSi}$, **M 115.2**, **b 90-92°/atm**, **b 91.3-91.6°/atm**, d_4^{20} **0.850** n_D^{20} **1.43943**. Purify it by repeated fractionation as for the isothiocyanate below. [Eaborn *J Chem Soc* 3077 1950, DOI: 10.1039/JR9500003077; *Beilstein* **4** III 1861, **4** IV 4011.]

Trimethylsilyl isothiocyanate [2290-65-5] $\text{C}_4\text{H}_9\text{NSSi}$, **M 131.3**, **m -49°, -33°**, **b 142.6-143.1°/759mm**, **143.8°/760mm**, n_D^{20} **1.4809**. The ^1H NMR spectrum should have only one peak; if not, purify it by repeated fractionation in an all-glass system using a 50cm (4mm internal diameter) column without packing. [Anderson *J Am Chem Soc* **69** 3049 1947, DOI: 10.1021/ja01204a036; Fehér & Blümcke *Chem Ber* **90** 1934 1957, DOI: 10.1002/cber.19570900933; Neidlein & Hege *Synthesis* **50** 1975, DOI: 10.1055/s-1975-23663; *Beilstein* **4** III 1861, **4** IV 4011.] It reacts with aziridines and cyclohexene oxide, without a catalyst, to provide *N*-substituted *trans*-2-amino-1-isothiocyanates or *trans*-2-hydroxy-1-isothiocyanates [Prusinowska & Gawronski *Synth Commun* **39** 2795 2009, DOI: 10.1080/00397910802691890].

(Trimethylsilyl)methanol [3219-63-4] $\text{C}_4\text{H}_{12}\text{OSi}$, **M 104.2**, **b 120-122°/754mm**, **122-123°/768mm**, d_4^{20} **0.83** n_D^{20} **1.4176**. If the NMR indicates impurities (should have only two signals), then dissolve it in Et_2O , shake this with aqueous 5N NaOH, $\text{M H}_2\text{SO}_4$, saturated aqueous NaCl, dry (MgSO_4) and distil it using an efficient column at atmospheric pressure. The **3,5-dinitrobenzoate** has **m 70-70.5°** (from 95% EtOH). [Huang & Wang *Acta Chem Sin* **23** 291 1957, http://sioc-journal.cn/Jwk_hxxb/EN/; cf. *Chem Abstr* **52** 19911 1958, Seyferth *J Am Chem Soc* **81** 1844 1959, DOI: 10.1021/ja01517a018; and Speier et al. *J Am Chem Soc* **70** 1117 1948, DOI: 10.1021/ja01183a074; *Beilstein* **4** III 1844, **4** IV 2876.]

(Trimethylsilyl)methylamine (aminomethyl trimethylsilane) [18166-02-4] $\text{C}_4\text{H}_{13}\text{NSi}$, **M 103.2**, **b 101.6°/735mm**, d_4^{20} **0.77**, n_D^{20} **1.416**. A possible contaminant is hexamethyldisiloxane. It should have two ^1H NMR signals in CDCl_3 ; if not, dissolve it in $^*\text{C}_6\text{H}_6$, shake it with 15% aqueous KOH, separate, dry (Na_2SO_4), filter, evaporate and distil it using a still of *ca* 10 theoretical plates. The water azeotrope has **b 83°/735mm**; hence it is important to dry the extract well. The **hydrochloride** has **m 198-199°** (from MeOH or Me_2CO). [Noll et al. *J Am Chem Soc* **73** 3867 1951, DOI: 10.1021/ja01152a092; *Beilstein* **4** IV 3878.]

(Trimethylsilyl)methyl chloride (chloromethyltrimethylsilane, silico-neopentyl chloride) [2344-80-1] $\text{C}_4\text{H}_{11}\text{ClSi}$, **M 122.7**, **b 97.1°/734mm**, **98.2-98.7°/747mm** d_4^{25} **0.979**, n_D^{20} **1.4180**. This chloride is prepared by stirring a mixture of TMS (174g, 2 moles, Me_4Si see [75-76-3]) in dry CCl_4 (150ml) and PCl_5 (3g) [use suitably placed traps of Dry-ice/ Me_2CO to prevent loss of TMS] under a reflux condenser and irradiated with a 450 watt GE sunlight lamp while dry chlorine is bubbled through for 4 hours at a rate of 0.5 mole per hour (check weight of flask occasionally). The mixture is then fractionated using a 15-plate glass helix packed column to give the **trimethylsilylmethyl chloride** (53g, 0.44 mole), **b 97.1°/734mm**, **polychlorinated tetramethylsilane** (93g) and recovered TMS (62g, 0.7 mole). The **chloride** can be analysed by placing ~0.2g in a gelatin capsule in a Parr bomb and fused with Na_2O_2 (15g) and sucrose (1g). The melt is treated with H_2O , acidified with HNO_3 and the chloride ion is determined by titration using the Volhard method. [Whitmore & Sommer *J Am Chem Soc* **68** 481 1946, DOI: 10.1021/ja01207a036; *Beilstein* **4** IV 3877.] It has also been prepared by reaction of chlorodimethyl-chloromethyl-silane and MeMgBr in Et_2O [Whitmore et al. *J Am Chem Soc* **69** 1976 1947, DOI: 10.1021/ja01200a043; Roedel *J Am Chem Soc* **71** 269 1949, DOI: 10.1021/ja01169a068]. [*Beilstein* **4** IV 3877.] It is readily converted to **trimethylsilylmethylmagnesium chloride** (see [13170-43-9] above) in ~90% yield. It reacts with *n*-BuLi in pentane to give **trimethylsilylmethylLi** [Sommer et al. *J Am Chem Soc* **71** 2746 1949, DOI: 10.1021/ja01176a043], which reacts with anhydrous cerium (III) chloride to form **trimethylsilylmethyl cerium dichloride**. The latter reacts with acyl halides to provide bis- β -silylethyl tertiary alcohols that efficiently undergo a trimethylchlorosilane-promoted **Peterson reaction** to generate allylsilanes in high overall yields [Anderson & Fuchs *Synth Commun* **17** 621 1987, DOI: 10.1080/00397918708075736]. It is useful also for Peterson olefination and homologation of aldehydes and ketones via 1,2-epoxysilanes [Lee et al. *Tetrahedron* **45** 5877 1989, DOI: 10.1016/S0040-4020(01)89114-6]. **Trimethylsilylmethyl bromide** [18243-41-9] has **M 167.1**, **b 115.5°/atm**, d_4^{25} **1.17**, n_D^{20} **1.444** [*Beilstein* **4** IV 3878], and **trimethylsilylmethyl iodide** [4206-67-1] has **M 214.1**, **b 139-141°/atm**, d_4^{25} **1.433**, n_D^{20} **1.491** [*Beilstein* **4** IV 3878], and both should be stored in the dark.

Trimethylsilylmethyl phenylsulfone (phenyltrimethylsilylmethylsulfone) [17872-92-3] $\text{C}_{10}\text{H}_{16}\text{O}_2\text{SSi}$, **M 228.4**, **m 28-32°**, **b 121°/0.01mm**, **160°/6mm**, n_D^{20} **1.5250**. Fractionate the sulfone at high vacuum and recrystal-

lise it from pentane at -80° . If too impure (cf. IR), dissolve it in CH_2Cl_2 (ca 800ml for 100g), wash this with 2M aqueous NaOH (2 x 200ml), brine, dry, evaporate and distil it. [Craig et al. *JCS Perkin Trans 1* 1949 1985, DOI: 10.1039/P19850001949; IR and NMR: Cooper *J Am Chem Soc* **76** 3713 1954, DOI: 10.1021/ja01643a035.]

1-Trimethylsilyloxy-1,3-butadiene [6651-43-0] $\text{C}_7\text{H}_{14}\text{OSi}$, M 142.3, b $131^{\circ}/760\text{mm}$ (mixture of isomers), **49.5°/25mm** (*E*-isomer), d_4^{20} 0.8237, n_D^{20} 1.447. Purify the butadiene by fractional distillation, and collect the fractions with the required ^1H NMR. Store it under N_2 — it is a flammable and moisture-sensitive liquid. [Caseau et al. *Bull Soc Chim Fr* 16658 1972, Belge Patent 670,769, *Chem Abstr* **65** 5487d 1966.] Used in the stepwise Diels-Alder reaction with 4,6-dinitrobenzofuroxan [Linder et al. *Org Lett* **1** 6 2012, PMID: 22126093].

1-(Trimethylsilyloxy)cyclopentene [19980-43-9] $\text{C}_8\text{H}_{16}\text{OSi}$, M 156.3, b $45^{\circ}/11\text{mm}$, $75-80^{\circ}/20-21\text{mm}$, d_4^{20} 0.878, n_D^{20} 1.441. If too impure as seen by the NMR spectrum, then dissolve it in 10 volumes of pentane, shake with cold NaHCO_3 (3 x 500ml), then 1.5M HCl (200ml) and aqueous NaHCO_3 (200ml) again, dry (Na_2SO_4), filter, evaporate and distil it through a short Vigreux column. Its ^1H NMR (CDCl_3) has δ at 0.21 (s, 9H), 1.55 (m, 2H), 1.69 (m, 2H), 2.05 (br d, 4H) and 4.88 (br s, 1H). GLC in a 6ft x 1/8inch with 3% SP2100 on 100-120 mesh Supelcoport column should give one peak. Store dry. [For the *cyclohexene* analogue [6651-36-1] $\text{C}_9\text{H}_{18}\text{OSi}$, M 170.3, b $64-65^{\circ}/15\text{mm}$, $165^{\circ}/\text{atm}$, see Varghese et al. *Org Synth Coll Vol* **8** 460 1993, DOI: 10.15227/orgsyn.067.0141.]

2-(Trimethylsilyloxy)furan [61550-02-5] $\text{C}_7\text{H}_{12}\text{O}_2\text{Si}$, M 156.3, b $34-35^{\circ}/9-10\text{mm}$, $42-50^{\circ}/17\text{mm}$, $40-42^{\circ}/25\text{mm}$, d_4^{20} 0.950, n_D^{20} 1.436. Fractionally distil the furan using a short path column. Its ^1H NMR in CCl_4 has δ at 4.90 (dd, $J = 1.3\text{Hz}$, 3H), 6.00 (t, $J = 3\text{Hz}$, 4H) and 6.60 (m, 5H). [Yoshii et al. *Heterocycles* **4** 1663 1976, DOI: 10.3987/R-1976-10-1663.]

4-Trimethylsilyloxy-3-penten-2-one (*cis*) (acetylacetone enol trimethylsilyl ether) [13257-81-3] $\text{C}_8\text{H}_{16}\text{O}_2\text{Si}$, M 172.3, b $66-68^{\circ}/4\text{mm}$, $61-63^{\circ}/5\text{mm}$, $205.1^{\circ}/760\text{mm}$, d_4^{20} 0.917, n_D^{20} 1.452. Fractionally distil the enone, and store it in glass ampoules which are sealed under N_2 . It hydrolyses readily in contact with moisture giving, as likely impurities, hexamethyldisiloxane and 2,4-pentanedione. It is a silylating agent. [West *J Am Chem Soc* **80** 3246 1958, DOI: 10.1021/ja01546a018; *Beilstein* **4** IV 4003.]

1-(Trimethylsilyl)-2-phenylacetylene (1-phenyl-2-trimethylsilylacetylene) [78905-09-6, 2170-06-1] $\text{C}_{11}\text{H}_{14}\text{Si}$, M 174.3, b $45-46^{\circ}/0.1\text{mm}$, $67^{\circ}/5\text{mm}$, $87.5^{\circ}/9\text{mm}$, d_4^{20} 0.8961, n_D^{20} 1.5284. Dissolve the acetylene in Et_2O , wash with H_2O , dry and fractionate it through a Todd column. [Benkeser & Hickner *J Am Chem Soc* **80** 5298 1958, DOI: 10.1021/ja01552a072.]

3-(Trimethylsilyl)-1-propyne (trimethyl[propargyl]silane) [13361-64-3] $\text{C}_8\text{H}_{12}\text{Si}$, M 112.3, b $99-100^{\circ}/760\text{mm}$, d_4^{20} 0.7581, n_D^{20} 1.4091. Fractionally distil the propyne, and add 2,6-di-*tert*-butyl-*p*-cresol ($\sim 0.5\%$) to stabilise it. [For polymerisation see Matsuda et al. *J Am Chem Soc* **105** 7473 1983, DOI: 10.1021/ja00363a061; use for the three-carbon elongation of aldehydes see Maeta & Suzuki *Tetrahedron Lett* **33** 5969 1992, DOI: 10.1016/S0040-4039(00)61102-4; Petrov et al. *Doklady Acad Nauk USSR* **93** 293 1953, cf. *Chem Abstr* **48** 13616 1954, *Beilstein* **4** IV 3938.]

2-Trimethylsilylpyridine [13737-04-7] $\text{C}_6\text{H}_{13}\text{NSi}$, M 151.3, b $47-49^{\circ}/5\text{mm}$, $69^{\circ}/14.5\text{mm}$, $74^{\circ}/21\text{mm}$, d_4^{25} 0.9113, n_D^{20} 1.489. Purify it by distillation in a vacuum, but if it is discoloured then dissolve it in Et_2O or C_6H_6 , wash it with H_2O , dry over Na_2SO_4 , filter, evaporate and distil the residue. It is more readily hydrolysed than the 3- or 4-trimethylsilyl isomers and the relative rates of hydrolysis in $\text{H}_2\text{O}:\text{MeOH}:\text{EtOH}$ are 740:120:1; pyridine and trimethylsilanol or the alkoxytrimethylsilane being liberated. Acids inhibit the reaction but bases have little effect. [Anderson et al. *J Chem Soc B* 450 1968, DOI: 10.1039/J29680000450; synthesis from 2-chloropyridine and $\text{Mg}/\text{Me}_3\text{SiCl}/\text{HMPT}$ then $\text{H}_2\text{O}/\text{NaHCO}_3$ in 62% yield: Effenberger & Häbich *Justus Liebigs Ann Chem* **1979** 842 1979, DOI: 10.1002/jlac.197919790613; Itami *Tetrahedron* **57** 5045 2001, DOI: 10.1016/S0040-4020(01)00348-9.]

3-Trimethylsilylpyridine [13779-37-2] $C_6H_{13}NSi$, M 151.3, b 94°/30mm, d_4^{25} 0.9113, n_D^{20} 1.4913. Purify it as for the 2-isomer above except that less care needs to be taken as it hydrolyses relatively (extremely) slowly. [Anderson et al. *J Chem Soc B* 450 1968, DOI: 10.1039/J29680000450.]

4-Trimethylsilylpyridine [18301-46-7] has, b 107°/48mm, d_4^{25} 0.9113, n_D^{20} 1.4868. Purify it as for the 2-isomer above except that less care needs to be taken as it hydrolyses relatively (extremely) slowly. [Anderson et al. *J Chem Soc B* 450 1968, DOI: 10.1039/J29680000450.]

1-Trimethylsilyl-1,2,4-triazole [18293-54-4] $C_5H_{11}N_3Si$, M 141.3, b 74°/12mm, 85.6°/760mm, d_4^{20} 0.99, n_D^{20} 1.4604. Fractionally distil it at atmospheric pressure in an inert atmosphere because it is moisture sensitive. [Birkofer et al. *Chem Ber* 93 2804 1960, DOI: 10.1002/cber.19600931207.]

Trimethylsilyl trifluoromethane (TMSCF₃, trifluoromethyl trimethylsilane, Ruppert's reagent) [81290-20-2] $C_4H_9F_3Si$, M 142.2, b 54-55°/atm, 55-55.5°/atm, d_4^{20} 0.962, n_D^{20} 1.332. Purify the silane by distilling it from trap to trap in a vacuum of 20mm using a bath at 45° and Dry-ice/Me₂CO bath for the trap. The liquid in the trap is then washed with ice cold H₂O (3x), the top layer is collected, dried (Na₂SO₄), and the liquid is decanted and fractionated through a helices-packed column at atmospheric pressure. ¹H, ¹³C, ¹⁹F, and ²⁹Si NMR can be used for assessing the purity of fractions. [Ruppert et al. *Tetrahedron Lett* 25 2195 1984, DOI: 10.1016/S0040-4039(01)80208-2; Krishnamurti et al. *J Org Chem* 56 984 1991, DOI: 10.1021/jo00003a017; Beilstein 4 IV 3892.] TMSCF₃ adds the trifluoromethyl nucleophile to aldehydes and ketones [Parkash et al. *J Org Chem* 71 6806 2006, DOI: 10.1021/jo060835d].

2,8,9-Trimethyl-2,5,8,9-tetraaza-1-phosphabicyclo[3.3.3]undecane (Verkade Superbase, 2,8,9-trimethyl-pro-azaphosphatrane) [120666-13-9] $C_9H_{21}N_4P$, M 216.3, m 110-115°, pKa 26.8 (DMSO). The free superbase can be prepared in three ways. When tris(dimethylamino)phosphine (8.8g, 54mmol, toxic see [1608-26-0]) and tris[2-(methylamino)-ethyl]amine (10.0g, 53mmol), see [65604-89-9] are dissolved in dry xylene and refluxed for 21 days, then the solvent is removed *in vacuo*, a residual thick oil is obtained. On heating this oil at 105°/0.05mm the *pro-azaphosphatrane* (53g, 24.5mmol, 46%) sublimes out as a colourless waxy solid.

In a second preparation the *trimethyl-pro-azaphosphatrane hydrochloride* (0.87g, 3.4mmol, see [138800-17-6] below) dissolved in MeCN (10ml) is added dropwise to a suspension of freshly sublimed *tert*-BuOK (0.41g, 3.7mmol) in MeCN (20ml), stirred for 30 minutes, the solvent is removed *in vacuo*, the residue is extracted with hexane (2 x 30ml), evaporated, and the residue is sublimed at 60°/0.01mm to give the pure *superbase* (80%) as a white solid. [Schmidt et al. *Z Anorg Allg Chem* 578 75 1989, DOI: 10.1002/zaac.19895780109.]

In a third preparation the *trimethyl-azaphosphatrane hydrochloride* is heated slowly with a large excess of anhydrous NaOH under vacuum. No reaction occurs below 200°, but above this temperature sudden sublimation of the free superbase occurs. Extraction of the sublimate, and/or the reaction mixture, with *C₆H₆ followed by evaporation of the solvent provided pure *free base* (53% yield). Sometimes charring occurs during this reaction resulting in lower yields. [Lensink et al. *J Am Chem Soc* 111 3478 1959, DOI: 10.1021/ja00191a081.]

The *free base* is an unusually strong Lewis base. It deprotonates phenol (pKa 10), protonated 'Proton Sponge' [1,8-(bismethylamino)naphthalene HI (pKa 12.3, Alder et al. *JCS Chem Commun* 723 1968, DOI: 10.1039/C19680000723), diethyl malonate (pKa 13, Pearson & Dillon *J Am Chem Soc* 75 2439 1953, DOI: 10.1021/ja01106a048), H₂O (measured pKa at 25° 13.9965, pKa at 20° 14.1669, Harned & Robinson *Trans Farad Soc* 36 973 1940, DOI: 10.1039/TF9403600973; or pKa 15.74 ? Reeve et al. *Can J Chem* 57 2747 1979, DOI: 10.1139/v79-444), but does not react with *tert*-BuOH (pKa 16.5, Reeve et al. *Can J Chem* 57 2747 1979, DOI: 10.1139/v79-444), to give the protonated base as the only product. In the presence of 1 equivalent of H₂O an equilibrium mixture of *free base to cation* of ca 4:1 is obtained (from NMR studies) [Lensink et al. *J Am Chem Soc* 111 3478 1989, DOI: 10.1021/ja00191a081]. The *un-methylated analogous free base* is not very stable although it *hydrochloride* is quite stable and it is a *stronger base* than the 2,8,9-trimethyl- or the respective 2,8,9-tribenzyl- derivative. The upper limit of the pKa of the trimethyl compound in DMSO is ~26.8 (by ¹³P NMR) [Laramay & Verkade *J Am Chem Soc* 112 9421 1990, DOI: 10.1021/ja00181a070]. The *free base* has IR (Nujol) with ν_{max} at 1332 s, 1303 m, 1244 s, 1226 s, 1197 m, 1145 s, 1128 s, 1053 s, 1004 s, 960 w, 887 m, 850 s, 767 w, 650 s and 634 s cm⁻¹; the ¹H NMR (300MHz, CD₃CN, TMS) has δ at 2.60 (d, 9H, ³J_{PH} = 11.0 Hz, CH₃) and 2.76 (br s, 12H, CH₂); the ¹³C NMR (75MHz, C₆D₆, TMS) has δ at 37.2 (²J_{PC} = 41.0 Hz, CH₃), 49.4 (d, ²J_{PC} = 6.7 Hz, N_{eq}CH₂) and 51.3 (s, N_{ax}CH₂); the ³¹P NMR (122MHz, C₆D₆, 85% H₃PO₄) has δ at 120.8; and the HRMS has *m/z* 216.15088 (calc for M is 216.15039). [Schmidt et al. *Z Anorg Allg Chem* 578

75 1989, DOI: 10.1002/zaac.19895780109; Lensink et al. *J Am Chem Soc* **111** 3478 1989, DOI: 10.1021/ja00191a081; Laramay & Verkade *J Am Chem Soc* **112** 9421 1990, DOI: 10.1021/ja00181a070; Laramay & Verkade *Z Anorg Allg Chem* **605** 163 1991, DOI: 10.1002/zaac.19916050120; Schmidt et al. *Inorg Chem* **29** 2214 1990, DOI: 10.1021/ic00337a008]. In addition to being easily protonated (see following entry), the free base has been readily converted to P-substituted adducts (**five-coordinated phosphatranes**), where the P-H is replaced by P-X, where -X is -O (by using Me_3SiO_2), -S (by using S_8), -Se (by using Se), $-\text{N}_3\text{P}$ (by using PhN_3), $-\text{NPh}$ (by using PhN_3 and heating), $-\text{CS}_2$ (by using CS_2) and $-\text{PtCl}_2$ [by using $(\text{Et}_2\text{S})_2\text{-PtCl}_2$] in 60 to >80% yields [Schmidt et al. *Z Anorg Allg Chem* **578** 75 1989, DOI: 10.1002/zaac.19895780109]. **Note** that the difference between the **pro-phosphatranes** and the **phosphatranes** is that the latter possess an apical $\text{N}\rightarrow\text{P}$ bond as evidenced by the shorter bond distance in X-Ray diffraction analyses and the large decrease in the $\delta^{31}\text{P}$ chemical shifts in the NMR spectra.

2,8,9-Trimethyl-2,5,8,9-tetraaza-1-phosphabicyclo[3.3.3]undecane hydrochloride (Verkade Superbase HCl, 2,8,9-trimethylazaphosphatrane HCl) [138800-17-6] $\text{C}_9\text{H}_{21}\text{N}_4\text{P}\cdot\text{HCl}$, **M 252.7, m 132-136°**. Like the preceding free base it is prepared in different ways — two are described here. A solution of PCl_3 (1.22g, 8.85mmol) in CH_2Cl_2 (5.0ml) is added to a solution of $\text{P}(\text{NMe})_3$ (2.89g, 17.7mmol, toxic see [1608-26-0]) in CH_2Cl_2 (5.0ml), then cooled to 5°, and TREN (3.88g, 26.5mmol, see [4097-89-6]) in CH_2Cl_2 (25ml) is added during ~15 minutes. The precipitate formed is filtered off, washed with CH_2Cl_2 (25ml) and dried to give the pure **hydrochloride** (5.58g, ~100%). Crystals for X-Ray diffraction can be grown from concentrated MeOH solutions. The **2,8,9-tribenzyl hydrochloride** can be similarly prepared in 89% yield by using tris-(*N*-benzylaminoethyl)amine in place of TREN. [Laramay & Verkade *Z Anorg Allg Chem* **605** 163 1991, DOI:10.1002/zaac.19916050120].

In the **second preparation** tris[2-(methylamino)-ethyl]amine (1.67g, 11.4mmol), see [65604-89-9] in CH_2Cl_2 (20ml) is added with stirring to a mixture of $\text{ClP}(\text{NMe}_2)_2$ (1.76g, 11.4mmol, TOXIC) and Et_3N (1.5g, 15mmol) in CH_2Cl_2 (30ml) during 5 minutes, then the mixture is stirred at ~25° for 1 hour, the volatiles are evaporated off *in vacuo* to give the **hydrochloride** quantitatively. It can be recrystallised from hexane/ CHCl_3 (82% yield) in colourless crystals. When the hydrochloride salt is treated with AgBF_4 in CH_2Cl_2 the **HB F_4 salt** is obtained quantitatively. The salts are very difficult to deprotonate (see preceding entry). The X-ray crystal structure shows a short intra-annular bond between the apical N atom and the P atom with a robust P—H bond, the positive charge being on the apical N atom. [Laramay & Verkade *J Am Chem Soc* **112** 9421 1990, DOI: 10.1021/ja00181a070; Lensink et al. *J Am Chem Soc* **111** 3478 1989, DOI: 10.1021/ja00191a081.] It has ^1H NMR (300MHz, CDCl_3 , TMS) with δ at 2.61 (d, 9H, $^3J_{\text{PH}} = 17.4$ Hz, CH_3) and 3.03 (dt, 6H, $\text{N}_{\text{ax}}\text{CH}_2$, $^3J_{\text{PH}} = 11.0$ Hz, $^3J_{\text{HH}} = 6.2$ Hz), 3.58 (dt, 6H, $\text{N}_{\text{eq}}\text{CH}_2$, $^3J_{\text{PH}} = 4.7$ Hz, $^3J_{\text{HH}} = 6.2$ Hz), 5.20 (d, 1H, $^1J_{\text{PH}} = 6.2$ Hz); the ^{13}C NMR (75MHz, CDCl_3) has δ at 34.4 (d, $^2J_{\text{PC}} = 17.1$ Hz, CH_3), 41.3 (d, $^2J_{\text{PC}} = 6.1$ Hz, $\text{N}_{\text{ax}}\text{CH}_2$) and 47.3 (d, $^2J_{\text{PC}} = 7.3$ Hz, $\text{N}_{\text{eq}}\text{CH}_2$); the ^{31}P NMR (122MHz, CDCl_3 , 85% H_3PO_4) has δ at -10.6 [Lensink et al. *J Am Chem Soc* **111** 3478 1989 DOI: 10.1021/ja00191a081].

Trimethyl vinyl silane [754-05-2] $\text{C}_5\text{H}_{12}\text{Si}$, **M 100.2, b 54.4°/744mm, 55.5°/767mm, d_4^{25} 0.6865, n_D^{25} 1.3880**. If the ^1H NMR spectrum shows impurities, then dissolve it in Et_2O , wash it with aqueous NH_4Cl solution, dry over CaCl_2 , filter, evaporate and distil it at atmospheric pressure in an inert atmosphere. It is used as a copolymer and may polymerise in the presence of free radicals. It is soluble in CH_2Cl_2 . [Nagel & Post *J Org Chem* **17** 1379 1952, DOI: 10.1021/jo50010a016; Beilstein **4** IV 3922.] A useful reagent for preparing trimethylsilylethylsulfoxides, which in the presence of fluoride ions (e.g with TBAF), provide a convenient source of sulfenate anions ($\text{R}'\text{SO}^-$) that react with RX to form the corresponding sulfoxides ($\text{R}'\text{SOR}$) [Foucoin et al. *Synthesis* 1315 2007, DOI: 10.1055/s-2007-966017], and in the efficient O-silylation of a variety of alcohols using Rh(I)/HCl catalyst at room temperature [Park & Jun *Org Lett* **9** 4073 2007, DOI: 10.1021/ol701909e]

2,4,6-Trinitrobenzenesulfonic acid hydrate (TNBS, picrylsulfonic acid) [2508-19-2] $\text{C}_6\text{H}_3\text{N}_3\text{O}_9$, **M 293.2, m 180°, λ_{max} 240nm (ϵ 650 $\text{M}^{-1}\text{cm}^{-1}$), d^{25} 0.955g/ml. $\text{pK}_{\text{Est}} \sim <0$** . It is also available as 0.1M and 5%w/v solutions in H_2O . Recrystallise TNBS from 1M HCl , or a mixture of EtOH (50ml), H_2O (30ml) and concentrated HCl (70ml) for 65g of acid, and dry it at 100°. It is a strong oxidising acid, keep away from reducing agents otherwise it can detonate. The **diethanolamine salt** had **m 182-183°** [Golumbic *J Org Chem* **11** 518 1946, DOI: 10.1021/jo50010a016]. [Beilstein **11** III 161.]

Tri(4-nitrophenyl)phosphate [3871-20-3] $C_{18}H_{12}N_3O_{10}P$, **M 461.3**, **m 155-156°, 156°, 156-158°, 157-159°**. This phosphate has been recrystallised from AcOH, dioxane, AcOEt and Me₂CO and dried it in a vacuum over P₂O₅. [Ketelaar & Gersmann *J Am Chem Soc* **72** 5777 1950, DOI: 10.1021/ja01168a532; Moffatt & Khorana *J Am Chem Soc* **79** 3741 1957, DOI: 10.1021/ja01571a035.]

Tri(*n*-octyl)phosphine (TOP) [4731-53-7] $C_{24}H_{51}P$, **M 370.6**, **m 30°, 48°, b 175°/0.3mm, 194-195°/1mm, 240°/10mm, 291°/50mm, 284-291°/50mm, 442°/760mm, 445°/760mm, 493°/760mm, d²⁵ 0.831g/ml, n_D²⁵ 1.666, pK_{Est} ~8.2**. Among various syntheses, it is also prepared by reaction of di-*n*-octylmagnesium and PCl₃, and purified by fractional distillation at as high a vacuum as possible. It reacts with PCl₃ (275°/1hr) to form *n*-octylphosphorous dichloride, a useful synthetic intermediate. **TOP** has basic properties [cf. pK_a of *n*-Bu₃P = 8.33, see Henderson & Streuli *J Am Chem Soc* **82** 5791 1960, DOI: 10.1021/ja01507a008; cf. compare NH₃, PH₃ with AsH₃; Davies & Addis *J Chem Soc* 1622 1937, DOI: 10.1039/JR9370001622]. [For synthesis from TOPCl₂ see Horner et al. *Chem Ber* **92** 2088 1959, DOI: 10.1002/cber.19590920920; for synthesis from 1-octene and PH₃ see Rauhut et al. *J Org Chem* **26** 5138 1961, DOI: 10.1021/jo01070a087; Pass et al. *Monatsh Chem* **90** 792 1959, DOI: 10.1007/BF00905667; *Beilstein* **4** IV 3438].

It is a useful solvent in extraction (mineral acids) processes, in formulation of flame retardants, in formulating fumigants and used to coat nanomaterials for manufacturing semiconductors. Tri-*n*-octylphosphine (TOP) alone has been used both as solvent and stabiliser, and is required in a simple method which is suitable for the large-scale preparation of high-quality CdS nanorods [for *Trioctylphosphine as Both Solvent and Stabiliser to Synthesise CdS Nanorods* see Chen et al. *Nanoscale Res Lett* **4**(10) 1159 2009, DOI: 10.1007/s11671-009-9375-x, PMCID: PMC2893852]. **IRRITATES** skin, eyes lungs and may cause burns — work in an efficient fume cupboard and use protective clothing.

Tri(*n*-octyl)phosphine oxide (TOPO) [78-50-2] $C_{24}H_{51}OP$, **M 386.7**, **m 50-52°, 52-54°, 59.5-60°, b 201-202°/2mm, 411.2°/760mm, pK_a 8.80 in MeNO₂**. Mason, McCarty and Peppard [*J Inorg Nuclear Chem* **24** 967 1962, DOI: 10.1016/0022-1902(62)80214-0] purified the oxide by stirring a 0.1M solution in *benzene with an equal volume of 6M HCl at 40° in a sealed flask for 48 hours, then washed the *benzene solution successively with water (twice), 5% aqueous Na₂CO₃ (three times) and water (six times). The *benzene and water were then evaporated under reduced pressure at room temperature. Zingaro and White [*J Inorg Nucl Chem* **12** 315 1960, DOI: 10.1016/0022-1902(60)80378-8] treated a petroleum ether solution of the oxide with aqueous KMnO₄ (to oxidise any phosphinous acids to phosphinic acids), then with sodium oxalate, H₂SO₄ and HCl (to remove any manganese compounds). The petroleum ether solution was slurried with activated alumina (to remove phosphinic acids), filtered, evaporated and the residue was recrystallised from petroleum ether or cyclohexane at -20°. It can also be recrystallised from EtOH. [Prepared in >95% yield by oxidation of TOP with SO₃ in CH₂Cl₂; Olah et al. *J Org Chem* **48** 1670 1983, DOI: 10.1021/jo00158a034; *Beilstein* **4** IV 3466.] Alternatively, crystallisation can be performed by dissolving TOPO (400g) in dry MeCN (3L) by heating and stirring, filtering, and the colourless clear filtrate is cooled to ~20° overnight, the colourless plates are collected by filtration and dried *in vacuo* for 24 hours (yield 90-95%). {¹H}³¹P NMR (162MHz, toluene-*d*₆) has δ at 40.1. [Owen et al. *J Am Chem Soc* **130** 12279 2008, DOI: 10.1021/ja804414f]. Its applications include extraction of carboxylic acids, metals and hydrogen bonding organic compounds [review: Watson & Rickelton *Solvent Extr Ion Exch* **10** 879 1992, DOI: 10.1080/07366299208918141]. For preparing CdSe, CdS and CdTe nanomaterials see Chapter 7.

Tri(*neo*-pentyl) phosphate [14540-59-1] $C_{15}H_{33}O_4P$, **M 308.4**. Crystallise it from hexane. [See Cherbuliez in *Organo Phosphorus Compounds* (Kosolapoff & Maier eds) Wiley Vol **6** pp 211-577 1973.]

Triphenylarsine [603-32-7] $C_{18}H_{15}As$, **M 306.2**, **m 58-61°, 61°, 60-62°, b 360°/atm, 373°/760mm**. Recrystallise Ph₃As from EtOH or aqueous EtOH [Dahlinger et al. *JCS Dalton Trans* 2145 1986, DOI: 10.1039/DT9860002145; Boere et al. *J Am Chem Soc* **109** 7781 1987, DOI: 10.1021/ja00259a029]. [Shriner & Wolf *Org Synth Coll Vol* **4** 910 1963, DOI: 10.15227/orgsyn.030.0095; *Beilstein* **16** H 829, **16** I 431, **16** II 407, **16** III 921, **16** IV 1139.] Useful compound in metal coordination chemistry. **HIGHLY TOXIC**.

Triphenylarsine oxide [1153-05-5] $C_{18}H_{15}AsO$, **M 322.2**, has **m 191-193°, 194-196° (anhydrous), b 468°/atm, d 1.45 g/cm³, pK²⁵ 0.99** (from H₀ scale in H₂SO₄), **2.10** (in H₂O). The anhydrous oxide crystallises from *C₆H₆, whereas the *monohydrate* separates from EtOH with **m 118° (m 114-117° was also reported)**. [*Beilstein* **16** H 846, **16** I 433, **16** II 433, **16** III 1022, **16** IV 1176.] **HIGHLY TOXIC**.

Triphenyl borane (borane triphenyl, triphenyl boron) [960-71-4] $C_{18}H_{15}B$, M 242.1, m 134-140°, 137°, 139-141°, 142-142.5°, 145°, 147.5-148°, 151°, b 203°/15mm. Recrystallise the borane three times from Et_2O or $*C_6H_6$ under N_2 and dry it at 130°. It can be distilled in a high vacuum at 300-350° and has been distilled (b 195-215°/~15mm) in vacuum using a bath temperature of 240-330°. N_2 is introduced into the apparatus before dismantling. It forms complexes with amines. [Nielsen et al. *Chemistry and Industry* 1069 1957, Wittig et al. *Justus Liebigs Ann Chem* 563 110 1949, DOI: 10.1002/jlac.19495630113; Bent & Dorfman *J Am Chem Soc* 57 1259 1935, DOI: 10.1021/ja01310a025; *Beilstein* 16 IV 1623.]

Triphenyl phosphate [115-86-6] $C_{18}H_{15}O_4P$, M 326.3, m 49.5-50°, b 245°/11mm, 370°/atm, d_4^{25} 1.2055 g/cm³. Crystallise the phosphate from EtOH or petroleum ether (b 60-80°)/EtOH. [Cox & Westheimer *J Am Chem Soc* 80 5441 1958, DOI: 10.1021/ja01553a031; Krishnakumar & Sharma *Synthesis* 558 1983, DOI: 10.1055/s-1983-30424; Cherbuliez in *Organo Phosphorus Compounds* (Kosolapoff & Maier eds) Wiley Vol 6 pp 211-577 1973, *Beilstein* 6 III 658, 6 IV 720.] It has been used as an effective ligand with phase transfer catalysts [Nakoji et al. *J Org Chem* 67 7418 2002, DOI: 10.1021/jo0260645]. Also useful as a plastisier and a fire retardant.

Triphenylphosphine [603-35-0] $C_{18}H_{15}P$, M 262.3, m 77-78°, 79°, 79-81°, 80.5°, 80-81°, b >360°(377°, in inert gas), d_4^{25} 1.194, d_4^{80} 1.075 (liq), pK^{25} 2.73. The phosphine crystallises from hexane, MeOH, diethyl ether, CH_2Cl_2 /hexane or 95% EtOH. Dry it at 65°/<1mm over $CaSO_4$ or P_2O_5 . Chromatograph it through alumina using (4:1) *benzene/ $CHCl_3$ as eluent. [Blau & Espenson et al. *J Am Chem Soc* 108 1962 1986, DOI: 10.1021/ja00268a039; Buchanan et al. *J Am Chem Soc* 108 1537 1986, DOI: 10.1021/ja00267a025; Randolph & Wrighton *J Am Chem Soc* 108 3366 1986, DOI: 10.1021/ja00272a035; Asali et al. *J Am Chem Soc* 109 5386 1987, DOI: 10.1021/ja00252a014.] It has also been crystallised twice from petroleum ether and 5 times from Et_2O /EtOH to give crystals with m 80.5°. Alternatively, dissolve it in concentrated HCl, and upon dilution with H_2O it separates because it is weakly basic, it is then crystallised from EtOH/ Et_2O . It recrystallises unchanged from AcOH. [Forward et al. *J Chem Soc Suppl.* p121 1949, Muller et al. *J Am Chem Soc* 78 3557 1956, DOI: 10.1021/ja01596a002.] **3Ph₃P.4HCl** crystallises out when HCl gas is bubbled through an Et_2O solution, it has m 70-73°, but recrystallises very slowly and is deliquescent. **Ph₃P.HBr** [6399-81-1] has m 196° (dec). The **hydriodide**, made by adding Ph_3P to hydriodic acid, is not hygroscopic and decomposes at ~100°. The **chlorate (1:1) salt** has m 165-167°, but decomposes slowly at 100°. All salts hydrolyse in H_2O to give Ph_3P [IR, UV: Sheldon & Tyree *J Am Chem Soc* 80 2117 1958, DOI: 10.1021/ja01542a024; pK : Henderson & Streuli *J Am Chem Soc* 82 5791 1960, DOI: 10.1021/ja01507a008; Kosolapoff, *Organophosphorus Compounds*, Wiley 1950]. [*Beilstein* 16 IV 951.] § It is also available commercially on a support of polystyrene cross-linked with 2% divinylbenzene as the free base [39319-11-4], loading of ~3.2 nmol/g, 100-200 mesh], and as the **hydrobromide** [loading 2.5-3.0 mmol Br, 200-400 mesh].

Triphenylphosphine dibromide [1034-39-5] $C_{18}H_{15}Br_2P$, M 422.1, has m 235°, 245-255°(dec). The dibromide recrystallises from MeCN/ Et_2O . Although it has been recrystallised from EtOH, this is not recommended as it converts alcohols to alkyl bromides since it has brominating power. It deteriorates on keeping, and it is best to prepare it afresh. [Anderson & Frenor *J Am Chem Soc* 86 5037 1964, DOI: 10.1021/ja01076a084; Horner et al. *Justus Liebigs Ann Chem* 626 26 1959, DOI: 10.1002/jlac.19596260104; Fieser 1 1247; *Beilstein* 16 III 864.] The **diiodide** [6396-07-2] $C_{18}H_{15}I_2P$, M 516.1, has m 210-220°. [*Beilstein* 16 IV 1013.]

Triphenylphosphine oxide [791-28-6] $C_{18}H_{15}OP$, M 278.3, has m 152.0°, 156-157°, pK_{Est} ~ -2.10 (aqueous H_2SO_4), pK^{25} 2.9 (in MeNO₂). It crystallises from absolute EtOH and is dried *in vacuo*. The **gold chloride complex** has m 177.5-178.5°. [Addison & Sheldon *J Chem Soc* 2705 1956, DOI: 10.1039/JR9560002705; Cox & Westheimer *J Am Chem Soc* 80 5441 1958, DOI: 10.1021/ja01553a031; *Beilstein* 16 III 864, 16 1011.]

Triphenyl phosphite [101-02-0] $C_{18}H_{15}O_3P$, M 310.3, m 16-20°, 21-23°, b 181-189°/1mm, 183-184°/1mm, d_4^{20} 1.183. Its ethereal solution is washed successively with aqueous 5% NaOH, distilled water and saturated aqueous NaCl, then dried with Na_2SO_4 and distilled under vacuum after evaporating the diethyl ether. [Walsh *J Am Chem Soc* 81 3023 1959, DOI: 10.1021/ja01521a028; Verkade & Coskren in *Organo Phosphorus Compounds* (Kosolapoff & Maier eds) Wiley Vol 6 pp 211-577 1973, Fieser 1 1249; *Beilstein* 6 IV 695.] It has been used as an effective ligand with phase transfer catalysts [Nakoji et al. *J Org Chem* 67 7418 2002, DOI: 10.1021/jo0260645].

Triphenylphosphorylidene acetaldehyde (formylmethylenetriphenylphosphorane) [2136-75-6] $C_{20}H_{17}OP$, **M 304.3**, **m 185-187°**, **186-187°(dec)**. Recrystallise it from Me_2CO , or dissolve it in *C_6H_6 , wash with $NaOH$, dry ($MgSO_4$), evaporate, and crystallise the residue from Me_2CO . It can be prepared from its precursor, **formylmethyltriphenylphosphonium chloride** (which crystallises from $CHCl_3/EtOAc$), by treatment with Et_3N and extraction with *C_6H_6 . [Trippett & Walker *J Chem Soc* 1266 1961, DOI: 10.1039/JR9610001266] A useful Wittig reagent for the two-carbon homologation of aldehydes and ketones to form α,β -unsaturated carbonyl compounds [Wang & Pagenkopf *Org Lett* 9 3703 2007, DOI: 10.1021/ol701797e; Nagamitsu et al. *J Org Chem* 72 2744 2007, DOI: 10.1021/jo062089i].

Triphenylphosphorylidene acetonitrile (cyanomethylenetriphenylphosphorane) [16640-68-9] $C_{20}H_{16}NP$, **M 301.3**, has **m 189-195°**, **195-196°**. The phosphorane is prepared from (cyanomethyl)triphenylphosphonium chloride (**m 278-279°**, ν_{max} at 2090 cm^{-1} for $-CN$, from $ClCH_2CN$ and Ph_3P in $MeCN$) by treatment with dilute $NaOH$ in H_2O , followed by isolation and recrystallisation from $EtOAc$. It has ν_{max} at 2174 cm^{-1} for $-C=C-CN$. Like the preceding phosphorane it is a useful Wittig reagent for the two-carbon homologation of aldehydes and some ketones to form α,β -unsaturated nitriles [Trippett & Walker *J Chem Soc* 3874 1959, DOI: 10.1039/JR9590003874; cf. Ohno et al. *J Org Chem* 72 4378 2007, DOI: 10.1021/jo0700528].

(Triphenylphosphoranylidene)ketene [Bestmann ylide (triphenylphosphoranylidene)ethenone, ketenylidene(triphenyl)phosphorane] [15596-07-3] $C_{20}H_{15}OP$, **M 302.3**, **m 167-173°**, **173°**. This versatile reagent is prepared in a dry inert atmosphere (N_2 or argon) by first mixing $NaNH_2$ (19.5g, 0.5mole) in toluene (1.3L, dried by passage through alumina) and bis(trimethylsilyl)amine (81g, 105ml, 0.5mol, toxic and corrosive), and heated ($70-80^\circ$ bath temperature) under a reflux condenser until a clear and colourless solution of sodium hexamethyldisilazane (see hexamethyl disilazane sodium salt [1070-89-9]) is obtained, and evolution of NH_3 ceased (2-4 hours). **Note** that repeated purging with argon or N_2 reduces the reaction time. The condenser is removed and under argon or N_2 , (methoxycarbonylmethylene)triphenylphosphorane (167g, 0.5mol, [2605-67-6]) is added in large portions, and the mixture is again heated ($70-80^\circ$ bath temperature) with stirring for 24-30 hours until the colour of the solution assumes a bright yellow hue [and the IR band of a film of aliquots between KBr plates, with ν_{max} at 1616 cm^{-1} of the starting ylide has disappeared, and is replaced by a strong band of the product with ν_{max} at 2090 cm^{-1} of constant intensity, indicating completion of reaction]. While still hot, and under argon or N_2 , the mixture is filtered rapidly [through a 20/40 standard taper vacuum filtration adapter and a 350ml fritted funnel of coarse porosity open to the atmosphere through a 2 cm thick layer of basic Al_2O_3 (III) and a 1cm Celite pad on top, and occasional scraping the top of the pad] into a 2L 24/40 single necked flask. The filtration requires 30-60 minutes and the funnel temperature should be kept between $30-50^\circ$ to avoid crystallisation. The filtrate is flushed with inert gas and evaporated (rotavap) to dryness. The residue is recrystallised from hot dry toluene (1g/5ml) followed by cooling at -20° overnight. While the mixture is cold, the solids are filtered off, washed with cold toluene (3 x 100ml at 0°). The combined filtrates are evaporated in the same way at low temperature as before to give a second crop. The combined crops are dried to constant weight at high vacuum (0.01mm) to give pure **triphenylphosphoranylidene)ketene** (106.8-114.9g, 74-76% and ~99.97% pure by 1H NMR) as a pale yellow, flaky powder with **m 173°**. It is remarkably stable, it can be stored at $\sim 25^\circ$ under argon or N_2 for months and can be handled at $\sim 25^\circ$ without obvious deterioration. It has IR (KBr) with ν_{max} at 2090 (s), 1625 (m), 1436 (m) and 1110 (m) cm^{-1} ; the 1H NMR (400MHz, $CDCl_3$) has δ at 7.44-7.54 (m, 6H), 7.55-7.63 (m, 3H) and 7.64-7.77 (m, 6H); the ^{13}C NMR (100MHz, $CDCl_3$) has δ at -10.5 (d, $^3J_{P-C} = 185.4\text{ Hz}$, C^α), 128.8 (d, $^3J_{P-C} = 12.9\text{ Hz}$, C-ortho), 129.6 (d, $^1J_{P-C} = 98.5\text{ Hz}$, C-ipso), 132.2 (s, C-para), 132.3 (s, C-meta) and 145.6 (d, $^2J_{P-C} = 63.0\text{ Hz}$, C^β); and the ^{31}P NMR (162MHz, $CDCl_3$) has δ at 6.0. Possible detectable impurities are starting ester ylide and toluene detected by 1H NMR spectroscopy which have singlets at δ 3.60 and 2.36 respectively. It is a nucleophilic-only C_2 -building block that reacts with a range of electrophiles in a variety of ways. It also undergoes cycloaddition reactions at the polar $P-C^\alpha$ bond as well as at the $C^\alpha=C^\beta$ double bond, and multi-component and domino reactions [Schobert *Org Synth* 82 140 2005, DOI: 10.1522/orgsyn.082.0140.]

Triphenyl silane [789-25-3] $C_{18}H_{16}Si$, **M 260.4**, **m 43-45°**, **45°**, **b 148-151°/1mm**, **152°/2mm**. Purify it by recrystallisation from $MeOH$. [Gilman & Zuech *J Am Chem Soc* 81 5925 1959, DOI: 10.1021/ja01531a021; Westermarck *Acta Chem Scand* 9 947 1955, DOI: 10.3891/acta.chem.scand.09-0947; IR: Kaplan *J Am Chem Soc* 76 5880 1954, DOI: 10.1021/ja01651a105; Beilstein 16 II 605, 16 III 1199, 16 IV 1369.]

Triphenylsilanol (hydroxytriphenylsilane) [791-31-1] $C_{28}H_{16}OSi$, M 276.4, m 150-153°, 151-153°, 154-155°, 156°, b 389°/760mm. It is purified by dissolving in petroleum ether, passing through an Al_2O_3 column, eluting thoroughly with CCl_4 to remove impurities and then eluting the silanol with MeOH. Evaporation gives crystals with m 153-155°. It can be recrystallised from petroleum ether, CCl_4 or from *benzene or Et_2O /petroleum ether (1:1). It has also been recrystallised by partial freezing from the melt to constant melting point. [George & Gilman *J Am Chem Soc* **81** 3288 1959, DOI: 10.1021/ja01522a034; IR: Tatlock & Rochow *J Org Chem* **17** 1555 1952, DOI: 10.1021/jo50012a001 and Richards & Thompson *J Chem Soc* 124 1949, DOI: 10.1039/JR9490000124; *Beilstein* **16** IV 1480.]

Triphenyl vinyl silane [18666-68-7] $C_{18}H_{15}Si$, M 286.5, m 58-59°, 57-59.5°, 67-68°, 69°, 68-71°, b 190-210°/3mm. The vinyl-silane has been recrystallised from EtOH, 95% EtOH, $EtOH/*C_6H_6$, petroleum ether (b 30-60°) and Et_2O , and has been distilled under reduced pressure. [Cason & Brooks *J Am Chem Soc* **74** 4582 1952, DOI: 10.1021/ja01138a039; Nagel & Post *J Org Chem* **17** 1379 1952, DOI: 10.1021/jo50010a016; *Beilstein* **16** IV 1371.]

Tri(*n*-propyl) borate [688-71-1] $C_9H_{21}BO_3$, M 188.1, b 64°/9mm, 175-177°/atm, d_4^{20} 0.857, n_D^{20} 1.395. Dry the ester over sodium and then distil it, preferably in a vacuum. (cf. tributyl borate.) [Charnley et al. *J Chem Soc* 2288 1952, DOI: 10.1039/JR9520002288; *Beilstein* **1** IV 1436.]

Tri(*iso*-propyl) borate [(*iso*-PrO)₃B, boric acid tri-*iso*-propyl ester, Boron *iso*-propoxide] [5419-55-6] $C_9H_{21}BO_3$, M 188.1, has m -59°, b 75°/76mm, 90°/120mm, 139-141°/760mm, d_4^{25} 0.815, n_D^{20} 1.376. The borate ester is prepared in 85% yield from *iso*-propanol (250ml) and $NaBH_4$ (6.84g) followed by dropwise addition of AcOH (0.18g) over a period of 11 minutes, and then refluxing for 4 hours (fume cupboard and a 'Dry-ice' condenser as 16.15L of H_2 are released, CARE due to its flammability). Then fractionate the *ester* through a Widmer column. [Brown et al. *J Am Chem Soc* **78** 3613 1956, DOI: 10.1021/ja01596a015.] It is moisture sensitive. The ester is a good reagent for borylation, e.g. *ortho*-borylating 1-substituted naphthalenes, by reacting with 1-halo- or cyano- naphthalenes (after treatment with LiTMP) to form 2-boryl esters which are later used in Pd-catalysed cross-coupling reactions [Lysén et al. *Synthesis* 3478 2006, DOI: 10.1055/s-2006-950239]. [*Beilstein* **1** H 363, **1** II 382, **1** III 1468, **1** IV 1488.]

Tri(quinol-8-yl) phosphate [52429-99-9] $C_{27}H_{18}N_3O_4P$, M 479.4, m 193-197°, 202-203°, 638°/760mm. Purify the phosphate by recrystallisation from dimethylformamide. The purity is checked by paper chromatography, R_F 0.90 [*i*-PrOH/saturated $(NH_4)_2SO_4/H_2O$, 2:79:19 as eluent], its IR (KBr) has ν_{max} at 1620, 1570 (C=C, C=N) and 1253 (P=O) cm^{-1} . [Takaku et al. *Bull Chem Soc Jpn* **47** 779 1974, DOI: 10.1246/bcsj.47.779.]

Tri-*o*-tolylphosphine [6163-58-2, 1038-95-5] $C_{21}H_{21}P$, M 304.4, m 123-125°, 125-126°, 127-129°, 129-130°, pK_{Est} ~1.0. Like PPh_3 , it crystallises from hexane, MeOH, diethyl ether, CH_2Cl_2 /hexane or 95% EtOH. Dry it at 65°/1mm over $CaSO_4$ or P_2O_5 . Purify further by chromatography through alumina using (4:1) *benzene/ $CHCl_3$ as eluent. Of the phosphine ligands studied, $P(o-Tol)_3$ (6 mol%) was the best in the $Ru_3(CO)_{12}$ (2 mol%) catalysed *N*-alkylation of primary amines and secondary alcohols (e.g. hexylamine with 1-phenylethanol to give *N*-octyl 1-phenethylamine) in high yields at 90-110° [Tillack et al. *Tetrahedron Lett* **47** 8881 2006, DOI: 10.1016/j.tetlet.2006.10.042]. The Heck reaction between aryl halides and *n*-butyl acrylates to provide substituted styrenes in ~80 to >90% yields at ~120° is catalysed by $Pd(OAc)_2$ (2 mol%) and $P(o-Tol)_3$ (8 mol%) without extensive P-C bond cleavage [Herrmann et al. *J Mol Catal A: Chem* **103** 133 1995, DOI: 10.1016/1381-1169(95)00140-9]. [*Beilstein* **16** III 835.]

Tris(2-biphenyl) phosphate [132-28-5] $C_{36}H_{27}O_4P$, M 554.6, m 115.5-117.5°. Crystallise it from MeOH containing a little acetone. (cf. triphenyl phosphate.)

Tris(2,4-di-*tert*-butylphenyl)phosphite [31570-04-4] $C_{42}H_{63}O_3P$, M 646.9, has m 181-184°, 187°. If the ester is suspect, e.g. partly hydrolysed, dissolve it in Et_2O , wash the solution successively with aqueous 5% NaOH, H_2O and saturated aqueous NaCl, then shake with charcoal, filter, dry with Na_2SO_4 , filter again, evaporate and add petroleum ether when the volume has decreased considerably to crystallise the ester. [Verkade & Coskren in *Organophosphorus Compounds* (Kosolapoff & Maier eds) Wiley Vol 6 pp 211-577 1973.] It is used as a processing stabiliser for polymers.

Tris(1,2-dioxyphenyl)cyclotriphosphazene {trispiro[1,3,5,2,4,6-triazatriphosphorine]-2,2':-2,4'':2,6'''-tris(1,3,2)benzodioxaphosphole} [311-03-5] $\text{C}_{18}\text{H}_{12}\text{O}_6\text{N}_3\text{P}_3$, **M 459.0**, **m 244-245°**, **245°**, **245-246°**. Recrystallise this phosphazine from C_6H_6 or chlorobenzene, then triple sublime it (175°/0.1mm, 200°/0.1mm, 230°/0.05mm). Its UV has λ_{max} nm (log ϵ) at 276 (3.72), 271 (3.79) 266sh (3.68) and 209 (4.38) in MeCN. Its IR has ν_{max} at 1270 (O-Ph), 1220 (P=N), 835 (P-O-Ph) and 745 (Ph) cm^{-1} . [Alcock *J Am Chem Soc* **86** 2591 1964, DOI: 10.1021/ja01067a015; for cyclophosphazene clathrates see Alcock et al. *J Am Chem Soc* **98** 5120 1976, DOI: 10.1021/ja00433a012; for mobility in a solid benzene-cyclophosphazene inclusion compound see Meirovitch et al. *J Phys Chem* **88** 1522 1984, DOI: 10.1021/j150652a016.]

Tris(2-ethylhexyl) phosphate (TEHP, tri-isooctylphosphate, 'trioctyl' phosphate. [78-42-2, 25103-23-5] $\text{C}_{24}\text{H}_{51}\text{O}_4\text{P}$, **M 434.6**, **m -75°**, **-74°**, **b 186°/1mm**, **215°/4mm**, **219°/5mm**, **450°/760mm**, **d_{25}^{20} 0.92042**, **n_D^{20} 1.44464**. TEHP, in an equal volume of diethyl ether, is shaken with aqueous 5% HCl, and the organic phase is filtered to remove traces of pyridine (used as a solvent during manufacture) as its hydrochloride. This layer is shaken with aqueous Na_2CO_3 , then water, and the ether is distilled off at room temperature. The ester is then filtered, dried for 12 hours at 100°/15mm, and again filtered, then shaken intermittently for 2 days with activated alumina (100g/L). It is decanted through a fine sintered-glass disc (with exclusion of moisture), and distilled under vacuum. [French & Muggleton *J Chem Soc* 5064 1957, DOI: 10.1039/JR9570005064.] *Benzene can be used as a solvent (to give 0.4M solution) instead of ether. Its IR has ν_{max} at 1702, 1701, 481 and 478 cm^{-1} [Bellamy & Becker *J Chem Soc* 475 1952, DOI: 10.1039/JR9520000475]. The **uranyl nitrate** salt is purified by crystallisation from hexane [Siddall & Dukes *J Am Chem Soc* **81** 790 1959, DOI: 10.1021/ja01513a007; Siddall *J Am Chem Soc* **81** 4176 1959, DOI: 10.1021/ja01525a015]. [Beilstein **1** IV 1786.]

Tris(hydroxymethyl)phosphine (phosphinidynetrimethanol) [2767-80-8] $\text{C}_3\text{H}_9\text{O}_3\text{P}$, **M 124.1**, **m 48-56°**, **55°**, **b 111-113°/2.5mm**, **311.7°/760mm**. **n_D^{20} 1.5497**, **pK 5.5 (half neutralisation point, glass electrode)**. It is obtained in two ways by neutralising precisely with THPC {*tetrakis(hydroxymethyl)phosphonium chloride*, [124-64-1]}. In the *first* method, NaOH (1.6g, 40mmol) in H_2O (25ml) is added rapidly to THPC (7.6g, 40mmol, see above [124-64-1]) in H_2O (50ml) under N_2 at room temperature in a closed system connected to a glass buret. No evolution of gas should be observed on stirring for 20 hours indicating that the stoichiometric amount of alkali has been added. Any liberation of gas (H_2) would indicate that excess of alkali had been added which would liberate H_2 with the formation of the corresponding phosphine oxide (see next entry, and Hoffman *J Am Chem Soc* **52** 2995 1930, DOI: 10.1021/ja01370a065). The solvent is then removed *in vacuo* at 50-60°, EtOH (under N_2) is added, the solid NaCl is filtered off, and the EtOH is removed *in vacuo* (45°) to give the **free base** (4.8g, 97%) as a clear viscous liquid which can be distilled at high vacuum in an atmosphere of N_2 (N_2 bleed). An IR (film) should indicate the absence of P=O and the presence of primary OH and P- CH_2 bands [Grayson *J Am Chem Soc* **85** 79 1963, DOI: 10.1021/ja00884a016].

In the *second* method, under N_2 flushing and stirring, a solution of THPC (25g, 131mmol) in distilled H_2O (100ml), into which is immersed a glass pH electrode, is treated with ~ 150ml (0.21 equivalents) of 20-50 mesh Dowex-1 x 8 resin (in the OH^- form) during 45 minutes until the pH was ~8.3-8.5. The resin is filtered off, washed with H_2O , the combined aqueous solutions are evaporated (rotavap) at 65-70°, then under high vacuum for 30 minutes, then further at -30° to give the **free base** as a viscous liquid which formed a waxy solid. Its IR (film between NaCl plates) should have no bands at ν_{max} 1052 (of THPC), or 1043 or 1135 (for P=O), but intense broad absorption at 1010 cm^{-1} . The ^1H NMR (60MHz, D_2O , internal TMS, or DSS at 0) has δ for CH_2 at 4.15 ($J_{\text{PCH}} = 5.5\text{Hz}$, $J_{^{13}\text{CH}} = \sim 148\text{Hz}$); and the ^{31}P NMR (40MHz, D_2O , external 85% H_3PO_4) has δ at -25.8. [Ellzey et al. *J Org Chem* **37** 3453 1972, DOI: 10.1021/jo00795a015.] With *n*-butyl iodide it formed ***n*-butyl phosphonium iodide** which gave, on treatment with sodium tetraphenylboron, the ***n*-butyl tris(hydroxymethyl)phosphonium tetraphenylboron complex**, **m 145-146°(dec)** after drying and recrystallising from $\text{Me}_2\text{CO}/\text{C}_6\text{H}_6$; and the **methiodide** gave with Ph_4BNa , the **methyl tris(hydroxymethyl)phosphonium tetraphenylboron complex**, **m 170-171°(dec)** after drying and recrystallising from $\text{Me}_2\text{CO}/\text{C}_6\text{H}_6$ [Grayson *J Am Chem Soc* **85** 79 1963, DOI: 10.1021/ja00884a016.]

Tris(hydroxymethyl)phosphine oxide [1067-12-5] $\text{C}_3\text{H}_9\text{O}_4\text{P}$, **M 137.1**, has **m 50-52°**, **54-55° (69° and 70° have also been reported)**. The oxide is prepared by slowly adding about 130% excess (over an equimolar amount) of 20-50 mesh Dowex-1 x 8 resin (in the OH^- form) to an aqueous solution of tris(hydroxymethyl)phosphine (25.0g, see preceding entry) which led to vigorous evolution of H_2 . After standing overnight the resin is filtered off, and the filtrate is evaporated (rotavap, vacuum) to give the **oxide**

(16.1g, 88%) which is recrystallised from absolute EtOH to yield hygroscopic crystals (11.9g). Store in a dry atmosphere. The IR has strong bands at ν_{\max} 1043 and 1134 cm^{-1} ; the ^1H NMR (60MHz, D_2O , internal TMS, or DSS at 0) has δ for CH_2 at 4.20 ($J_{\text{PCH}} = 3.1\text{Hz}$, $J_{^{13}\text{CH}} = \sim 146\text{Hz}$); and the ^{31}P NMR (40MHz, D_2O , external 85% H_3PO_4) has δ at -48.7. [Ellzey et al. *J Org Chem* **37** 3453 1972, DOI: 10.1021/jo00795a015; Anteunis et al. *Bull Soc Chim Belg* **74** 622 1965, DOI: 10.1002/bscb.19650741114.] The **tribenzoate**, prepared by boiling THPC with excess of aqueous NaOH solution until no further evolution of H_2 occurred followed by treatment with a slight excess of PhCOCl , gave fine needles **m 111°** upon recrystallisation from MeOH [Hoffman *J Am Chem Soc* **43** 1684 1921, DOI: 10.1021/ja01440a035].

Trisodium 8-hydroxy-1,3,6-pyrenetrisulfonate (Pyranine Solvent Green 7) [6358-69-6] $\text{C}_{16}\text{H}_7\text{Na}_3\text{O}_{10}\text{S}_3$, **M 523.4**, **m >300(dec)**, **CI 59040**, λ_{\max} **403nm**. Purify the salt by chromatography through an alumina column, and elute with *n*-propanol/water (3:1, v/v). Recrystallise it from aqueous acetone (5:95, v/v) using decolorising charcoal. [Beilstein **1** III 565.] **IRRITANT**.

Trisodium 1,3,6-naphthalenetrisulfonate [5182-30-9] $\text{C}_{10}\text{H}_5\text{Na}_3\text{O}_9\text{S}_3$, **M 434.3**. The **free acid** is obtained by passing the salt through an ion-exchange column and converting it to the lanthanum salt by treatment with La_2O_3 . This salt is crystallised twice from hot water. [The much lower solubility of $\text{La}_2(\text{SO}_4)_3$ and its retrograde temperature dependence allows a good separation from sulfate impurity]. The lanthanum salt is then passed through an appropriate ion-exchange column to obtain the free acid, the sodium or potassium salt. (The sodium salt is **hygroscopic**.) [Atkinson et al. *J Am Chem Soc* **83** 1570 1961, DOI: 10.1021/ja01468a008.] It can also be recrystallised from aqueous acetone [Okahata et al. *J Am Chem Soc* **108** 2863 1986, DOI: 10.1021/ja00271a013].

Tris(2,2,2-trifluoroethyl) phosphite [370-69-4] $\text{C}_6\text{H}_6\text{F}_9\text{O}_3\text{P}$, **M 328.1**, **b 130-131°/743mm**, **128.8±40°/760mm**, d_4^{20} **1.487**, n_D^{20} **1.324**. Fractionate the phosphite through a 10inch Helipak column [Krogh et al. *J Org Chem* **19** 1124 1954, DOI: 10.1021/jo01372a018]. [Beilstein **1** IV 1371.]

Tris(trimethylsilyl)silane (TTMSS) [1873-77-4] $\text{C}_3\text{H}_{10}\text{Si}_4$, **M 248.7**, **b 73°/5mm**, **82-84°/12mm**, d_4^{20} **0.808**, n_D^{20} **1.4900**. Purify it by fractional distillation and taking the middle cut. Store it under N_2 or argon as it is **PYROPHORIC** and is an **IRRITANT**. [Chatgililoglu et al. *J Org Chem* **53** 3641 1988, DOI: 10.1021/jo00250a051; Chatgililoglu *Acc Chem Res* **25** 188 1992, DOI: 10.1021/ar00016a003; NMR: Gilman et al. *J Organomet Chem* **4** 163 1965, DOI: 10.1016/S0022-328X(00)84384-3; Dickhaut & Giese *Org Synth* 141 2003, DOI: 10.1002/0471264180.os070.19] It is a radical-based reducing agent as in the dehydrohalogenation reactions of halides to the respective hydrocarbons [Balestri et al. *J Org Chem* **56** 678 1991, DOI: 10.1002/0471264180.os070.19; for a review see Boxer et al. *Aldrichimica Acta* **42** 3 2009].

Tri(*p*-tolyl) phosphate [78-32-0, 20756-92-7, 1330-78-5 (isomeric tritolyl phosphate mixture)] $\text{C}_{21}\text{H}_{21}\text{O}_4\text{P}$, **M 368.4**, **m 77°**, **74-78°**, **b 232-234°/atm**, d^{25} **1.16484**, n_D^{20} **1.56703**. Dry the ester with CaCl_2 , percolate it through a column of alumina, then distil it under a vacuum. Alternatively, pass it through a packed column of alumina at 150°, with a counter-current stream of nitrogen, under reduced pressure, to remove residual traces of volatile impurities. It also crystallises from petroleum ether (b 60-80°). [Cherbuliez in *Organic Phosphorus Compounds* (Kosolapoff & Maier eds) Wiley-Interscience Vol **6** pp 454-457 1973.]

Tropaeolin 00 {4-[(4-anilinophenyl)azo]benzenesulfonic acid Na salt} [554-73-4] $\text{C}_{18}\text{H}_{14}\text{N}_3\text{NaO}_3\text{S}$, **M 375.4**, $\text{pK}_{\text{Est}(1)} \sim 2.3$, $\text{pK}_{\text{Est}(2)} \sim 5.8$, $\text{pK}_{\text{Est}(3)} \sim 10.3$. Recrystallise it twice from water [Kolthoff & Gus *J Am Chem Soc* **60** 2516 1938, DOI: 10.1021/ja01277a068]. It is an indicator which is red at pH 1.4 and orange-yellow at pH 3.2. [Beilstein **16** II 171.]

Vinyl chlorosilane [75-94-5] $\text{C}_2\text{H}_3\text{Cl}_3\text{Si}_4$, **M 161.5**, **m -95°**, **b 17.7°/46.3mm**, **82.9°/599.4mm**, **92°/742mm**, **91-91.5°/atm**, d_4^{20} **0.12717**, n_D^{20} **1.435**. Fractionally distil the chlorosilane at atmospheric pressure. It is water sensitive and is stored in the dark and it is likely to polymerise. [Müller & Schnurrbusch *Chem Ber* **91** 1805 1958, DOI: 10.1002/cber.19580910903; Munkelt & Müller *Chem Ber* **92** 1012 1959, DOI: 10.1002/cber.19590920503; Polarography: Abrahamson & Reynolds *Anal Chem* **24** 1827 1952,

DOI: 10.1021/ac60071a033; *Beilstein* **4** IV 4258.]

3-Vinylphenylboronic acid ([3-ethenylphenyl]-dihydroxyborane) [15016-43-0] $\text{C}_8\text{H}_9\text{BO}_2$, **M 148.0**, **m 141-147°**, **145-160°**, $\text{pK}_{\text{Est}(1)} \sim 8.8$. This binder for paper or glass crystallises in white plates from H_2O (**m 144-145°**), and its IR has ν_{max} at 919 and 995 ($\text{CH}_2\text{CH}=\text{CH}_2$), 1350 (B-O) and 3220 (OH) cm^{-1} . The *dibromo derivative* crystallises in white needles from H_2O with **m 196-197.5°**. [Dale & Rush *J Org Chem* **27** 2598 1962, DOI: 10.1021/jo01054a071; *Beilstein* **16** III 1279, **16** IV 1677.]

Vinylphosphonic acid [1746-03-8] $\text{C}_2\text{H}_5\text{O}_3\text{P}$, **M 108.0**, **m 41-45°**, $d_{25}^{25} 1.389$, $\text{pK}_1^{20} 3.48$, $\text{pK}_2^{20} 8.54$ (50% EtOH). This fireproofing agent, and ingredient for making polymers, is obtained as a syrup on hydrolysing vinylphosphonyl dichloride with cold H_2O and solidifies on prolonged drying over $\text{P}_2\text{O}_5/\text{KOH}$. When distilled at 235-240°/0.0006mm, it gives the *anhydride* ($d_4^{20} 1.304$, $n_D^{20} 1.5874$) [Kabachnik & Medvedi *Izvest Akad Nauk SSSR, Ser Khim* 868 1953, *Chem Abstr* **54** 10834 1960]. It is best kept as the *sodium salt* (**m 350°**) which precipitates when a solution of EtOH containing NaOEt (from 2g of Na) is added to vinylphosphonic acid (3.2g), and is recrystallised from EtOH (5.1g, quantitative). The *p-anisidinium salt* forms mauve prisms **m 250°** (from EtOH/Et₂O). The *dimethyl ester*, [4645-32-3] **M 136.1**, $d_4^{20} 1.1405$, $n_D^{20} 1.4330$, has **b 72.5°/10mm** and **197-202°/760mm**. [Kabachnik et al. *J Gen Chem USSR* (Engl Trans) **33** 375 1963, Svara et al. *Ullmann's Encyclopedia of Industrial Chemistry*, 2008 on line, Wiley-VCH, DOI: 10.1002/14356007.a19_545.pub2; *Beilstein* **4** IV 3560.]

Xylenol Orange {3*H*-2,1-benzoxathiol-3-ylidene-bis-[(6-hydroxy-5-methyl-*m*-phenylene)-methyl-nitrilo]-tetraacetic acid, S,S-dioxide} [1611-35-4] $\text{C}_{31}\text{H}_{32}\text{N}_2\text{O}_{13}\text{S}$, **M 672.6**, **m 210°(dec)**, $\epsilon_{578} 6.09 \times 10^4$ (pH 14), $\epsilon_{435} 2.62 \times 10^4$ (pH 3.1), $\text{pK}_1 -1.74$, $\text{pK}_2 -1.09$ (aqueous $\text{H}_2\text{SO}_4\text{-HNO}_3$), $\text{pK}_3 2.58$, $\text{pK}_4 3.23$, $\text{pK}_5 6.46$, $\text{pK}_6 10.46$, $\text{pK}_7 12.28$. It is generally contaminated with starting material (cresol red) and semi-xylenol orange. Purify it by ion-exchange chromatography using DEAE-cellulose, eluting with 0.1M NaCl solution, which will give the sodium salt [3618-43-7], Cresol Red, semi-xylenol orange and iminodiacetic acid bands elute first. This method will give the *sodium salt* [3618-43-7] $\text{C}_{31}\text{H}_{28}\text{N}_2\text{Na}_4\text{O}_{13}\text{S}$, **m 195°(dec)**, of the dye. To obtain the *free acid*, dissolve the salt in H_2O and acidify it with AcOH. Filter it off, wash it with H_2O and dry it first in air and then in a vacuum desiccator over P_2O_5 in the dark [Sato et al. *Anal Chim Acta* **94** 217 1977, DOI:10.1016/S0003-2670(01)83654-0]. Useful indicator for metal ion titrations, also as a metallochromic ligand employed in chelation ion chromatography of trace metal ions [Paull & Haddad *Trends Analyt Chem* **18** 107 1999, DOI:10.1016/S0165-9936(98)00104-6; Gay et al. *Analyt Biochem* **273** 142 1999, DOI: 10.1006/abio.1999.4207]. [*Beilstein* **19** II 111, **19** III/IV 1135, **19/3** V 461.]

Zinc diethyldithiocarbamate [14324-55-1] $\text{C}_{10}\text{H}_{20}\text{N}_2\text{S}_4\text{Zn}$, **M 361.9**, **m 172-183°**, **178-181°**, $d 1.462$, $\text{pK}^{25} 3.04$ (for $\text{Et}_2\text{NCS}_2^-$). Crystallise this herbicide several times from hot toluene or from hot CHCl_3 by addition of EtOH. It also crystallises from xylene, **m 180°**. [*Beilstein* **4** II 613.] **TOXIC**.

Zinc dimethyldithiocarbamate (Ziram) [137-30-4] $\text{C}_6\text{H}_{12}\text{N}_2\text{S}_4\text{Zn}$, **M 305.8**, has **m 151-152°**, **248-250°**, $\text{pK}^{25} 3.36$ (for $\text{Me}_2\text{NCS}_2^-$ in H_2O), $\text{pK}^{20} 1.23$ (octanol/ H_2O). Crystallise this herbicide several times from hot toluene or from hot CHCl_3 by addition of EtOH. It has low solubility (65mg/L) in H_2O at 25°. [*Beilstein* **4** III 149, **4** IV 234.]

Zinc ethylenebis(dithiocarbamate) (Zineb) [142-14-3 monomer, 9006-42-2 polymer, 12122-67-7] $\text{C}_4\text{H}_8\text{N}_2\text{S}_4\text{Zn}$, **M 275.7**. Crystallise this herbicide several times from hot toluene or from hot CHCl_3 by addition of EtOH. It is a skin irritant. [*Beilstein* **4** III 149, **4** IV 234.]

Zinc phenol-*o*-sulfonate (8H₂O) (Phenozin) [1300-55-6, 127-82-2] $\text{C}_{12}\text{H}_{10}\text{O}_8\text{S}_2\text{Zn}$, **M 411.7**. Phenozin crystallises from warm water by cooling to 0°. It effloresces in dry air, and loses all its H_2O at 120°. Its solubility in H_2O is 63% at 20° and 250% at boiling point, and in EtOH it is 55% at 50°. [*Beilstein* **11** IV 574.]

Zinc trifluoromethanesulfonate (zinc triflate) [54010-75-2] $(\text{CF}_3\text{SO}_3)_2\text{Zn}$, **M 363.5**, **m >300°**. This zinc salt should be dried at 125° for 2 hours at 3mm before use. It is soluble in CH_2Cl_2 but insoluble in petroleum ether. It is a good catalyst for the preparation of dithioketals [Corey & Shimoji *Tetrahedron Lett* **24** 169 1983, DOI: 10.1016/S0040-4039(00)81357-X], and is a Lewis acid catalyst in silylations [Jiang & Zhu (2005). *Tetrahedron Letters* **46** 517 2005, DOI: 10.1016/j.tetlet.2004.10.175]. [*Beilstein* **16** H 392.]

CHAPTER 4

PURIFICATION OF INORGANIC AND METAL-ORGANIC CHEMICALS

INTRODUCTION

The most common method of purification of inorganic species is by recrystallisation, usually from water. However, especially with salts of weak acids or of cations other than the alkaline and alkaline earth metals, care must be taken to minimise hydrolysis. This can be achieved, for example, by recrystallising acetates in the presence of dilute acetic acid. Nevertheless, there are many inorganic chemicals that are too insoluble or are hydrolysed by water so that no general purification method can be given. It is convenient that many inorganic substances have large temperature-solubility coefficients for their solubility in water, i.e. large amounts dissolve in boiling water and on cooling to <25° large amounts of crystals separate. In cases where the temperature-solubility coefficients are small, crystallisation can be achieved by slow partial solvent evaporation.

Organo-metallic compounds, on the other hand, behave very much like organic compounds, e.g. they can be redistilled and may be soluble in organic solvents. A note of **caution** should be made about handling organo-metallic compounds, e.g. arsines, because of their **potential toxicities**, particularly when they are volatile. Generally the suppliers of such compounds provide details about their safe manipulation. These should be read carefully and adhered to closely. If in any doubt, always assume that the materials are lethal and treat them with utmost care. The same **safety precautions** about the handling of substances as stated in Chapter 3 should be followed here (see Chapter 1).

For information on **ionization (pK)** see Chapter 1, and Chapter 3. In order to avoid repetition, the literature (or predicted) pK values of anionic and/or cationic species are usually reported at least once, and in several cases are entered for the free acid or free base; e.g. Na₂SO₄ will have a pK value for Na⁺ at the entry for NaOH and the pK values for SO₄²⁻ at the entries for H₂SO₄. When the pK values of the organic counter-ions are not given in this chapter, as in case of sodium benzoate, the reader is referred to the value(s) in Chapter 3, 'Aromatic Compounds', e.g. of benzoic acid.

Abbreviations of titles of periodicals are defined as in the Chemical Abstracts Service Source Index (CASSI). References to Fieser & Fieser's *Reagents for Organic Synthesis* will be shortened to Fieser throughout, e.g. Fieser **2** 254, **11** 88, etc. All temperatures are in degrees Centigrade unless otherwise stated. Other abbreviations are self evident.

Benzene, which has been used as a solvent successfully and extensively in the past for reactions and purification by chromatography and crystallisation, is now considered a **very dangerous substance**, so it has to be used with extreme care. It is important that an alternative solvent to benzene (e.g. toluene, toluene/petroleum ether, or a petroleum ether to name a few) should be used first. However, if benzene has to be used, then all operations have to be performed in well-ventilated fumehoods and precautions taken to avoid inhalation and contact with skin and eyes. Whenever benzene is mentioned in the text, an asterisk e.g. *C₆H₆ or *benzene, is inserted to remind the user that special precaution should be adopted.

Organic dyes which are *not* complexed or are salts of metals are included in 'Miscellaneous As, B, P.....' in Chapter 3 (use the CAS Registry Numbers to find them). Commercially available polymer-supported reagents are indicated with § under the appropriate reagent.

This chapter is subdivided into two sections: the **Purification of Inorganic Compounds** and the **Purification of Metal-Organic Compounds** which includes ammonium and metal salts of organic acids.

INORGANIC COMPOUNDS

Alumina (aluminium oxide) (neutral) [1344-28-1] Al_2O_3 , **M 102.0 (anhydrous)**. Stir the oxide with hot 2M HNO_3 , either on a steam bath for 12 hours (changing the acid every hour) or three times for 30 minutes, then wash it with hot distilled water until the washings have pH 4, and follow by three washings with hot MeOH. The product is dried at 270° [Angyal & Young *J Am Chem Soc* **81** 5251 1959, DOI: 10.1021/ja01528a055]. For the preparation of alumina for chromatography see Chapter 1. [For α , β and γ Al_2O_3 see Becher in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 822-823 1963 and Wagner in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1662 1965, for chromatography see Fieser **1** 19 and in later volumes.] Also commercially available in various physical forms and dryness.

Aluminium ammonium sulfate dodecahydrate [7784-26-1] $\text{AlNH}_4(\text{SO}_4)_2 \cdot 12\text{H}_2\text{O}$, **M 453.3, m 93°, m 95°, pK_1^{25} 4.89, pK_2^{25} 5.43, pK_3^{25} 5.86 (Al^{3+} aquo), pK_4^{25} 11.22 [aluminate $\text{Al}(\text{OH})_4^-$** . Crystallise it from hot H_2O and cool in ice. When the melt is heated, it loses NH_3 and H_2SO_4 , and gives pure alumina at red heat. Solubility (%) in H_2O is 3.9% (0°), 15.0 (20°) and 135 (100°). Crystals for X-ray studies were easily grown in aqueous solution and were ground into spheres about 0.15mm in diameter [crystal structure: Larson & Cromer *Acta Cryst* **22** 793 1967, DOI: 10.1107/S0365110X67001586].

Aluminium bromide [7727-15-3] AlBr_3 , **M 266.7, m 97°, b 114°/10mm, d_4^{18} 3.205**. Reflux it and then distil it from pure aluminium chips in a stream of nitrogen into a flask containing more of the chips. It is then redistilled under vacuum into ampoules [Tipper & Walker *J Chem Soc* 1352 1959, DOI: 10.1039/JR9590001352]. Anhydrous conditions are essential, and the white to very light brown solid distillate can be broken into lumps in a dry-box (under nitrogen). It fumes in moist air. [Becher in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 812-813 1963, Fieser **1** 22 and in 7 later volumes.]

Aluminium caesium sulfate dodecahydrate [7784-17-0 ($12\text{H}_2\text{O}$), 14284-36-7] $\text{AlCs}(\text{SO}_4)_2 \cdot 12\text{H}_2\text{O}$, **M 352.1 (anhydr), 568.2, m 110°(dec)**. Recrystallise it from hot water (3ml/g).

Aluminium chloride (anhydrous) [7446-70-0] AlCl_3 , **M 133.3, m 192.6°, d_4^{17} 2.465**. Sublime it several times in an all-glass system under nitrogen at 30-50mm pressure. It has also been sublimed in a stream of dry HCl and has been subjected to a preliminary sublimation through a section of granular aluminium metal [for manipulative details see Jensen *J Am Chem Soc* **79** 1226 1957, DOI: 10.1021/ja01562a051]. It fumes in moist air. Used in Friedel-Crafts reactions. It is also commercially available as a 0.1M solution in nitrobenzene [7446-70-0], in functionalised silica gel (70-120mesh, with a 1.5mmol/g loading) for use as a Lewis acid and Friedel-Crafts reactions, and as $\text{AlCl}_3 \cdot \text{THF}$ complex [192656-42-1] as a 0.5M solution in THF

Aluminium fluoride (anhydrous) [7784-18-1 (anhydr), 32287-65-3 (H_2O), 15098-87-0 ($3\text{H}_2\text{O}$)] AlF_3 , **M 84.0 (anhydr), m 1292°(subliming), d^{25} 3.1g/cm³(anhydr)**. The technical material may contain up to 15% alumina, and minor impurities such as aluminium sulfate, cryolite, silica and iron oxide. Reagent grade AlF_3 (hydrated) contains only traces of impurities, but its water content is variable (and may be up to 40%). It can be dried by calcining at 600-800° in a stream of dry air (some hydrolysis occurs), followed by vacuum distillation at low pressure in a graphite system, heated to approximately 925° (condenser at 900°) [Henry & Dreisbach *J Am Chem Soc* **81** 5274 1959, DOI: 10.1021/ja01529a003]. Its solubility in H_2O is 0.5%. [Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 225-226 1963, Aigueperse et al. 'Fluorine Compounds, Inorganic' in *Ullmann's Encyclopedia of Industrial Chemistry* Wiley-VCH, Weinheim, 2005, DOI: 10.1002/14356007.a11_307]

Aluminium nitrate nonahydrate [7784-27-2 ($9\text{H}_2\text{O}$); 13473-90-0 (anhydr)] $\text{Al}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$, **M 375.1, m 72.8°, 73°, 73.5°, b 135°/atm, d^{25} 1.72g/cm³**. Crystallise the nitrate from dilute HNO_3 , and dry it by passing dry nitrogen through the crystals for several hours at 40°. After 2 recrystallisations of ACS grade, it had S, Na and Fe at 2.2, 0.01 and 0.02 ppm, respectively.

Aluminium potassium sulfate dodecahydrate (alum) [7784-24-9] $\text{AlK}(\text{SO}_4)_2 \cdot 12\text{H}_2\text{O}$, **M 474.4, m 92°, d^{25} 1.757g/cm³**. Crystallise it from weak aqueous H_2SO_4 (ca 0.5ml/g). Its solubility (%) in H_2O is 5.7 (0°), 12.0

(20°) and 136.9 (100°). Crystals for X-ray studies were easily grown in H₂O solution and were ground into spheres about 0.15mm in diameter [crystal structure: Larson & Cromer *Acta Cryst* **22** 793 1967, DOI: 10.1107/S0365110X67001586].

Aluminium rubidium sulfate dodecahydrate [7784-29-4, 1350-57-9 (anhydr)] **AlRb(SO₄)₂ · 12H₂O**, **M 520.8, 304.6 (anhydr)**, **m 99-109°, 100°(dec)**, **d 1.90g/cm³**. Crystallise the double salt from aqueous H₂SO₄ (ca 2.5ml/g). Crystals for X-ray studies were easily grown in aqueous solution and were ground into spheres about 0.15mm in diameter. It is insoluble in EtOH. [Prepn: *Gmelin's, Aluminium* (8th edn) **35B** p 525-527 1934; crystal structure: Larson & Cromer *Acta Cryst* **22** 793 1967, DOI: 10.1107/S0365110X67001586.]

Aluminium sulfate (anhydrous) [10043-01-3] **M 342.2(anhydr)**, **m 765°(dec)**, **d²⁵ 2.71g/cm³**, **Al₂(SO₄)₃ 14-18 H₂O** [17927-65-0], **d²⁵ 1.69g/cm³**, **Al₂(SO₄)₃ 18 H₂O** [7784-31-8]. It crystallises from hot dilute H₂SO₄ (1 ml/g) on cooling in ice. When a solution of alumina (Al₂O₃) in concentrated H₂SO₄ is slowly cooled, Al₂(SO₄)₃ 17 or 18H₂O deposits as a crystalline mass. Al₂(SO₄)₃ 17H₂O is the stable form in equilibrium with its saturated aqueous solution at 25° [Smith *J Am Chem Soc* **64** 41 1942, DOI: 10.1021/ja01253a012]. This is purified by dissolving it in a small volume of H₂O and adding EtOH until the sulfate readily crystallises from the oily supersaturated solution. It forms Al₂(SO₄)₃ 16H₂O between 0-112°. On gradual heating, the hydrate melts, giving the anhydrous salt at ca 250°. Several hydrates up to 27H₂O have been described. Further heating to red heat (~ 600-800°) causes decomposition to Al₂O₃ + SO₃ + SO₂ and O₂ [Cobb *J Soc Chem Ind* **29** 250 1910]. The ACS reagent is Al₂(SO₄)₃ 18H₂O (98+%).

Ammonia (gas) [7664-41-7] **NH₃**, **M 17.0**, **pK²⁵ 9.25**. Major contaminants are water, oil and non-condensable gases. Most of these impurities are removed by passing the ammonia through a trap at -22° and condensing it at -176° under vacuum. Water is removed by distilling the ammonia into a tube containing a small lump of sodium. Also dry it by passage through porous BaO, or over alumina followed by glass wool impregnated with sodium (prepared by soaking the glass wool in a solution of sodium in liquid ammonia and evaporating off the ammonia). It can be rendered oxygen-free by passage through a solution of potassium in liquid ammonia. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 460-463 1963.] **AMMONIA (gas, liquid or aqueous solution) is very irritating and should not be inhaled in any quantity as it can lead to olfactory paralysis (temporary or permanent).**

Ammonia (liquid) [7664-41-7] **NH₃**, **M 17.0**, **m -77.8°, b -33.35°, n_D²⁰ 1.325**, **d₄²⁰ 0.597**, **d⁻⁷⁹ 0.817g/ml**. Dry the liquid, and store it, with sodium in a steel cylinder, then distil and condense it by means of liquid air, the non-condensable gases being pumped off. In order to obtain liquid NH₃ from a cylinder, turn the cylinder upside-down (i.e. with the valve at the bottom, use a metal stand to secure it in this position; a special stand can be constructed for it) and lead a plastic tube from the tap to a measuring cylinder placed in an efficient fume cupboard which is kept running. Turn the tap on and allow the ammonia to be released. At first, gas and liquid will splatter out (make sure that the plastic tube is secure), but soon the liquid will drip into the measuring cylinder. The high latent heat of evaporation will cool the ammonia so that the liquid will remain cool and not boil vigorously. If the ammonia is required dry, the necessary precautions should be taken, i.e. the gas is allowed to flow through tubes packed with coarse CaO pellets. **AMMONIA (gas, liquid or aqueous solution) is very irritating and should not be inhaled in any quantity as it can lead to olfactory paralysis (temporary or permanent).**

Ammonia (aqueous) [7664-41-7] **NH₃ · xH₂O**, **M 17.0 + H₂O**, **d₄²⁰ 0.90 (saturated, 27% w/v, 14.3 N)**, **pK²⁵ 9.25** (pK_b at 25° = 4.75, i.e. pK_a = 14.00-4.75 = 9.25, or K_b = 1.81 × 10⁻⁵). Obtained metal-free by saturating distilled water, in a cooling bath, with ammonia (from a cylinder) gas. *Alternatively*, isothermal distillation can be used by placing a dish of concentrated aqueous ammonia and a dish of pure water in an empty desiccator and leaving to equilibrate for several days. **AMMONIA (gas, liquid or aqueous solution) is very irritating and should not be inhaled in any quantity as it can lead to olfactory paralysis (temporary or permanent).**

Ammonium bisulfate (ammonium hydrogen sulfate) [7803-63-6] **(NH₄) HSO₄**, **M 115.1**, **m 121-145°, ~147°, d₄²⁰ 1.79**, **pK²⁵ 1.96 (HSO₄⁻)**. It crystallises from water at room temperature (1ml/g) on adding EtOH

and cooling. $(\text{NH}_4)\text{HSO}_4$ is formed as deliquescent rhombic crystals on cooling a solution of $(\text{NH}_4)_2\text{SO}_4$ in hot concentrated H_2SO_4 but EtOH decomposes it to $(\text{NH}_4)_3\text{H}(\text{SO}_4)_2$. *Alternatively*, if the salt is slightly acidic, it is extracted (Soxhlet) with dry Et_2O (no reaction) until the runnings are neutral. This is found to be the most efficient way of purifying $(\text{NH}_4)\text{HSO}_4$, **m 145°**, free from traces of H_2O and H_2SO_4 . It is *extremely* hygroscopic and should be stored in glass bottles in desiccators. When $(\text{NH}_4)\text{HSO}_4$ is shaken with 5 to 7 times its weight of EtOH (dried over lime or CaC_2 and distilled with **b 77.1-77.2°/726mm**) for 18 hours, filtered quickly then extracted with Et_2O (Soxhlet) until the washings are neutral (litmus), a salt with the formula $(\text{NH}_4)_3\text{H}(\text{SO}_4)_2$ is produced. This salt is not deliquescent and decomposes before melting. [Dunncliff *J Chem Soc* **123** 476 1923, DOI: 10.1039/CT9232300476; *Gmelin's, Ammonia* [8th Ed.] **23** 405-406 1936.]. When powdered $(\text{NH}_4)_2\text{SO}_4$ is heated in a Pt dish below 100°, it loses NH_3 , and at 300° it is completely converted to fused $(\text{NH}_4)\text{HSO}_4$, **m 140°**, but >300° it decomposes to SO_2 and N_2 [Smith *J Soc Chem Ind* **14** 629 1895, **15** 3 1896]. [Dunncliff *J Chem Soc* **123** 731 1923, DOI: 10.1039/CT9232300731.] Used in hair waving and as a source of weak acid.

Ammonium bromide [12124-97-9] NH_3HBr , **M 98.0**, **m 450°(sublimes)**, **d₄²⁰ 2.43**. It crystallises from 95% EtOH and is slightly hygroscopic.

Ammonium chloride [12125-02-9] NH_3HCl , **M 53.5**, **m 338°(sublime point, without melting)**, **d₄²⁰ 1.53**. Crystallise it several times from conductivity water (1.5ml/g) between 90° and 0°. It sublimes. The salt is fully ionised in aqueous solution, i.e. $K \approx \infty$. A 1M aqueous solution of NH_4Cl has a pH of ~4.7, i.e. is acidic. After one crystallisation, ACS grade has: metal(ppm) As (1.2), K (1), Sb (7.2), V (10.2). [Becher in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 812 1963.]

Ammonium chromate [7788-98-9] $(\text{NH}_4)_2\text{CrO}_4$, **M 152.1**, **m 185°(dec)**, **d₄²⁰ 1.81**, **pK₁²⁵ 0.74**, **pK₂²⁵ 6.49 (for H_2CrO_4)**. Crystallise it from weak aqueous ammonia (ca 2.5ml/g) by cooling from room temperature. It loses NH_3 on heating to form ammonium dichromate (below). [Prepn: *Gmelin's, Chromium* (8th edn) **52B** p 707-712 1962.] Used as a reagent in analytical chemistry, in textile and wool dyeing, and in sensitising photographic gelatin. **POISONOUS**.

Ammonium dichromate [7789-09-5] $(\text{NH}_4)_2\text{Cr}_2\text{O}_7$, **M 252.1**, **m 170°(dec)**, **d₄²⁰ 1.26**. It crystallises from weak aqueous HCl (ca 1ml/g). It decomposes rapidly on heating. (Possible **carcinogen**, **irritates skin** possibly forming **chrome sores**, and is **POISONOUS**)

Ammonium dihydrogen arsenate [13462-93-6] $(\text{NH}_4)\text{H}_2\text{AsO}_4$, **M 159.0**, **Sp gr 2.311g/ml**, **n 1.577**, **m 300°(dec)**. Crystallise it from water (1ml/g). **POISONOUS**.

Ammonium dihydrogen orthophosphate [7722-76-1] $(\text{NH}_4)\text{H}_2\text{PO}_4$, **M 115.0**, **m 190°(dec)**, **d₄²⁰ 1.80**. Crystallise it from water (0.7ml/g) between 100° and 0°. It is slightly soluble in EtOH and insoluble in Me_2CO . With NaHCO_3 (a baking powder), it is used in fermentation and culture media. It is a fire retardant for paper and related fibrous materials.

Ammonium ferric sulfate dodecahydrate [7783-83-7 (12 H_2O), 10138-04-2 (anhydrous)] $(\text{NH}_4)\text{Fe}(\text{SO}_4)_2 \cdot 12\text{H}_2\text{O}$, **M 482.2**, **m ~37°**, **39-41°**, **d₄²⁰ 1.71**. Crystallise it from aqueous ethanol. Its solubility is 1M in dilute HCl , and 1.24g/100ml in H_2O .

Ammonium ferrous sulfate hexahydrate [Mohr's salt] [7783-85-9 (6 H_2O), 10045-89-3 (anhydrous)] $(\text{NH}_4)_2\text{Fe}(\text{SO}_4)_2 \cdot 6\text{H}_2\text{O}$, **M 392.1**, **284.05(anhydr)**, **m 100°(dec)**, **d₄²⁰ 1.86**. A solution in warm water (0.67g/ml) is cooled rapidly to 0°, and the resulting light bluish-green monoclinic crystals are filtered at the pump, washed with cold distilled water and pressed between sheets of filter paper to dry it. The solubility at 25° is 0.36g/ml. It separates as an almost white powder when a saturated aqueous solution is diluted with EtOH .

Ammonium hexachloroiridate (IV) [16940-92-4] $(\text{NH}_4)_2\text{IrCl}_6$, **M 441.0**, **d²⁵ 2.86g/ml**. It is precipitated several times from aqueous solution by saturation with ammonium chloride. This removes any palladium and rhodium. It is then washed with ice-cold water and dried over conc H_2SO_4 in a vacuum desiccator. If osmium

or ruthenium is present, it can be removed as the tetroxide by heating with conc HNO_3 , followed by concentrated HClO_4 , until most of the acid has been driven off. (This treatment is repeated.) The near-dry residue is dissolved in a small amount of water and added to excess NaHCO_3 solution and bromine water. On boiling, iridic (but not platinic) hydroxide is precipitated. It is dissolved in HCl and precipitated several times, then dissolved in HBr and treated with HNO_3 and HCl to convert the bromides to chlorides. Saturation with ammonium chloride and cooling precipitates ammonium hexachloroiridate which is filtered off and purified as above [Woo & Yost *J Am Chem Soc* **53** 884 1931, DOI: 10.1021/ja01354a008]. This commercially important iridium compound is the most common complex of Ir(IV) [Renner et al. 'Platinum group metals and compounds' in *Ullmann's Encyclopedia of Industrial Chemistry*, Wiley 2002, DOI: 10.1002/14356007.a21_075].

Ammonium hexacyanoferrate II hydrate [14481-29-9] $(\text{NH}_4)_4[\text{Fe}(\text{CN})_6] \cdot x\text{H}_2\text{O}$, **M 284.1, m dec on heating**. The pale yellow *trihydrate* powder can be washed with 10% aqueous NH_3 , filtered, then washed several times with EtOH and Et_2O , and dried at room temperature. It decomposes in a vacuum above 100° and should be stored away from light and under N_2 . In light and air it decomposes by losing NH_3 . [Lux in *Handbook of Preparative Inorganic Chem* (Ed. Brauer) Academic Press Vol **II** p 1509 1965; *Gmelin's, Iron* (8th edn) **59B** p 1024 1932.] Used as a spot test reagent for Ca [Feigl, *Tüpfelanalyse*, Anorganischer Teil Akadem Verlag, Frankfurt/Main, p228 1960].

Ammonium hexafluorophosphate [16941-11-0] NH_4PF_6 , **M 163.0, d_4^{18} 2.181, $\text{pK}_1^{25} \sim 0.5$, pK_2^{25} 5.12 (for fluorophosphoric acid $\text{H}_2\text{PO}_3\text{F}$)**. It crystallises from H_2O in square plates and decomposes on heating before melting. Its solubility in H_2O at 20° is 74.8% w/v, and it is very soluble in Me_2CO , MeOH , EtOH and MeOAc , but is decomposed by boiling mineral acids. It does **not** etch glass. [Lange & Müller *Chem Ber* **63** 1058 1930, DOI: 10.1002/cber.19300630510; Woyski et al. *Inorg Synth* **3** 111 1951, DOI: 10.1002/9780470132340.ch29; Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 195 1963.]

Ammonium hexafluorosilicate [16919-19-0] $(\text{NH}_4)_2\text{SiF}_6$, **M 178.1, d^{25} 2.01g/ml, pK_2 1.92 (for H_2SiF_6)**. Crystallise the salt from water (2ml/g). After 3 recrystallisations, the Technical grade salt has Li, Na, K and Fe at 0.3, 0.2, 0.1 and 1.0 ppm respectively. Used as a flux in soldering and for etching glass. [*Gmelin's, Ammonium* (8th edn) **23** p414-415 1936.]

Ammonium hypophosphite (ammonium phosphinate)[7803-65-8] $\text{NH}_4 \cdot \text{H}_2\text{PO}_2$, **M 83.0, m 155-160°**. Crystallise it from hot EtOH . Its solubility in H_2O at $\sim 25^\circ$ is 1g/ml, and it liberates PH_3 (TOXIC) at $\sim 240^\circ$.

Ammonium iodate [13446-09-8] NH_4IO_3 , **M 192.9, m 150° (dec), d^{25} 3.309g/ml, pK^{25} 0.79 (IO_3^-)**. Ammonium iodate crystallises from water (8ml/g) on cooling from 100° to 0° .

Ammonium iodide [12027-06-4] NH_4I , **M 144.9, sublimes with dec at $\sim 405^\circ$, d_4^{20} 2.51**. The iodide crystallises from EtOH on addition of ethyl iodide, and is very *hygroscopic*. Store it in a tightly stoppered bottle, in the dark. Its solubility is 0.4g/ml in MeOH , 0.27g/ml in EtOH , 0.67g/ml in glycerol, and 1.67g/ml in H_2O at 25° , and 2.0g/ml in H_2O at $\sim 100^\circ$. [Schmeisser in *Handbook of Preparative Inorganic Chem* (Ed. Brauer) Academic Press Vol **I** p 289-290 1963.]

Ammonium magnesium chloride hexahydrate [60314-43-4 ($12\text{H}_2\text{O}$), 12125-06-3 (anhydrous)] $\text{NH}_4\text{Cl MgCl}_2$, **M 256.8, 148.7(anhydr)**. It crystallises from water (6ml/g) by partial evaporation in a desiccator over KOH (deliquescent).

Ammonium magnesium sulfate hexahydrate [20861-69-2] $\text{NH}_4\text{MgSO}_4 \cdot 6\text{H}_2\text{O}$, **M 246.4, 138.4(anhydr)**. It crystallises from water (1ml/g) between 100° and 0° .

Ammonium manganous sulfate hexahydrate (diammonium manganese II sulfate hexahydrate) [7785-19-5, 13566-22-8] $\text{NH}_4\text{SO}_4 \cdot \text{NH}_4\text{MnSO}_4 \cdot 6\text{H}_2\text{O}$, **M 391.3, 138.4(anhydr)**. It crystallises from water (2ml/g) by partial evaporation in a desiccator.

Ammonium molybdate [13106-76-8] $(\text{NH}_4)_2\text{MoO}_4$, **M 196.0, pK_1^{25} 0.9 (proton addition), pK_2^{25} 3.57, pK**

²⁵**4.08 (for H₂MoO₄)**. Crystallise the salt from water (2.5ml/g) by partial evaporation in a desiccator. When a solution of MoO₃ in excess of hot concentrated NH₃ is cooled, **normal** ammonium molybdate (NH₄)₂MoO₄ crystallises out. However, when this solution is made to evaporated which allows the pH to drop to 6, the **common** hydrated ammonium **paramolybdate** (NH₄)₆Mo₇O₂₄·4H₂O {or 3[(NH₄)₂O]·7MoO₃·4H₂O, [12054-85-2] M 1235.9} crystallises out. This was an **old formula** which was confirmed by Sturdivant. [Sturdivant *J Am Chem Soc* **59** 630 1937, DOI: 10.1021/ja01283a010; Grüttner & Jauder in *Handbook of Preparative Inorganic Chem* (Ed. Brauer) Academic Press Vol II p 1711 1965.]

Ammonium nickel (II) sulfate hexahydrate [7785-20-8 (6H₂O), 15699-18-0 (anhydrous)] (NH₄)₂Ni(SO₄)₂·6H₂O, M 395.0, m 58-89°, d₄²⁰ 1.923. Crystallise this salt from water (3ml/g) on cooling from 90° to 0°.

Ammonium nitrate [6484-52-2] NH₄NO₃, M 80.0, m 165°(moist salt), 169°, 169.6°, b 210°/atm(dec explosively), d₄²⁰ 1.725. It is crystallised twice from distilled water (1ml/g) by adding EtOH, or from warm water (0.5ml/g) by cooling in an ice-salt bath. Its solubility in H₂O is 1.18g/ml (0°), 1.5g/ml (20°), 2.3g/ml (40°), 4.1g/ml (60°), 5.76g/ml (80°) and 10.2g/ml (100°). Dry it in air, then under vacuum. After 3 recrystallisations of ACS grade, it contained Li and B at 0.03 and 0.74 ppm, respectively. It is **deliquescent**. [Early & Lowry *J Chem Soc* **115** 1387 1919, DOI: 10.1039/CT9191501387; **121** 963 1922, DOI: 10.1039/CT9222100963; Hendricks et al. *J Am Chem Soc* **54** 2766 1932, DOI: 10.1021/ja01346a020.] It is a high nitrogen fertiliser in agriculture, a useful explosive in mining and ammunition and should be **handled with care**.

Ammonium perchlorate [7790-98-9] NH₄ClO₄, M 117.5, d₄²⁰ 1.95, pK²⁵ -2.4 to -3.1 (for HClO₄). It is recrystallised twice from distilled water (2.5ml/g) between 80° and 0°, and dried in a vacuum desiccator over P₂O₅. Drying at 110° might lead to slow decomposition to the chloride. Useful in pyrotechnic materials and rocket propellants. [Jacobs & Whitehead *Chem Rev* **69** 551-50 1969, DOI: 10.1021/cr60260a005; *Gmelin's, Ammonium* (8th edn) **23** p 196-200 1936.] **POTENTIALLY EXPLOSIVE**.

Ammonium persulfate (APS, ammonium peroxydisulfate) [7727-54-0] (NH₄)₂S₂O₈, M 228.2, m dec when heated wet liberating oxygen, d₄²⁰ 1.98. Recrystallise it at room temperature from EtOH/water. It is freely soluble in H₂O: 0.5 (0.11)g/ml at 20°, and 2g/ml at 100°, also soluble in MeOH, but less so in EtOH and Me₂CO, and insoluble in most other organic solvents. Stable when pure and dry, but decomposes when moist evolving O₂ and O₃, and gradually loses NH₃ on exposure to air. It is a bleacher and a free radical initiator being used for polymerising acrylamide gels. [Feher in *Handbook of Preparative Inorganic Chem* (Ed. Brauer) Academic Press Vol I p 190 1963, Fieser **1** 952, **2** 348, **3** 238, **5** **5** 15, **6**, 20, **12** 33.]

Ammonium reineckate (Reineckate salt) [13573-16-5] NH₄[Cr(NH₃)₂(SCN)₄], M 336.4 (anhydrous), m 268-272°(dec), 270-273°(dec), d 1.49g/cm³. Crystallise this dark red solid from water, between 30° and 0°, while working under artificial light. Solutions of reineckate salt (aqueous or alcoholic) decompose slowly at room temperature in the dark (~2 weeks) and more rapidly at higher temperatures or in diffuse sunlight. The solutions are blue in colour and liberate HCN (**POISONOUS**). Store it dry in the dark under a vacuum. [Dakin *Org Synth Coll Vol* **2** 555 1943, DOI: 10.15227/orgsyn.015.0074.] Used to precipitate primary, secondary amines and some amino acids, e.g. crystalline products with proline and hydroxyproline, and gives a red colour or precipitate with Hg²⁺ compounds.

Ammonium selenate [7783-21-3] (NH₄)₂SeO₄, M 179.0, d₄²⁰ 2.19, m dec on heating. Crystallise the selenate from water at room temperature by adding EtOH and cooling. Its solubility in H₂O is 117% at 7° and 197% at 100°. [King *J Phys Chem* **41** 797 1937, DOI: 10.1021/j150384a003.] Used for mothproofing.

Ammonium sulfamate [7773-06-0] NH₄NH₂SO₃, M 114.1, m 132-135°, dec at 160°. Crystallise it from water at room temperature (1ml/g) by adding EtOH and cooling. [Sisler et al. *Inorg Synth* **2** 179 1946, DOI: 10.1002/9780470132333.ch53.]

Ammonium sulfate [7783-20-2] (NH₄)₂SO₄, M 132.1, m 230°(dec), 280°(dec), d₄²⁰ 1.77. Crystallise it twice from hot water containing 0.2% EDTA to remove metal ions, then finally from distilled water. Dry it in a

desiccator for 2 weeks over $\text{Mg}(\text{ClO}_4)_2$. After 3 recrystallisations, ACS grade had Ti, K, Fe, Na at 11, 4.4, 4.4, 3.2 ppm respectively. Extensively used in H_2O or aqueous buffer at various concentrations to fractionate, precipitate and/or crystallise proteins, purification of antibodies and crystallography of nucleic acids and proteins.

Ammonium tetrafluoroborate [13826-83-0] $\text{NH}_4 \text{BF}_4$, M 104.8, m 220°(sublimes), $d^{25}_{25} 1.871 \text{ g/cm}^3$, $\text{pK}^{25}_{25} 2.77$ (for HBF_4). Crystallise it from conductivity water (1m/g) between 100° and 0°.

Ammonium thiocyanate (ammonium rhodanide) [1762-95-4] $\text{NH}_4 \text{SCN}$, M 76.1, m 138°(dec), 149°(dec), $\text{pK}^{25}_{25} -1.85$ (for HSCN), 149. Crystallise it three times from dilute HClO_4 to give material optically transparent at wavelengths longer than 270nm. It has also been crystallised from absolute MeOH or from acetonitrile, and is soluble in H_2O (1.28g/1ml at 0°), EtOH, Me_2CO and liquid NH_3 . It is a useful source of the SCN anion. Used for preparing herbicides, transparent artificial resins, matches, rustproofing materials, textile dyeing and printing, stabilising agents in photography, separation of Hf and Zr, titrimetric analysis, and detection of Cu, Ag, Zn, Pb and Hg which form precipitates that can be extracted into organic solvents. [A.F.Wells *Structural Inorganic Chemistry* 5th edn, OUP Oxford UK 1984, ISBN 9780198553700.]

Ammonium tungstate (VI) [11120-25-5, 89127-99-1] $\text{H}_{40}\text{N}_{10}\text{O}_{41}\text{W}_{12} = (\text{NH}_4)_{10} \text{W}_{12}\text{O}_{41}$, M 3042.6, $[(\text{NH}_4)_2 \text{WO}_4]$, M 283.9, $\text{pK}^{25}_1 2.20$, $\text{pK}^{25}_2 3.70$ (for tungstic acid, H_2WO_4). It crystallises as the *pentahydrate* from warm water on adding EtOH and cooling. It decomposes at 600° to give *tungsten(VI) oxide* (WO_3) which yields elemental *tungsten* on heating in an atmosphere of H_2 . The metal is used in the manufacture of tungsten alloys. The tungstate anion can be complex, forming a white tungsten-oxygen cage, e.g. *ammonium paratungstate* which has the $[\text{H}_2\text{W}_{12}\text{O}_{42}]^{10-}$ anion. [Greenwood & Earnshaw, *Chemistry of the Elements*, 2nd ed. B-H, Oxford, pp 1012–1014 1997, ISBN 0080379419.]

Ammonium (meta) vanadate [7803-55-6] $\text{NH}_4 \text{VO}_3$, M 117.0, $d^{20}_{10} 2.326$. Wash the salt with H_2O until free from Cl^- ions and dry it in air. It is soluble in H_2O (5.18g/100ml at 15°, 10.4g/100ml at 32°) but is more soluble in dilute NH_3 . It crystallises from conductivity water (20ml/g). When heated at relatively low temperature, it loses H_2O and NH_3 to give vanadium oxide (V_2O_5), and at 210° it forms lower oxides. [Baker et al. *Inorg Synth* 3 117 1950, DOI: 10.1002/9780470132340.ch30.] Its solubility in H_2O is 0.52% (15°), 1% (32°) and 1.6% (50°). After washing the technical grade salt with H_2O , it had Na, Mn and U at 0.06, 0.2 and 0.1 ppm, respectively. [Brauer in *Handbook of Preparative Inorganic Chem* (Ed. Brauer) Academic Press Vol II p 1272-1273 1965.]

Antimony (V) pentafluoride [7783-70-2] SbF_5 , M 216.7, m 7.0°, 8.3°, b 141°, 150°, 148-150°, $d^{20}_4 2.99$, $\text{pK}^{25}_{25} 2.55$ [for $\text{HSb}(\text{OH})_6 = \text{Sb}(\text{OH})_6^- + \text{H}^+$]. Purify it by vacuum distillation, preferably in a quartz apparatus, and store it in quartz or aluminum bottles. It is a *hygroscopic* viscous liquid which reacts *violently* with H_2O and is hydrolysed by alkalis. It is **POISONOUS** and attacks the skin. [Woolf & Greenwood *J Chem Soc* 2200 1950, DOI: 10.1039/JR9500002200; Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 200 1963.]

Antimony trichloride [10025-91-9] SbCl_3 , M 228.1, m 73°, 73.4°, b 220.3°/760mm, 283°/atm, $\text{pK}^{25}_1 1.4$, $\text{pK}^{25}_2 11.0$ (11.8), $\text{pK}^{25}_3 12.95$ (for Sb^{3+} aquo). Dry the trichloride over P_2O_5 or by mixing it with toluene or xylene and distilling (water is carried off with the organic solvent), then distil it twice under dry nitrogen at 50mm, and sublime it twice in a vacuum into ampoules and seal. It can be crystallised from CS_2 and is *deliquescent*. It fumes in moist air and is decomposed by H_2O with precipitation of the basic chloride, but forms a clear solution in dilute HCl. It is soluble in AcOH (4.4g/g at 25°), Me_2CO (5.4g/g at 18°), PhCOCl (1.7g/g at 25°), 36.7%w/w HCl (7.9g/g at 20°), *p*-cymene (1.5g/g at 30°). It is a useful reagent in the **Carr-Price test** for detecting vitamin A and related carotenoids producing a blue complex which can be evaluated colorimetrically. [Patnaik *Handbook of Inorganic Chemicals*. McGraw-Hill, 2002, ISBN 0-07-049439-8.]

Antimony trifluoride [7783-56-4] SbF_3 , M 178.8, m 235°(dec), 292°, b 376°/atm, $d^{20}_4 4.379$. It crystallises from MeOH to remove oxide and oxyfluoride, then it is sublimed under vacuum in an aluminium cup on to a water-cooled copper condenser. Its solubility is 443g/100g in H_2O at 20° and 562g/100g in H_2O at 30° with

partial hydrolysis. Store it in a glass or steel vessel. [Woolf *J Chem Soc* 273 279 1955, DOI: 10.1039/JR9550000273; Kwasnik in *Handbook of Preparative Inorganic Chem* (Ed. Brauer) Academic Press Vol I p 199 1963].

Antimony triiodide [7790-44-5] SbI_3 , **M 502.5, m 167°, 168°, 170°, b 401°/atm, 420°/atm.** It sublimes under vacuum as a ruby-red solid with an orange vapour. It hydrolyses to the yellow oxyiodide (SbOI) with H_2O . It is insoluble in CCl_4 but is soluble in aqueous KI , HCl , EtOH , Me_2CO , and CS_2 . [Bailar et al. *Inorg Synth* I 104 1939, DOI: 10.1002/9780470132326.ch36; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 614 1963.]

Antimony trioxide [1309-64-4] Sb_2O_3 , **M 291.5, m 656°, b 870°/10mm, 1424°/atm, 1550°/atm.** Dissolve the trioxide in the minimum volume of dilute HCl , filter, and add six volumes of water to precipitate the basic antimonous chloride (free from Fe and Sb_2O_5). The precipitate is redissolved in dilute HCl , and added slowly, with stirring, to a boiling solution (containing a slight excess) of Na_2CO_3 . The oxide is filtered off, washed with hot water, then boiled and filtered. The process is repeated until the filtrate gives no test for chloride ions. The product is dried in a vacuum desiccator [Schuhmann *J Am Chem Soc* 46 52 1924, DOI: 10.1021/ja01666a008]. After one crystallisation (precipitation?), the oxide from a Chinese source had: metal (ppm) Al (8), Ag (0.2), As (56), Cr (6), Ge (0.4), Mn (0.2), Na (16), Ni (2.2) Pb (2.4), Sn (0.4) and V (32). It sublimes in a vacuum at 400°, being yellow on heating and pale buff in colour on cooling. Used in pigments, enamels, glass, flame-proofing canvas and as a mordant. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 615-616 1963.]

Aqua regia. This is prepared by adding slowly concentrated HNO_3 (1 volume) to concentrated hydrochloric acid (3 volumes) in a glass container. This mixture is used to dissolve metals, including noble metals and alloys, as well as minerals and refractory substances. It is done by suspending the material and boiling (**EFFICIENT FUME CUPBOARD — EYE PROTECTION**) to dryness and repeating the process until the residue dissolves in H_2O . If the aqua regia is to be stored for long periods it is advisable to dilute it with one volume of H_2O which will prevent it from releasing chlorine and other chloro and nitrous compounds which are objectionable and toxic. Store it cool in a fume cupboard. However, it is good laboratory practice to prepare it freshly and dispose of it down the fume cupboard sink with copious amounts of water.

Argon [7440-37-1] Ar , **M 39.9481 (atomic weight standard), m -189.34°, b -185.848°, d 1.784g/L (at STP, 0°/101.325 kPa), 1.3954g/L (liquid at b.p.)** Argon is rendered oxygen-free by passage over reduced copper at 450°, or by bubbling through alkaline pyrogallol and H_2SO_4 , then dried with CaSO_4 , $\text{Mg}(\text{ClO}_4)_2$, or Linde 5A molecular sieves. Other purification steps include passage through *Ascarite* (**CARE: asbestos** impregnated with sodium hydroxide), through finely divided uranium at about 800° and through a -78° cold trap.

Alternatively, the gas is passed over CuO pellets at 300° to remove hydrogen and hydrocarbons, over Ca chips at 600° to remove oxygen and, finally, over titanium chips at 700° to remove nitrogen. It has also been purified by freeze-pump-thaw cycles and by passage over sputtered sodium [Arnold & Smith *JCS Faraday Trans* 2 77 861 1981, DOI: 10.1039/F29817700861]. Argon is inert and **non-toxic** and is 2.5 times more soluble in H_2O than N_2 . It comprises 0.934% by volume and 1.288% by mass of the Earth's crust. Being 38% denser than air, it is considered a **dangerous asphyxiant** in confined spaces. This inert gas is a useful shield in metal arc welding, a gas in fluorescent tubes and light bulbs and a commonly used carrier gas in GLC. [Haynes ed. *CRC Handbook of Chemistry and Physics* (92nd ed.) Boca Raton, FL, CRC Press p. 4.121 2011, ISBN 1439855110; Hwang et al. 'Noble Gases' in Kirk Othmer's *Encyclopedia of Chemical Technology*, Wiley, pp 343–383 2005, DOI: 10.1002/0471238961.0701190508230114.a01.]

Arsenic acid (arsenic pentoxide hydrate, arsenic V oxide hydrate, orthoarsenic acid) [12044-50-7] $\text{As}_2\text{O}_5 \cdot x\text{H}_2\text{O}$, **M 229.8 + $x\text{H}_2\text{O}$, d²⁵ 4.32g/ml, pK₁²⁵ 2.26, pK₂²⁵ 6.76, pK₃²⁵ 11.29 (H_3AsO_4).** The acid crystallises from concentrated solutions of boiling concentrated HNO_3 as rhombic crystals. Dry it in a vacuum to give the **hemihydrate** (hygroscopic). Heating above 300° yields As_2O_5 . [Simon & Thaler *Z Anorg Allgem Chem* 246 19 1941, DOI: 10.1002/zaac.19412460104; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 601 1963.] **POISONOUS.**

Arsenic tribromide [7784-33-0] AsBr_3 , M 314.6, m 31.1° , b $89^\circ/11\text{mm}$, $221^\circ/760\text{mm}$, d_4^{20} 3.67. Distil it under vacuum. It hydrolyses in H_2O (fumes in moist air), but less readily than AsCl_3 . It has a characteristically **high refractive index** (~ 2.3) as well as a **very high diamagnetic susceptibility**. **POISONOUS**. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 597 1963.]

Arsenic trichloride (butter of arsenic) [7784-34-1] AsCl_3 , M 181.3, m -16° , b $25^\circ/11\text{mm}$, $130.0^\circ/\text{atm}$, d_4^{20} 2.2, n_D^{20} 1.006. Reflux the trichloride with arsenic for 4 hours, then fractionally distil it. The middle fraction is stored with sodium wire for two days, then again distilled [Lewis & Sowerby *J Chem Soc* 336 1957, DOI: 10.1039/JR9570000336]. It fumes in moist air forming the solid hydroxy-chloride $[\text{AsCl}(\text{OH})_2]$ and is readily hydrolysed by H_2O to form arsenious acid. Useful in the ceramic industry. **POISONOUS**. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 596 1963.]

Arsenic triiodide [7784-45-4] AsI_3 , M 455.6, m 146° , b $400^\circ/\text{atm}$, d_4^{25} 4.688. It crystallises from acetone and sublimes below 100° . It is very slowly hydrolysed by H_2O (much more slowly than the chloride), e.g. 0.083g dissolves in H_2O (1ml) to give a yellow solution from which it can be recovered unchanged within 5hrs, by extraction into C_6H_6 , toluene or xylene. However, in time the aqueous acidic solution will form HI and As_2O_3 . **POISONOUS**. [Bailar et al. *Inorg Synth* 1 103 1939, DOI: 10.1002/9780470132326.ch36; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 597-598 1963.]

Arsenic III oxide (arsenic trioxide, arsenous oxide) [1327-53-3] As_2O_3 , M 197.8, three forms: m $\sim 200^\circ$ (amorphous glass), m 275° (sealed tube, octahedral, common form, sublimes $> 125^\circ$ without fusion but melts under pressure), and m $\sim 312^\circ$, b $465^\circ/\text{atm}$, pK_1^{20} 9.27, pK_2^{20} 13.54, pK_3^{20} 13.99 (for H_3AsO_3). It crystallises in an **octahedral form** (common form) from H_2O or from dilute HCl (1:2), and is then washed, dried and sublimed ($193^\circ/760\text{mm}$). Analytical reagent grade material is suitable for use as an analytical standard after it has been dried at 105° for 1-2 hours or has been left in a desiccator for several hours over concentrated H_2SO_4 . Alternatively, As_2O_3 (15g) is dissolved by heating in a mixture of H_2O (60ml) and HCl (90g, s.g. 1.1), and crystallisation occurs on cooling, accompanied by brilliant flashes of light.

The **amorphous** form is a colourless transparent glass (m 200°) which is obtained when the vapour is slowly condensed below the vapourisation temperature, and should be kept in a sealed tube because it changes to the **octahedral** form (m 275°) in the presence of moisture. [Rushton & Daniels *J Am Chem Soc* 48 384 1926, DOI: 10.1021/ja01413a011.]

A third **monoclinic** form, is obtained by heating the oxide in a sealed tube at 400° (the vitreous, amorphous form remains at the bottom of the tube) with the **monoclinic** form subliming onto the intermediate part of the tube at 200° (m 312°), and the **octahedral** form deposits at the top of the tube. The transition temperature between the last two forms is $\sim 250^\circ$. **POISONOUS (particularly the vapour, handle in a ventilated fume cupboard)**. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 600 1963; Grund et al. 'Arsenic and Arsenic Compounds' *Ullmann's Encyclopedia of Industrial Chemistry*, Weinheim: Wiley-VCH, 2005, DOI: 10.1002/14356007.a03_113.pub2.]

Barium (metal) [7440-39-3] Ba M 137.3, m 725° , 727° , b $1640^\circ(1537^\circ)/760\text{mm}$, d_4^{20} 3.56(3.76). Barium is cleaned by washing with diethyl ether to remove adhering paraffin, then filed in an argon-filled glove box, washed first with ethanol containing 2% concentrated HCl, then with dry ethanol. It is dried in a vacuum and stored under argon [Addison et al. *J Chem Soc* 3868 1962, DOI: 10.1039/JR9620003868]. It has also been purified by double distillation under 10mm of argon pressure. Useful as a catalyst for making graphitic nanofilaments [Liang et al. *Nanotech in Catalysis* 2 543 2004].

Barium bromide dihydrate [7791-28-8 ($2\text{H}_2\text{O}$), 10553-31-8 (anhydrous)] $\text{BaBr}_2 \cdot 2\text{H}_2\text{O}$, M 333.2, m at 75° loses first H_2O and at 120° it loses the second H_2O and melts at 847° , 857° , b $1835^\circ/\text{atm}$, d $4.78\text{g}/\text{cm}^3$. It crystallises from H_2O (0.92g/ml at 0°) by partial evaporation in a desiccator. The **anhydrous salt** is readily prepared by drying the dihydrate at 200° . Drying at temperatures of 120° gives a product which is either a lower hydrate or a different crystalline modification of the anhydrous material. The **anhydrous salt** is **deliquescent** and should be handled in a drybox. [Brackett et al. *J Phys Chem* 67 2132 1963, DOI: 10.1021/j100804a038.] [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 930 1963.]

Barium chlorate monohydrate [10294-38-9 (*hydrate*), 13477-00-4 (*anhydrous*)] $\text{Ba}(\text{ClO}_3)_2 \cdot \text{H}_2\text{O}$, **M 322.3, m 414°, d²⁵ 3.18g/cm³**. It crystallises from H_2O (1ml/g) between 100° and 0°, and loses H_2O at 120°. [Schmeisser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 314 1963.]

Barium chloride dihydrate [10326-27-9] $\text{BaCl}_2 \cdot 2\text{H}_2\text{O}$, **M 244.3, m ~120°(dec, hydrate); M 208.2, 963° (anhydrous), d²⁵ 3.856g/cm³**. It is crystallised twice from water (2ml/g) and dried in an oven to constant weight. The solubilities of the *hydrate* (% of anhydrous wt) in H_2O are 31.6 at 0°, 35.7 at 20° and 58.7 at 100°. Drying the dihydrate at 200° gives a mixture of *monoclinic* and *orthorhombic* forms. Conversion of the *monoclinic* form to the more stable *orthorhombic* modification is very slow (> 1 week). Attempted preparation of a pure sample of the *orthorhombic* form from the dihydrate by heating under vacuum at 60° overnight provided a *cubic* form which is stable at 925-960°. The *cubic* form does not revert appreciably to the *orthorhombic* form at room temperature, but at 200° the conversion is complete in 2 days. Thus the three forms can be obtained for X-ray diffraction. The *anhydrous* salts are *deliquescent* and should be handled in a drybox. [Brackett et al. *J Phys Chem* **67** 2132 963, DOI: 10.1021/j100804a038.]

Barium dithionate dihydrate [13845-17-5] $\text{Ba}(\text{S}_2\text{O}_6) \cdot 2\text{H}_2\text{O}$, **M 333.5, m >150° loses SO_2 , d²⁵ 4.54g/cm³, pK²⁵ 0.49 (for $\text{H}_2\text{S}_2\text{O}_6$, theory pK₁ -3.4, pK₂ -0.2)**. Purify by successive crystallisations from water at room temperature by evaporation. Its solubility in H_2O is 7.9% (0°), 15.7% (20°) and 19.9% (30°). [Pfanstiel et al. **2** 167 1946, DOI: 10.1002/9780470132333.ch50; Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 397 1963; crystal structure: Rausell-Colom & Garcia-Blanco *Acta Cryst* **21** 672 1966, DOI: 10.1107/S0365110X66003682.]

Barium ferrocyanide hexahydrate [13821-06-2] $\text{Ba}_2[\text{Fe}(\text{CN})_6]$, **M 594.8, m 80°(dec), pK₃²⁵ 2.57, pK₄²⁵ 4.35 (for ferrocyanide)**. Prepare the complex salt by boiling Prussian Blue with the equivalent of $\text{Ba}(\text{OH})_2$ solution, or FeSO_4 and $\text{Ba}(\text{CN})_2$, and it crystallises in yellow *monoclinic* crystals from hot water (100ml/g). The hydrate loses most of its H_2O on heating at 40° becoming colourless, and decomposes at 80° evolving HCN . [Grat-Cabanac *Bull Soc Chim Fr* 1743 1956.]

Barium fluoride [7787-32-8] BaF_2 , **M 175.3, m 1353°, 1368°, b 2260°, d₄²⁰ 4.83**. Wash it well with distilled H_2O and dry it in a vacuum. Its solubility in H_2O is 1.6g (10°), 1.6g (20°) and 1.62g (30°) per L, and is soluble in mineral acids and aqueous NH_4Cl . It may be stored in glass bottles. [Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 234 1963.]

Barium hydroxide octahydrate [12230-71-6] $\text{Ba}(\text{OH})_2 \cdot 8\text{H}_2\text{O}$, **M 315.5, m 78°, pK₁²⁵ 13.13, pK₂²⁵ 13.36**. It crystallises from water (1ml/g) and readily absorbs CO_2 from air. It effloresces to the *monohydrate* [22326-55-2] **M 189.4**. It dehydrates to $\text{Ba}(\text{OH})_2$ [17194-00-2] **M 171.3**, in dry air at 100°. An aqueous solution (*baryta water*) absorbs CO_2 to form a white precipitate of BaCO_3 . This base catalyses the β -elimination of phosphoserine residues; and a 0.3N solution of $\text{Ba}(\text{OH})_2$ [17194-002] **M 171.3**, has been used for precipitating proteins in turbid fluids like whole blood in assays based on Somogyi-Nelson's method. [Hatanaka & Kobara *Agricultural and Biological Chemistry* **44** 2943 1980, DOI: 10.1080/00021369.1980.10864408.]

Barium hypophosphite monohydrate [14871-79-5] $\text{Ba}(\text{H}_2\text{PO}_2)_2 \cdot \text{H}_2\text{O}$, **M 285.4**. Prepared by heating white phosphorus with $\text{Ba}(\text{OH})_2$ and purified by precipitating it from aqueous solution (3ml/g) on adding EtOH . Its solubility in H_2O is 28.6% at 17° and 33.3% at 100°. [Klement in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 557 1963.]

Barium iodate monohydrate [7787-34-0] $\text{Ba}(\text{IO}_3)_2 \cdot \text{H}_2\text{O}$, **M 487.1 (anhydr), m 130°(loses H_2O), 476°(dec)**. The iodate is purified by recrystallisation from a large volume of hot water. Its solubility in H_2O is 1g/3.35L at 25° and 1g/0.625L at the boiling point. [Lambert et al. *Inorg Synth* **7** 13 1963, DOI: 10.1002/9780470132388.ch4.]

Barium iodide dihydrate [7787-33-9 ($2\text{H}_2\text{O}$), 13718-50-8 (*anhydrous*)] $\text{BaI}_2 \cdot 2\text{H}_2\text{O}$, **M 427.2, m 740°(dec)**. The *dihydrate* salt crystallises from water (0.5ml/g) by partial evaporation in a desiccator. The *anhydrous* salt is

best obtained by heating the dihydrate to 400° in vacuum with no evidence of more than one modification (compare with BaCl₂). The monohydrate can apparently be obtained by heating the dihydrate at 500° in the presence of HI, or under vacuum at 150-200°. The *anhydrous* salt is *deliquescent* and should be handled in a drybox. [Brackett et al. *J Phys Chem* **67** 2132 963, DOI: 10.1021/j100804a038.] **POISONOUS.**

Barium manganate (VI) [7787-35-1] **BaMnO₄, M 256.3, d₄²⁰ 3.77.** Wash the salt (blue-green crystals) with cold conductivity H₂O by decantation until the supernatant gives a faint test for Ba²⁺. Remove excess H₂O in a vacuum (IMPORTANT), then heat at 100° and the last traces of H₂O are removed in a vacuum desiccator over P₂O₅. Store it over KOH. It disproportionates in hot H₂O or dilute acid into Ba(MnO₂)₂ and MnO₂ (see below). It is a mild oxidant, and is used in some wall paints. [Schlesinger & Siems *J Am Chem Soc* **46** 1965 1924, DOI: 10.1021/ja01674a001; Nyholm et al. *Inorg Synth* **11** 56 1968, DOI: 10.1002/9780470132425.ch11.]

Barium nitrate [10022-31-8] **Ba(NO₃)₂, M 261.4, m 593°(dec).** Crystallise it twice from water (4ml/g) and dry it overnight at 110°. It decomposes at higher temperatures to give mostly the oxide and the peroxide with only a little of the nitrite. **POISONOUS.** [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 941 1963.] The salt is used for making BaO, produces a green light in flares and pyrotechnics, and in fluorescent tubes.

Barium nitrite monohydrate [7787-38-4] **Ba(NO₂)₂·H₂O, M 247.4, m 217°(anhydr).** Barium nitrite monohydrate crystallises in yellow-white hexagonal crystals from water (1ml/g) on cooling in an ice-salt bath, and is insoluble in EtOH. Although stable in air it melts at ~184° with partial hydrolysis and concentrated aqueous solutions undergo hydrolysis at 100°. It may be dehydrated *in vacuo* over P₂O₅, and the *anhydrous* salt has d° 3.52g/ml at 0° and melts at 217° then decomposes at ~325°. **POISONOUS.**

Barium oxide (calcined baryta) [1304-28-5] **BaO, M153.3, m 1923°, b ~2000°/atm, d²⁵ 5.72g/cm³.** Best to prepare it by heating pure BaCO₃ with C, or other pure barium salts, e.g. Ba(NO₃)₂. It is a white powder which is used to absorb H₂O and CO₂ from gases (e.g. air, O₂, N₂). It does not become sticky with moisture, but dissolves in H₂O *exothermically* to form Ba(OH)₂ (*baryta water*, alkaline) which is used to absorb CO₂. Solubility in H₂O is 0.035g/1ml (20°) and 0.91g/1ml (100°). It is soluble in EtOH but insoluble in Me₂CO and liquid NH₃, and the porous grade is used for drying solvents that are not affected by alkali. IRRITATES skin, eyes, should not be inhaled and is harmful to aquatic and other organisms. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 933 1963.]

Barium perchlorate [13465-95-7] **Ba(ClO₄)₂, M 336.2(anhydr), m 505°, d²⁵ 3.2g/cm³. pK²⁵ -2.4 to -3.1 (for HClO₄).** Recrystallise the perchlorate twice from water (solubility is 66.5g/100ml at 25°). The hydrate has [15318-52-2, xH₂O]. [Schmeisser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 320 1963.]

Barium permanganate [7787-36-2] **Ba(MnO₄)₂, or BaMn₂O₈, M 375.2, m 200°(dec), d²⁵ 3.77g/cm³.** Obtained by disproportionation of BaMnO₄ in dilute H₂SO₄, the BaSO₄ and MnO₂ are filtered off and the filtrate is evaporated to give dark violet to brown rhombic crystals. The reaction is very slow because the solubility of the manganate is low. Dry in a vacuum. It is poorly soluble in cold H₂O (but 62.5g/100ml at 30°), and decomposes in EtOH. [Lux in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 1462 1963.] The salt is a mild versatile oxidising agent used under aprotic and non-aqueous solutions [Firouzabadi et al. *Tetrahedron* **46** 6869 1990, DOI: 10.1016/S0040-4020(01)87874-1].

Barium sulfate [7722-43-7] **BaSO₄, M 233.4, m 1345°, >1580°, d²⁵ 4.50g/cm³.** Wash the sulfate five times by decantation with hot distilled water, dialyse it against distilled water for one week, then freeze-dry and dry in an oven at 105° to constant weight (~12 hours). It is *practically insoluble* in H₂O (solubility is 0.0024g/L at 25°), an analytical product of *sulfate determination*, but is soluble in concentrated H₂SO₄. Used as a catalyst support, in making various kinds of paper whiter, in white pigments, and because of its high density it is useful as a radiocontrast agent for X-ray imaging.

Barium tetrathionate [82203-66-5] **BaS₄O₆, M 361.6, m 100-110°(forming BaSO₄, SO₂ and S).** Purify the

tetrathionate by dissolving it in a small volume of water and precipitating it with EtOH below 5°. After drying, the salt is stored in the dark at 0°. Aqueous solutions are stable, but alcoholic solutions in the presence of acid decompose it to give barium thiosulfate. [see Frehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 399-401 1963 for potassium tetrathionate K₂S₄O₄.]

Barium thiocyanate dihydrate [2092-17-3] Ba(SCN)₂·2H₂O, M 253.5(anhydr), pK²⁵ -1.85 (for HSCN). It is crystallised from water (2.5ml/g) by partial evaporation in a desiccator. It is *deliquescent*. [Herstein et al. *Inorg Synth* 3 24 1950, DOI: 10.1002/9780470132340.ch5.]

Barium thiosulfate [35112-53-9] BaS₂O₃, M 249.5, m 220°(dec), pK₁²⁵ 0.6, pK₂²⁵ 1.74 (for H₂S₂O₃). It is very slightly soluble in water and is washed repeatedly with chilled water and dried in air at 40°. Analytical standard in iodometry.

Beryllium carbonate [744998-97-8, 13106-47-3] BeCO₃, M 69.0, m 54°, b 100°/atm (dec). The commercial carbonate contains more than 1% of impurities. It is best purified by converting it to the basic acetate (see [1332-52-1]) which should be sublimed, converted into the nitrate with HNO₃ see below [13587-99-4]) and evaporated in a Pt dish until free from excess acid. The residual nitrate salt is dissolved in a small volume of H₂O and enough ammonium carbonate solution is added to redissolve the beryllium carbonate that precipitates and gives a clear solution. This is then evaporated in a Pt dish until course BeCO₃ separates. It is washed thoroughly with pure H₂O, then distilled EtOH and dried *in vacuo*. It has low solubility in H₂O (0.36g/100ml at 25°). Be(OH)₂, BeO and Be salts are generally insoluble in NH₃ but dissolve in **aqueous** (NH₄)₂CO₃ solution to form the carbonate. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 893 1963.] Beryllium compounds are potentially **carcinogenic**.

Beryllium chloride [7787-47-5] BeCl₂, M 79.9, m 399°, 405°, b 488°/atm, d²⁵ 1.89. The anhydrous chloride is obtained by heating Be powder or chips (or BeO + C) with Cl₂ gas in a furnace at 300-350° for *ca* 40 hours when anhydrous BeCl₂ sublimes out as a white to faintly yellow crystalline mass or orthorhombic crystals. It is purified further by sublimation at ~300° *in vacuo*. It is very hygroscopic and reacts exothermically with H₂O which becomes acidic. The solid should be stored in a tightly stoppered vessel and handled in a dry atmosphere much like anhydrous AlCl₃ and has similar catalytic properties to it. It is insoluble in *C₆H₆ and toluene but is soluble EtOH, Et₂O, pyridine and CS₂ with which it probably **complexes**. The conductivity of fused BeCl₂ is ~0.001 of that of fused NaCl implying that 1 molecule in 1000 is ionised. [Tannenbaum et al. *Inorg Synth* 5 22 1957, DOI: 10.1002/9780470132364.ch7; Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 889 1963.] Beryllium compounds are potentially **carcinogenic**.

Beryllium hydroxide [13327-32-7] Be(OH)₂, M 43.0, d²⁵ 1.89g/cm³. The hydroxide exists in two forms, a metastable **α-form** and a stable **β-form**. The former is obtained in amorphous form by precipitating a Be salt with NH₃ in the absence of CO₂ and allowing it to age by prolonged heating (~ 24 hours) with NH₄OH solution. **β-Be(OH)₂** is made by boiling a saturated solution of the amorphous α-form in 10N aqueous NaOH until a permanent turbidity is attained. On slow cooling fine crystals (regular double pyramids) of the β-form separate, and are purified by washing with warm H₂O until the washings are no longer alkaline, then drying at 80°. The mother liquours can be reused in the above process. It is very slightly soluble in H₂O and dilute NaOH, but both forms are amphoteric and are soluble in hot concentrated aqueous NaOH solutions and in acids. It dehydrates on strong heating to BeO. Several sodium beryllates are known. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 894, 895 1963.] Beryllium compounds are potentially **carcinogenic**.

Beryllium nitrate [13587-99-4; 7787-55-5 (3H₂O)] Be(NO₃)₂, M 133.0, m ~60° (for 3H₂O), b 142°/atm (dec), d²⁵ 1.56g/cm³. The nitrate does not crystallise easily and is **very deliquescent**. It can be prepared from the metal by digesting it with conc HCl until most of it has dissolved, filter any insoluble material through a 0.45 micron acetate filter. Then at low boil, add concentrated HNO₃ with low heat to reduce to a slurry. Repeat with careful addition of HNO₃ as many times as necessary (3-6 additions) until the slurry tests negative for chloride ions with 0.1N aqueous AgNO₃. Then isolate the nitrate as described below [J. Papa personal communication. 2011, see end of Chapter 1]. *Alternatively* it is obtained by heating recrystallised, then sublimed, basic beryllium

acetate $\{\text{Be}_4\text{O}(\text{OAc})_6$, see [1332-52-1]}, with pure concentrated HNO_3 in a degassed pyrex container until the volatiles are removed. Heat well below 1100° , so as not to decompose it to give pure BeO . It picks up moisture to give the white to pale yellow *trihydrate*. Store it in a well stoppered bottle in a cool place. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 893 1963.] Beryllium compounds are potentially **carcinogenic**.

Beryllium oxide (beryllia) [1304-56-9] BeO , M 25.0, m 2507° , 2530° , b $4120^\circ/\text{atm}$, d^{25} 3.01. It is prepared by calcining the nitrate at 1100° {see $\text{Be}(\text{NO}_3)_2$ above [13587-99-4]}, and this oxide always contains $\sim 0.35\text{ml}$ of gas (N_2 , O_2) per gram of oxide. Alternatively, freshly prepared beryllium carbonate is calcined in a Pt boat placed in an electric furnace at 900° . BeO is amphoteric, dissolving slowly in acids and in alkalis but if heated strongly it becomes refractory and difficult to dissolve except in hydrofluoric acid. It is *almost insoluble* in H_2O (0.002g/ml at 25°). Like ceramics it is a good electrical insulator, but it is a good conductor of heat like some metals. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 893 1963.] Beryllium compounds are potentially **carcinogenic**.

Beryllium potassium fluoride [7787-50-0] BeK_2F_4 , M 163.4 m $\sim 350^\circ$. It crystallises from hot water (25ml/g). Beryllium compounds are potentially **carcinogenic**.

Beryllium sulfate tetrahydrate [7787-56-6] $\text{BeSO}_4 \cdot 4\text{H}_2\text{O}$, M 177.1, m $\sim 100^\circ(\text{dec})$, d^{25} 1.713, pK_1^{25} 3.2, $\text{pK}_2^{25} \sim 6.5$ (Be^{2+}). It crystallises from a concentrated solution of weak aqueous H_2SO_4 (with partial evaporation), and is dried in air. It is quite soluble in H_2O . The *tetrahydrate* loses H_2O at $\sim 100^\circ$ to give the *dihydrate*. [cf. *Gmelin's Beryllium* (8th edn) 26 169 1930.] Beryllium compounds are potentially **carcinogenic**.

Bismuth [7440-69-9] Bi, M 209.0, m $271-273^\circ$, 271.44° , $271.04 \pm 0.06^\circ$, b $1450^\circ/\text{atm}$, $1560^\circ/\text{atm}$, d^{25} 9.8g/cm^3 . Melt it in an atmosphere of dry helium, then filter it through dry Pyrex wool to remove any bismuth oxide present [Mayer et al. *J Phys Chem* 64 238 1960, DOI: 10.1021/j100831a013].

Precipitated bismuth prepared by reduction of BiCl_3 in HCl with hypophosphorous acid, filtering it off, washing with H_2O and drying *in vacuo*, is a very fine dull grey powder with particles size $< 0.015\text{mm}$ ($< 15\text{microns}$) which is easily dispersed in H_2O . Bi dissolves in dilute mineral acids to give the respective salts *without* liberation of H_2 , whereas the red-hot metal reacts with H_2O to form Bi_2O_3 and hydrogen. A poor conductor of electricity, its resistance increases in a magnetic field, and it forms alloys with other metals, e.g. with Fe.

Bismuth trichloride [7787-60-2] BiCl_3 , M 315.3, m $230-232^\circ$, 233.6° , b $447^\circ/\text{atm}$, pK^{25} 1.58 for hydrolysis ($\text{Bi}_3^+ = \text{BiOH}_2^+ + \text{H}^+$). Sublime the trichloride under high vacuum (or $\sim 450^\circ/\text{atm}$), or dry it under a current of HCl gas, followed by fractional distillation, once under HCl and once under argon. [Mayer et al. *J Phys Chem* 64 238 1960, DOI: 10.1021/j100831a013; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 621 (BiCl_3), 622 (BiOCl) 1963.] It is *deliquescent*, and hydrolyses in H_2O or aqueous EtOH to give BiOCl [7787-59-9] M 260.4, d 7.72, which melts at low red heat. It dissolves in EtOH , Me_2CO , EtOAc , HCl and HNO_3 . BiCl_3 is a source of bismuth salts and catalysts for organic reactions.

Boric acid (orthoboric acid, boracic acid) [10043-35-3] H_3BO_3 , M 61.8, m $160^\circ(\text{dec})$, 171° , pK^{25} 9.23. Crystallise the acid three times from H_2O (3ml/g) between 100° and 0° , after filtering through sintered glass. Dry it to constant weight over metaboric acid in a desiccator. It is steam volatile. After two recrystallisations of ACS grade, it had Ag at 0.2 ppm. Its solubility (%) in H_2O is 2.66 at 0° , 4.0 at 12° and 24 at 80° . At 100° it loses H_2O to form *metaboric acid* (HBO_2). When it is heated to redness or slowly to 200° , or over P_2O_5 *in vacuo*, it dehydrates to *boric anhydride* (B_2O_3) [1303-82-6] to give a white hard glass or crystals with m $\sim 294^\circ$. The glass softens on heating and liquefies at red heat. It is an astringent, a fungicide and an antibacterial. [McCulloch *J Am Chem Soc* 59 2650 1937, DOI: 10.1021/ja01291a050; Kelley *J Am Chem Soc* 63 1137 1941, DOI: 10.1021/ja01849a072; Briscoe et al. *J Chem Soc* 70 1926, DOI: 10.1039/JR9262900070; Conti *J Soc Chem Ind* 44 343T 1925.]

Boron trichloride (trichloroborane) [10294-34-5] BCl_3 , M 117.2, m -107° , b $0^\circ/476\text{mm}$, $12.5^\circ/\text{atm}$, d^0 1.3728, d_4^{12} 1.35, (also available as a 1.0M solution in toluene, *p*-xylene, CH_2Cl_2 , heptane or hexanes). Purify it (from chlorine) by passage through two mercury-filled bubblers, then fractionally distil it under a slight

vacuum. In a more extensive purification the nitrobenzene addition compound is formed by passage of the gas over nitrobenzene in a vacuum system at 10°. Volatile impurities are removed from the crystalline yellow solid formed by pumping at -20°, and the BCl₃ is recovered by warming the addition compound at 50°. Passage through a trap at -78° removes entrained nitrobenzene, the BCl₃ finally condensing in a trap at -112° [Brown & Holmes *J Am Chem Soc* **78** 2173 1956, DOI: 10.1021/ja01591a042]. Alternatively, purify it by condensing it into a trap cooled in acetone/Dry-ice, where it is pumped for 15 minutes to remove volatile impurities. It is then warmed, recondensed and again pumped. [Gamble et al. *Inorg Synth* **3** 27 1950, DOI: 10.1002/9780470132340.ch6.]

It fumes in moist air. **TOXIC, do not breath in the vapours.** It forms addition compounds with Et₂O; and with dimethyl sulfide it forms **Me₂S·BCl₃** [5523-19-3] **M 179.3**, which has **m 88-90°**, and is a convenient solid form of it (also commercially available as a 1.0M solutions in CH₂Cl₂, heptane, hexanes, toluene and *p*-xylene) — **beware** of the foul odour of Me₂S; work in an efficient fume cupboard and absorb the sulfide vapours by flushing them with a stream of N₂ into lead acetate solution which will precipitate insoluble black PbS. [Gamble et al. *Inorg Synth* **3** 27 1950, DOI: 10.1002/9780470132340.ch6; Gerrard & Leppert *Chem Rev* **58** 1081 1958, DOI: 10.1021/cr50024a003.] Boronated aluminas (highly acidic) prepared from BCl₃ effect region- and diastereoselectivity in Diels-Alder reactions [McGinnis et al. *J Org Chem* **61** 3496 1996, DOI: 10.1021/jo951077m]. High-temperature synthesis of two layered graphite-like materials with p-type semiconductor properties was achieved by the interaction of MeCN with BCl₃, and acrylonitrile with BCl₃, in a H₂ and N₂ atmosphere in a quartz tube at 1000° [Kawaguchi *Chem Mater* **8** 1197 1996, DOI: 10.1021/cm950471y]

Boron trifluoride [7637-07-2] **BF₃**, **M 67.8**, **b -101°/760mm**. The usual impurities-bromine, BF₅, HF and non-volatile fluorides are readily separated by distillation. Brown and Johannesen [*J Am Chem Soc* **72** 2934 1950, DOI: 10.1021/ja01163a035] passed BF₃ into benzonitrile at 0° until the latter was saturated. Evacuation to 10⁻⁵mm then removed all traces of SiF₄ and other gaseous impurities. [A small amount of the BF₃-benzonitrile addition compound sublimes and is collected in a U-tube cooled to -80°]. The pressure is raised to 20mm by admitting dry air, and the flask containing the BF₃ addition compound is warmed with hot water. The BF₃ that evolves is passed through a -80° trap (to condense any benzonitrile) into a tube cooled in liquid air. The addition compound with anisole can also be used. BF₃ can be dried by passing it through H₂SO₄ saturated with boric oxide. It fumes in moist air. [It is commercially available as a 1.3M solution in MeOH or PrOH.] [Booth et al. *Inorg Synth* **I** 21 1939, DOI: 10.1002/9780470132326.ch8; Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 219-222 1963.] The following are **BF₃ complexes** that are among the commercially available sources of the reagent: **BF₃·2H₂O** [13319-75-0] **M 103.8**, **d²⁵ 1.636g/ml**; **BF₃·2AcOH** [373-61-5] **M 187.9**, **d²⁵ 1.353g/ml**; **BF₃·MeCN** [420-16-6] **M 108.9**, **d²⁰ 0.87-0.88g/ml**; **BF₃·Me₃COMe** [123334-27-0] **M 155.9**, **d²⁵ 1.381g/ml**; **BF₃·(n-Bu)₂O** [593-04-4] **M 198.0**, **d²⁵ 0.99g/ml**; **BF₃·Et₂O** [109-63-7] **M 141.9**, **d²⁵ 1.15g/ml**, **m -58°**, **b 126-129°/atm**; **BF₃·Et₂NH** [75-23-0] **M 112.9**, **m 85-89°**; **BF₃·MeOH** [373-57-9] **M 99.9**, **d²⁵ 1.203g/ml**, **b 59°/4mm**; **BF₃·Me₂O** [353-42-4] **M 113.9**, **d²⁵ 1.239g/ml**, **m -15°**, **b 126-127°/atm**; **BF₃·Me₂S** [353-43-5] **M 129.9**, **d²⁵ 1.235g/ml**; **BF₃·H₃PO₄** [13669-76-6] **M 165.8**, **d²⁵ 1.84g/ml**, **b 147°/atm**; **BF₃·n-PrOH** [762-48-1] **M 127.9**, **d²⁵ 1.379g/ml**; **BF₃·THF** [462-34-0] **M 139.9**, **d²⁵ 1.268g/ml**, **m -123°**, **b 180°/atm**. **TOXIC**.

Bromine [7726-95-6] **Br₂**, **M 159.8**, **m -7.2°**, **b 58.8°/atm**, **59°/atm**, **d₄²⁰ 3.102**, **n_D²⁰ 1.661**. Reflux the brown liquid with solid KBr and distil; dry the distillate by shaking it with an equal volume of conc H₂SO₄, then redistil it. The H₂SO₄ treatment can be replaced by direct distillation from BaO or P₂O₅. A more extensive purification [Hildenbrand et al. *J Am Chem Soc* **80** 4129 1958, DOI: 10.1021/ja01549a004] is to reflux about 1L of bromine for 1 hour with a mixture of 16g of CrO₃ in 200ml of conc H₂SO₄ (to remove organic material). The bromine is distilled into a clean, dry, glass-stoppered bottle, and chlorine is removed by dissolving *ca* 25g of freshly fused CsBr in 500ml of the bromine and standing overnight. To remove HBr and water, the bromine is then distilled back and forth through a train containing alternate tubes of MgO and P₂O₅. [Schmeisser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 275 1963.] Also available commercially are **bromine solution** [7726-95-6] 1.0M in **trimethyl phosphate**, **volumetric standard** bromine 0.0950-0.1050 N in H₂O, and **bromine on polymer support** on Amberlite IRA-900 Br₃⁻ form (loading: 1.2-1.8.mmol/g active bromine (Br₃⁻) [Cacchi et al. *Synthesis* 64 1979, DOI: 10.1055/s-1979-28560; Bongini et al. *Synthesis* 143 1980, DOI: 10.1055/s-1980-28952; Smith et al. *JCS Perkin Trans 1* 1877 1992, DOI: 10.1039/P19920001877].

HIGHLY TOXIC.

Bromine pentafluoride [7789-30-2] BrF_5 , **M 174.9**, **m** -60.5° , **b** 41.3° , **40.25** $^\circ$, **d**₄²⁵ **2.466**. Purify it *via* its KF complex, as described for chlorine trifluoride. [Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 158-159 1963.] While working on a metal vacuum line excess fluorine is removed by passage through a trap at -134° , the reactor is heated to 60° at the end to drive off any product adsorbed on the solids. BrF_5 is then transferred to a metal storage cylinder. It is a **powerful oxidant**, reacts violently with H_2O to give HBrO_3 and HF , and is an extremely effective **fluorinating agent**. Its IR has ν_{max} at 587, 644 and 683 cm^{-1} [Hyde & Boudakian *Inorg Chem* **7** 2648 1968, DOI: 10.1021/ic50070a039; for IR and Raman see Begun et al. *J Chem Phys* **42** 2236 1965, DOI: 10.1063/1.1696273.] **HIGHLY TOXIC.**

Cadmium [7440-43-9] **Cd**, **M 112.4**, **m** **320.9** $^\circ$, **321.1** $^\circ$, **b** **765** $^\circ$ /atm, **767** $^\circ$ /atm, **d**₄²⁰ **8.642**. Any oxide contaminant is removed by filtering the molten metal, under vacuum, through quartz wool. Its solubility in Hg is 5.2% (18°), and it is soluble in mineral acids. [Wagenknecht & Juza in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **11** p 1092 1965.] All Cadmium compounds are considered **carcinogenic** or potentially **carcinogenic**.

Cadmium bromide [7789-42-6 (anhydrous), 13464-92-1 ($4\text{H}_2\text{O}$)] $\text{CdBr}_2 \cdot 4\text{H}_2\text{O}$, **M 344.3**, **m** **566** $^\circ$, **b** **863** $^\circ$, **963** $^\circ$, **d**₄²⁰ **5.192**. Crystallise it from water (0.57g/ml at 10° , 1.26g/ml at 100°) between and 0° , and dry it at 110° . It forms the **monohydrate** below 36° and the **tetrahydrate** above 36° . It is **hygroscopic**. [Wagenknecht & Juza in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1096 1965.] **Carcinogenic.**

Cadmium chloride [10108-64-2, 7790-78-5 ($2.5\text{H}_2\text{O}$), 654054-66-7 ($x\text{H}_2\text{O}$)] CdCl_2 , **M 183.3**(anhydr), **m** **568** $^\circ$, **b** **960** $^\circ$ /atm, **d**₄²⁰ **4.06**. Crystallise it from water (1ml/g) by addition of EtOH and cooling. [Pray et al. *Inorg Synth* **5** 153 1957, DOI: 10.1002/9780470132364.ch43; Wagenknecht & Juza in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1093 1965.] **Carcinogenic.**

Cadmium fluoride [7790-79-6] CdF_2 , **M 150.4**, **m** **>1000** $^\circ$, **1049** $^\circ$, **b** **1748** $^\circ$ /atm, **d**₄²⁰ **6.35**. Crystallise it by dissolving it in hot water (25ml/g at 25°) at 60° , filtering, then cooling. It is soluble in mineral acids and HF but insoluble in EtOH and liquid NH_3 . [Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 243 1963.] **Carcinogenic.**

Cadmium iodide [7790-80-9] CdI_2 , **M 366.2**, **m** **388** $^\circ$, **b** **787** $^\circ$ /atm, **d**₄²⁰ **5.66**. Crystallise it from ethanol (2ml/g) by partial evaporation, or from H_2O [solubility is 78.7g/100ml (0°), 84.7g/100ml (20°), 125.0g/100ml (100°)]. The salt is also soluble in Et_2O , Me_2CO and in NH_3 , and turns yellow on exposure to light. It is used in the production of phosphors, in lithography, in photography, in electroplating, in analytical chemistry and as a nematocide. [Wagenknecht & Juza in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1096 1965.] **Carcinogenic.**

Cadmium nitrate tetrahydrate [10022-68-1, 10325-94-7 (anhydrous)] $\text{Cd}(\text{NO}_3)_2 \cdot 4\text{H}_2\text{O}$, **M 308.5**, **m** **59.5** $^\circ$ (hydrate), **b** **132** $^\circ$ /760mm, **d**₂₅²⁵ **3.6g/cm³** (anhydrous), **d**₂₅²⁵ **2.45g/cm³** ($4\text{H}_2\text{O}$). Crystallise the nitrate from water [solubility is 1.1g/ml (0°), 1.3g/ml (18°), 1.4g/ml (30°), 3.2g/ml (60°)] by cooling in an ice-salt bath. The salt is also soluble in Et_2O , EtOAc, Me_2CO , ROH, dilute acids and in NH_3 . Used as a flash in photography and emulsions; and for colouring glass and porcelain. [Gmelin's *Cadmium* (8th edn) **33** pp 76-78 1925, Suppl p 446 1959.] **Carcinogenic.**

Cadmium potassium iodide [13601-63-3] KCdI_3 , **M 532.2**. Crystallise it from ethanol by partial evaporation. **Carcinogenic.**

Cadmium selenide [1306-24-7] CdSe , **M 191.4**, **m** **1268** $^\circ$, **d**₂₅²⁵ **5.816g/cm³**, **n**_D²⁵ **2.5**. It is prepared by heating Cd in a stream of H_2Se (foul odour, work in a well ventilated fume cupboard), then subliming it under H_2 at dull red heat. Alternatively, CdSe is precipitated from an aqueous CdSO_4 solution and H_2Se (or a soluble metal

selenide) as for CdS below, and is almost insoluble in water (hence wash well with H₂O). It is a white to brown salt which turns red on exposure to sunlight. It exists in three forms: an unstable *sphalerite cubic form*, which is converted into the *wurzite form* (hexagonal) with a transition starting at 300° and completing at 700°; and a *rock-salt cubic* form only seen at high pressures. [Wagenknecht & Juza in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II p 1099 1965.] Bulk crystalline CdSe is prepared by the High-Pressure Vertical Bridgman (HPVB) method using high pressures (10-100atm) of inert gas [in for *Device for crystal growth by vertical direct crystallization of melt* see Kulakov & Fadeev Pat. USSR N 774286] or high-pressure vertical zone melting (HPVB) of crystalline selenide [for *HPVB and HPVZM shaped growth of CdZnTe, CdSe and ZnSe crystals* see Kolesnikov et al. *Proc SPIE* **4787** 93-104 2002]. For CdSe nanoparticle and quantum dots see Chapter 7. It is used in phosphors, semiconductors, photoelectric cells etc.

Cadmium sulfide [1306-23-6] CdS, **M 144.5**, **m 1750°**, **d²⁵ 4.826g/cm³**, **n_D²⁵ 2.529**. CdS is prepared by precipitation from an aqueous 0.1M solution of CdSO₄, Cd(NO₃)₂, CdCl₂, CdBr₂ or CdI₂ at 30° or 100° by adding S²⁻ ions in the form of H₂S or a sulfide salt, e.g. Na₂S. The colour of the precipitate varies from yellow to orange to red depending on the original salt used and on whether the original solution is neutral or acidic, e.g. by addition of a few drop of the respective concentrated acid, e.g. H₂SO₄, HNO₃, HCl, HBr or HI respectively. At the lower or higher temperature the cubic or the hexagonal form separate. This is a test for Cd in qualitative analysis. The salt is collected, washed well with H₂O until no salts filter through, then washed with EtOH, Et₂O and is dried in air at room temperature or preferably in a vacuum. Commercial salts should be washed free from other salts as stated and dried. The differences in colour are due to particle size. When used as a pigment (**Cadmium Yellow, CI pigment yellow 37**, cf. Ryan et al. US Pat 07/104,716 to Raychem Corp 2 Oct 1987) the dried salt is ground and calcined for an hour or so, but note that the temperature must be kept below 980°, as it sublimes above it. Its solubility in H₂O is 0.13w/w% at 25°, and it liberates H₂S in strong acid. **Carcinogenic**. It is a useful pigment for paints, plastics, is piezoelectric, used in solar cells when combined with other metals (e.g. Cu₂S), and for the preparation of nanomaterials see entry in Chapter 7. [Milligan *J Phys Chem* **38** 797 1934, DOI: 10.1021/j150357a009; Wagenknecht & Juza in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II pp 1098-1099 1965.] **TOXIC**.

Cadmium sulfate [7790-84-3 (for 3CdSO₄ 8H₂O), 15244-35-6] (2.67 H₂O), 10124-36-4 (anhydrous)] CdSO₄, **M 208.4** (anhydrous), **769.5** (hydrate), **m 40°** (8H₂O), **105°** (H₂O), **1000°** (anhydr), **d²⁵ 4.69g/cm³** (anhydr), **d²⁵ 3.79g/cm³** (H₂O), **3.08g/cm³** (8H₂O), **n_D²⁵ 1.565**. The sulfate crystallises from distilled water as a *hydrate* by partial evaporation in a desiccator. It gives the *monohydrate* on heating at 80°. Its solubility in H₂O is 0.75g/ml (0°), 0.76g/ml (25°), and 0.54g/ml (99°) for the *anhydrous salt*; 0.77g/ml (25°) for the *monohydrate salt*, and the *octahydrate salt* is freely soluble.] It is insoluble in EtOH, Me₂CO or EtOAc. It forms a white precipitate of Cd(OH)₂ with aqueous NH₃ which dissolves in excess of NH₃ to form soluble [Cd(NH₃)₄]SO₄. [Gmelin's *Cadmium* (8th edn) **33** p 121 1925, Suppl pp 609-610 1959.] **Carcinogenic**.

Cadmium telluride [1306-25-8] CdTe, **M 240.0**, **m 1041°**, **b 1130°/atm**, **d²⁵ 6.20g/cm³**, **n_D²⁵ 2.67**. It is prepared by heating stoichiometric amounts of Te and Se (both purified by sublimation *in vacuo*) in a crucible at 500° under a pressure of 40atm of N₂. On completion of the reaction, the temperature is raised above the melting point (1050°) and then decreased slowly as well as the N₂ pressure when an ingot of polycrystalline CdTe is obtained. This is purified by recrystallising from the melt under 1atm pressure of Cd. Zone melting under such conditions gives spectroscopically purer brown-black crystals [For *Preparation and Electrical Properties of CdTe Single Crystals* see Kröger & Nobel *J Electronic & Control* **1** 190 1955, DOI: 10.1080/00207215508961407; for prep and phys props of H₂Te see Dennis & Anderson *J Am Chem Soc* **36** 882 1914, DOI: 10.1021/ja02182a012] CdTe is insoluble in H₂O, but decomposes slowly on prolonged exposure to moist air. Thin CdTe films, prepared from pure Cd and Te in an evacuated Vycor tube, can be deposited by condensation across the shot gaps between Aquadag electrode in Dewar tubes. [Kretschmar & Schilberg *J Appl Phys* **28** 865 1957, DOI: 10.1063/1.1722876.] Used in photoconductors and solar cells, and for the preparation of nanomaterials see entry in Chapter 7. **Carcinogenic**.

Calcium [7440-70-2] CdSO₄, **M 40.1**, **m 842°**, **845°**, **b 1484°/atm**, **d²⁵ 1.55g/cm³**, **d⁸⁴⁵ 1.378g/cm³** (liquid). Clean the silver white soft metal by washing it with ether to remove adhering paraffin, file the surface in an argon-filled glove box, and wash it with ethanol containing 2% of conc HCl. Then wash it with dry ethanol, dry

it in a vacuum and store it under pure argon [Addison et al. *J Chem Soc* 3868 1962, DOI: 10.1039/JR9620003868]. Ca sublimates below 800° at high vacuum, oxidises slowly in moist air and reacts with cold H₂O slowly.

Calcium bromide monohydrate [62648-72-0, 71626-99-8 (*x*H₂O), 7789-41-5 (anhydrous)] **CaBr₂ · H₂O**, **M 217.9**, **d₄²⁰ 3.35**. Crystallise the bromide from EtOH or Me₂CO. It loses H₂O on heating, is anhydrous at 750°, then it loses Br₂ at higher temperatures. It is *deliquescent*. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 930 1963.]

Calcium carbonate (limestone, chalk) [471-34-1] **CaCO₃**, **M 100.1**, **m 1339° (calcite), 825° (aragonite)**, **n_D²⁵ 1.59**, **d₂₅²⁵ 2.711g/cm³ (calcite), d₂₅²⁵ 2.83g/cm³ (aragonite)**, **pK₂₅²⁵ 9.0**. Pure CaCO₃ is produced from pure quarried *marble* (as for food or pharmaceuticals). It is *quite insoluble* in H₂O (0.013g/cm³ at 25°, pK_{sp} 3.3x10⁻⁹) and is best prepared from pure CaO, adding distilled H₂O [to give Ca(OH)₂] and bubbling CO₂ through the solution, collecting the white precipitate, washing thoroughly with H₂O until the filtrate is no longer alkaline and dried *in vacuo* to give *precipitated calcium carbonate (PCC)* [Ropp in *Encyclopedia of the alkaline earth compounds* Elsevier pp 359–370. ISBN 9780444595508]. The carbonate exists in equilibrium with CaO and CO₂ and is best calcined above ~898° at atmospheric pressure (101kPa or 760mm) when CO₂ is *outgassed* and pure CaO (quicklime) is formed. It effervesces with dilute mineral acids to give CO₂. [Fieser **1** 103, **2** 57, **4** 67, **5** 89; Rohleder & Kroker *Calcium Carbonate: From the Cretaceous Period Into the 21st Century* Springer Science & Business Media. 2001, ISBN 3-7643-6425-4.]

Calcium chloride (anhydrous) [10043-52-4] **CaCl₂**, **M 111.0**, **m 772°**, **b >1600°**, **d₄¹⁵ 2.15**. It is available as fused granules or cubic crystals. It is very *hygroscopic*, very soluble in H₂O (exothermic), and EtOH. Store it in a tightly closed container. Used for drying many organic solvents (except MeOH and EtOH) and extracts e.g. ethereal, chloroform extracts. Also for removing EtOH (added to inhibit formation of phosgene) from CHCl₃. Useful desiccant. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 931 1963.] For drying agent properties see Chapter 1.

Calcium chloride dehydrate [10035-04-8] **CaCl₂ · 2H₂O**, **M 147.0**, **m 175°(dehydrates), 772°(dec)**. Crystallise it from ethanol. It is *hygroscopic*. It loses H₂O at 200° so it can be dried at high temperatures to dehydrate it. The *hexahydrate* [7774-34-7] has **m 30°** and **d 1.67**. [Fieser **1** 104, **4** 368.]

Calcium dithionite (calcium hydrosulfite) [13812-88-9, 15512-36-4] **CaS₂O₄**, **M 168.2**, **m dec on heating**, **d₂₅²⁵ 2.83g/cm³**. Crystallise it from water, or water followed by acetone and dry it in air at room temperature. Reducing agent which may *ignite* in contact with moist air. If spilled into excess H₂O (at least 5 fold) half the theoretical yield of SO₂ gas will evolve in ~7minutes. Reaction with oxidising agents could be violent. [Development of the Table of Initial Isolation and Protective Distances for the 2008 Emergency Response Guidebook, ANL/DIS-09-2, D.F. Brown et al. Argonne National Laboratory, Argonne, Illinois, June 2009.]

Calcium hexacyanoferrate (II) undecahydrate (yellow prussiate of lime) [13821-08-4] **Ca₂[Fe(CN)₆] · 11H₂O**, **M 490.1**, **292.1(anhydr)**. Recrystallise it three times from conductivity H₂O and dry it in air to constant weight over the partially dehydrated salt. [James *Trans Faraday Soc* **45** 855 1949, DOI: 10.1039/TF9494500855.] Alternatively, the Ca salt can be purified by precipitation with absolute EtOH in the cold (to avoid oxidation) from an air-free saturated aqueous solution. The pure lemon yellow crystals are centrifuged, dried in a vacuum desiccator first over dry charcoal for 24 hours, then over partly dehydrated salt and stored in a dark glass stoppered bottle. No deterioration occurs after 18 months. No trace of Na, K or NH₄ ions can be detected in the salt from the residue after decomposition of the salt with conc H₂SO₄. Analyses indicate 11mols of H₂O per mol of salt. The solubility in H₂O is 36.45g (24.9°) and 64.7g (44.7°) per 100g of solution. [Farrow *J Chem Soc* 49 1926, DOI: 10.1039/JR9262900049.]

Calcium hydroxide [1305-62-0] **Ca(OH)₂**, **M 74.1**, **m loses H₂O on heating**, **d₂₅²⁵ 2.24g/cm³**, **pK₂₅²⁵ 12.7 (for Ca²⁺)**. Heat analytical grade calcium carbonate at 1000° during 1 hour. Allow the resulting oxide to cool and add slowly to water. Heat the suspension to boiling, cool and filter through a sintered glass funnel of medium

porosity (to remove soluble alkaline impurities). Dry the solid at 110° and crush it to a uniformly fine powder. Store it in a tightly stoppered bottle as it absorbs CO₂ from air to form CaCO₃. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 934 1963.]

Calcium iodate [7789-80-2 (anhydrous), 10031-33-1 (6H₂O)] **Ca(IO₃)₂**, **M 389.9**, **m >540°**, **d²⁵ 4.520g/cm³**, **pK²⁵ 0.79 (for HIO₃)**. Crystallise it from water (100ml/g at 100° and 100ml/0.1g at 0°) [Bahl & Singh *J Indian Chem Soc* **17** 397 1940]. Used as iodine supplement, in lotions and ointments as antiseptic and deodorant.

Calcium iodide x-hydrate [71626-98-7 (xH₂O), 10102-68-8 (anhydrous)] **CaI₂**, **M 293.9 (anhydr)**, **m 740°, 779°, b 1100°/atm**, **d²⁵ 3.956g/cm³**. Dissolve the salt in acetone, which is then diluted and evaporated. This drying process is repeated twice, then the CaI₂ is recrystallised from acetone/diethyl ether and stored over P₂O₅. It is very *hygroscopic* when anhydrous and is light sensitive [Cremlyn et al. *J Chem Soc* 520 528 1958, DOI: 10.1039/JR9580000520]. The yellow *hexahydrate* has **m 42°**. It is soluble in H₂O, MeOH, EtOH and Me₂CO but insoluble in Et₂O.

Calcium nitrate tetrahydrate [13477-34-4 (4H₂O), 35054-52-5 (xH₂O), 10124-37-5] **Ca(NO₃)₂**, **M 236.1(4H₂O)**, **164.1(anhydr)**, **m 42.7°, 45°(hydrate, dec)**, **561°(anhydrous)**, **d²⁵ 2.504g/cm³(anhydr)**, **d²⁵ 1.896g/cm³(4H₂O)**, **pK²⁵ 6.0**. Crystallise the nitrate four times from water [solubility: 1.9g/ml (0°), 1.3g/ml (20°), 3.6g/ml (100°) for 4H₂O; 1.2g/ml (20°), 2.7g/ml (40°) for anhydrous salt] by cooling in a CaCl₂-ice freezing mixture. The *tetrahydrate* is dried over concentrated H₂SO₄ and stored over P₂O₅, to give the *anhydrous salt*. It is *deliquescent*. Its solubility in EtOH is 0.5g/ml (20°) and 0.6g/ml (40°); and 1.3g/ml (10°), 1.4g/ml (40°) and 1.4g/ml (60°) in MeOH; and 0.013g/ml (20°) in Me₂CO. It is soluble in NH₃ but almost insoluble in HNO₃. After 3 recrystallisations of ACS grade salt, it had Co, Fe, Mg, Sr and Zn at 0.2, 1.0, 0.02, 10 and 0.02 ppm resp. [Bassett & Taylor *J Chem Soc* **105** 1926 1914, DOI: 10.1039/CT9140501926.]

Calcium nitrite dihydrate [13780-06-8 (30% w/w aqueous solution)] **Ca(NO₂)₂ · 2H₂O**, **M 168.1(2H₂O)**, **132.1(anhydr)**, **m 390° (dec on heating)**, **d_D²⁰ 2.22**, **n_D²⁰ 1.3946**. Crystallise it from hot water (1.4ml/g) by adding ethanol and cooling to give the *hydrate*. It is *deliquescent*. [Ray & Ogg *J Am Chem Soc* **79** 265 1957, DOI: 10.1021/ja01559a003.] It has many applications, e.g. rust inhibitor of steel, antifreeze, large scale hydraulic, heavy oil detergent.

Calcium permanganate tetrahydrate [10118-76-0 (anhydrous)] **Ca(MnO₄)₂ · 4H₂O**, **M 350.0 (for 4H₂O)**, **277.9(anhydr)**, **m 140°(dec)**, **d²⁵ 2.49g/cm³**. Crystallise the purple solid from water [3.3g/ml (14°), 3.4g/ml (25°)] by partial evaporation in a desiccator. It is *deliquescent*. It is soluble in NH₄OH and decomposes in EtOH. *Note* that it loses oxygen more readily than the potassium salt. It is an oxidising agent.

Calcium sulfate dihydrate [10101-41-4] **CaSO₄ · 2H₂O**, **M 172.2**, **m 150°(dec)**, **d₄²⁰ 2.32**, **pK²⁵ 7.3**, **K_{sp} 3.14 x 10⁻⁵ mol².L⁻²**. It loses only part of its H₂O at 100-150° (see below). It is soluble in H₂O and very slowly soluble in glycerol. It is insoluble in most organic solvents.

Calcium sulfate hemihydrate [10034-76-1] **CaSO₄ · 0.5H₂O**, **M 145.2**. Its solubility in H₂O is 0.2parts/100 at 18.75°. It dehydrates completely >650°. Dry it below 300° to give a solid with estimated pore size *ca* 38% of volume. *Anhydrous* CaSO₄ (**Drierite**) [7778-18-9] **M 136.1**, **d²⁵ 2.96g/cm³**, **m 1460°**, **pK²⁵ 10.4**, **K_{sp} 4.93 x 10⁻⁵ mol².L⁻²**, has a high affinity for H₂O and will absorb 6.6% of its weight of H₂O to form the *hemihydrate* (**gypsum**). It sets to a hard mass with H₂O; hence it should be kept in a tightly sealed container. The solubility of gypsum in H₂O is unusual: 0.176% at 0°, 0.209% at 30°, 0.210% at 40°, 0.204% at 50° and 0.200% at 60°. [Hulett *J Am Chem Soc* **27** 49 1905, DOI: 10.1021/ja01979a008; James & Partington *J Chem Soc* **107** 1019 1915, DOI: 10.1039/CT9150701019; Namba *J Soc Chem Ind* **40** 2797 1920, D.R. Linde (ed.) *CRC Handbook of Chemistry and Physics* 83rd Edition, CRC Press, 2002]

Calcium thiosulfate (hypo, hydrosulfite) [10124-41-1] **CaS₂O₃**, **M 152.2**, **m 43-49°**, **pK₁²⁵ 0.6**, **pK₂²⁵ 1.74 (for H₂S₂O₃)**. Recrystallise the thiosulfate from water below 60° in a N₂ atmosphere, followed by drying with EtOH and Et₂O. Store it in a refrigerator. The *hexahydrate* can decompose spontaneously at 43-49°. Store it in a cool closed container. A ~6% aqueous solution is a useful sulfur fertiliser. [Pethybridge & Taba *JCS Faraday*

Trans I **78** 1331 1982, DOI: 10.1039/F19827801331.]

Carbon dioxide [124-38-9] CO_2 , M 44.0, m -56.6° , sublimates at -78.5° , triple point 5.1atm at -0° , d 1562kg/m^3 (solid at -78.5°), d 770kg/m^3 (liquid at 56atm and 20°), d 1.977kg/m^3 (gas at 1atm and -56.6° ; 1.67 times that of air), pK_1^{25} 6.35, pK_2^{25} 10.33 (for carbonic acid: H_2CO_3). Pass the gas over CuO wire at 800° to oxidise CO and other reducing impurities (such as H_2), then over copper dispersed on Kieselguhr at 180° to remove O_2 . Drying it at -78° removes the water vapour. Final purification is by vacuum distillation at liquid nitrogen temperature to remove non-condensable gases [Anderson et al. *J Chem Soc* 3498 1962, DOI: 10.1039/JR9620003498]. Sulfur dioxide contaminant can be removed at 450° using silver wool combined with a plug of platinised quartz wool. Halogens are removed by using Mg, Zn or Cu, heated to 450° . [Glemser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 647 1963, Fieser **3** 40, and in 5 later volumes.] At temperature and pressures above the critical point (31.1° at 7.38MPa), CO_2 is a **supercritical fluid** in which chemical reactions can be performed. Solid CO_2 is commercially available as **Dry-Ice**, and in organic solvents is used in cooling baths (see Table 8, Materials for Cooling Baths in Chapter 1).

Carbon disulfide [75-15-0] CS_2 , M 76.1, m -112 to -111° , -110.8° , b 46.3° , d 1.264 , n_D^{20} 1.627. Shake it for 3 hours with three portions of KMnO_4 solution (5g/L), twice for 6 hours with mercury (to remove sulfide impurities) until no further darkening of the interface occurs, and finally with a solution of HgSO_4 (2.5g/L) or cold, saturated HgCl_2 . Dry it with CaCl_2 , MgSO_4 , or CaH_2 (with further drying by refluxing over P_2O_5), followed by fractional distillation in diffuse light. **Alkali metals cannot be used as drying agents.** It has also been purified by standing with bromine (0.5ml/L) for 3-4 hours, shaking rapidly with KOH solution, then copper turnings (to remove unreacted bromine), and drying with CaCl_2 . CS_2 is highly **TOXIC** has a **foul odour** and is highly **FLAMMABLE**. *Work in a good fumehood.*

Small quantities of CS_2 have been purified (including removal of hydrocarbons) by mechanical agitation of a 45-50g sample with a solution of 130g of sodium sulfide in 150ml of H_2O for 24 hours at $35-40^\circ$. The aqueous sodium thiocarbonate solution is separated from unreacted CS_2 , then precipitated with 140g of copper sulfate in 350g of water, with cooling. After filtering off the **copper thiocarbonate**, it is decomposed by passing steam into it. The distillate is separated from H_2O and distilled from P_2O_5 . [Ruff & Golla *Z Anorg Chem* **138** 17 1924, DOI: 10.1002/zaac.19241380103; Fieser **1** 114, **5** 94, **6** 95; *Beilstein* **3** IV 395.] Available commercially in various grades of purity including *Spectroscopic grades*.

Carbon monoxide [630-08-0] CO , M 28.0, m -205° , -200° , b $-191.5^\circ/\text{atm}$, d 1.282g/cm^3 , d 789kg/m^3 (liquid), d 1250kg/m^3 (at $0^\circ/1\text{atm}$), d 1145kg/m^3 (at $25^\circ/1\text{atm}$), n_D^{25} 1.0003364. Iron carbonyl is a likely impurity in CO stored under pressure in steel tanks. It can be decomposed by passing the gas through a hot porcelain tube at $350-400^\circ$. Passage through alkaline pyrogallol solution removes oxygen (and CO_2). Removal of CO_2 and water are effected by passage through soda-lime followed by $\text{Mg}(\text{ClO}_4)_2$ or P_2O_5 and collected over Hg. Carbon monoxide can be condensed and distilled at -195° . It is sparingly soluble in H_2O (27.6g/L at 25°), but is readily absorbed by a solution of CuCl in HCl to give the white crystalline **adduct** **$\text{CuCl}\cdot\text{CO}\cdot 2\text{H}_2\text{O}$** . It burns in air with a bright blue flame but a mixture of 2 volumes of CO and 1 volume of O_2 explode when kindled, although in a small jar the combustion is not violent. **HIGHLY POISONOUS** gas as it reacts with haemoglobin to form **bright red carboxyhaemoglobin** which is stable and not readily decomposed by oxygen. The **TOXIC LEVEL** of CO is 50ppm ($\sim 55\text{mg/m}^3$). **The ANTIDOTE should be at hand and always available in laboratories using CO; and staff should be trained to administer it.** [Gilliland et al. *Inorg Synth* **2** 81 1946, DOI: 10.1002/9780470132333.ch22; Glemser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 645-646 1963.]

Carbonyl bromide (bromophosgene) [593-95-3] COBr_2 , M 187.8, b $64.5^\circ/760\text{mm}(\text{dec})$, d $2.52\text{g/cm}^3(15^\circ)$. Purify it by distillation from Hg and from powdered Sb to remove free bromine, then distil it in a slight vacuum to remove volatile SO_2 (the major impurity) [Carpenter et al. *JCS Faraday Trans 2* 384 1977, DOI: 10.1039/F2977300384]. **TOXIC.**

Carbonyl sulfide [463-58-1] COS , M 60.1, m -138° , b $-47.5^\circ/\text{atm}$ $-50.2^\circ/\text{atm}$, d 2.51g/L . Purify the **sulfide-like smelling** gas by scrubbing it through three consecutive fritted washing flasks containing conc NaOH at 0° (to remove HCN), and then through concentrated H_2SO_4 (to remove CS_2) followed by a mixture of NaN_3

and NaOH solution; or passed through traps containing saturated aqueous lead acetate, then through a column of anhydrous CaSO_4 . Then it is freeze-pumped repeatedly and distilled through a trap packed with glass wool and cooled to -130° (using an *n*-pentane slurry). It liquefies at $0^\circ/12.5\text{mm}$. Use stainless steel containers. The gas is stored over conc H_2SO_4 , because in the presence of moisture and bases it decomposes to CO and H_2S . [Glemser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 654 1963.] **TOXIC**

Ceric (IV) ammonium nitrate [16774-21-3] $\text{Ce}(\text{NH}_4)_2(\text{NO}_3)_6$, **M 548.2**, **pK₁²⁵ -1.15**, **pK₂²⁵ -0.72**, **pK₃²⁵ 1.68**, **pK₄²⁵ 2.29 (for aquo Ce^{4+})**. Ceric ammonium nitrate (125g) is warmed with 100ml of dilute HNO_3 (1:3 v/v) and 40g of NH_4NO_3 until it dissolves, and filtered through a sintered-glass funnel. The orange-red solid which separates on cooling in ice is filtered off on a sintered funnel (at the pump) and air is sucked through the solid for 1-2 hours to remove most of the nitric acid. Finally, the solid is dried at $80-85^\circ$. A primary standard in **iodometry** [Smith & Fly *Anal Chem* **21** 1233 1949, DOI: 10.1021/ac60034a027; Smith *Talanta* **10** 709 1963, DOI: 10.1021/ac60034a027]. This nitrate is also commercially available as a silica gel supported oxidising agent [Hwu et al. *J Org Chem* **65** 5077 2000, DOI: 10.1021/jo000024o].

Cerium(III) sulfate [cerous sulfate,] [13454-94-9 anhydrous; 10450-59-6 $8\text{H}_2\text{O}$; 13550-47-5 $12\text{H}_2\text{O}$] $\text{Ce}_2(\text{SO}_4)_3$, **M 568.4** (anhydr), **m 920° (dec)**, **d₄²⁵ 2.89**, **pK²⁵ 9.29 (for hydrolysis of Ce^{3+})**. To prepare cerous sulfate, cerium oxide [CeO_2 , 0.3g] or chloride is dissolved in hot 6N H_2SO_4 (20ml), filtered through a sintered glass funnel and allowed to crystallise over concentrated H_2SO_4 in a vacuum dessicator. The crystals are filtered off, washed twice with H_2O (10ml) and once with EtOH (10ml), and dried in air for 4 hours to provide the **pentahydrate**. Alternatively, a neutral or slightly acidic (with H_2SO_4) aqueous solution of the sulfate is treated with $\frac{3}{4}$ of its volume of EtOH. The salt is thus obtained rapidly and quantitatively without the evaporation stage. The **anhydrous** sulfate is formed by heating any sulfate hydrate salt at $400-500^\circ$, but is **hygroscopic**. The hydrate loses most of its H_2O at *ca* 250° , and above 650° it loses SO_3 also to give the basic salt. [Wendlandt *J Inorg Nucl Chem* **7** 51 1958, DOI: 10.1016/0022-1902(58)80026-3; Wetzel in *Handbook of Preparative Inorganic Chem* (Ed. Brauer) Academic Press Vol **11** p 1156 1965.] When a fairly concentrated solution of the anhydrous salt in ice H_2O , is allowed to evaporate at $40-50^\circ$, the **octahydrate salt** separates as colourless rhombic pyramids [Koppel *Z Anorg Chem* **41** 377 1904, DOI: 10.1002/zaac.19040410123]. The **dodecahydrate** is obtained in fine needles when a clear fairly concentrated solution of cerous sulfate is allowed to evaporate over concentrated H_2SO_4 in a desiccator in a refrigerator [Koppel *Z Anorg Chem* **41** 377 1904, DOI: 10.1002/zaac.19040410123]. [Vanino *Handbuch der Präparativen Chemie* Vol **I**, 3rd Edn, pp 754-755, Stuttgart 1925.]

Cerium(IV) sulfate [ceric sulfate,] [13590-82-4 anhydrous] $\text{Ce}(\text{SO}_4)_2$, **M 332.2**, **d₄²⁵ 3.01**, **pK₁²⁵ -1.15**, **pK₂²⁵ -0.72**, **pK₃²⁵ 1.68**, **pK₄²⁵ 2.29 (for hydrolysis of Ce^{4+})**. Ceric sulfate is prepared by heating pure ceric oxide (CeO_2) with an excess of concentrated H_2SO_4 for an hour, cooling, adding glacial acetic acid, stirring, allowing to settle and decanting off. The process is repeated several times and the yellow orthorhombic crystals of the sulfate are filtered off (sintered glass), washed with glacial acetic acid and dried in a vacuum desiccator over NaOH/KOH and/or soda lime. [Vanino *Handbuch der Präparativen Chemie* Vol **I**, 3rd Edn, pp 754-755, Stuttgart 1925, Meyer & Aufrecht *Chem Ber* **37** 140 1904, DOI: 10.1002/cber.19040370124.] It is a strong **oxidising agent** (oxidation potential of a 1 to 8N H_2SO_4 solution at 25° is 1.42 ± 0.1 volts), and like ceric ammonium sulfate is used in volumetric analysis. Solutions in dilute H_2SO_4 are standardised by titration with pure arsenious oxide, iron or ferrous ammonium sulfate. Although it is its own end-point indicator (Ce^{4+} ions are yellow and Ce^{3+} ions are colourless), the end point is not as sharp as that of permanganate, and it is best to use 0.005M *N*-phenylanthranilic acid as indicator (colour change from yellow-green to purple at end point) [J. Mendham, R.C. Denney, J.D. Barnes and M.J.K. Thomas, *Vogel's Quantitative Chemical Analysis*, 6th Edn, Prentice Hall, Harlow, 2000, ISBN 0582226287].

Ceric sulfate tetrahydrate is obtained thus: pure ceric ammonium nitrate [16774-21-3] in a small volume of H_2O is treated with ammonia, $\text{Ce}(\text{OH})_4$ separates, is collected and is dissolved in concentrated H_2SO_4 . This solution is evaporated to dryness and the residue is crystallised from H_2O (perhaps a small amount of H_2SO_4 should be added to avoid hydrolysis). A small volume of H_2O should be used as boiling with a large volume of H_2O leads to separation of the basic salt. The yellow-orange powder or orthorhombic crystals lose H_2O to give the anhydrous salt at $180-200^\circ$, but decomposes $>350^\circ$ to give the basic salt CeOSO_4 . [Vanino *Handbuch der Präparativen Chemie* Vol **I**, 3rd Edn, pp 755-756, Stuttgart 1925, Muthmann & Stützel *Chem Ber* **33** 1763 1900,

DOI: 10.1002/cber.19000330255.]

Cesium bromide [7787-69-1] CsBr , **M 212.8**, **m 636°**, **b ca 1300°/atm**, **d₄²⁰ 4.44**. It is very soluble in H_2O , soluble in EtOH but insoluble in Me_2CO . Dissolve it in the minimum volume of H_2O , filter and precipitate it by adding Me_2CO . Filter off the solid and dry it at 100°. Also recrystallise it from water [1.06g/ml (15°), 1.24g/ml (25°)] by partial evaporation in a desiccator. Used as a beam splitter in spectrometers.

Cesium carbonate [534-17-8] Cs_2CO_3 , **M 325.8**, **m 610°(dec)**, **792°(at red heat)**. Crystallise it from ethanol (10ml/g) by partial evaporation. [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 988 1963.] It promotes efficient *O*-alkylation of alcohols to produce alkyl carbonates, and is used for the aerobic oxidation of arylalkanes to aryl ketones [Park et al. *Synthesis* 3617 2006, DOI: 10.1055/s-2006-950189].

Cesium chloride [7647-17-8] CsCl , **M 168.4**, **m 645°**, **b 1303°/atm**, **d₄²⁰ 3.99**. It is soluble in H_2O but can be purified by crystallisation from H_2O [solubility in g percent: 162.3(0.7°), 182.2(16.2°) and 290(at bp 119.4°)] and dried in high a vacuum. It is soluble in EtOH and is *deliquescent*; keep it in a tightly closed container. [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 951-955 1963.] For further purification of CsCl , a concentrated aqueous solution of the practically pure reagent is treated with an equivalent weight of I_2 and Cl_2 is bubbled into the solution until precipitation of CsCl_2I is complete. Recrystallisation yields a salt that is free from other alkali metals. It is then decomposed to pure CsCl on heating. [Harned & Schupp *J Am Chem Soc* **52** 3886 1930, DOI: 10.1021/ja01373a019.] It can also be recrystallised from acetone/water, or from water (0.5ml/g) by cooling in a CaCl_2 /ice bath. Dry it at 78° under vacuum. It is a very useful salt for making solutions for the separation of RNA and DNA by density gradient centrifugation [Maniatis et al. *Molecular Cloning: A Laboratory Manual* Cold Spring Harbor Laboratory Press (Cold Spring Harbor) 1989, ISBN 0-87969-309-6].

Cesium chromate [56320-90-2] Cs_2CrO_4 , **M 381.8**, **d²⁵ 4.237g/cm³**, **pK₁²⁵ 0.74**, **pK₂²⁵ 6.49 (for H_2CrO_4)**. Crystallise the yellow chromate from water (0.71g/ml at 13°) by partial evaporation in a desiccator. [Boer et al. *Z Anorg Allgem Chem* **191** 113 1930, DOI: 10.1002/zaac.19301910114; Hein & Herzog in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **11** p 1389 1963.]

Cesium fluoride [13400-13-0] CsF , **M 151.9**, **m 682°**, **703°**, **b 1251°/atm**, **d²⁵ 4.115g/cm³**, **n_D²⁵ 1.477**. Crystallise it from aqueous solution by adding ethanol. The solubility in H_2O is 3.67g/ml at 18°. CsF chains with one or two atom thickness can be grown inside carbon nanotubes [Senga & Suenaga *Nature Commun* **6** 7943 2015, DOI: 10.1038/ncomms8943]. The strong Si—F bond makes CsF (soluble in dioxane, THF and DMF) a useful reagent for de-silylation reactions [see Smith et al. *Org Synth Coll Vol* **10** 107 2002, DOI: 10.15227/orysyn.078.0082]. CsF is a base in *Suzuki cross-coupling* of *ortho*-substituted biaryls [Li et al. *J Orgmet Chem* **691** 5688 2006, DOI: 10.1016/j.jorganchem.2006.09.023], and is employed as a reagent for nucleophilic fluorination of primary halides and sulfonates in protic media, e.g. *tert*-butyl and *tert*-pentyl alcohols [Kim et al. *J Am Chem Soc* **128** 16394 2006, DOI: 10.1021/ja0646895].

Cesium iodide [7789-17-5] CI , **M 259.8**, **m 621°**, **b~1280°/atm**, **d₄²⁰ 4.51**, **n_D²⁵ 1.739**. Crystallise from warm water (1ml/g) by cooling to -5°. Like CsF , it has been grown in carbon nanotubes [Senga et al. *Nature Materials* **13** 1050 2014, DOI: 10.1038/nmat4069], and used in phosphor screens for imaging, scintillators, calorimeters and various particle detectors [Kalivas et al. *Appl Phys A* **78** 915 2004, DOI: 10.1007/s00339-003-2089-5].

Cesium nitrate [7789-18-6] CsNO_3 , **M 194.9**, **m 414°(dec)**, **d₄²⁰ 3.65**. It crystallises from water [solubility: 0.2g/ml (~10°), 2.0g/ml (100°)] between 100° and 0°. After 1 crystallisation of 99.9% grade salt, it had K, Na and Se at 0.8, 0.4 and 0.2 ppm respectively. [Watt et al. *Inorg Synth* **4** 5 1953, DOI: 10.1002/9780470132357.ch2.]

Cesium perchlorate [13454-84-7] CsClO_4 , **M 232.4**, **m 250°(dec)**, **d²⁵ 3.327g/cm³**, **n_D²⁵ 1.4887**, **pK²⁵ -2.4 to -3.1 (for HClO_4)**. Crystallise it from water (4ml/g) between 100° and 0°. It decomposes above 250° to CsCl . Strong oxidant.

Cesium sulfate [10294-54-9] Cs_2SO_4 , M 361.9, m 1005°, 1010°, d_4^{20} 4.243. Crystallise it from water (0.5ml/g) by adding ethanol and cooling. It is very soluble in H_2O (1.7g/ml at 0°), but insoluble in most organic solvents. Used to prepare dense aqueous solutions for *density gradient* centrifugation of DNA. [Gmelin's, Cesium (8th edn) 25 pp 218-225 1938.]

Chlorine [7782-50-5] Cl_2 , M 70.9, m -101.5°, b -34.0°/atm, d_4^{20} 2.898. Pass the gas in succession through aqueous KMnO_4 , dilute H_2SO_4 , conc H_2SO_4 , and a drying tower containing $\text{Mg}(\text{ClO}_4)_2$. Or bubble it through water, dry it over P_2O_5 and distil it from bulb to bulb in a vacuum line. One volume of water dissolves 4.6 volumes of Cl_2 at 0°, 2.15 volumes at 20°, 1.22 volumes at 50° and 0.39 volumes at 90°. [Schmeisser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 272 1963.] **HIGHLY TOXIC.**

Chlorine trifluoride [7790-91-2] ClF_3 , M 92.5, m -76.34°, b 12.1°/atm, 11.75°/atm, d^{25} 4mg/cm³. Impurities include chloryl fluoride, chlorine dioxide and hydrogen fluoride. Passed it first through two U-tubes containing NaF to remove HF, then through a series of traps in which the liquid is fractionally distilled. It can be purified *via* the KF complex; KClF_4 , formed by adding excess ClF_3 to solid KF in a stainless steel cylinder in a dry-box and shaking overnight. After pumping out the volatile materials, pure ClF_3 is obtained by heating the bomb to 100-150° and condensing the evolved gas in a -196° trap [Schack et al. *Chem Ind (London)* 545 1967]. It attacks glass very vigorously, and reacts with H_2O violently to form HF, HCl and O_2 . **HIGHLY TOXIC.**

Chlorosulfonic acid (chlorosulfuric acid) [7790-94-5] ClSO_3H , M 116.5, m -80°, b 151-152°/750mm, d_4^{20} 1.753, n_D^{20} 1.4929, pK^{25} -5.9 (aqueous H_2SO_4). Distil the acid at atmospheric pressure in an all-glass apparatus, taking the fraction boiling at 156-158°. It reacts **EXPLOSIVELY** with water, **wear gloves and face protection**. [Cremlyn *Chlorosulfonic acid: A Versatile Reagent*, Royal Society of Chemistry UK, 2002, p 308, ISBN 0854044981, Fieser 1 140, 2 70, 6 121, Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 385 1963].

Chromium ammonium sulfate dodecahydrate [34275-72-4 (hydrate), 10022-47-6 ($12\text{H}_2\text{O}$), 13548-43-1 (anhydrous)] $\text{NH}_4\text{Cr}(\text{SO}_4)_2 \cdot 12\text{H}_2\text{O}$, M 478.4, m 94° loses $9\text{H}_2\text{O}$ then dehydrates at 300°, d_4^{20} 1.72. Crystallise the blue-violet double salt from a saturated aqueous solution at 55° by cooling slowly with rapid mechanical stirring. The resulting fine crystals are filtered on a Büchner funnel, partly dried on a porous plate, then equilibrated for several months in a vacuum desiccator over crude chromium ammonium sulfate (partially dehydrated by heating at 100° for several hours before use) [Johnston et al. *J Am Chem Soc* 75 3922 1953, DOI: 10.1021/ja01112a013].

Chromium (II) chloride (anhydrous) [10049-05-5] CrCl_2 , M 122.9, m 824°, d_4^{14} 2.75. It is obtained from the *dihydrate* by heating *in vacuo* at 180°. It is a very *hygroscopic* white powder which dissolves in H_2O to give a sky blue solution. It is stable in dry air but oxidises rapidly in moist air and should be stored in air tight containers. It sublimes at 800° in a current of HCl gas and should be cooled in the presence of HCl gas. *Alternatively*, it can be washed with air-free Et_2O and dried at 110-120°. [Burg et al. *Inorg Synth* 3 150 1950, DOI: 10.1002/9780470132340.ch40; Balthis et al. (4 H_2O) *Inorg Synth* 1 125 1939, DOI: 10.1002/9780470132326.ch46; Fieser 1 149, 2 76, 3 60, 4 144, 7 73, 9 511, 11 132, 12 136, 13 84, 14 94, 15 95, 16 93, 17 84; Hein & Herzog in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II pp 1336-1338 1965.] CrCl_3 is useful for the preparation of a very active *precatalyst* for ethylene polymerisation [Small et al. *Macromolecules* 37 4375 2004, DOI: 10.1021/ma035554b], promotes the condensation of aldehydes with trisubstituted chloroalkenes [Falek et al. *Tetrahedron Lett* 45 3039 2004, DOI: 10.1016/j.tetlet.2004.02.101], converts terminal olefins into *trans*-iodocyclopropanes in excellent yields when used with CHI_3 and tetra-alkyl ethylenediamine [Takai et al *J Organomet Chem* 692 520 2007, DOI: 10.1016/j.jorganchem.2006.06.047], and for the preparation of bis(indenyl)chromium dimer and related compounds [Heinemann et al. *Organometallics* 15 5462 1996, DOI: 10.1021/om960899m].

Chromium (III) chloride (anhydrous) [10025-73-7, 10060-12-5 ($6\text{H}_2\text{O}$)] CrCl_3 , M 158.4(anhydrous), m 1152°, b 1300°/atm dissociates, d^{25} 2.87g/cm³ (25°, anhydrous), d^{25} 1,760g/cm³ (hexahydrate), pK_1^{25} 3.95,

pK₂²⁵ 5.55, pK₃²⁵ 10.5 (for Cr³⁺). The chloride is purple when anhydrous and dark green when hexahydrated. Sublime the chloride in a stream of dry HCl. *Alternatively*, the impure chromic chloride (100g) is added to 1L of 10% aqueous K₂Cr₂O₇ and several millilitres of concentrated HCl, and the mixture is brought to a gentle boil with constant stirring for 10 minutes. (This removed a reducing impurity.) The solid is separated and washed by boiling with successive 1L lots of distilled water until the wash water no longer gives a test for chloride ion, then dry it at 110°. The **anhydrous** chloride is only slightly soluble in H₂O, dissolving slowly but more rapidly if a wetting agent is added. The **hexahydrate** has a solubility of 585g/L at ~25°, and X-ray analysis showed that it was **trans-dichlorotetraaquo chromium (III) chloride dihydrate 'Bjerrum-green'** [Dance & Freeman *Inorg Chem* **4** 1555 1965, DOI: 10.1021/ic50033a006]. [For preparation see Heisig et al. *Inorg Synth* **2** 193 1946, DOI: 10.1002/9780470132333.ch59; and Pray *Inorg Synth* **5** 153 1953, DOI: 10.1002/9780470132364.ch43]. [Poulsen & Garner *J Am Chem Soc* **81** 2615 1959, DOI: 10.1021/ja01520a005; Hein & Herzog in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1338 1965]. It has uses as a mordant, in tanning, as a waterproofing agent and corrosion inhibitor, as well as an olefin polymerising catalyst.

Chromium (III) nitrate nonahydrate [7789-02-8] Cr(NO₃)₃, M 400.2, m ~60° (dec >100°). The pure deep purple nonahydrate salt is best prepared freshly from pure recrystallised (3 times) chromium (VI) trioxide and pure nitric acid. A solution is prepared by dissolving 1g of CrO₃ in 3ml of H₂O and 2ml of pure HNO₃, and carefully, with stirring (behind a screen) pure MeOH (0.5-1.0ml) is added dropwise carefully (shield) with cooling to avoid a violent reaction. A bluish colour develops as Cr(VI) is reduced to Cr(III). When this solution is diluted to 130,000ppm of the salt, analysis detected by ICP/MS gave the following trace elements (ppm in brackets): Sr (33), Na (3.5), Mo, Ni, Cu, Si and Mg (0.13 each), Se, Zn, Al (0.026 each), Ti (1.5), Fe (4.9), Co (0.07), Sn (0.065), Ba (0.013 and W (0.078). Evaporation of the methanolic solution in high vacuum over CaCl₂ eventually yields pure deep violet rhombic crystals of chromium (III) nitrate **nonahydrate**. An aqueous solution of this salt becomes green on heating but reverts to the violet colour on cooling.

The pale green (blue-violet) **anhydrous chromium (III) nitrate**, [13548-38-4] **M 238.0, m dec >60°**, is best obtained pure by mixing a solution of chromium hexacarbonyl in CCl₄ with excess of N₂O₅ in CCl₄ under N₂ for 12 hours, whereby evolution of gasses occurs. Filter off the salt and wash it with CCl₄ in a closed system under N₂, and dry it *in vacuo*. It is very soluble in H₂O, EtOAc and Me₂SO but insoluble in *C₆H₆, CCl₄ and CHCl₃. The **deliquescent** powder reacts vigorously with Et₂O. [Addison & Chapman *J Chem Soc* 508 539 1964, DOI: 10.1039/JR9640000508.] **Cr(VI)** ions are **CARCINOGENIC** as they cause DNA breaks, and **Cr(III)** ions affect DNA synthesis.

Chromium (III) potassium sulfate dodecahydrate (chrome alum) [7788-99-0, 10141-00-6] CrK(SO₄)₂·12H₂O, M 499.4, m 89°, b 400°/atm, d²⁵ 1.38g/cm³, pK₁²⁵ 0.74, pK₂²⁵ 6.49 (for H₂CrO₄, chromic acid). Crystallise it from hot water (solubility: 0.24g/ml at 25°) by cooling (dark purple needles). Aqueous solutions are dark violet but turn green if heated above 50°. It is used in tanning leather and to harden photographic gel emulsions.

Chromium (III) oxide (chromium sesquioxide, Pigment Green 17, Chromia) [1333-82-0] Cr₂O₃, M 152.0, m 2435°, b 3000-4000°/atm, d²⁵ 5.22g/cm³, n_D²⁵ 2.551, CI 77288. This Cr(III) oxide is insoluble in H₂O, organic solvents, but is **amphoteric**, dissolving in acids to give the hydrated chromium cation [Cr(H₂O)₆]³⁺, and reacts in bases to give [Cr(OH)₆]³⁻ salts. In concentrated alkali it provides chromite ions. It is best prepared by decomposition of chromium salts, e.g. Cr(NO₃)₃ or ammonium dichromate [(NH₄)₂Cr₂O₇] (**exothermic**, a reaction with a low ignition temperature <200°, frequently used in '**volcano**' **demonstrations**). The oxide is a hard solid and useful as an abrasive in sharpening (stropping) the edges of knives. On heating it turns brown in colour but reverts to green on cooling. It is used as a pigment (Viridian or Chrome Green) in paints, for colouring glass, printing banknotes, and dyeing fabrics. [Gerd Anger et al. 'Chromium Compounds' in *Ullmann's Encyclopedia of Industrial Chemistry* Wiley-VCH, Weinheim, 2005, DOI: 10.1002/14356007.a07_067].

Chromium (IV) trioxide (chromic anhydride) [1333-82-0] CrO₃, M 100.0, m 197°, dec at 250° to Cr₂O₃, d²⁰ 2.70 (pK₁²⁵ 0.74, pK₂²⁵ 6.49, for H₂CrO₄, chromic acid). It forms red crystals from water between 100° and -5°, [solubility: 1.65g/ml (0°), 1.69g/ml (25°), 1.72g/ml (40°) and 1.98g/ml (100°)], or from water/concentrated HNO₃ (1:5). It separates when potassium or sodium dichromate are dissolved in concentrated H₂SO₄. Filter it

off using a sintered glass funnel. Dry it in a vacuum desiccator over NaOH pellets. It is dark purple under anhydrous conditions. It is a **hygroscopic**, powerful oxidant and can **ignite** with organic compounds. [Fieser **1** 44, **1** 144, **2** 72, **3** 54, **4** 96, **5** 140, **7** 70, **9** 115, **10** 99, **12** 131, **15** 92, **17** 86; Keyes et al. *Industrial Chemicals* (Lowenheim & Moran eds.) 4th edn J. Wiley pp 270-274 1975.] $\text{CrO}_3\text{-NH}_4\text{Cl}$ oxidises hydrobenzoin to the respective benzils in CHCl_3 in high yields at $\sim 25^\circ$ under ultrasound irradiation [Li & Sun *Lett Org Chem* **3**(11) 842 2006, DOI: 10.2174/157017806779116987]. It is a skin and pulmonary **IRRITANT**, and a **CANCER SUSPECT**.

Chromyl chloride [14977-61-8] CrO_2Cl_2 , **M 154.9**, **m -96.5°** , **b $115.7^\circ/\text{atm}$** , **$117^\circ/\text{atm}$** , **d²⁵ 1.911**. Purify it by distillation under reduced pressure. It hydrolyses violently with H_2O and is a powerful **oxidant** which explodes with P, and **ignites** in contact with S, NH_3 , EtOH and many organic compounds. [Freeman et al. 'Aldehydes by Oxidation of Terminal Olefins with Chromyl Chloride: 2,4,4-Trimethylpentanal' *Org Synth Coll.* Vol **6**, 1028 1988, DOI: 10.15227/orgsyn.051.0004.] Its formation is used as a specific test for chloride: if the sample containing chloride is mixed with $\text{K}_2\text{Cr}_2\text{O}_7$ and concentrated H_2SO_4 , the solution forms red CrO_2Cl_2 and red fumes are liberated — This does not happen when chloride is absent or in the presence of fluorides, bromides, iodides and cyanide. [Freeman 'Chromyl Chloride' in *Encyclopedia of Reagents for Organic Synthesis* J. Wiley & Sons, NY, L. Paquette ed. 2004, DOI: 10.1002/047084289.] **TOXIC**.

Claisen alkali (alkali Claisen). Prepare this from KOH (35g) in H_2O (25ml) and dilute it to 100ml with MeOH. **STRONGLY CAUSTIC**.

Cobaltous ammonium sulfate hexahydrate [13596-46-8, 13586-38-4] $(\text{NH}_4)_2\text{Co}(\text{SO}_4)_2 \cdot 6\text{H}_2\text{O}$, **M 395.5**, **m $120^\circ(\text{dec})$** , **d²⁵ 5.22g/cm³**. Crystallise it from boiling water (2ml/g) by cooling. Wash it with ethanol and dry it in a vacuum.

Cobaltous bromide hexahydrate [85017-77-2 ($x\text{H}_2\text{O}$), 7789-43-7 (anhydrous)] $\text{CoBr}_2 \cdot 6\text{H}_2\text{O}$, **M 326.9** ($6\text{H}_2\text{O}$), **M 218.7(anhydr)**, **m $47^\circ(\text{dec}, 6\text{H}_2\text{O})$** , **$678^\circ(\text{anhydr})$** , **d²⁰ 4.9**. Crystallise it from water (1ml/g) by partial evaporation in a desiccator (red-purple, hydrate). The **anhydrous** salt (bright green) is soluble in EtOH, Me_2CO , MeOAc to form blue-coloured solutions. [Glemser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1517 1965.]

Cobaltous chloride hexahydrate [7791-13-1 ($6\text{H}_2\text{O}$), 69098-14-2 ($x\text{H}_2\text{O}$), 7646-79-9 (anhydrous)] $\text{CoCl}_2 \cdot 6\text{H}_2\text{O}$, **M 237.9**, **m $87^\circ(\text{dec})$** , **d²⁰ 1.92**. A saturated aqueous solution at room temperature is fractionally crystallised by standing overnight. The first half of the material that crystallises in this way is used in the next crystallisation. The process is repeated several times, water being removed in a dry-box using air filtered through glass wool and dried over CaCl_2 [Hutchinson *J Am Chem Soc* **76** 1022 1954, DOI: 10.1021/ja01633a025]. It has also been crystallised from dilute aqueous HCl. The **hexahydrate m 86°** forms pink to red **deliquescent** crystals. It loses $4\text{H}_2\text{O}$ on heating at $52\text{-}56^\circ$ and forms the violet **dihydrate** which loses a further H_2O at 100° to form the violet **monohydrate** which loses the last H_2O molecule at $120\text{-}140^\circ$ to give the pale blue **anhydrous deliquescent** salt **m 735° (724°)** and **b 1049°** . A pink solution of CoCl_2 in H_2O becomes blue on heating to 50° or adding conc HCl which may precipitate the **mono** or **dihydrate**. The solid **dihydrate** gives a blue-purple solution with EtOH. **Note:** CoCl_2 in H_2O is a '**sympathetic ink**', i.e. writing with an aqueous solution is almost invisible on paper, but becomes blue on warming the paper. On cooling or standing, the writing becomes invisible again. The **anhydrous** salt is soluble in H_2O , EtOH, Et_2O , Me_2CO and pyridine. [Glemser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1515 1965.]

Cobaltous nitrate hexahydrate [10026-22-9] $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$, **M 291.0**, **m $\sim 55^\circ(6\text{H}_2\text{O})$** , **$100\text{-}105^\circ(\text{dec})$** , **d²⁰ 1.88**. Crystallise the red nitrate from water (1ml/g), or ethanol (1ml/g), by partial evaporation. After 3 crystallisations from H_2O it contains: metal (ppm) As (8), Fe (1.2), K (1), Mg (4), Mn (4), Mo (4), Na (0.6), Ni (18), Zn (1.6). The **hexahydrate** gives the pink **anhydrous** salt by the action of HNO_3 and N_2O_5 . The **hexahydrate** melts at $\sim 55^\circ$ to give a red liquid which decomposes on further heating at $100\text{-}105^\circ$ to form Co_3O_4 .

Cobaltous perchlorate hexahydrate [13478-33-6] $\text{Co}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$, **M 365.9**, **m 1534°** , **pK²⁵ -2.4 to -3.1** (for HClO_4). Crystallise the red solid from warm water (0.7ml/g) by cooling. Used in battery manufacture.

Cobaltous potassium sulfate [13596-22-0] $\text{CoK}_2(\text{SO}_4)_2$, **M 329.3**, d_4^{20} **2.22(anhydr)**. Crystallise it from water (1ml/g) between 50° and 0°, and dry it in a vacuum desiccator over conc H_2SO_4 . Red, monoclinic prismatic, crystals (*hexahydrate*) are obtained that lose H_2O on heating above 75° to form the reddish-brown *dihydrate* that is stable between 120-150°. It becomes *anhydrous* (violet solid) above 220°. [Gmelin's, *Cobalt* (8th ed.) **58** (part A) p 414 1932, and supplement pp 782-786 1961.]

Cobaltous sulfate heptahydrate [10026-24-1 ($7\text{H}_2\text{O}$), 60459-08-7 ($x\text{H}_2\text{O}$), 10124-43-3 (*anhydrous*)] $\text{CoSO}_4 \cdot 7\text{H}_2\text{O}$, **M 281.1**, **m** (see text), d_4^{20} **2.03**. Crystallise it three times from conductivity water (1.3ml/g) between 100° and 0° depending on which hydrate is required. The *heptahydrate* crystallises below 44° and is *efflorescent* with **m 97°**. Between 44° and 70° the monoclinic *hexahydrate* $\text{CoSO}_4 \cdot 6\text{H}_2\text{O}$ **m 41.5°** is formed, and above 70° the *monohydrate* $\text{CoSO}_4 \cdot \text{H}_2\text{O}$ **m 71°** is obtained. The pale reddish or lavender-coloured *anhydrous* salt is obtained by heating the hydrate above 250°, boiling with conc H_2SO_4 or by heating with $(\text{NH}_4)_2\text{SO}_4$.

Cupric ammonium chloride dihydrate [10534-87-9 (*hydrate*), 15610-76-1 (*anhydrous*)] $(\text{NH}_4)_2\text{CuCl}_4 \cdot 2\text{H}_2\text{O}$, **M 277.5**, **m 110-120°(anhydrous)** then **dec at higher temperature**, d_4^{20} **1.993**. Crystallise it from weak aqueous HCl (1ml/g). It crystallises out of a hot solution of CuCl_2 saturated with NH_3 gas.

Cupric (II) bromide [7789-45-9] CuBr_2 , **M 223.4**, **m 498°**, **b 900°/atm**, d_4^{20} **4.77**. Crystallise it twice by dissolving it in water (140ml/g), filtering to remove any Cu_2Br_2 , and concentrating under vacuum at 30° until crystals appear. The cupric bromide is then allowed to crystallise by leaving the solution in a vacuum desiccator containing P_2O_5 . [Hope et al. *J Chem Soc* 5226 1960, DOI: 10.1039/JR9600005226; Glemser & Sauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II p 1009 1965, Fieser **1** 161, **2** 84, **5** 158, **6** 138, **14** 100, **15** 100.]

Cupric (II) chloride [7447-39-4] CuCl_2 , **M 134.4**, **m 498°**, **620°**, **630°(dec)**. Crystallise the chloride from hot dilute aqueous HCl (0.6ml/g) by cooling in a CaCl_2 -ice bath. It is dehydrated by heating on a steam bath under vacuum. It is *deliquescent* in moist air but *efflorescent* in dry air. The *dihydrate* is emerald green but blue when free from solvent. Concentrated solutions are yellow-green in colour but are blue when free from solvent. Concentrated solutions are yellow-green and become yellow on adding conc HCl. A very dilute solution is pure **blue** due to $\text{Cu}(\text{H}_2\text{O})_4^{2+}$ [Donnan & Bassett *J Chem Soc* **81** 939 1902, DOI: 10.1039/CT9028100939]. CuCl_2 is very *deliquescent* and is soluble in MeOH or EtOH to give **green** crystals of $\text{Cu}(\text{ROH})_2\text{Cl}_2$. [Glemser & Sauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II p 1008 1965, Fieser **1** 163, **2** 84, **3** 66, **4** 105, **5** 158, **6** 139, **7** 79, **8** 119, **9** 123, **10** 106, **13** 85, **14** 100.] Used with Pd in the catalytic synthesis of 3-haloindoles *via* annulation [Tang et al. *Synthesis* 1841 2007, DOI: 10.1055/s-2007-983717.]

Cupric (II) nitrate trihydrate [10031-43-3 ($3\text{H}_2\text{O}$), 13478-38-1 ($6\text{H}_2\text{O}$), 19004-19-6 ($2.5\text{H}_2\text{O}$), 3251-23-8 (*anhydrous*)] $\text{Cu}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$, **M 241.6($3\text{H}_2\text{O}$)**, **m 26.4°($6\text{H}_2\text{O}$, dec)**, **114.5°($3\text{H}_2\text{O}$, dec)**, **256°(anhydr, dec)**, **b 170°/atm($3\text{H}_2\text{O}$, dec)**, d_4^{20} **2.0**. Crystallise it from weak aqueous HNO_3 (0.5ml/g) by cooling from room temperature. The *anhydrous* salt can be prepared by dissolving copper metal in a 1:1 mixture of liquid NO_2 and ethyl acetate and purified by sublimation [Evans et al. *JCS Faraday Trans 1* **75** 1023 1979, DOI: 10.1039/F19797501023]. The *hexahydrate* dehydrates to the *trihydrate* at 26°, and the *anhydrous salt* sublimes between 150 and 225°/high vacuum, but melts at **255-256°** and is *deliquescent*.

Cupric (II) perchlorate hexahydrate [10294-46-9 (*hydrate*), 13770-18-8] $\text{Cu}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$, **M 370.5**, **m 230-240°**, d^{25} **2.225g/cm³**, pK^{25} **-2.4 to -3.1 (for HClO_4)**. Crystallise it from distilled water. The *anhydrous salt* is hygroscopic.

Cupric (II) sulfate (blue vitriol, blue stone) [7758-98-7 (*anhydrous*), 7758-99-8 ($5\text{H}_2\text{O}$), 16448-28-5 ($3\text{H}_2\text{O}$), 23254-43-5 ($4\text{-}6\text{H}_2\text{O}$), 19086-18-1 ($7\text{H}_2\text{O}$)] $\text{Cu}(\text{SO}_4)_2 \cdot 5\text{H}_2\text{O}$, **M 159.6**, **m 110°(hydrate)**, **>560°(anhydr)**. After adding 0.02g of KOH to a litre of nearly saturated aqueous solution of the sulfate, it is left for two weeks, then the precipitate is filtered on to a fibreglass filter with pore diameter of 5-15 microns. The filtrate is heated to 90° and allowed to evaporate until some $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$ crystallises out. The solution is then filtered hot and cooled rapidly to give crystals which are freed from mother liquor by filtering under suction [Geballe & Giauque

J Am Chem Soc **74** 3513 1952, DOI: 10.1021/ja01134a017]. Alternatively, crystallise the sulfate from water (0.6ml/g) between 100° and 0°. The **pentahydrate** is slowly **efflorescent**, losing 2H₂O at 30°, two more H₂O are lost at 110° and a **white** anhydrous powder (desiccant) is obtained on heating above 250°. [Hoffman *Copper(II) Sulfate*, in *Encyclopedia of Reagents for Organic Synthesis*, John Wiley & Sons. 2001, DOI: 10.1002/047084289X.rc247].

Cuprous (I) bromide [7787-70-4] **CuBr**, **M 143.4**, **m 497°, 504°, b 1345°, d₄²⁰ 4.72**. Purify it as for cuprous iodide but using aqueous NaBr. [Keller et al. *Inorg Synth* **2** 1 1946, DOI: 10.1002/9780470132333.ch1; Glemser & Sauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1006 1965, Fieser **1** 165, **2** 9, **2** 90, **3** 7, **4** 108, **5** 163, **6** 143, **7** 79, **8** 116, **11** 140, **14** 99, **16** 96.]

Cuprous (I) bromide dimethylsulfide complex [54678-23-8] **CuBr·CH₃SCH₃**, **M 205.6**, **m 132°(dec)**. If it needs to be prepared afresh then wash CuBr (25g) with MeOH (4 x 50ml) to remove coloured impurities, and dry for 1 hour *in vacuo*, then dissolve the green-tinted white powder in Me₂S (60ml) [**foul odour**, use an efficient fume cupboard] and filter to remove insoluble impurities. Dilute the filtrate with hexane (200ml) and collect the colourless crystals by filtration under suction. Wash the light sensitive crystals five times with hexane and dry them under a stream of dry N₂ for 3 hours. Do **not** dry under vacuum, as loss of Me₂S would occur with deleterious effect on its reactions. If the sample is reasonably good then dissolve it in MeOH and crystallise it by diluting the solution with an equal volume of hexane, collect the crystals, wash with hexane and dry as stated previously. [Wuts *Synth Commun* **11** 139 1981, DOI: 10.1080/00397918108064294; Fieser **6** 225, **8** 117, **10** 104, **15** 100, **16** 96.] Catalyst used for the addition of alkyl, alkenyl and aryl Grignard reagents to Fullerenes in which 5 organic groups add to [60]fullerene, and 3 groups to [70]fullerene [Matsuo et al. *Org Synth* **83** 80 2006, DOI: 10.15227/orgsyn.083.0080].

Cuprous (I) chloride [7758-89-6] **CuCl**, **M 99.0**, **m 430°, b~1400°**. Dissolve it in strong HCl, precipitate it by diluting with water and filter it off. Wash the solid with ethanol and diethyl ether, then dry it and store it in a vacuum desiccator [Österlöf *Acta Chem Scand* **4** 374 1950, DOI: 10.3891/acta.chem.scand.04-0374]. Alternatively, to an aqueous solution of CuCl₂·2H₂O is added, with stirring, an aqueous solution of anhydrous sodium sulfite. The colourless product is dried at 80° for 30 minutes and stored under N₂. Cu₂Cl₂ can be purified by zone-refining [Hall et al. *JCS Faraday Trans 1* **79** 343 1983, DOI: 10.1039/F19837900343]. [Glemser & Sauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1005 1965, Fieser **1** 166, **2** 91, **3** 67, **4** 109, **5** 164, **6** 145, **7** 80, **8** 118, **11** 140, **12** 141, **13** 85, **15** 101.]

Cuprous (I) cyanide [544-92-3] **CuCN**, **M 89.6**, **m 474°, d₄²⁵ 2.92g/cm³**. Wash the cyanide thoroughly with boiling H₂O, then with EtOH. Dry it at 100° to a fine soft powder. It dissolves in excess alkali cyanide solutions to form the very soluble **complex ion Cu(CN)₄³⁻**. [Bassett & Corbett *J Chem Soc* **125** 1660 1924, DOI: 10.1039/CT9242501660; Barber et al. *J Chem Soc* **79** 1943, DOI: 10.1039/JR9430000079.] Applied in a widely useful **thiosulfate-assisted** preparation of new diamine-CuCN complexes [Stocker et al. *Inorg Chem* **38** 984 1999, DOI: 10.1021/ic970226b].

Cuprous (I) iodide [7681-65-4] **CuI**, **M 190.5**, **m 605°, b 1336°/atm, d₄²⁵ 5.63**. It can be freshly prepared by dissolving an appropriate quantity of CuI in boiling saturated aqueous NaI over 30 minutes. Pure CuI is obtained by cooling and diluting the solution with water, followed by filtering and washing sequentially with H₂O, EtOH, EtOAc, Et₂O and pentane, then drying *in vacuo* for 24 hours [Dieter et al. *J Am Chem Soc* **107** 4679 1985, DOI: 10.1021/ja00302a014]. Alternatively, wash it with H₂O, then EtOH and finally with Et₂O containing a little iodine. Traces of H₂O are best removed first by heating at 110° and then at 400°. Excess of I₂ is removed completely at 400°. It dissolves in Et₂O if an amine is present to form the **amine complex**. On heating it becomes red, then black, but changes to white on cooling. It is sparingly soluble in H₂O or alkali iodide solutions but readily soluble in NH₃ (which absorbs CO), and in cyanide or thiosulfate solutions. [Glemser & Sauer in *Handbook of Preparative Inorganic Chemistry* (Ed Brauer) Academic Press Vol **II** p 1007 1965, Bawn & Ledwith *Chem Ind (London)* 1180 1957, Fieser **1** 169, **2** 92, **3** 69, **5** 167, **6** 147, **7** 81, **8** 121, **10** 107, **11** 141, **12** 141, **16** 98, **17** 87.] Useful catalyst for the Ullmann type *N*-arylation of amines and amino acids [see Jiang et al. *Synlett* 1837 2007, DOI: 10.1055/s-2007-982564], and the preparation of sulfenamides [see Taniguchi *Synlett* 1917 2007, DOI: 10.1055/s-2007-984539], and some catalysed reactions by CuI have

been reviewed by Coeffard [see *Synlett* **15** 2456 2007, DOI: 10.1055/s-2007-986636].

Cuprous (I) thiocyanate [18223-42-2, 1111-67-7] CuSCN , **M 121.6**, **m 1084°**, **d**²⁵ **2.84g/cm³**. **pK²⁵ -1.85 (for HSCN)**. Purify the grey-greenish yellow solid as for cuprous iodide but using aqueous NaSCN. [Demmerle et al. *Ind Eng Chem* **42** 2 1950, DOI: 10.1021/ie50481a002; Newman *Analyst* **88** 500 1963, DOI: 10.1039/AN9638800500.]

Cyanamide (carbidiimide) [420-04-2] NCNH_2 , **M 42.0**, **m 43°, 45°, 46°**, **b 83°/0.5mm, 85-87°/0.5mm**, **pK₁²⁰ -0.36 (1.1 at 29°)**, **pK₂²⁰ 10.27**. Purify it by placing *ca* 15g in a Soxhlet thimble and extracting exhaustively (2-3 hours) with two successive portions of Et_2O (400ml, saturated with H_2O by shaking before use) containing two drops of 1N acetic acid. Two successive portions of Et_2O are used so that the NH_2CN is not heated for too long. Each extract is dried over Na_2SO_4 (30g), then combined and evaporated under reduced pressure. The NH_2CN may be stored unchanged at 0° in Et_2O solution in the presence of a trace of AcOH . Extracts from several runs may be combined and evaporated together. The residue from evaporation of an Et_2O solution is a colourless viscous oil which sets to a solid and can be recrystallised from a mixture of 2 parts of C_6H_6 and 1 part of Et_2O . Concentrating an aqueous solution of NH_2CN at high temperatures causes **EXPLOSIVE** polymerisation. [Kurzer & Lawson *Org Synth Coll Vol* **4** 645 1963, DOI: 10.15227/orgsyn.034.0067; Pinck et al. *Inorg Synth* **3** 39 1950, DOI: 10.1002/9780470132340.ch9; Soloway & Lipschitz *J Org Chem* **23** 613 1958, DOI: 10.1021/jo01098a603.] *Hygroscopic*. [Beilstein **3** IV 145.]

Cyanogen bromide [506-68-3] BrCN , **M 105.9**, **m 49-51°, 50-53°**, **b 60-62°/atm**, **d**²⁵ **2.02g/cm³**. *All operations with this substance should be performed in a very efficient fume cupboard—it is very POISONOUS and should be handled in small amounts. Fresh commercial material is satisfactory for nearly all purposes and does not need to be purified. It is a white crystalline solid with a strong cyanide odour. If it is reddish in colour and partly liquid or paste-like, then it is too 'far gone' to be purified, and fresh material should be sought.* It can be purified by distillation using small amounts at a time, and using a short wide-bore condenser because it readily solidifies to a crystalline white solid which may clog the condenser. *An appropriate gas mask should be used when transferring the molten solid from one container to another, and the operation should be done in an efficient fume cupboard.* The melting point (**m 49-51°**) should be measured in a sealed tube. [Hartman & Dreger *Org Synth Coll Vol* **2** 150 1948, DOI: 10.15227/orgsyn.011.0030.] Also commercially available are a 5.0M solution in MeCN (**d**²⁵ **1.903g/cm³**), and a 3.0M solution in CH_2Cl_2 (**d**²⁵ **1.443g/cm³**). BrCN is a useful reagent for peptide cleavage [Orlando et al. *Org Mass Spectrom* **28** 1395 1993, DOI: 10.1002/oms.1210281207], and for cleaving peptides bound to a support for sequencing by mass spectrometry [Anspach et al. *Appl Biochem Biotechnol* **44** 135 1944, DOI: 10.1007/BF02921651]. **POISONOUS**.

Cyanogen iodide [506-78-5] ICN , **M 152.9**, **m 146.7°, 146-147°, m 146.7°**, **d**²⁵ **1.84g/cm³**. *This compound is POISONOUS, and the precautions for cyanogen bromide (above) apply here.* The reagent (*ca* 5.9g) is dissolved in boiling CHCl_3 (15ml), filtered through a plug of glass wool into a 25ml Erlenmeyer flask. Cool to room temperature for 15 minutes, then place it in an ice-salt bath and cool to -10°. This cooling causes a small aqueous layer to separate as ice. The ice is filtered with the CNI , but melts on the filter and is also removed with the CHCl_3 used as washing liquid. The CNI which is collected on a sintered glass funnel is washed 3x with CHCl_3 (1.5ml at 0°) and freed from last traces of solvent by placing it on a watch glass and exposing it to the atmosphere in a good fume cupboard at room temperature for 1 hour to give colourless needles (*ca* 4.5g), **m 146-147°** (sealed capillary totally immersed in the oil bath). The yield depends slightly on the rapidity of the operation; in this way loss by sublimation can be minimised. If desired, it can be sublimed under reduced pressure at temperatures at which CNI is only slowly decomposed into I_2 and $(\text{CN})_2$. The vacuum will need to be renewed constantly due to the volatility of CNI . [Bak & Hillebert *Org Synth Coll Vol* **4** 207 1963, DOI: 10.15227/orgsyn.032.0029.]

Decaborane [17702-41-9] $\text{B}_{10}\text{H}_{14}$, **M 122.2**, **m 99.7-100°, b 100°/19mm, 213°/atm**, **d**²⁵ **0.94g/cm³**. Purify decaborane by vacuum sublimation at 80°/0.1mm, followed by crystallisation from methylcyclohexane, CH_2Cl_2 , or dry olefin-free-*n*-pentane, the solvent being subsequently removed by storing the crystals in a vacuum

desiccator containing CaCl_2 . It is soluble in H_2O but is slowly decomposed to give H_2 . It is soluble in alkali, and on acidification it liberates H_2 . **TOXIC**. [Greenwood in *Comprehensive Chemistry* (Ed Bailer et al.) Pergamon Press Vol 1 pp 818-837 1973.]

Deuterium (heavy hydrogen) [7782-39-0] D_2 , M 4.028, m $-254.43^\circ/\text{atm}$, b $-249.49^\circ/\text{atm}$, d $^{25}_4$ $0.169\text{g}/\text{cm}^3$, d $0.189\text{Kg}/\text{m}^3$ (at STP i.e. $0^\circ/101.325\text{kPa}$), d $162.4\text{kg}/\text{m}^3$ (liquid), critical temp -234.75° , critical pressure **128.5mm (triple point)**. Pass the gas over activated charcoal at -195° [MacIver & Tobin *J Phys Chem* **64** 451 1960, DOI: 10.1021/j100833a018]. Purify it also by diffusion through nickel [Pratt & Rogers, *JCS Faraday Trans I* **92** 1589 1976, DOI: 10.1039/F19767201589]. It is as flammable as H_2 . Always check deuterium for radioactivity to determine the amount of tritium in it (see D_2O below). [A Hydrogen Isotope of Mass 2, Urey et al. *Phys Rev* **39** 164 1932, DOI: 10.1103/PhysRev.39.164; D_2 Isotope effect: Wiberg *Chem Rev* **55** 713 1955, DOI: 10.1021/cr50004a004.]

Deuterium oxide [7789-20-0] D_2O , M 20, f $3.8^\circ/760\text{mm}$, b $101.4^\circ/760\text{mm}$, d $^{20}_4$ 1.105 , pK 20 **14.955**. Distil 'heavy water' from alkaline KMnO_4 [de Giovanni & Zamenhof *Biochem J* **87** 79 1963, DOI: 10.1042/bj0870079]. **NOTE that D_2O invariably contains tritiated water and will therefore be RADIOACTIVE; always check the radioactivity level of D_2O in a scintillation counter before using.** [For pKa of D_2O see Covington et al. *J Phys Chem* **70** 3820 1966, DOI: 10.1021/j100884a011.]

cis-Diamminedichloroplatinum(II) (Cisplatin) [15663-27-1] $\text{Pt}(\text{NH}_4)_2\text{Cl}_2$, M 300.1, m $270^\circ(\text{dec})$. Recrystallise it from dimethylformamide and check the purity by IR and UV-VIS spectroscopy. [Raudaschl et al. *Inorg Chim Acta* **78** L43 1983, DOI: 10.1016/S0020-1693(00)86472-5.] **HIGHLY TOXIC, SUSPECTED CARCINOGEN.**

Diammonium hydrogen orthophosphate [7783-28-0] $(\text{NH}_4)_2\text{HPO}_4$, M 132.1, m $155^\circ(\text{dec})$. Crystallise it from water (1ml/g) between 70° and 0° . Its solubility in H_2O is 59% at room temperature and 200% at the boiling point. It slowly evolves NH_3 and should be stored in a well-stoppered container. After one crystallisation, ACS grade salt had Fe, Mo, Na, Se and Ti at 1, 0.2, 1.4, 0.2 and 0.8ppm, respectively. [Gmelin's, Ammonium (8th edn) **23** pp422-426 1936.]

Dinitrogen tetroxide (nitrogen dioxide) [10544-72-6] N_2O_4 , M 92.0, m -11.2° , b $21.69^\circ/\text{atm}$, d $^{21}_4$ 1.44246 , n $^{21}_D$ **1.00112(liquid)**. Purify it by oxidation at 0° in a stream of oxygen until the blue colour changes to red-brown. Alternatively, distil it from P_2O_5 , then solidify it by cooling in a deep-freeze (at -78° , giving nearly colourless crystals). Oxygen can be removed by alternate freezing and melting cycles. **TOXIC VAPOUR**. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 488-489 1963.]

Disodium hydrogen orthophosphate (anhydrous) [7558-79-4] Na_2HPO_4 , M 142.0, m 250° , d $^{25}_4$ $1.7\text{g}/\text{cm}^3$, (see pK of H_3PO_4). Crystallise the salt twice from warm water, by cooling. Dry in air, then in an oven overnight at 130° . It should be dried before use as it is slightly *hygroscopic*. It forms di-, hepta- and deca-hydrates.

Ferric Bromide [10031-26-2] FeBr_3 , M 395.6, m $>130^\circ(\text{dec})$, $>200^\circ(\text{dec})$, d $^{25}_4$ $4.50\text{g}/\text{cm}^3$. Sublime it in a sealed tube with Br_2 at 120° - 200° . Decomposition to FeBr_2 and Br_2 occurs above 200° , or by boiling in H_2O . It is soluble in AcOH , EtOH , Et_2O and H_2O . Store it in a tightly stoppered bottle away from light. [Lux in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II p 1494 1965, Gregory & Thackrey *J Am Chem Soc* **72** 3176 1950, DOI: 10.1021/ja01163a102.]

Ferric chloride (anhydrous) [7705-08-0] FeCl_3 , M 162.2, m $>300^\circ(\text{dec})$, 304° , 306° , d $^{25}_4$ $2.898\text{g}/\text{cm}^3$. Sublime it at 200° in an atmosphere of chlorine. It is an 'iron-black' coloured powder with green iridescence. It is soluble in C_6H_6 , EtOH , Et_2O , Me_2CO and pyridine in which the *monomeric species* ' FeCl_3 ' exist whereas in the vapour state it exists in the *dimeric* ' Fe_2Cl_6 ' form. Decomposes to FeCl_2 and Cl_2 above 315° . It is soluble in H_2O (0.7g/ml at 0°), in Me_2CO (0.63/ml at 18°) in MeOH (very soluble), EtOH (0.83g/ml) and Et_2O (very

soluble). Store it in a weighing bottle inside a desiccator as it absorbs moisture from air to form the yellow **hexahydrate** (see next entry). [Tarr et al. *Inorg Synth* **3** 191 1950, DOI: 10.1002/9780470132340.ch51; Pray et al. *Inorg Synth* **5** 153 1957, DOI: 10.1002/9780470132364.ch43; Epperson et al. *Inorg Synth* **7** 163 1963, DOI: 10.1002/9780470132388.ch45; Fieser **1** 390, **2** 199, **3** 145, **4** 236, **5** 307, **7** 153, **8** 228, **10** 185.]

Ferric chloride hexahydrate [10025-77-1] $\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$, **M 270.3**, **m** 37°(dec), **b** 280-285°/atm(partial decomposition), **d**²⁵ 1.83g/cm³, **pK**₁²⁵ 2.83, **pK**₂²⁵ 4.59 (for hydrolysis of Fe^{3+}). An aqueous solution, saturated with the salt at room temperature, is cooled to -20° for several hours. Separation of the crystals is slow, even with scratching and seeding, and it is generally necessary to stir this overnight. The presence of free HCl retards crystallisation. [Linke *J Phys Chem* **60** 91 1956, DOI: 10.1021/j150535a022; Fieser **1** 390, **2** 199, **3** 145, **4** 236, **5** 307, **6** 259, **7** 153, **8** 228, **9** 222, **10** 185, **12** 230, **14** 164, **15** 158, **16** 190.]

Ferric nitrate nonahydrate [7782-61-8 ($9\text{H}_2\text{O}$), 13476-08-9 (H_2O), 10421-48-4 (anhydrous)] $\text{Fe}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$, **M 404.0**, **m** 47°(dec), **b** 125°/atm, **d**²⁵ 1.6429g/cm³($9\text{H}_2\text{O}$), **d**²⁵ 1.68g/cm³($6\text{H}_2\text{O}$). It crystallises from aqueous solutions of moderately strong HNO_3 as the pale violet **nonahydrate m** 40° and is soluble in EtOH and Me_2CO . With more concentrated aqueous solutions (containing some HNO_3), the **hexahydrate** crystallises out **m** 60.5°. The **anhydrous** salt is *slightly deliquescent* and decomposes at 47°. [Lambert & Thomson *J Chem Soc* **97** 2426 1910, DOI: 10.1039/CT9109702426; *Gmelin's, Iron* (8th edn) **59** Part B pp 161-172 1932.] Solutions of this nitrate are used by metalsmiths and jewellers for etching Ag and Ag alloys. $\text{Fe}(\text{NO}_3)_3$ on Silica Gel (15-20wt% salt) is a useful **oxidising reagent**.

Ferric perchlorate nonahydrate [13537-24-1 ($9\text{H}_2\text{O}$), 15201-61-3 ($x\text{H}_2\text{O}$)] $\text{Fe}(\text{ClO}_4)_3 \cdot 9\text{H}_2\text{O}$, **M 516.3**, **pK**²⁵ -2.4 to -3.1 (for HClO_4). Crystallise it twice from concentrated HClO_4 , the first time in the presence of a small amount of H_2O_2 to ensure that the iron is fully oxidised [Sullivan *J Am Chem Soc* **84** 4256 1962, DOI: 10.1021/ja00881a012]. Extreme care should be taken with this preparation because it is potentially **EXPLOSIVE**.

Ferric sulfate x-hydrate [10028-22-5 (anhydrous), 15244-10-7 (H_2O)] $\text{Fe}_2(\text{SO}_4)_3 \cdot x\text{H}_2\text{O}$, **M 399.9 + $x\text{H}_2\text{O}$** , **m** 480°(anhydr), 175°(H_2O), **d**²⁵ 3.097g/cm³(anhydr), **d**²⁵ 1.898g/cm³($5\text{H}_2\text{O}$). Dissolve the sulfate in the minimum volume of dilute aqueous H_2SO_4 and allow it to evaporate at room temperature until yellowish-white crystals start to form. Do not concentrate by boiling off the H_2O as basic salts will be formed. Various **hydrates** are formed; the common ones are the **nona** and **dodeca hydrates** which are violet in colour. The **anhydrous** salt is colourless and is quite **hygroscopic**, but it dissolves in H_2O slowly unless ferrous sulfate is added. [*Gmelin's, Iron* (8th edn) pp 439-462 1932.]

Ferrous bromide [20049-65-4, 7789-46-0] FeBr_3 , **M 215.7 + $x\text{H}_2\text{O}$** , **m** 684°, **b** 927°/atm, 934°/atm, **d**²⁵ 4.63. It crystallises from air-free H_2O to provide the **hexahydrate** as pale green to bluish-green rhombic prisms. On heating at 49° H_2O is lost and the **tetrahydrate** is formed. On further heating at 83° more H_2O is lost and the **dihydrate** is formed as a light yellow to dark brown **hygroscopic** powder. The ferrous iron in aqueous solutions of these salts readily oxidises to ferric iron. The salts should be stored over H_2SO_4 under N_2 in tightly closed containers. They have some solubility in MeOH, EtOH and THF. [Baxter *Z Anorg Chem* **38** 232 1904, DOI: 10.1002/zaac.19040380119; Kühn & Ernst *Z Anorg Allgem Chem* **317** 84 1962, DOI: 10.1002/zaac.19623170112; Lux in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1493 1965, Winter *Inorg Synth* **14** 99 1973, DOI: 10.1002/9780470132456.ch20.]

Ferrous chloride tetrahydrate [13478-10-9 ($4\text{H}_2\text{O}$), 16399-77-2 ($2\text{H}_2\text{O}$)] $\text{FeCl}_2 \cdot 9\text{H}_2\text{O}$, **M 198.8**, **m** 105°($4\text{H}_2\text{O}$, dec), 120°($2\text{H}_2\text{O}$, dec), **d**²⁵ 3.16g/cm³(anhydr), **d**²⁵ 2.39g/cm³($2\text{H}_2\text{O}$), **d**²⁵ 1.93g/cm³($4\text{H}_2\text{O}$), **pK**₁²⁵ 6.7, **pK**₂²⁵ 9.3 (for aquo Fe^{2+}). A 550ml round-bottomed Pyrex flask is connected, *via* a glass tube fitted with a medium porosity sintered-glass disc, to a similar flask. To 240g of $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ in the first flask is added conductivity water (200ml), 38% HCl (10ml), and pure electrolytic iron (8-10g). A stream of purified N_2 gas is passed through the assembly, escaping through a mercury trap. The salt is dissolved by heating which is continued until complete reduction has occurred. By inverting the apparatus and filtering (under N_2 pressure) through the sintered glass disc, unreacted iron is removed. After cooling and crystallisation, the unit is again inverted, and the crystals of ferrous chloride are filtered free from mother liquor by applied N_2 pressure. Partial

drying by overnight evacuation at room temperature gives a mixed hydrate which, on further evacuation on a water bath at 80°, loses water of hydration and absorbed HCl (with vigorous effervescence) to give a white powder of $\text{FeCl}_2 \cdot 2\text{H}_2\text{O}$ (see below). [Gayer & Wootner *J Am Chem Soc* **78** 3944 1956, DOI: 10.1021/ja01597a021; $(2\text{H}_2\text{O})$ Gayer et al. *Inorg Synth* **5** 179 1957, DOI: 10.1002/9780470132364.ch48; Kovacic & Brace 'Iron(II) Chloride' *Inorg. Synth.* **6** 172 1960, DOI: 10.1002/9780470132371.ch54]

Ferrous chloride [7758-94-3] FeCl_2 , **M 126.8, m 674°, b 1023°, d²⁵ 3.16**. It sublimes in a stream of HCl at *ca* 700°, or in H_2 below 300°. Its vapour pressure at 700° is 12mm. It forms white **hygroscopic** rhombohedral crystals with a green tint which oxidise in air to FeCl_3 and Fe_2O_3 . It is soluble in H_2O , EtOH Me_2CO but insoluble in Et_2O . The **tetrahydrate** is pale green to pale blue in colour and loses $2\text{H}_2\text{O}$ at 105–115°. The **dihydrate** loses H_2O at 120°. [**Anhydrous** FeBr_2 can be obtained by carefully dehydrating the **tetrahydrate** in a stream of HBr and N_2 , and it can be sublimed under N_2 .] The ferrous iron in aqueous solutions of these salts readily oxidises to ferric iron. (See above.) [Kovacumic & Brace *Inorg Synth* **6** 172 1960, DOI: 10.1002/9780470132371.ch54; Lux in *Handbook of Preparative Inorganic Chemistry* (Ed Brauer) Academic Press Vol **II** p 1491 1965.]

Ferrous perchlorate hexahydrate [13933-23-8, 1350-69-9, 335159-18-7] $\text{Fe}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$, **M 362.8, pK²⁵ -2.4 to -3.1 (for HClO_4)**. Crystallise it from HClO_4 . [**CARE, see ferric perchlorate above.**]

Ferrous sulfate heptahydrate (green vitriol) [7782-63-0 ($7\text{H}_2\text{O}$), 13463-43-9 ($x\text{H}_2\text{O}$), 17375-41-6 (H_2O), 7720-78-7 (anhydrous)] $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$, **M 278.0($4\text{H}_2\text{O}$), 151.9(anhydr), m ~60–64°($7\text{H}_2\text{O}$, dec), 300°(H_2O , dec), 680°(anhydr, dec), d²⁵ 1.895g/cm³($7\text{H}_2\text{O}$), d²⁵ 1.934g/cm³($6\text{H}_2\text{O}$), d²⁵ 2.15g/cm³($5\text{H}_2\text{O}$), d²⁵ 3.0g/cm³(H_2O), d²⁵ 3.65g/cm³(anhydrous)**. Crystallise the sulfate from 0.4M H_2SO_4 , or precipitate it from an aqueous solution with EtOH. It is **efflorescent** in dry air, and is converted to the **tetrahydrate** at 57°, then to the **monohydrate** (white-yellow) at 65° [or by heating the **heptahydrate** (blue-green crystals) in a vacuum at 140°; the **anhydrous** salt forms white crystals]. At 650° it decomposes to Fe_2O_3 , SO_2 and SO_3 . The solubility in H_2O is 0.45g/ml(77°) and 0.36g/ml(90°) for the **monohydrate**, 0.66g/ml(0°), 0.21g/ml(10°), 0.29g/ml(25°), 0.40g/ml(40°) and 0.51g/ml(54°) for the **heptahydrate**; in ethylene glycol it is 0.06g/ml(20°); but it is insoluble in EtOH. It forms a brown-black complex, $\text{FeSO}_4 \cdot \text{NO}$, with nitric oxide and is used in a qualitative test for nitrates ('**brown ring**' test). Used medically as an iron supplement. [Wildermuth et al. 'Iron Compounds' in *Ullmann's Encyclopedia of Industrial Chemistry* Wiley-VCH, Weinheim, 2005.]

Fluorine [7782-41-4] F_2 , **M 38.0, m -219.7°, b -188.1°/atm, d 1.696g/L(STP: 0°/101.325kPa)**. Pass the gas through a bed of NaF at 100° to remove HF and SiF_4 . [For description of stills used in fractional distillation, see Greenberg et al. *J Phys Chem* **65** 1168 1961, DOI: 10.1021/j100825a017; Stein et al. *Purification of Fluorine by Distillation*, Argonne National Laboratory, ANL-6364 1961 (from Office of Technical Services, US Dept of Commerce, Washington 25); Aigueperse, et al., Ullmann (ed.) 'Fluorine Compounds, Inorganic' *Ullmann's Encyclopedia of Industrial Chemistry* Wiley-VCH Weinheim **15** 397–441 2000, DOI: 10.1002/14356007, ISBN 3527306730.] **HIGHLY TOXIC use strictest precautions — do not inhale.**

Fluoroboric acid (tetrafluoroboric acid, fluoboric acid) [16872-11-0] HBF_4 , **M 87.8, b 130°(dec), d²⁵ 1.4g/cm³(of 48% aqueous solution), pK²⁵ -4.9**. Crystallise fluoroboric acid several times from conductivity water. It can be stored in a glass vessel at room temperature. It is available commercially as ~48% aqueous solution. It is most useful for preparing **tetrafluoroborate salts** which are generally insoluble. For example, addition of the acid to aryldiazonium salt solutions precipitates the more stable aryldiazonium tetrafluoroborate salts which can be washed with H_2O to remove impurities, followed by EtOH and Et_2O , and stored for short periods of time before further use. It is a catalyst for preparing acetals. [Mathers et al. *J Am Chem Soc* **37** 1515 1915, DOI: 10.1021/ja02171a012; Wamser *J Am Chem Soc* **70** 1209 1940, DOI: 10.1021/ja01183a101; Sharp *Adv Fluorine Chem* **1** 68-128 1960, Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 221-222 1963.] **STRONG IRRITANT and is TOXIC.**

Gallium (metallic liquid) [7440-55-3] Ga, **M 69.7, m 29.78°, b 2403°/atm, d^{29.6} 5.904, d^{29.8} 6.095**.

Dissolve the metal in dilute HCl and extract it with Et₂O. Bubbling H₂S through the solution removes many metals, and a second extraction with Et₂O frees Ga further from metal impurities, except for Mo, Th(III) and Fe which are largely removed by precipitation with NaOH. The solution is then electrolysed in 10% NaOH with a Pt anode and cathode (2-5A at 4-5V) to deposit Ga, In, Zn and Pb, from which Ga was obtained by fractional crystallisation of the melt [Hoffman *J Res Nat Bur Stand* **13** 665 1934]. Ga is also purified by heating to boiling in 0.5-1M HCl, then heating to 40° in water and pouring the molten Ga with water under vacuum through a glass filter (30-50 μ pore size), to remove any unmelted metals or oxide film. The Ga is then fractionally crystallised from the melt under water. [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 837 1963.]

Gallium (III) Chloride [13450-90-3] GaCl₃, M 176.1, m 77.8°, b 133°/100mm, 197.7°/700mm, d₄²⁰ 2.47, pK₁²⁵ 2.91, pK₂²⁵ 3.70, pK₃²⁰ 4.42 (for Ga³⁺). The pure compound can be obtained by redistillation in a stream of Cl₂ or Cl₂/N₂ followed by vacuum sublimation or zone refining. It forms colourless needles which give *gallium dichloride* [Ga(GaCl₄), m 172.4°] on heating. It dissolves in H₂O with liberation of heat. It is soluble in Et₂O and can be extracted from an HCl solution with Et₂O. Also commercially available in a 0.5M solution in pentane. [Laubengayer & Schirmer *J Am Chem Soc* **62** 1578 1940, DOI: 10.1021/ja01863a070; Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 846 1963.] GaCl₃ has been used as a mediator and catalyst in organic bond formation reactions [Barman *Synlett* 2440 2003, DOI: 10.1055/s-2003-43340]; and with NaN₃ [Grocholl et al. *Chem Mater* **13** 4290 2001, DOI: 10.1021/cm010342j] and Li₃N [Xie et al. *Science* **272** 1926 1996, DOI: jstor.org/stable/2890619] was used in lower-temperature solvothermal techniques for preparing nanocrystalline GaN.

Gallium (III) nitrate nonahydrate [69365-72-6] Ga(NO₃)₃, M 255.7, 417.9, m ca 65°. Recrystallise the nitrate from H₂O (solubility is 295g/100ml at 20°). It forms a white *deliquescent*, colourless powder soluble in H₂O, absolute EtOH and Et₂O. It loses HNO₃ upon heating at 40°. Addition of Et₂O to a warm ethanolic solution (40-50°) of Ga(NO₃)₃·9H₂O precipitates Ga(OH)₂NO₃·Ga(OH)₃·2H₂O. If the salt has partly hydrolysed, dissolve it in concentrated HNO₃, reflux, dilute with H₂O and concentrate on a sand bath. Wash the solid several times by adding H₂O and evaporating until there is no odour of acid. Dilute the residue to a Ga concentration of 26g/100ml. At this concentration, *spongy* Ga(NO₃)₃·xH₂O separates from the viscous solution. After standing for several days the crystals are collected and dried in a stream of dry air first at room temperature, then at 40°. *Dehydration* is complete after 2 days. Recrystallise it from H₂O and dry it at water pump vacuum at room temperature. [Reimmann & Tanner *Z Naturforsch* **20B** 71 1965, Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 856 1963; Hendsbee et al. '[Ga(H₂O)₆](NO₃)₃·3H₂O' *Acta Cryst E* **65** i65 2009, DOI: 10.1107/S1600536809028086/mg2076sup1.cif.]

Gallium (III) sulfate [13494-91-2 (anhydrous), 13780-42-2 (hydrate)] Ga₂(SO₄)₃, M 427.6(anhydr), m 105-110°, d₂₅ 3.86g/cm³(xH₂O), Recrystallisation from H₂O gives the 16-18H₂O hydrate (solubility at 20° is 170g/100ml). Alternatively, dissolve it in 50% H₂SO₄ and evaporate (60-70°), cool and precipitate it by adding EtOH/Et₂O. On heating at 165° it provides the *anhydrous* salt, which is a white *hygroscopic* solid. [Reimmann & Tanner *Z Naturforsch* **20B** 71 1965.]

Germanium [7440-56-4] Ge, M 72.6, m 937°, 925-975°, b 2700°/atm, 2830°/atm, d₄²⁰ 5.3. Copper contamination on the surface and in the bulk of single crystals of Ge can be removed by immersion in molten alkali cyanide under N₂. The Ge is placed in dry K and/or Na cyanide powder in a graphite holder in a quartz or porcelain boat. The boat is then inserted into a heated furnace which, after a suitable time, is left to cool to room temperature. At 750°, a 1mm thickness of metal requires about 1 minute, whereas 0.5cm needs about half hour. The boat is removed from the furnace, and the solid samples are taken out with plastic-coated tweezers, carefully rinsed in hot water and dried in air [Wang *J Phys Chem* **60** 45 1956, DOI: 10.1021/j150535a012; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 712 1963]. **Care with the use of cyanide.**

Germanium (IV) oxide (germanium dioxide) [1310-53-8] GeO₂, M 104.6, m 1080°(soluble form), d₂₅ 6.239; m 1116°(insoluble form) d₂₅ 4.228, pK₁²⁵ 9.02, pK₂²⁵ 12.82 (for germanic acid H₂GeO₃). The oxide (GeO₂) is usually prepared by hydrolysing redistilled GeCl₄ and igniting it in order to remove H₂O and chloride.

It can be further purified by dissolving in hot H₂O (solubility is 4g/L cold) evaporating and drying the residual crystalline solid. When the *soluble form* (which is produced in H₂O at 355°) is heated for 100 hours, it is converted to the *insoluble form*. This form is stable at temperatures up to 1033°, and fusion at 1080° for 4 hours causes complete de-vitrification and it reverts to the *soluble form*. [Müller & Blank *J Am Chem Soc* **46** 2358 1924, DOI: 10.1021/ja01676a002; Dennis & Laubengayer *J Am Chem Soc* **47** 1945 1925, DOI: 10.1021/ja01684a504; Laubengayer & Morton *J Am Chem Soc* **54** 2303 1932, DOI: 10.1021/ja01345a019; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 706 1963.] The oxide is used for the synthesis of ultra-high refractive index and ultra-low loss wave guides [Imoto *Optromics* **260** 116 2004], and as a building block for inorganic nanowires [Rao et al. *J Mater Chem* **14** 440 2004, DOI: 10.1039/B310387K],

Germanium (IV) tetrabromide [13450-92-5] **GeBr₄**, **M392.2**, **m 26.1°**, **b 185.9°/atm**, **d₄²⁹ 3.123**. Purify it by simple distillation or fractionation depending on purity. It is soluble in EtOH, CHCl₃, *C₆H₆ and Et₂O. It *fumes* in moist air and is readily hydrolysed by water. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 718 1963]. **LACHRYMATORY**.

Germanium tetrachloride [10038-98-9] **GeCl₄**, **M 214.4**, **m -49.5° (α)**, **-52.0° (β)**, **b 83.1°/760mm**, **86.5°/760mm corr**, **d₄²⁰ 1.84**. Traces of Cl₂ and HCl can be removed from the liquid by blowing dry air through it for a few hours at room temperature or by shaking it with Hg or Hg₂Cl₂ and then fractionating it in a vacuum. It decomposes on heating at 950°. It has a sharp penetrating odour and *fumes* in moist air to give a chalky coat of GeO₂. It is slowly hydrolysed by H₂O to give GeO₂, but distils from concentrated HCl. [Foster et al. *Inorg Synth* **2** 109 1946, DOI: 10.1002/9780470132333.ch30; Dennis & Hance *J Am Chem Soc* **44** 299 1922, DOI: 10.1021/ja01423a008; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 715 1963.] **LACHRYMATORY**.

Glass Powder (100-300 mesh). Washed with 10% HNO₃, water and dry in air.

Glass Spheres[65997-17-3] **SiO₂**, **M 60.08**, (mean particle size 9-13μm), **d²⁵ 1.1g/ml**. These are hollow spherical non-porous glass beads with an alkalinity of <0.5 meq/g. If too alkaline for the purpose then wash with 10% HNO₃, water and dry in air.

Gold powder [7440-57-5] **Au**, **M 196.97**, **m 1064.79°**, **b 2808° (2700°)/atm**, **d₄²⁰ 19.3**, **resistivity at 0° is 2.05 μΩ-cm**. Massive gold reacts very slowly and it is best converted to the powder that is more readily reactive. While working in a well vented fume cupboard, solid gold (10g, 51mmol) is dissolved in *Aqua Regia*, made from concentrated hydrochloric acid (12M HCl, 50ml) and concentrated nitric acid (16M HNO₃, 13ml), by heating to 90° which will hasten dissolution, but will require topping up with *Aqua Regia*. Alternatively, the suspension is allowed to stand at ~25° until the solid has dissolved. The solution is evaporated to ca 10ml, concentrated hydrochloric acid (30ml) is added and evaporation is repeated. The solution is diluted with H₂O (20ml), filtered through a medium sintered-glass funnel, and the filter is washed with small volumes of H₂O. The filtrate and washings, which contain **HAuCl₄**, are heated to boiling and hydroquinone (10g, 91mmol) in hot H₂O (100ml) is slowly added. The mixture is kept at 90° for 1 hour (sufficient time to reduce all the HAuCl₄ to Au), cooled and filtered through an extraction thimble. The thimble is placed in a Soxhlet apparatus and extracted with MeOH. After 15 minutes, when the circulating MeOH in the thimble is colourless, the thimble is removed and the contents are collected and dried in air to give Au powder quantitatively. [Block et al. *Inorg Synth* **4** 14 1953, DOI: 10.1002/9780470132357.ch4.] Also available in slugs, mica, beads and wires of various thicknesses, diameters or lengths.

Gold (III) bromide (gold tribromide) [10294-28-7] **AuBr₃ · xH₂O**, **M 436.7**, **m 150°(dec)**. Purify it by adding pure Br₂ to the dark powder, securely stopper the container, warm a little and shake while keeping away from light for ca 48 hours. Remove the stopper and place it over NaOH until free Br₂ is no longer in the apparatus (48-60 hours). The bright yellow needles of the tribromide are stable over NaOH in the dark. It is soluble in H₂O and in EtOH where it is slowly reduced. Keep it in a cooled, closed container and protect it from light as decomposition causes free gold to be formed. *Auribromic acid* can be obtained by adding the calculated amount of concentrated HBr to AuBr₃ (actually Au₂Br₆) until all dissolves, whereby the acid crystallises out as

HAuBr₄·5H₂O; a *deliquescent* solid soluble in EtOH with **m ca 27°**, and store it as above. [Gibson & Colles *J Chem Soc* 2407 1931, DOI: 10.1039/JR9310002407; Burawoy & Gibson *J Chem Soc* 217 1935, DOI: 10.1039/JR9350000217; Burawoy & Gibson *J Chem Soc* 219 1935, DOI: 10.1039/JR9350000219.]

Gold (I) chloride [aurous chloride, Au(I)Cl] [10294-29-8] AuCl, M 232.4, m 289°(dec), d²⁵ 7.57. It is best prepared by heating AuCl₃·3H₂O, or HAuCl₄ (see next entry) at 100° in a high vacuum until the vapour pressure drops considerably when most of the H₂O has been expelled. As the HAuCl₄ liquefies again in this process much spluttering will occur. The solid residue (mostly anhydrous AuCl₃) is then heated to 156° (bromobenzene bath) and further to 170-205° when decomposition to AuCl is complete. Another preparation includes heating AuCl₃ in air at 185° [Thomsen *J Prakt Chem* 13 337 1876, DOI: 10.1002/prac.18760130120], but purer salt is obtained by heating AuCl₃ in a stream of dry HCl at 175° [Diemer *J Am Chem Soc* 35 552 1913, DOI: 10.1021/ja02194a006]. It forms pale yellow crystals which are not deliquescent, slightly soluble in cold EtOH, dissolve in alkali chloride solutions to form chloroaurates(I), e.g. KAuCl₂, but it decompose in H₂O to give gold and hydrolysed Au(III) species. [Blitz & Wein *Z Anorg Allgem Chem* 148 192 1925, DOI: 10.1002/zaac.19251480121; Glemser & Sauer *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer Academic Press Vol II p 1055 1965.)

Gold (III) chloride (hydrate) [13453-07-1 (anhydrous), 16961-25-4 (3H₂O), 16903-35-8 (30w% in dil HCl)] AuCl₃ (anhydrous), HAuCl₄·xH₂O, M 339.8 + xH₂O, x ~3, m 229°, b 354°(dec), d₄²⁰ 3.9. It is obtained as a dark red crystalline mass by dissolving Au in *aqua regia* and evaporating. When sublimed at 180°, the *anhydrous* crystals are ruby red. The anhydrous salt is *hygroscopic*, soluble in H₂O but sparingly soluble in EtOH and Et₂O. *Aurichloric acid* (chloroauric acid, HAuCl₄) is formed when AuCl₃ is dissolved in HCl. [Diemer *J Am Chem Soc* 35 552 1913, DOI: 10.1021/ja02194a006; Block *Inorg Chem* 4 14 1953, Glemser & Sauer *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II p 1056 1965.] The trihydrate is commonly used for the preparation of **Au nanoparticles** [Bhattacharjee et al. *J Phys Chem B* 110 (13) 6768 2006, DOI: 10.1021/jp056675b; Wang et al. *Biomacro* 7(4) 1203 2006, DOI: 10.1021/bm060030f]; and for preparing stable nanosized and undervatised *colloidal Au* particles in solution [Fink et al. *Chem Mater* 10 922 1998, DOI: 10.1021/cm970702w].

Gold (I) cyanide [506-65-0] AuCN, M 223.0, m dec on heating, d²⁵ 7.14g/ml. The lemon yellow powder is sparingly soluble in H₂O and EtOH but soluble in aqueous NH₃. It is obtained by heating H[Au(CN)₂] at 110°. Wash it well with H₂O and EtOH and dry it at 110°. It has an IR band at ν_{max} 2239 cm⁻¹ typical for C≡N stretching vibration. [Glemser & Sauer *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II p 1064 1965.] **CARE:** may evolve HCN.

Gold (I) iodide [10294-31-2] AuI, M 323.9, m 120°(dec), d₄²⁰ 8.25. It has been prepared by heating gold and iodine in a tube at 120° for 4 months. Since it decomposes to Au and I₂ in the presence of UV light and heat, then the main impurity is Au. The salt is therefore purified by heating it at 120° with I₂ for several weeks. The crystals should be kept dry and in a cool place in the dark. [Weiss & Weiss *Z Naturforsch* 11B 604 1956.]

Gold (III) oxide hydrate [1303-58-8] Au₂O₃·xH₂O, AuH₃O₃, M 441.9 + xH₂O, evolves O₂ at 110°, pK₁²⁵ <11.7, pK₂²⁵ 13.36, pK₃²⁵ >15.3 [for Au(OH)₃]. The most probable impurities are SO₄²⁻ and Cl⁻ ions. Dissolve the brown powder in strong boiling KOH solution (ca 5M) and precipitate (**care**) with excess of 3N H₂SO₄. Then shake and centrifuge, resuspend in H₂O and repeat the washing several times until free from SO₄ and Cl ions. This gives a *wet oxide* that is dried in air, but decomposes to free gold in sunlight. It is advisable to **keep it wet** as it decomposes on drying (analyse wet sample). Store it away from light in the presence of H₂O vapour. It evolves O₂ at 110° to provide gold, being completed at 250°. It is insoluble in H₂O but soluble in HCl and concentrated HNO₃. [Roseveare & Buehrer *J Am Chem Soc* 49 1221 1927, DOI: 10.1021/ja01404a011.] The *wet oxide* solutions are used to decorate porcelain, and the decorations become gold in colour when fired.

Graphite (black lead, mineral carbon, plumbago) [7782-42-5] C, M 12.01, m 3652-3697°, d²⁵ 2.09, 2.23. This carbon allotrope is obtained by mining (e.g. in Shrilanka, Canada). It is normally soft, forms black/grey scales, and although even small crystals of it are rare it is composed of crystallised carbon containing traces of

SiO₂, Fe and other minerals. Treat graphite with hot 1:1 HCl. Then filter, wash and the dried powdered is heated in an evacuated quartz tube at 1000° until a high vacuum is obtained. Cool this and store it in an atmosphere of helium [Craig et al. *J Phys Chem* **60** 1225 1956, DOI: 10.1021/j150543a019; Ubbelohde & Lewis *Graphite and Its Crystals* Oxford 1960, Holliday et al. *Comprehensive Inorganic Chemistry* Bailar Jr et al. eds, **vol 1** 1250-1294 1973]. Also available in flakes, rods, powder and nanofibers. It has been used in numerous applications viz: strengthened with Fe and SiO₂ in 'lead' pencils, polishing, cements, pigments, lubricants, matches, with NaNO₃ in explosives, commutator brushes, arc lamps, anodes, cathode coatings, electroplating, in nuclear piles as moderator etc. See also entry on *Graphene* in Chapter 7.

Helium [7440-59-7] He, M 4.0, m -272.2°/26atm, b -268.93°/atm, d^{-270.3} 0.147. Dry the *inert* gas by passing it through a column of Linde 5A molecular sieves and CaSO₄, then through an activated-charcoal trap cooled in liquid N₂, to adsorb N₂, argon, xenon and krypton. Also pass it over CuO pellets at 300° to remove hydrogen and hydrocarbons, over Ca chips at 600° to remove oxygen, and then over titanium chips at 700° to remove N₂ [Arnold & Smith *JCS Faraday Trans 2* **77** 861 1981, DOI: 10.1039/F29817700861]. Its solubility in 100ml of H₂O is 0.94ml at 25°, 1.05ml at 50° and 1.21ml at 75°.

Hexachloroplatinic acid hydrate (chloroplatinic acid, platinum IV chloride solution) [16941-12-1] H₂PtCl₆ · xH₂O, M 409.8 + H₂O, m 60° (deliquescent solid). If it is to be purified, or regenerated from Pt recovered from catalytic hydrogenations, it should be dissolved in *aqua regia* followed by evaporation to dryness and dissolution in the minimum volume of H₂O. Then the aqueous solution is treated with saturated ammonium chloride until all the *ammonium hexachloroplatinate* [16919-58-7] separates. This (NH₄)₂PtCl₆ is filtered off and dried at 100°. Igniting this salt gives **Pt sponge**; dissolve the Pt sponge in *aqua regia*, boil to dryness, dissolve the residue in concentrated HCl, boil to dryness again and repeat the process. Protect it from light. [Wichers *J Am Chem Soc* **43** 1268 1921, DOI: 10.1021/ja01439a008; Adams et al. *Org Synth Coll Vol* **1** 463, 466 1941, DOI: 10.15227/orgsyn.008.0092; Bruce *J Am Chem Soc* **58** 687 1936, DOI: 10.1021/ja01295a501.]

Hexammine cobalt(III) chloride [10534-89-1] [Co(NH₃)₆]Cl₃, M 267.5, m 217°, d²⁵ 1.71g/ml. It crystallises from warm water (8ml/g) on cooling. It is soluble in NH₃. [Bjerrum et al. *Inorg Synth* **2** 216 1946, DOI: 10.1002/9780470132333.ch69.]

Hexammine ruthenium(III) chloride [14282-91-8] [Ru(NH₃)₆]Cl₃, M 309.6. Crystallise it twice from 1M HCl. In the presense of glucose oxidase (GOD) it is used as an electron mediator for the detection and estimation of glucose which is oxidised to D-glucono-δ-lactone.

Hydrazine (anhydrous) [302-01-2] NH₂NH₂, M 32.1, m 1.4°, 1.5-2.0°, b 47°/26mm, 56°/71mm, 113-113.5°/atm, n_D²⁰ 1.470, d²⁵ 1.032g/ml, pK₁²⁵ -0.88, pK₂²⁵ 8.11. Hydrazine hydrate is dried by refluxing with an equal weight of KOH pellets for 3 hours, then distilled from fresh solid NaOH or BaO in a current of dry N₂. Use stainless steel or copper equipment. Hydrazine and its hydrates have **VERY IRRITATING** and **TOXIC** vapours and should be used in an efficient fume cupboard. Store in a well-stoppered vessel, preferably under N₂. It is a useful **reducing agent**. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 469-472 1963, Fieser **1** 434, **2** 211, **3** 153, **4** 248, **5** 327, **6** 280, **7** 170, **8** 245, **9** 236, **11** 255, **12** 241, **13** 144, **16** 174.]

Hydrazine hydrate (N₂H₄ · x H₂O) [10217-52-4] NH₂NH₂ · xH₂O, M 32.05(anhydr) + x 18.02, usually 50-60% N₂H₄ in H₂O, d²⁵ 1.029g/ml. Hydrated hydrazine can be obtained as above and diluted as required. Solutions containing various amounts of H₂O (e.g. 35 wt%), and a 1M solution in THF [302-01-2] are also available commercially.

Hydrazine monohydrate (N₂H₄ · H₂O) [7803-57-8] NH₂NH₂ · H₂O, M 50.1, m -51.7°, b 120.1°/atm, d²⁵ 1.032g/ml, n_D²⁰ 1.428. It is best obtained by heating hydrazine sulphate (200g), NaOH (160g) and H₂O (75ml, exothermic) in a copper flask under reflux for 1.5 hours then distilled off (using a flame to remove all the hydrazine). The distillate (175ml) is a clear liquid which contains ~40-45% of N₂H₄. **Note** that hydrazine

attacks glass, rubber and cork, and stainless steel equipment should be used. The percentage of hydrazine is determined by titration with standard acid (methyl orange indicator) or against standard iodine (starch indicator). **Hydrazine monohydrate** should contain 64% of N_2H_4 . The ~40-45% solution may be concentrated by mixing it (144ml) with xylene (230ml) and distilling it through an efficient fractionating column (e.g. Hempel column). All the xylene passes over with about 85ml of H_2O . On distilling the residue, hydrated hydrazine (50ml) is obtained containing 80-85% of N_2H_4 . This can be diluted with conductivity H_2O to 64% N_2H_4 to give the **monohydrate**. **Anhydrous** hydrazine has also been obtained from mixtures of hydrazine monohydrohalides and NH_4Cl in four different ways [Sisler et al. *J Am Chem Soc* **76** 3914 1954, DOI: 10.1021/ja01644a013]. Hydrazine and its hydrates have **VERY IRRITATING** and **TOXIC** vapours and should be used in an efficient fume cupboard. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 469-472 1963, Fieser **1** 434, **2** 211, **3** 153, **4** 248, **5** 327, **6** 280, **7** 170, **9** 236, **11** 255, **12** 241, **13** 144, **16** 174.]

Hydrazine dihydrochloride [5341-61-7] $\text{NH}_2\text{NH}_2 \cdot 2\text{HCl}$, **M 105.0**, **m 198°**, **200°(dec)** **d**²⁵ **1.42g/ml**. It is recrystallised from aqueous EtOH and dried under vacuum over CaSO_4 .

Hydrazine monohydrochloride [2644-70-4] $\text{NH}_2\text{NH}_2 \cdot \text{HCl}$, **M 68.5**, **m 89°**, **91-92.5°**, **92.5°**. Prepare it by dropwise addition of cold concentrated HCl to cold liquid hydrazine in equimolar amounts. The crystals are harvested from water and are twice recrystallised from absolute MeOH and dried under a vacuum [Kovack et al. *J Am Chem Soc* **107** 7360 1985, DOI: 10.1021/ja00311a024]. Alternatively, $\text{NH}_2\text{NH}_2 \cdot 2\text{HCl}$ is heated and the low melting product recrystallised from H_2O to give the **monohydrochloride m 92.5°** [Sisler et al. *J Am Chem Soc* **76** 3914 1954, DOI: 10.1021/ja01644a013.]

Hydrazine sulfate [10034-93-2] $\text{NH}_2\text{NH}_2 \cdot \text{H}_2\text{SO}_4$, **M 130.1**, **m 254°**. Crystallise it from H_2O . Its solubility in H_2O is 3% at room temperature, but is very soluble in hot H_2O . It is a suspected **carcinogen**. [Adams & Brown *Org Synth Coll Vol* **1** 309 1941, DOI: 10.15227/orgsyn.002.0037; Audrieth et al. *Inorg Synth* **I** 90 1939, DOI: 10.1002/9780470132326.ch31.]

Hydrazoic acid (hydrogen azide, triazoic acid) [7782-79-8] H_3N , **M 43.0**, **m -80°**, **b 37°/atm**, **d**²⁵ **1.09g/ml**, **pK**²⁵ **4.72**. The free acid is **HIGHLY EXPLOSIVE** and **POISONOUS**, so it is prepared and used in solution, and in an **efficient fume cupboard**. It is a very useful reagent for Schmidt and related reactions [Wolff *Organic Reactions* **3** 307 1946, DOI: 10.1002/0471264180.or003.08]. Solutions in C_6H_6 or CHCl_3 are prepared by making a paste from NaN_3 (65g, 1mole) in warm H_2O (65ml), and added to C_6H_6 or CHCl_3 (400ml), cooled to 0°, and concentrated H_2SO_4 (26.6mls, 0.5mol) is dropped into the stirred mixture while carefully controlling the temperature between 0° and 5°. The organic layer is separated, dried (Na_2SO_4), and stored in a cold room. Its concentration is determined by shaking a small aliquot with ten times its volume of H_2O in a glass-stoppered flask, and titrating it against standard alkali. It can then be diluted to the desired concentration. [Audrieth et al. *Inorg Synth* **1** 77 1939, DOI: 10.1002/9780470132326.ch26; Frost et al. *J Am Chem Soc* **55** 3516 1933, DOI: 10.1021/ja01336a003; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 472-474 1963.] The pure acid is a mobile, pungent explosive liquid, and **poisonous** compound, causing irritation of membranes, headaches, palpitation, lowering blood pressure and ataxia — **handle with great care**.

Hydriodic acid [10034-85-2] **HI**, **M 127.9**, **b 127°(aqueous azeotrope)**, **d**₄²⁰ **1.701**, **pK**²⁵ **-8.56**. Iodine can be removed from aqueous HI, probably as the amine hydrogen triiodide, by three successive extractions using a 4% solution of Amberlite LA-2 (a long-chain aliphatic amine) in CCl_4 , toluene or petroleum ether (10ml per 100ml of acid). [Davidson & Jameson *Chem Ind (London)* 1686 1963.] Extraction with tributyl phosphate in CHCl_3 or other organic solvents is also suitable. Alternatively, a De-acidite FF anion-exchange resin column in the OH-form using 2M NaOH, then into its I-form by passing dilute KI solution through, can be used. Passage of an HI solution under CO_2 through such a column removes polyiodide. The column can be regenerated with NaOH. [Irving & Wilson *Chem Ind (London)* 653 1964]. The earlier method was to reflux with red phosphorus and distil in a stream of N_2 . The colourless product is stored in ampoules in the dark [Bradbury *J Am Chem Soc* **74** 2709 1952, DOI: 10.1021/ja01131a005; Heisig et al. *Inorg Synth* **1** 157 1939, DOI: 10.1002/9780470132326.ch54]. It fumes in moist air. **HARMFUL VAPOURS**.

Hydrobromic acid [10035-10-6] **HBr**, **M 80.9**, **b 125°(aqueous azeotrope, 47.5% HBr)/atm**, **d**₄²⁰ **1.38** (34%

HBr, pK^{25} -8.69. A solution of aqueous HBr *ca* 48% (w/w, constant boiling) is purified by distilling twice with a little red phosphorus, and the middle half of the distillate is taken. (The azeotrope at 760mm contains 47.8% (w/w) HBr.) [Hetzer et al. *J Phys Chem* **66** 1423 1962, DOI: 10.1021/j100814a012]. Free bromine can be removed by Irvine and Wilson's method for HI (see above), except that the column is regenerated by washing with an ethanolic solution of aniline or styrene. Hydrobromic acid can also be purified by aerating with H₂S, distilling and collecting the fraction boiling at 125-127°. [Heisig & Andur *Inorg Synth* **1** 149(155) 1939, DOI: 10.1002/9780470132326.ch53.] Also commercially available as a ~33% HBr solution in AcOH [37348-16-6], M 80.1, d_4^{20} 1.40g/ml. **HARMFUL VAPOURS**.

Hydrochloric acid (muriatic acid) [7647-01-0] **HCl**, M 36.5, b 108.6°(aqueous azeotrope, 20.2% HCl), d_4^{20} 1.09(20%), pK^{25} -6.1. It is readily purified by fractional distillation as the constant boiling point acid, following dilution with H₂O. The constant-boiling fraction contains 1 mole of HCl in the following weights of distillate at the stated pressures: 179.555g (730mm), 179.766g (740mm), 179.979 (750mm), 180.193 (760mm), 180.407 (770mm). [Foulk & Hollingsworth *J Am Chem Soc* **45** 1220 1923, DOI: 10.1021/ja01658a016.] **Toxic**.

Hydrofluoric acid [7664-39-3] **HF**, M 20.0, b 112.2°(aqueous azeotrope, 38.2% HF), d_4^{20} 1.15 (47-53% HF), pK^{25} 3.21. It is freed from lead (Pb *ca* 0.002ppm) by co-precipitation with SrF₂, by addition of 10ml of 10% SrCl₂ solution per kilogram of the concentrated acid. After the precipitate has settled, the supernatant is decanted through a filter in a hard-rubber or paraffin lined-glass vessel [Rosenqvist *Am J Sci* **240** 356 1942, DOI: 10.2475/ajs.240.5.356]. An advantage of the method of Coppola and Hughes [*Anal Chem* **24** 768 1952, DOI: 10.1021/ac60064a047] is that the temperature of distillation (in a polyethylene still of special design) is 70° to 85°, and so boiling and entrapment of fine droplets by the vapour do not occur. Pure aqueous HF solutions (up to 25M) can be prepared by isothermal distillation in polyethylene, polypropylene or platinum apparatus. Thus: place in a polyethylene washing bowl two 250ml polyethylene beakers. Place in one beaker technical grade HF (150ml, 35M, 75% v/v). In the second beaker place de-mineralised water (150ml). Another washing bowl similar to the first one is then placed upside down on top of the first bowl making sure of a close fit. The de-mineralised water will absorb the HF vapour so that after a while pure HF will condense in the H₂O. After 2 days the pure HF is 12M, and by replenishing the technical acid beaker, pure 25M HF can be obtained. When ultra pure HF (50% v/v) obtained in this way was subjected to spectrochemical analysis, the concentrations of the 23 elements recorded were less than 10⁻³ µg/ml. [Kwestroo & Visser *Analyst* **90** 297 1965, DOI: 10.1039/AN9659000297]. It attacks glass and is used for etching glass. **HIGHLY TOXIC do not inhale**.

Hydrogen [1333-74-0] **H₂**, M 2.02, m -259.1°, b -252.9°/atm, d 0.08988g/L (gas at STP: 0°/101.325kPa), 0.07099g/L (liquid at b), d 0.0763g/ml (solid), 1 mole occupies 22.4L at STP. It is usually purified by passing through a suitable absorption train of tubes. Carbon dioxide is removed with KOH pellets, soda-lime or NaOH pellets. Oxygen is removed with a 'De-oxo' unit or by passage over Cu heated to 450-500° and Cu on Kieselguhr at 250°. Passage over a mixture of MnO₂ and CuO (Hopcalite) oxidises any CO to CO₂ (which is removed as above). Hydrogen can be dried by passage through dried silica-alumina at -195°, through a dry-ice trap followed by a liquid-N₂ trap packed with glass wool, through CaCl₂ tubes, or through Mg(ClO₄)₂ or P₂O₅. Other purification steps include passage through a hot palladium thimble [Masson *J Am Chem Soc* **74** 4731 1952, DOI: 10.1021/ja01139a001], through an activated-charcoal trap at -195°, and through a non-absorbent cotton-wool filter or small glass spheres coated with a thin layer of silicone grease. **Potentially VERY EXPLOSIVE in air or with O₂**.

Hydrogen bromide (anhydrous) [10035-10-6] **HBr**, M 80.9, m -67°, b -66.8°/atm. Dry it by passing it through Mg(ClO₄)₂ towers. This procedure is **hazardous** [Stross & Zimmermann *Ind Eng Chem News* **17** 70 1939, Publication date: January 20, 1939]. *Alternatively*, shake it with mercury, distil it through a -78° trap and condense it at -195°/10⁻⁵mm. It *fumes* in moist air. **HARMFUL VAPOURS**. It is soluble in H₂O. A constant boiling aqueous solution of HBr has b 126°/760mm, and its HBr concentration is 47.4% (see hydrobromic acid above). It is soluble in AcOH. [Schneider & Johnson *Inorg Synth* **1** 149 1939, DOI: 10.1002/9780470132326.ch53; Schmeisser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 282-286 1963.]

Hydrogen chloride [7647-01-0] **HCl**, M 36.5, m -114°, b -85°/760mm, d^{25} 1.2g/ml. Pass it through concent-

rated H_2SO_4 , then over activated charcoal and silica gel. It fumes in moist air. Hydrogen chloride in gas cylinders contains ethylene, 1,1-dichloroethane and ethyl chloride. The latter two may be removed by fractionating the HCl through a trap cooled to -112° . Ethylene is difficult to remove. HCl fumes in moist air. **HARMFUL VAPOURS.** Its solubility in H_2O is 82% at 0° . A constant boiling aqueous solution (azeotrope) has **b 108.6°/760mm** with an HCl concentration of ~20%, and is called *Hydrochloric acid (muriatic acid)* (see above). [Schmeisser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 280-282 1963.]

Hydrogen cyanide (anhydrous) [prussic acid, Blausäure (German)] [74-90-8] HCN, M 27.0, m -14 to -15° , b 25.7°/760mm, d^{25}_D 0.687g/ml, n^{20}_D 1.2675, pK^{25} 9.21 (aqueous acid). HCN is prepared from NaCN and H_2SO_4 , and dried by passage through H_2SO_4 and over CaCl_2 , then distilled in a vacuum system and degassed at 77°K before use [Arnold & Smith *JCS Faraday Trans 2* 77 861 1981, DOI: 10.1039/F29817700861]. Cylinder HCN may contain stabilisers against explosive polymerisation, together with small amounts of H_3PO_4 , H_2SO_4 , SO_2 , and water. It can be purified by distillation over P_2O_5 , then frozen in Pyrex bottles at Dry-ice temperature for storage. [Zeigler *Org Synth Coll Vol* 1 314 1941, DOI: 10.15227/orgsyn.007.0050; cf. Glemser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 658-660 1963.] Liquid HCN, like liquid ammonia, evaporates very slowly since the latent heat of evaporation is high, and keeps it in the liquid state because the temperature of the liquid is lowered to below its boiling point. **Note** that some individuals (genetic variants) cannot smell this odour (the HCN molecule does not span across their olfactory sensors) so should take great care when working with compounds that liberate HCN. Earlier chemists with this olfactory deficiency smoked tobacco pipes near the equipment because the HCN vapours sucked through the pipes produced a foul taste in the mouth. Cyanides, and particularly HCN are extremely poisonous, readily absorbed in the mouth; the **lethal dose** is of the order of 150-300mg, depending on the individual. It is a respiratory inhibitor, blocks electron transport by acting on mitochondrial cytochrome oxidase, leading to anaerobic metabolism and lactic acidosis. The face and body become blue, hence the german name. **EXTREMELY POISONOUS; all due precautions should be taken.**

Hydrogen fluoride (anhydrous, hydrofluoric acid) [7664-39-3] HF, M 20.0, m -83.55° , b $19.5^\circ/\text{atm}$, d^{25} 1.15g/ml, pK_a 3.17. It can be purified by trap-to-trap distillation, followed by drying over CoF_2 at room temperature and further distillation. *Alternatively*, it can be absorbed on NaF to form NaHF_2 which is then heated under vacuum at 150° to remove volatile impurities. The HF is regenerated by heating at 300° and is stored with CoF_3 in a nickel vessel, being distilled as required. (Water content should be *ca* 0.01%.) To avoid contact with base metals, use can be made of nickel, polychlorotrifluoroethylene and gold-lined fittings [Hyman et al. *J Am Chem Soc* 79 3668 1957, DOI: 10.1021/ja01571a016]. An aqueous solution is *hydrofluoric acid* (see above). It is **HIGHLY TOXIC and attacks glass.**

Hydrogen iodide (anhydrous, hydriodic acid) [10034-85-2] HI, M 127.9, m -50.8° , b -35.36° , d^{25} 1.701g/ml, d 2.85g/ml (-47°), pK_{Est} -10 (H_2O), pK_a 2.8 (MeCN) After removal of free iodine from aqueous HI, the solution is frozen, then covered with P_2O_5 and allowed to melt under vacuum. The gas evolved is dried by passing through P_2O_5 on glass wool. It can be freed from iodine contamination by repeated fractional distillation at low temperatures. It **fumes** in moist air, and an aqueous solution is *hydriodic acid* (see above). **HARMFUL VAPOURS, CORROSIVE.**

Hydrogen peroxide [7722-84-1] H_2O_2 , M 34.0, d^{20}_4 1.110, pK^{25} 11.65. The 30% material has been steam distilled using distilled water. Gross and Taylor [*J Am Chem Soc* 72 2075 1950, DOI: 10.1021/ja01161a055] made 90% H_2O_2 approximately 0.001M in NaOH and then distilled it under its own vapour pressure, keeping the temperature below 40° , the receiver being cooled with a Dry-ice/isopropyl alcohol slush. The 98% material has been rendered *anhydrous* by repeated fractional crystallisation in all-quartz vessels. **EXPLOSIVE IN CONTACT WITH ORGANIC MATERIAL.** Available as a 3, 30-32, 30 (ACS grade), 35 and 50 wt% in H_2O (some stabilised with ~200ppm of MeCN), as well as in various semiconductor PURANAL (Honeywell) grades.

Hydrogen peroxide urea adduct (UHP, urea hydrogen peroxide 1:1 complex, carbamide peroxide, Debrox, Hyperol) [124-43-6] $\text{CO}(\text{NH}_2)_2 \cdot \text{H}_2\text{O}_2$, M 94.1, m. $85-90^\circ(\text{dec})$, $90^\circ(\text{dec})$. It is a safe alternative to

H₂O₂ in various oxidation reactions. It is commercially available in tablets ('rapidly soluble', equivalent to ~30% H₂O₂) or as a white powder (with 15-17% active oxygen). It is usually used without purification after assaying for active oxygen. This is done by titration with potassium permanganate or by iodometry, i.e. titration of liberated iodine when glacial acetic acid containing Fe³⁺ and NaI are added. It can be recrystallised from 30% H₂O₂ in a molar ratio of ~2:3 by heating in a pyrex dish for a few minutes at ~60°, cooling and allowed to crystallise slowly by evaporation in a crystallising dish. It forms elongated white needles, but if the solution is seeded just before crystallisation and shaken gently for a few seconds, then small plates are formed. Preferably collect the crystals by centrifugation at low temperature and dry them at 0° *in vacuo*. When dry, it is stable at room temperature and it has been reported that the available oxygen content had not decreased noticeably after 12 months. However, it is best to store it dry at low temperature. It is soluble in organic solvents e.g. EtOH, Et₂O, CHCl₃, CH₂Cl₂ and Me₂CO with slow decomposition, and its solubility in H₂O is 40% where it also decomposes slowly. It decomposes slowly at 40-60°/20mm and at 55-70°/760mm in air, but decomposition appears to accelerate above 80°. It is very useful (and in many cases superior to *p*-chloroperbenzoic acid) in the oxidation of alkenes, (epoxides), aromatic hydrocarbons (to phenols), ketones (Baeyer-Villiger), sulfides (to sulfones) and N-heterocycles (to N-oxides) when using 5 to 10 molar ratios of oxidant in the presence of acetic or trifluoroacetic anhydrides. Care should be used with this reagent as it is **potentially explosive**. [Lu et al. *J Am Chem Soc* **63** 1507 1941, DOI: 10.1021/ja01161a055; Cooper et al. *Synlett* 533 1990, DOI: 10.1055/s-1990-21156; *Beilstein* **3 H** 54, **3 I** 25, **3 II** 45, **3 III** 105, **3 IV** 102.]

Hydrogen sulfide [7783-06-4] H₂S, M **34.1**, m **-59.6°**, **-60°**, b **-80°/atm**, pK₁²⁵ **7.05**, pK₂²⁵ **12.89**. Wash it through water, then pass the gas through a train of tubes containing saturated Ba(OH)₂ (2x), water (2x), and dilute HCl [Goates et al. *J Am Chem Soc* **73** 707 1951, DOI: 10.1021/ja01146a063]. It is available in gas cylinders. **HIGHLY POISONOUS**.

Hydroxylamine [7803-49-8] NH₂OH, M **33.0**, m **33.1°**, b **56.5°/22mm**, d₄²⁰ **1.226**, pK_a²⁰ **5.96** (pK_b **7.97**). Crystallise it from *n*-butanol at -10°, collect it by vacuum filtration and wash it with cold diethyl ether. Also available as a 50wt % solution in H₂O. **Harmful vapours, and potentially explosive**. [Hurd et al. *Inorg Synth* **1** 87 1939, DOI: 10.1002/9780470132326.ch30; Semon in *Org Synth Coll Vol* **1** 318 1932, DOI: 10.15227/orgsyn.003.0061.]

Hydroxylamine hydrochloride [5470-11-1] NH₂OH·HCl, M **69.5**, m **151°**, **155-157°(dec)**, d₄²⁵ **1.67g/cm³**. Crystallise the salt from aqueous 75% ethanol or boiling methanol, and dry it under vacuum over CaSO₄ or P₂O₅. It has also been dissolved in a minimum of water and saturated with HCl; after three such crystallisations, it is dried under a vacuum over CaCl₂ and NaOH. Its solubility at 20° is 85% in H₂O, 6% in EtOH and 12% in MeOH. [Hurd et al. *Inorg Synth* **1** 87 1939, DOI: 10.1002/9780470132326.ch30; Semon in *Org Synth Coll Vol* **1** 318 1941, DOI: 10.15227/orgsyn.003.0061; Fieser **1** 478, **2** 217, **7** 176, **9** 245, **11** 257, **15** 170.] It is a monoamine oxidase (MAO) inhibitor, inhibits platelet aggregation and is used in a simple assay for superoxide dismutase [Elstner & Heupel *Anal. Biochem.* **70** 616 1976, DOI: 10.1016/0003-2697(76)90488-7].

Hydroxylamine sulfate [10039-54-0] (NH₂OH)₂·H₂SO₄, M **164.1**, m **130°**, **170°(dec)**, d₄²⁵ **1.88g/cm³**. Crystallise it from boiling H₂O by cooling to 0° (solubility: 0.6g/ml at 20°). Strong **reducing agent**, converts aldehydes and ketones to oximes, acid chlorides to hydroxamic acids, used as a catalyst, swelling agent and a co-polymerisation inhibitor.

Hydroxylamine-O-sulfonic acid [2950-43-8] H₂NOSO₃H, M **113.1**, m **210-211°**, **215°(dec)**, pK_a⁴⁵ **1.48**. Stir the solid vigorously with anhydrous Et₂O and filter it off using large volumes of dry Et₂O. Drain dry at the pump for 5 minutes and then for 12-14 hours in a vacuum. Store it in a vacuum desiccator over concentrated H₂SO₄. Determine the purity by oxidation of iodide to I₂. It must be stored in a dry atmosphere at 0-4°. It decomposes slowly in H₂O at 25° and more rapidly above this temperature. [Matsuguma et al. *Inorg Synth* **5** 122 1957, DOI: 10.1002/9780470132364.ch32; Fieser **1** 481, **2** 217, **3** 156, **4** 256, **5** 343, **6** 290, **8** 250, **9** 245, **10** 207, **15** 170.]

Hydroxyurea [127-07-1] NH₂CONHOH, M **76.1**, m **70-72° (unstable form)**, m **133-136°**, **141° (stable form)**, pK_a²⁵ **10.6**. Recrystallise hydroxyurea from absolute EtOH (10g in 150ml). Note that the rate of solution

in boiling EtOH is slow (15-30 minutes). It should be stored in a cool dry place, but some decomposition could occur after several weeks. [Deghenghi *Org Synth Coll Vol* **5** 645 1973, DOI: 10.15227/orgsyn.040.0060.] The solubility in H₂O is 50mg/ml at ~25° and it can be crystallised from Et₂O. [Kofod *Acta Chem Scand* **10** 256 1956, DOI: 10.3891/acta.chem.scand.10-0256; Beilstein **3** IV 170.]

Hypophosphorous acid (Phosphinic acid) [6303-21-5] H₃PO₂, M 66.0, m 26.5°, d₄³⁰ 1.217, 1.13 and 1.04 for 50, 30-32, and 10% aqueous solutions resp, pK²⁵ 1.31 (H₃PO₂). Phosphorous acid is a common contaminant of commercial 50% hypophosphorous acid. Jenkins and Jones [*J Am Chem Soc* **74** 1353 1952, DOI: 10.1021/ja01125a514] purified this material by evaporating about 600ml in a 1L flask at 40°, under reduced pressure (in N₂), to a volume of about 300ml. After the solution was cooled, it was transferred to a wide-mouthed Erlenmeyer flask which was stoppered and left in a Dry-ice/acetone bath for several hours to freeze (if necessary, with scratching of the wall). When the flask was then left at ca 5° for 12 hours, about 30-40% of it liquefied, and was again filtered. This process was repeated, then the solid was stored over Mg(ClO₄)₂ in a vacuum desiccator in the cold. Subsequent crystallisations from *n*-butanol by dissolving it at room temperature and then cooling in an ice-salt bath at -20° did not appear to purify it further. The **free acid** forms **deliquescent** crystals m 26.5° and is soluble in H₂O and EtOH. The NaH₂PO₂ salt can be purified through an anion exchange resin [Klement *Z Anorg Allgem Chem* **260** 267 1949, DOI: 10.1002/zaac.19492600408.]

Indium [7440-74-6] In, M 114.8, m 155-158°, 156.6°, b 2000°/atm, d₄²⁰ 7.31. Before use, the metal surface is cleaned with dilute HNO₃, followed by thorough washing with water and an alcohol rinse. [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 856 1963.] The metal is commercially available as rods, bars, wire, foil, granular form and powder in a large variety of sizes and purity.

Indium (III) chloride [10025-82-8 (anhydrous)] InCl₃, M 211.2, m 586°, b 800°/atm, d²⁵ 3.46g/cm³, pK₁²⁵ 3.54, pK₂²⁵ 4.28, pK₃²⁵ 5.16 (for aqueous In³⁺). The anhydrous salt forms yellow **deliquescent** crystals which can be sublimed at 500-600° in the presence of Cl₂/N₂ (1:1) [does not melt]. It is resublimed in the presence of Cl₂/N₂ (1:10) and finally heated to 150° to expel excess Cl₂. It is soluble in THF and H₂O (exothermically) and should be stored in a tightly closed container. [Baxter & Alter *J Am Chem Soc* **55** 1943 1933, DOI: 10.1021/ja01332a023.] It is a Lewis acid catalyst, e.g. in Friedel-Crafts and Diels-Alder reactions, and is the **most available soluble salt of Indium** [Araki & Hirashita 'Indium trichloride' in *Encyclopedia of Reagents for Organic Synthesis* Paquette ed 2004, Wiley NY, DOI:10.1002/047084289]. It forms a **tetrahydrate** [22519-64-8] M 293.1, and a solution in H₂O (2 mol/mol) is also available commercially.

Indium (III) oxide [1312-43-2] In₂O₃, M 277.6, d₄²⁰ 7.18, m sublimes at 850°. Wash it with H₂O and dry it below 850°. It volatilises at 850° and dissolves in hot mineral acids to form salts. Store it away from light because it darkens due to the formation of free In.

Indium sulfate [13464-82-9] In₂(SO₄)₃, M 517.8, d²⁵ 3.44g/cm³. Crystallise the white-grey powder from dilute aqueous H₂SO₄. It is **hygroscopic**; store it in a well-stoppered vessel.

Indium (III) sulfate pentahydrate [17069-79-3] In₂(SO₄)₃·5H₂O, M 607.9, d₄²⁰ 3.44. Dissolve the salt in strong H₂SO₄ and slowly evaporate at ca 50°. Wash the crystals with glacial AcOH and then heat them in a furnace at a temperature of 450-500° for 6 hours. Its solubility in H₂O is 529g/L at 20°. The **pentahydrate** is converted to an **anhydrous hygroscopic** powder on heating at 500° for 6 hours; but heating above this temperature over N₂ yields the oxide-sulfate. Evaporation of neutral aqueous solutions provides basic sulfates. [Baxter & Alter *J Am Chem Soc* **55** 1943 1933, DOI: 10.1021/ja01332a023; Hattox & Vries *J Am Chem Soc* **58** 2126 1936, DOI: 10.1021/ja01302a012.] The **hydrate is a complex** in which the H₂O in the complex exchanges with water at the rate of 10,000,000/sec, so the NMR cannot detect the difference between complexed and uncomplexed ion [Wolfram et al. *Physical Chemistry Chemical Physics* **6** 5145 2004, DOI: 10.1039/b407419j].

Iodic acid [7782-68-5] HIO₃, M 175.9, m 118°(dec), d₄²⁰ 4.628, pK²⁵ 0.75, 0.79. Dissolve iodic acid in the minimum volume of hot dilute HNO₃, filter and evaporate in a vacuum desiccator until crystals are formed. Collect the crystals and wash them with a little cold H₂O, and dry them in air in the dark. It is soluble in H₂O:

269g/100ml at 20° and 295g/100ml at 40°. It is soluble in dilute EtOH and darkens on exposure to light. It is converted to $\text{HIO}_3 \cdot \text{I}_2\text{O}_5$ on heating at 70°, but at 220° complete conversion to HIO_3 occurs. [Lamb et al. *J Am Chem Soc* **42** 1636 1920, DOI: 10.1021/ja01453a014; Bray & Caulkins *J Am Chem Soc* **53** 44 1931, DOI: 10.1021/ja01352a007.]

Iodine [7553-56-2] I_2 , **M 253.8**, **m 113.7°**, **b 184.3°/atm**, **d²⁵ 4.93g/cm³(solid)**, **d 3.960 (liquid at 120°)**. The almost black crystals with violet vapour are usually purified by vacuum sublimation. Preliminary purifications include grinding with 25% by weight of KI, blending with 10% BaO and subliming, subliming with CaO, grinding to a powder and treating with successive portions of H_2O to remove dissolved salts, then drying, and recrystallising from *benzene. Barrer and Wasilewski [*Trans Faraday Soc* **57** 1140 1961, DOI: 10.1039/TF9615701140] dissolved I_2 in concentrated KI and distilled it, then steam distilled it three times and washed it with distilled H_2O . Organic material is removed by sublimation in a current of O_2 over platinum at about 700°, the iodine being finally sublimed under vacuum. It has a high vapour pressure, and Iodine crystals sometimes sublime on to the sides and top of its stoppered container. [Fieser **1** 495, **2** 200, **2** 220, **3** 159, **4** 258, **5** 346, **6** 293, **7** 179, **8** 256, **9** 248, **10** 210, **11** 201, **11** 261, **12** 253, **13** 148, **14** 181, **15** 172, **16** 182, **17** 153.] **HARMFUL VAPOURS.** I_2 is slightly soluble in H_2O (0.0013moles/L at 25°), but more so in aqueous KI solution where it is in equilibrium with KI_3 . It is soluble in organic solvents to varying amounts at 35° (g/100g solvent) solubilities are: * C_6H_6 (17.9), CCl_4 (6.2), heptane (4.29), CS_2 (21.6), 2,2-dimethylbutane (1.99), cyclohexane (3.90), EtOH (24.60), *p*-xylene (20.14), perfluoroheptane (0.019) and at 25°: * C_6H_6 (14.1), CS_2 (14.5), EtOH (21.4), Et_2O (25.2), cyclohexane (2.27) and heptane (1.70), and CCl_4 (16.4) [Hildebrand & Jenks *J Am Chem Soc* **42** 2180 1920, DOI: 10.1021/ja01456a008; Hildebrand et al. *J Am Chem Soc* **72** 1017 1950, DOI: 10.1021/ja01158a096]. It is a good and commonly used topical antiseptic.

Iodine monobromide [7789-33-5] IBr , **M 206.8**, **m 42°**, **42-50°**, **d²⁵ 4.416g/cm³**. The brown-black crystals are purified by repeated fractional crystallisation from its melt. The vapour dissociates on heating [Yost et al. *J Amer Chem Soc* **55** 552 1933, DOI: 10.1021/ja01329a016; Schmeisser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 291-292 1963]. A useful electrophile in a new preparation of polyketides synthons [Liu et al. *Org Lett* **8** 5393 2006, DOI: 10.1021/ol0623318].

Iodine monochloride [7790-99-0] ICl , **M 162.4**, **m 27.2°(α-form)**, **13.9°(β-form)**, **b 97.4°/atm**, **d²⁵ 3.24g/cm³**. Recrystallise repeatedly from its melt at low temperatures. The black crystals melt to a red-brown liquid. [Cornog et al. *Inorg Synth* **I** 165 1939, DOI: 10.1002/9780470132326.ch56.] A useful **electrophile** in a new preparation of polyketides synthons [Liu et al. *Org Lett* **8** 5393 2006, DOI: 10.1021/ol0623318].

Iodine pentafluoride [7783-66-6] IF_5 , **M 221.9**, **m -9.3°**, **-8.0°**, **b 97.85°/760mm**, **d²⁵ 3.250g/cm³**. Rogers et al. [*J Am Chem Soc* **76** 4843 1954, DOI: 10.1021/ja01648a022] removed dissolved iodine from IF_5 by agitating with a mixture of dry air and ClF_3 in a Fluorothene beaker using a magnetic stirrer. The mixture is transferred to a still, and the more volatile impurities are pumped off as the pressure is reduced below 40mm. The still is gradually heated (kept at 40mm) to remove the ClF_3 before IF_5 distilled. Stevens [*J Org Chem* **26** 3451 1961, DOI: 10.1021/jo01067a103] pumped IF_5 under vacuum from its cylinder, trapping it at -78°, then allowing it to melt in a stream of dry N_2 . It **reacts vigorously** with H_2O to form HF, is a strong fluorinating agent, is strongly oxidative; and forms nitriles with primary amines after hydrolysis with H_2O [Stevens *J Org Chem* **31** 2025 1966, DOI:10.1021/jo01344a539]. **HARMFUL VAPOURS.** [Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 159-160 1963, Booth & Pinkston Jr. 'The Halogen Fluorides' *Chem Rev* **41** 421-439 1947, DOI: 10.1021/cr60130a001.]

Iodine trichloride [865-44-1] ICl_3 , I_2Cl_6 , **M 233.3**, **m 33°**, **b 77°(dec)**. Purify ICl_3 by sublimation at room temperature. At 77° it decomposes to ICl and Cl_2 . It is stable as a 20% to 35% in hydrochloric acid. A strong chlorinating and oxidising agent. **Irritant, toxic vapours.** [Booth et al. *Inorg Synth* **1** 167 1939, DOI: 10.1002/9780470132326.ch57.] It is an interhalogen compound and is in fact a **dimer $\text{Cl}_2\text{ICl}_2\text{ICl}_2$** [Boswijk & Wiebenga 'The crystal structure of I_2Cl_6 (ICl_3)' *Acta Cryst* **7** 417 1954, DOI: 10.1107/S0365110X54001260]

Iridium [7439-88-5] **Ir**, **M 192.2**, **m 2450°**, **b ~4500°/atm**, **4130°/atm**, **d²⁵ 22.65g/cm³**. Iridium is a silver

white hard solid which oxidises on the surface in air. Scrape the outer tarnished layer until silver clear and store it under paraffin. It is stable to acids but dissolves in *aqua regia*. Available as a black powder, foil, and wire. [Gilchrist *Chem Rev* **32** 277 1943, DOI: 10.1021/cr60103a002.]

Iridium (IV) chloride hydrate (hexachloroiridic acid) [16941-92-7 ($6\text{H}_2\text{O}$), 207399-11-9 ($x\text{H}_2\text{O}$)] $\text{IrCl}_4 \cdot x\text{H}_2\text{O}$, **M 334.0**+ H_2O . If it contains nitrogen, then repeatedly concentrate a concentrated HCl solution until free from nitrogen, and dry free from HCl in a vacuum over CaO until crystals are formed. The olive-green solid (yellow at $\sim 700^\circ$) is very *hygroscopic*. [Woo & Yost *J Am Chem Soc* **53** 884 1931, DOI: 10.1021/ja01354a008; Grube in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p1592 1965.]

Iron (wire) [7439-89-6] **Fe**, **M 55.9**, **m 1535°**, **1538°**, **b 2750°/atm**, **2862°/atm**, **d 25 7.86g/cm 3** . Clean it in concentrated HCl, rinse in de-ionised water, then reagent grade acetone and dry it under vacuum. It reacts with mineral acids to give the respective iron salts and hydrogen. Available as chips, granular form, powder, rods, and wire. [Fieser **1** 519, **2** 229, **3** 167, **5** 357, **9** 257, **12** 263, **14** 184, **17** 157.]

Lanthanum [7439-91-0] **La**, **M 138.9**, **m 920°**, **b 3464°/atm**, **3470°/atm**, **d 25 6.19g/cm 3** . It is a shiny metal that slowly tarnishes in air due to oxidation. It slowly decomposes by H_2O in the cold and more rapidly on heating to form the hydroxide. The metal is cleaned by scraping off the tarnished areas until the shiny metal is revealed and stored under oil or paraffin. It burns in air at 450° . It exists in three forms: α -form, β -form and γ -form with transition temperatures of 310° and 864° , respectively. [Spedding et al. *Ind Eng Chem* **44** 553 1952, DOI: 10.1021/ie50507a034.]

Lead (II) bromide [10031-22-8] **PbBr $_2$** , **M 367.0**, **m 371°**, **373°**, **b 892°/atm**, **d 25 6.66g/cm 3** . Crystallise it from water containing a few drops of HBr (25ml of water per gram PbBr_2) between 100° and 0° . A neutral solution is evaporated at 110° , and the crystals that separate are collected by rapid filtration at 70° and dried at 105° (to give the *monohydrate*). Its solubility in H_2O is 0.5% (at $\sim 10^\circ$) and 5% (at $\sim 100^\circ$), (solubility product at 20° is 7.9×10^{-5}). To prepare the *anhydrous* bromide, the hydrate is heated for several hours at 170° and then in a Pt boat at 200° in a stream of HBr and H_2 . Finally it is fused [Clayton et al. *JCS Faraday Trans 1* **76** 2362 1980, DOI: 10.1039/F19807602362]. **POISONOUS**.

Lead (II) chloride [7758-95-4] **PbCl $_2$** , **M 278.1**, **m 501°**, **b 950°/atm**, **d 25 5.85g/cm 3** . Crystallise it from distilled water at 100° (33ml/g) after filtering through sintered-glass and adding a few drops of HCl, by cooling. After three crystallisations the solid is dried under vacuum or under anhydrous HCl vapour by heating slowly to 400° . The solubility in H_2O is 0.07% at $\sim 10^\circ$, and 0.43% at $\sim 100^\circ$, (solubility product at 20° is 2.4×10^{-4}). **POISONOUS**.

Lead (II) iodide [10101-63-0] **PbI $_2$** , **M 461.0**, **m 402°**, **b 954°/atm**, **d 25 6.16g/cm 3** . It crystallises from a large volume of water. The solubility in H_2O is 1.1% at $\sim 10^\circ$, and 3.3% at $\sim 100^\circ$. **POISONOUS**.

Lead monoxide [1317-36-8] **PbO**, **M 223.2**, **m 886°**, **888°**, **b 1477°/atm**, **d 25 9.53g/cm 3** . Higher oxides are removed by heating under vacuum at 550° with subsequent cooling under vacuum. It is red at room temperature but becomes yellow at high temperatures ($\sim 480^\circ$) reversibly. It has low solubility in H_2O (0.017g/L at 25°), but is soluble in concentrated alkalis, HCl and NH_4Cl . [Ray & Ogg *J Am Chem Soc* **78** 5994 1956, DOI: 10.1021/ja01604a010; Kwestroo et al. *J Inorg Nucl Chem* **29** 39 1967, DOI: 10.1016/0022-1902(67)80141-6.] **POISONOUS**.

Lead sulfide (Galena) [1314-87-0] **PbS**, **M 239.3**, **m 1112°**, **d 25 7.5g/ml**. The sulfide is a black powder which is almost insoluble in H_2O ($4.9\text{g} \times 10^{-11}\text{g/L}$ at $\sim 25^\circ$; solubility product at 20° is 4.2×10^{-29}) meaning that traces of lead can be detected in solution by bubbling H_2S through it or adding aqueous H_2S solution to it. Thus, lead can be detected in the supernatant of an aqueous solution of precipitated lead chloride (cf. above, solubility product at 20° is 2.4×10^{-4}). If PbS is required pure then it is best prepared by precipitating it by bubbling washed H_2S through a solution of pure recrystallised lead acetate, filtering off the sulfide, washing it thoroughly with H_2O and drying it in a vacuum. *Note* that the precipitate is often red in the presence of hydrochloric acid

due to the initial formation of **red lead sulfochloride** ($PbS.PbCl_2$) which decomposes on dilution, or by further passage of H_2S to form the black sulfide [Lenher *J Am Chem Soc* **23** 680 1901, DOI: 10.1021/ja02035a008]. Although PbS has a high melting point it will sublime under N_2 above 860° . It dissolves in boiling dilute HNO_3 with precipitation of S ; and with concentrated HNO_3 it is completely converted to the insoluble white $PbSO_4$ (solubility product at 20° is 2.3×10^{-8}).

Lead sulfide paper test for H_2O_2 . The paper is prepared by soaking filter paper in aqueous lead acetate solution and exposing it to a small amount of H_2S then drying it in a vacuum desiccator, and storing the brown paper in a stoppered container. A drop of a neutral or slightly acidic H_2O_2 solution on the paper will show up as a white spot on a brown background. The white spot is from $BaSO_4$ formed by oxidation. The sensitivity for H_2O_2 is $0.07\mu g$ with a concentration limit of 1 in 700,000.

Lead nitrate [10099-74-8] $Pb(NO_3)_2$, **M 331.2, m 270° (dec), d 25 $4.53g/cm^3$** . Precipitate it twice from a hot (60°) concentrated aqueous solution by adding HNO_3 . The precipitate is sucked dry on a sintered-glass funnel, then transferred to a crystallising dish which is covered by a clock glass and left in an electric oven at 110° for several hours [Beck et al. *Trans Faraday Soc* **55** 331 1959, DOI: 10.1039/TF9595500331]. Its solubility in H_2O is 0.38g/ml (0°), 0.52g/ml (20°), 1.27g/ml (100°); in EtOH it is 0.04g/100ml ($\sim 25^\circ$), in MeOH it is 1.3g/100ml ($\sim 25^\circ$), and virtually insoluble in HNO_3 . After two recrystallisations of ACS grade, no metals above 0.001ppm were detected. **POISONOUS.**

Lithium (metal) [7439-93-2] **Li, M 6.9, m 180.5° , b $1342^\circ/atm$, d 25 $0.534g/cm^3$** . After washing with petroleum ether to remove storage oil, lithium is fused at 400° and then forced through a 10-micron stainless-steel filter with argon pressure. It is again melted in a dry-box, skimmed, and poured into an iron distillation pot. After heating under a vacuum to 500° , cooling and returning it to the dry-box for a further cleaning of its surface, the lithium is distilled at 600° using an all-iron distillation apparatus [Gunn & Green *J Am Chem Soc* **80** 4782 1958, DOI: 10.1021/ja01551a008].

Lithium aluminium hydride [16853-85-3] $LiAlH_4$, **M 37.9, m 125° (dec)**. Extract it with Et_2O , and, after filtering, the solvent is removed under vacuum. The residue is dried at 60° for 3 hours, under high vacuum [Ruff *J Am Chem Soc* **83** 1798 1961, DOI: 10.1021/ja01469a006]. It is a strong **reducing agent**. Store it in aliquots in a strictly dry atmosphere, and use in these aliquoted quantities. **It IGNITES in the presence of a small amount of water and reacts with it EXPLOSIVELY.** [Becher in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 805 1963; Fieser **1** 581, **2** 242, **3** 176, **4** 291, **5** 382, **6** 325, **7** 196, **8** 286, **9** 274, **10** 236, **11** 289, **12** 272, **13** 158, **14** 190, **15** 184.] Also commercially available as a 1.0M solution in Et_2O , 0.5M solution in 2-methoxyethyl ether, 0.5M solution in ethylene glycol dimethyl ether, 1.0M and 2.0M solution in THF, and as a $\sim 3.5M$ suspension in THF/toluene.

Lithium amide [7782-89-0] $LiNH_2$, **M 23.0, m $380-400^\circ$, d 17.5 1.178 , d 25 $1.178g/cm^3$** . Purify it by heating at 400° while NH_3 is passed over it in the upper of two crucibles (the upper crucible is perforated). The $LiNH_2$ will drip into the lower crucible through the holes in the upper crucible. The product is cooled in a stream of NH_3 . Protect it from air and moisture, store it under N_2 in a clear glass bottle sealed with paraffin. Store it in small quantities so that all the material is used once the bottle is opened. If the colour of the amide is yellow, it should be destroyed as it is likely to have oxidised and to **EXPLODE**. On heating above 450° it is decomposed to Li_2NH , which is stable up to $750-800^\circ$. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 463 1963, Greenlee et al. *Inorg Synth* **2** 128 1953, DOI: 10.1002/9780470132333.ch38.]

Lithium azide [19597-69-4] LiN_3 , **M 49.0, m $115-298^\circ$ (dec), pK 25 4.72 (HN_3)**. Digest $\sim 1g$ with 15ml of 96% EtOH at 35° , filter and dry it in air at temperatures below 80° . Store it in a cool place and treat it as **potentially explosive**. Its solubility in H_2O is 66.4% at 16° , and 20.26% at 16° in EtOH. [Hofmann-Bang *Acta Chem Scand* **11** 581 1957, DOI: 10.3891/acta.chem.scand.11-0581; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 581 1963.] Also available as a 20wt % solution in H_2O . **Poisonous.**

Lithium borohydride [16949-15-8] $LiBH_4$, **M 21.8, m 268° , 275° (dec), b 380° (dec), d 25 $0.666g/cm^3$** . It is crystallised from Et_2O , and pumped free of ether at $90-100^\circ$ during 2 hours [Schaeffer et al. *J Am Chem Soc* **78**

729 1956, DOI: 10.1021/ja01585a011]. Store it dry as it decomposes slowly in moist air. [Becher in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 775 1963, Fieser 1 603, 4 296, 11 293, 12 276, 14 191, 15 186.] Also commercially available as a 2.0M solution in THF.

Lithium bromide [7550-35-8] **LiBr**, **M 86.8**, **m 550°**. Crystallise it several times from water or EtOH, then dry it under high vacuum for 2 days at room temperature, followed by drying at 100°. Its solubility in H₂O is 167% at ~20°, and 250% at ~100°. It is *deliquescent* and should be stored in a tightly stoppered vessel. [Fieser 1 604, 2 245, 4 297, 13 332.]

Lithium carbonate [554-13-2] **Li₂CO₃**, **M 73.9**, **m 552°, 618°, d²⁵ 2.11g/cm³ (powder)**. Crystallise it from water. Its solubility decreases as the temperature is raised. The solubility in H₂O is 1.3% at ~10°, and 0.7% at ~100°. [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 987 1963, Calby et al. *Inorg Synth* 1 1 1939, DOI: 10.1002/9780470132326.ch1; Fieser 1 606, 3 183, 4 192, 3 396.]

Lithium chloride [7447-41-8] **LiCl**, **M 42.4**, **m 600°, 605°, 723°, d²⁵ 2.068g/cm³, n_D²⁴ 1.662**. Crystallise the white powder from water (1ml/g) or MeOH and dry it for several hours at 130°. Other metal ions can be removed by preliminary crystallisation from hot aqueous 0.01M disodium EDTA. It has also been crystallised from concentrated HCl, fused in an atmosphere of dry HCl gas, cooled under dry N₂ and pulverised in a dry-box. Runner & Wagner [*J Am Chem Soc* 74 2529 1952, DOI: 10.1021/ja01130a022] precipitated it with ammonium carbonate, washed it with Li₂CO₃ five times by decantation and finally with suction, then dissolved it in HCl. The LiCl solution is evaporated slowly with continuous stirring in a large evaporating dish, the dry powder being stored (while still hot) in a desiccator over CaCl₂. It has the following solubilities in H₂O: 0.68g/ml (0°), 0.84g/ml (25°), 1.23g/ml (100°); in EtOH: 0.14g/ml (0°), 0.25g/ml (30°), 0.23g/ml (60°); in HCOOH: 0.27g/ml (25°); in Me₂CO: 0.012g/ml (20°), 0.008g/ml (25°), 0.006g/ml (50°); in liquid NH₃: 0.0054g/ml (-34°) and 0.032g/ml (25°). [Fieser 1 609, 2 246, 4 298, 5 677, 13 332, 16 194.] It forms a *hydrate* [85144-11-2] LiCl·xH₂O [Fieser 13 332, 16 194.] **TOXIC, affects the nervous system.**

Lithium dimethylaminoborohydride (in 1M solution in THF) [53042-33-4] **Li(Me₂N)BH₃**, **M 64.8**, **d²⁵ 0.882g/cm³, n_D²⁰ 1.423**. This reagent is prepared under a N₂ atmosphere by reacting 1mol of borane (BH₃) or borane-dimethylsulfide (BMS) in THF with 1 mol of Me₂NH (or other primary or secondary amine) at ~25° for 1 hour followed by addition of just under 1mol of *n*BuLi at 0° slowly, and the temperature allowed to rise to ~25° while stirring, and stirred for a further hour. These solutions are kept in vials with a *sure-fit* cap and are stable for at least 9 months. The solid reagent can be obtained by removing the solvent at 0°/1mm. **Note** that care should be taken that a slight excess of *n*BuLi will cause the solid to be highly *pyrophoric*. The solid or solution retain their activity when kept under nitrogen as well as in a dry air atmosphere and are as stable as NaBH₄. It is however better to store the reagent as a solution. These **LABs (lithium aminoborohydrides)** are selective, air stable reducing agents capable of reducing a variety of functional groups [Fisher et al. *J Org Chem* 59 6378 1994, DOI: 10.1021/jo00100a046; Review: Pasumansky et al *Aldrichimica Acta* 38 61 2005, Saikai *Synlett* 995 2007, DOI: 10.1055/s-2007-973864]. They are also capable of transferring the amine group as in their reactions with halopyridines [Thomas et al *Org Lett* 5 3867 2003, DOI: 10.1021/ol035430j] and primary alkyl methanesulfonates [Thomas et al. *Org Lett* 3 3915 2001, DOI: 10.1021/ol0167659]

Lithium fluoride [7789-24-4] **LiF**, **M 25.9**, **m 842°, 845°, 848°, b 1676°, 1681°, d₄²⁰ 2.640**. Possible impurities are LiCO₃, H₂O and HF. These can be removed by calcining it at red heat, then pulverising it with a Pt pestle and storing it in a paraffin bottle. Its solubility in H₂O is 0.27% at 18°. It volatilises between 1100-1200°. [Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 235 1963].

Lithium hydride [7580-67-8] **LiH**, **M 7.95**, **m 680°, d₄²⁰ 0.76-0.77**. It should be a white powder; otherwise replace it. It darkens rapidly on exposure to air and is decomposed by H₂O to give H₂ and LiOH, and reacts with lower alcohols. One gram in H₂O liberates 2.8L of H₂ (could be explosive). [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 987 1963, Fieser 4 304, 13 165.]

Lithium hydroxide monohydrate [1310-66-3 (H₂O), 1310-65-2 (anhydrous)] **LiOH·H₂O**, **M 42.0**, **m 470°**

(dec), 471°, d 1.51, pK²⁵ 13.82. It crystallises from water (3ml/g) as the *monohydrate*. It *dehydrates* at 150° in a stream of CO₂-free air, and sublimes at 220° with some decomposition [Cohen *Inorg Synth* **5** 1 1957, DOI: 10.1002/9780470132364.ch1; Bravo & Groot *Inorg Synth* **7** 1 1963, DOI: 10.1002/9780470132388.ch1].

Lithium iodate [13765-03-2] LiIO₃, M 181.9, m 420-450°, d²⁵ 4.487g/cm³, n_D²⁰ 1.8875. Crystallise it from water and dry it in a vacuum oven at 60°. Its solubility in H₂O is 0.89g/ml (10°), 0.83g/ml (25°), 0.78g/ml (40°), and 0.73g/ml (75.6°). [Zachariasen & BartaLars 'Crystal Structure of Lithium Iodate'. *Physical Review Letters* **37** 1626 1931, DOI: 10.1103/PhysRev.37.1626.]

Lithium iodide [10377-51-2, 85017-80-7 (hydrate)] LiI, M 133.8, m 73° (3H₂O), 446°, 469°, b 1171°, d²⁵ 3.49g/cm³. Crystallise it from hot water (0.5mL/g) by cooling in a CaCl₂/ice/EtOH or from an acetone/Dry-ice bath. Dry it under a vacuum over P₂O₅ for 1 hour at 60° and then at 120°. It is *deliquescent* and should be stored in dark tightly stoppered vessels. [Fieser **1** 615, **4** 304, **5** 410, **7** 208, **9** 283, **10** 245, **11** 300, **12** 282, **13** 322.]

Lithium nitrate [7790-69-4] LiNO₃, M 68.9, m 253°, 255°, 264°, b 600°/atm(dec), d²⁵ 2.38g/cm³, n_D²⁰ 1.735. It crystallises from water [solubility: 0.52g/ml (20°), 0.9/ml (28°), 2.3g/ml (100°)] or EtOH. Dry it at 180° for several days by repeated melting under vacuum. If it is crystallised from water, and keeping the temperature above 70°, formation of *trihydrate* is avoided. The *anhydrous* salt is dried at 120° and stored in a vacuum desiccator over CaSO₄. After the 99% pure salt was recrystallised 3 times, it contained: metal (ppm) Ca (1.6), K (1.1), Mo (0.4), Na (2.2). [Donnan & Burt *J Chem Soc* **83** 335 1903, DOI: 10.1039/CT9038300335.]

Lithium nitrite monohydrate [13568-33-7] LiNO₂·H₂O, M 71.0 m 222°. Crystallise it from water by cooling from room temperature. The salt is very soluble in EtOH, so dissolve it in EtOH and allow to evaporate slowly which leaves white crystals on the side of the flask. On addition of a small amount to the saturated solution will provide larger needle-shaped crystals of the *monohydrate*. [Ball & Abram 'The nitrites of thallium, lithium, caesium, and rubidium' *JCS Transactions* **103** 2130 1913, DOI: 10.1039/CT9130302130.]

Lithium perchlorate [7791-03-9, 13453-78-6 (2H₂O)] LiClO₄, M 106.4, m 236°, pK²⁵ -2.4 to -3.1 (for HClO₄). Crystallise it from water or 50% aqueous MeOH. It is rendered *anhydrous* by heating the *trihydrate* at 170-180° in an air oven. It can then be recrystallised twice from acetonitrile and again dried under vacuum [Kosower & Mohammad *J Am Chem Soc* **93** 2713 1971, DOI: 10.1021/ja00740a022]. **SKIN IRRITANT.**

Lithium sulfate (anhydrous) [10377-48-7] Li₂SO₄, M 109.9, loses H₂O at 130° and m 845°, 859°, d²⁵ 2.22g/cm³(anhydr). Crystallise it from H₂O (4ml/g) by partial evaporation, and dry it above 130° *in vacuo*.

Lithium tetrafluoroborate [14283-07-9] LiBF₄, M 93.7, m 293-300°(dec), d²⁵ 0.852g/cm³(1.0M in MeCN), pK²⁵ 13.82 (Li⁺), pK²⁵ -4.9 (for HBF₄). Dissolve it in THF just below its solubility, filter from insoluble material and evaporate it to dryness in a vacuum below 50°. Wash the residue with dry Et₂O, and pass dry N₂ gas over the solid and finally heat it in an oven at 80-90°. Its solubility in Et₂O is 1.3g in 100ml at 25°; in THF it is 71g in 100ml at 25°. It is *hygroscopic* and is an **IRRITANT**. [Elliott et al. *J Am Chem Soc* **74** 5211 1952, DOI: 10.1021/ja01140a507; **75** 1753 1953.]

Lithium thiocyanate (lithium rhodanide) [556-65-0, 123333-85-7 (xH₂O)] LiSCN, M 65.0, pK²⁵ -1.85 (for HSCN). It crystallises from H₂O as the *dihydrate*, but on drying at 38-42° it gives the *monohydrate*. It can be purified by allowing an aqueous solution to crystallise in a vacuum over P₂O₅. The crystals are collected, dried out *in vacuo* at 80°/P₂O₅ in a stream of pure N₂ at 110°. [Coates & Taylor *J Chem Soc* 1245 1936, DOI: 10.1039/JR9360001245; *Beilstein* **3** III 251.]

Magnesium [7439-95-4] Mg, M 24.3, m 648°, 651°, b 1091°/atm, 1100°/atm, d₄²⁰ 1.739, d₄⁶⁵⁰ 1.584 g/cm³(liquid). It slowly oxidises in moist air and tarnishes. If too dark in colour, do not use it. The shiny solid should be degreased by washing with dry Et₂O, dry it *in vacuo* and keep it in a N₂ atmosphere. It can be activated by stirring it in Et₂O containing a crystal of I₂ then filtering it off, before drying and storing.

[Gmelin's Magnesium (8th edn) **27A** 121 1937; Amundsen et al. 'Magnesium'. *Ullmann's Encyclopedia of Industrial Chemistry* 2002 Wiley-VCH. DOI:10.1002/14356007.a15_559; Fieser **1** 627, **2** 254, **3** 189, **3** 254, **4** 315, **5** 419, **6** 351, **7** 218, **10** 251, **11** 307, **12** 290, **13** 170, **16** 198.] It is used in flares as it burns with a very bright light; and used for making light alloys.

Magnesium bromide (anhydrous) [7789-48-2] **MgBr₂**, **M 184.1**, **m 711°**, **d₄²⁰ 3.72**, **d²⁵ 3.72g/cm³**. Crystallise it from EtOH or H₂O (3.3g/ml). Dry it in a vacuum at ~150°, or heat the *hydrate* in a stream of HCl. It is very *deliquescent*. It forms *hexahydrate* crystals [13446-53-2] with **d²⁵ 2.0g/cm³**. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 909 1963.]

Magnesium chloride hexahydrate [7791-18-6 (6H₂O), 7786-30-3 (anhydrous)] **MgCl₂·6H₂O**, **M 203.3 (6H₂O)**, **95.2 (anhydr)** **m ~100°(dec)**, **pK₁²⁵ 10.3**, **pK₂²⁵ 12.2 (for Mg²⁺ hydrolysis)**. Crystallise it from hot water (3.3g/ml) by cooling. Dry it in a vacuum at ~175°, or heat the *hydrate* in a stream of HCl. When the hydrate is heated above 180° it is hydrolysed to the oxychloride (Mg₂OCl₂). It is *deliquescent*; store it in a well-stoppered vessel. [Bryce-Smith & Hunt *Inorg Synth* **6** 9 1960, DOI: 10.1002/9780470132371.ch4; Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 908 1963.]

Magnesium iodate tetrahydrate [13446-17-8 (4H₂O), 7790-32-1] **Mg(IO₃)₂·4H₂O**, **M 446.2**, **m 210°(dec)**. Crystallise from water (0.2g/ml) between 100° and 0°.

Magnesium iodide [10377-58-9] **MgI₂**, **M 278.1**, **m 634°**, **637°(dec)**, **d²⁵ 4.43g/cm³**. Crystallise it from water (0.8g/ml) by partial evaporation in a desiccator. It is *deliquescent* and should be stored in the dark. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 910 1963.]

Magnesium nitrate hexahydrate [13446-18-9] **Mg(NO₃)₂·6H₂O**, **M 256.4**, **m 89°(dec)**, **~95°(dec)**. Crystallise the nitrate from water (2.5mlg) by partial evaporation in a desiccator. It is *deliquescent* and is soluble in EtOH. After two recrystallisations, ACS grade salt has: metal (ppm) Ca (6.2), Fe (8.4), K (2), Mo (0.6), Na (0.8), Se (0.02).

Magnesium perchlorate (Anhydron, Dehydrite) [10034-81-8 (anhydrous)] **Mg(ClO₄)₂**, **M 223.2(anhyd)**, **m >250°**, **pK²⁵ -2.4 to -3.1 (for HClO₄)**. Crystallise it from water to give the *hexahydrate* **M 331.3** [13346-19-0]. Coll et al. [*J Am Chem Soc* **81** 1284 1959, DOI: 10.1021/ja01515a005] removed traces of unspecified contaminants by washing it with small portions of Et₂O and drying in a vacuum (**CARE**). The anhydrous salt is commercially available as an ACS reagent, and is as efficient a dehydrating agent as P₂O₅ and is known as 'Dehydrite' or 'Anhydron'. [Willard & Smith *J Am Chem Soc* **44** 2255 1922, DOI: 10.1021/ja01431a022; and Smith et al. *Ind Eng Chem* **16** 20 1924, DOI: 10.1021/ie50169a006.] It is *hygroscopic*; keep it in a tightly closed container. It is **EXPLOSIVE in contact with organic materials, and is a SKIN IRRITANT**.

Magnesium sulfate (anhydrous) [7487-88-9 (anhydrous), 14168-73-1 (H₂O), 17830-18-1 (6H₂O)] **MgSO₄**, **M 120.4**, **m 1127°**. Crystallise it from warm H₂O (1g/ml) by cooling. Dry the *heptahydrate* (*Epsom salt*) [10034-99-8 (7H₂O)] **M 246.5**, at ~250° until it loses 25% of its weight. Its solubility in H₂O is 36% at 20°, 55% at 60° and 74% at 100°; above 110° the solubility decreases with rise of temperature. Store it in a sealed container. [Fieser **1** 634, **4** 425, **5** 421.] Used as a purgative, in general anesthesia, local inflammation and septic wounds

Manganous(II) bromide (anhydrous) [13446-03-2 (anhydrous), 10031-20-6 (4H₂O)] **MnBr₂**, **M 214.8**, **m 695° (4H₂O)** [10031-20-6] **M 286.8**, **m 64°(dec)**. It forms rose-red *deliquescent* crystals which are soluble in EtOH. The H₂O is removed by heating at 100° then in HBr gas at 725°, or dry it in an atmosphere of N₂ at 200°.

Manganous(II) chloride tetrahydrate [13446-34-9 (4H₂O), 38639-72-4 (2H₂O), 7773-01-5 (anhydrous)] **MnCl₂·4H₂O**, **M 197.9(4H₂O)**, **125.8(anhydr)**, **m 58°**, **87.5°**, **650°**, **654°(anhydrous)**, **b 1190°/atm (anhydrous)**, **1225°/atm**, **d²⁵ 2.977g/cm³(anhydr)**, **d²⁵ 2.27g/cm³(2H₂O)**, **d²⁵ 2.01g/cm³(4H₂O)**. It crystallises from water (0.3ml/g) on cooling. The red-rose *tetrahydrate* melts at ~52° and forms the *dihydrate salt* which loses all its H₂O at 198° to give MnCl₂. It is soluble in EtOH. Another *hydrate* [73913-06-1 (xH₂O)]

has **m 122°** and **d**²⁵ 1.193g/cm³. It is soluble in EtOH and pyridine but insoluble in Et₂O. [Reidies 'Manganese Compounds', *Ullmann's Encyclopedia of Industrial Chemistry* 2002 Weinheim: Wiley-VCH, DOI:10.1002/14356007.a16_123].

Manganous sulfate monohydrate [10034-96-5 (H₂O), 10101-68-5 (4H₂O), 15244-36-7 (xH₂O), 7785-87-7 (anhydrous)] **MnSO₄ · H₂O**, **M 169.0**, **m 710°(anhydr)**, **27°(4H₂O)**, **b 850°/atm(anhydr)**, **d**²⁵ **3.25g/cm³(anhydr)**, **d**²⁵ **2.95g/cm³(H₂O)**, **d**²⁵ **2.107g/cm³(4H₂O)**. Crystallise it from water (0.9ml/g) at 54-55° by evaporating about two-thirds of the water. It *dehydrates* above 400°. The anhydrous salt is soluble in EtOH but insoluble in Et₂O. [Reidies 'Manganese Compounds' *Ullmann's Encyclopedia of Chemical Technology* 2007 Wiley-VCH, Weinheim, DOI:10.1002/14356007.a16_123.]

Mercuric bromide [7789-47-1] **HgBr₂**, **M 360.4**, **m 236°**, **b 320°/atm**, **322°/atm**, **d**²⁵ **6.03g/cm³**. Crystallise it from hot saturated ethanolic solution, dry and keep it at 100° for several hours under a vacuum, then sublime it. [Garrett *J Am Chem Soc* **61** 2744 1939, DOI: 10.1021/ja01265a055.] Its solubility in H₂O is 0.6% at 20°, and 22% at 100°; in EtOH it is 30% at 25°; and in MeOH it is 69.6% at 25°. [Wagenknecht & Juza *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1109 1965.] **POISONOUS**.

Mercuric chloride [7487-94-7] **HgCl₂**, **M 271.5**, **m 276°**, **b 304°/atm** **d**²⁵ **5.43g/cm³**, **n**_D²⁵ **1.859**, **pKa 3.2 (0.2M solution)**. Crystallise it twice from distilled water, dry it at 70° and sublime it under high vacuum. Its solubility in H₂O is 4.3% at ~0°, 6.6% at ~10° and 54% at ~100°. It is soluble in EtOH, Me₂CO, EtOAc, and is extracted into Et₂O (solubility: 4g/100ml at ~25°) from an aqueous solution. It is **very POISONOUS** and 0.2-0.4g is fatal. [Simon 'Mercury, Mercury Alloys, and Mercury Compounds' in *Ullmann's Encyclopedia of Industrial Chemistry* 2006, Wiley-VCH, Weinheim, DOI: 10.1002/14356007.a16_269.pub2]. The antidote is immediate administration of white of egg as an emetic.

Mercuric cyanide [592-04-1] **Hg(CN)₂**, **M 252.6**, **m 320°(dec)**, **d**²⁵ **3.996g/cm³**. Crystallise it from water. The solubility in H₂O is 8% at ~20° and 33% at ~100°; in EtOH it is 8% at ~20° and in MeOH it is 25% at ~20°. [Blitz *Z Anorg Allgem Chem* **170** 161 1928, DOI: 10.1002/zaac.19281700123.] **POISONOUS**.

Mercuric iodide (red) [7774-29-0] **HgI₂**, **M 454.4**, **m 259°(yellow >130°)**, **b ~350°(subl)**, **354°**, **d**₄²⁰ **6.3**. Crystallise it from MeOH or EtOH and wash it repeatedly with distilled water (solubility is 0.006% at ~25°). It has also been mixed thoroughly with excess 0.001M iodine solution, filtered, washed with cold distilled water, rinsed with EtOH and Et₂O, and dried in air. It changes colour reversibly to yellow at ~130°. [Friend *Nature* **109** 341 1922, DOI: 10.1038/109341b0.] **POISONOUS**.

Mercuric nitrate monohydrate [7783-34-8 (H₂O), 10045-94-0 (anhydrous)] **Hg(NO₃)₂ · H₂O**, **M 324.6(anhydr)**, **m 79°**, **d**²⁵ **4.30g/cm³**. This salt is prepared by reacting concentrated HNO₃ with Hg, and under these conditions the HNO₃ is an oxidising agent. The *anhydrous* salt is *deliquescent* and is soluble in a small volume of H₂O, from which the *monohydrate* can be obtained by slow evaporation at room temperature *in vacuo*. A 0.14N solution in H₂O (d²⁵ 1.025g/cm³) is also available commercially. However, with large volumes of H₂O, or on boiling, it is hydrolysed forming the insoluble basic salt. It is a strong *oxidising agent*, is light sensitive and is explosive in contact with EtOH in which it is insoluble. It is poisonous if absorbed through the skin. [Fieser **3** 197, **10**, 254, **11** 317.] **Caution as this is a poisonous substance**.

Mercuric oxide (yellow) [21908-53-2] **HgO**, **M 216.6**, **m 500°(dec)**, **d**²⁵ **11.14g/cm³**, **n**_{550nm}²⁵ **2.5**. Dissolve it in HClO₄ and precipitate it with NaOH solution. It is yellow when cold and changes to red at ~130° reversibly. Its solubility in H₂O is quite low: 0.0053g/100ml (25°) and 0.0395g/100ml (100°), and is insoluble in EtOH, Et₂O, Me₂CO and NH₃. [Fieser **1** 655, **2** 267, **4** 323, **5** 428, **6** 360, **7** 224, **8** 316, **9** 293, **12** 305.] **POISONOUS**.

Mercuric thiocyanate [592-85-8] **Hg(SCN)₂**, **M 316.8**, **m 165°(dec)**, **d**²⁵ **3.71g/cm³**, **pK_a²⁵ -1.85 (for HSCN)**. Recrystallise it from H₂O, and it can give various crystal forms depending on conditions. Its solubility in H₂O is 0.069% at 25°, but is more soluble at higher temperatures. It decomposes to Hg above 165°. **POISONOUS**. [Mason & Forgeng *J Phys Chem* **35** 1123 1930, DOI: 10.1021/j150322a017; Birckenbach & Kolb *Chem Ber* **68** 895 1935, DOI: 10.1002/cber.19350680532; *Beilstein* **3** IV 305.]

Mercurous nitrate dihydrate [7782-86-7 ($2\text{H}_2\text{O}$)] $\text{Hg}_2(\text{NO}_3)_2$, **M 561.2, m 70°(dec), d₄²⁵ 4.78g/cm³, pK_a²⁵ 2.68 (for Hg_2^{2+} hydrolysis).** Its solubility in H_2O containing 1% HNO_3 is 7.7%. Recrystallise it from a warm saturated solution of dilute HNO_3 and cool to room temperature slowly to give elongated prisms. Rapid cooling gives plates. The colourless crystals should be stored in the dark. **POISONOUS.** [For crystal structure see Grdenic *J Chem Soc* 1312 1956, DOI: 10.1039/JR9560001312]

Mercurous sulfate [7783-36-0] Hg_2SO_4 , **M 497.3, d₄²⁵ 7.56g/cm³.** The white-yellow powder is recrystallised from dilute H_2SO_4 , dried in a vacuum under N_2 , and stored in the dark. Its solubility in H_2O is 0.6% at 25°. It is hydrolysed by excessive washing with H_2O to form the greenish-yellow **basic salt** $\text{Hg}_2\text{SO}_4 \cdot \text{Hg}_2\text{O} \cdot \text{H}_2\text{O}$. Store it in the dark, as exposure to light decomposes it to Hg and HgSO_4 . **POISONOUS.**

Mercury [7439-97-6] **Hg, M 200.6, m -38.9°, b 126°/1mm, 184°/10mm, 261°/100mm, 356.9°/atm, d₄²⁰ 13.534 g/cm³.** After air has been bubbled through mercury for several hours to oxidise metallic impurities, it is filtered to remove coarser particles of oxide and dirt, then sprayed through a 4-ft column containing 10% HNO_3 . It is washed with distilled water, dried with filter paper and distilled under vacuum. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 8 1963.] A **mercury absorption kit**, available commercially, which consists of a bottle of **Mercurisorb** (100g, which absorbs its own wt of Hg in 5 minutes), a brush, a minishovel, disposable gloves, a container for adsorbed mixture, disposable pipettes and a compact case.

Molybdenum [7439-98-7] **Mo, M 95.9, m 2622°, b ~4825°, d₂₅²⁵ 10.3g/ml, resistivity of 5.0 $\mu\Omega\text{-cm}$ at 20°.** The dark-grey metal is commercially available in varying degrees of purity from 98 to 99.99%, as foil with thickness from 0.025mm (5.8g 150 x 150mm) to 1.0mm (25.6g, 50mm x 50mm or 102.4g, 100mm x 100 mm), or black powder from <150 μm to 1-2 μm particle size as well in nanopowder form of <100nm (BET) size. It is also available as wire of 1.0mm in diameter. The metal is stable in air but is oxidised at red heat to MoO_3 ; is unreactive towards dilute acids or alkalies but reacts with strong HNO_3 , H_2SO_4 , fused KClO_3 or KNO_3 , F_2 (at ~ 25°), Cl_2 and Br_2 (at red heat). [**CARE:** it is potentially quite toxic—particularly the nano sized particles.]

Molybdenum (VI) dichloride dioxide (MoO_2Cl_2) [13637-68-8] **MoO_2Cl_2 , M 198.8, d₂₅²⁵ 6.31g/cm³.** It is prepared from MoO_2 (40.0g, 313mmol) by placing it in a test tube with a wide side arm and a central tube which ends ~2cm above the surface of the oxide. The side arm is connected to a large round bottomed flask which vents through a bubbler containing H_2SO_4 . The apparatus is flushed with argon, and the tube containing the oxide is immersed in an oil bath which is kept at 140° for 12 hours to dry the oxide. The temperature of the tube is then raised to 160° and a stream of Cl_2 gas (pre-dried by bubbling through two H_2SO_4 traps) is allowed to flow over the hot oxide for 10 hours. The wide tube connecting the test tube and the flask is wrapped with a heating tape to encourage the MoO_2Cl_2 to sublime into the flask as it is formed. Stirring the contents of the flask (magnetic stirrer bar) hinders the clogging entry and exit tubes. MoO_2Cl_2 collects as ivory flakes that are quite pure for most purposes, but can be resublimed if necessary. All due **PRECAUTIONS** should be taken as chlorine is a **TOXIC GAS** and concentrated H_2SO_4 is used in the bubblers. [Schrock et al. *J Am Chem Soc* **112** 3875 1990, DOI: 10.1021/ja00166a023; Epperson et al. *Inorg Synth* **7** 163 1963, DOI: 10.1002/9780470132388.ch45.] Commercial fluffy MoO_2Cl_2 gives poor yields of substitution products (e.g. with ArNHSiMe_3), but the **THF complex $\text{MoO}_2\text{Cl}_2(\text{THF})_2$** [556907-19-8; 12081-12-8] is easier to handle, reacts in a similar way, and gives much higher yields of substitution products. It is readily prepared in solution by carefully adding solid MoO_2Cl_2 to dry THF at -30°. Addition of THF to MoO_2Cl_2 is too exothermic to keep under control. [Schrock et al. *J Am Chem Soc* **112** 3875 1990, DOI: 10.1021/ja00166a023.] Krauss & Huber prepared it by adding MoO_2Cl_2 (2.0g, 10mmol) in four portions to THF (7ml, 6.5g, 90mmol) while shaking. The yellow solution is diluted with light petroleum (5ml), filtered, the filtrate is diluted further with light petroleum (10ml) when the complex crystallises out. The supernatant is decanted off, the solid is washed several times with light petroleum (10ml lots), and dried at room temperature in a stream of N_2 to give analytically **pure $\text{MoO}_2\text{Cl}_2(\text{THF})_2$** (3.2g, 93%) as very pale yellow needles, **m > 50° (dec)**, that are soluble in THF, slightly soluble in Et_2O but insoluble in light petroleum [Krauss & Huber *Chem Ber* **94** 2864 1961, DOI: 10.1002/cber.19610941106]. It is very useful for preparing a variety of Mo complexes and catalysts.

Molybdenum hexafluoride [7783-77-9] **MoF_6 , M 209.9, m 17.5°, b 35°/760mm, d₄²⁰ 2.543.** Purify the hexa-

fluoride by low-temperature trap-to-trap distillation over pre-dried NaF. It is **hygroscopic**, fumes in moist air and is hydrolysed readily by H₂O. [Oppegard et al. *J Am Chem Soc* **82** 3835 1960, DOI: 10.1021/ja01500a011; Anderson & Winfield *JCS Dalton Trans* 337 1986, DOI: 10.1039/DT9860000337; Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 259 1963.] **Poisonous vapours, take necessary precaution.**

Molybdenum (IV) oxide [molybdenum (IV) dioxide] [18868-43-4] MoO₂, M 127.9, m 1100°(dec), d²⁵ 6.47g/cm³. It is prepared by grinding together ~10-15g of a mixture in a 2:1 ratio of MoO₃ (dried by heating in at 500° for 1 hour) and Mo metal powder in a quartz combustion tube under argon, or in an evacuated tube (CARE when opening), at 800° for 70 hours. It forms a brown-violet powder, oblong needles or thick platelets with a metallic lustre. It is insoluble H₂O, alkalies, HCl, HF, but only slightly soluble in hot HNO₃. [Conroy et al. *Inorg Synth* **14** 149 1973, DOI: 10.1002/9780470132456.ch31; **30** 105 1997, DOI: 10.1002/9780470132616.ch21; Herzog et al. in *Handbuch der Preparativen Anorganische Chemie* (Ed. Brauer) Enke Verlag Stuttgart Vol **3** p 1542 1981.] Beware of **TOXIC** vapours.

Molybdenum trichloride [13478-18-7] MoCl₃, M 202.3, m 1027°, d²⁰ 3.74. Boil it with 12M HCl, wash it with absolute EtOH and dry it in a vacuum desiccator. It is a brown-red powder soluble in H₂O, EtOH or Et₂O and gives a blue solution in concentrated H₂SO₄. [Hein & Herzog *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1404 1965, Larson et al. *Inorg Synth* **12**, 165 1970, DOI: 10.1002/9780470132432.ch29.]

Molybdenum trioxide [molybdenum (VI) oxide, MoO₃] [1313-27-5] MoO₃, M 143.9, m 795°, b 1155°/atm, d²⁰ 4.60. MoO₃ is prepared by adding HNO₃ to an aqueous solution of ammonium molybdate and stirring for several hours which precipitates H₂MoO₄. The acid is filtered off, washed with H₂O, heated at 150° for 1-2 hours and the hydrated oxide is dried at 450°. It recrystallises in rhombs from water (1g/50ml) between 70° and 0°, and is dried in air at ~500° for 1 hour before use. Its solubility in H₂O is 0.1% at 18°, and 2% at 70°. It is a white powder which turns yellow reversibly on heating. It sublimes readily at >780°/760mm in a quartz tube to give pure MoO₃. [Hein & Herzog *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1412 1965, Herzog et al. in *Handbuch der Preparativen Anorganische Chemie* (Ed. Brauer) Enke Verlag Stuttgart Vol **3** p 1544 1981.] It is used in the synthesis of Mo compounds, and of the ternary reduced molybdenum oxide Pr₄Mo₉O₁₈, which contains the previously unknown Mo₇, Mo₁₃, and Mo₁₉ clusters; and is a precursor for the preparation of fast ion conductors and superconductors [Arulraj et al. *Chem Mater* **14** 2492 2002, DOI: 10.1021/cm011239x ; Lacorre et al. *Nature* **404** 856 2000, DOI: 10.1038/35009069]. The oxide forms hydrates and alcoholates which have different IR spectra [Krauss & Huber *Chem Ber* **94** 2864 1961, DOI: 10.1002/cber.19610941106]. Beware of **TOXIC** vapours.

Molybdic acid [7782-91-4] H₂MoO₄, M 162.0, 180.0 (monohydrate), d²⁵ 3.124g/cm³ (for monohydrate), pK₁²⁵ 0.9 (proton addition), pK₂²⁰ 4.00, pK₃²⁰ 4.21. Treatment of an aqueous solution of 0.4N Na₂MoO₄ [1L, or of (NH₄)₂MoO₄ as in the preceding entry] at 60° with 30% HNO₃ (1L, prepared from 300ml of concentrated acid, d 1.42, in 1L of H₂O with cooling) and allowing to stand at 25° for several days deposited the theoretical amount of canary yellow monoclinic crystals which are collected, washed with ice water and dried *in vacuo* over H₂SO₄ for 2 weeks to give H₂MoO₄·H₂O. This is referred to as **α-molybdic acid** which on warming in H₂O at 70° is transformed to the white anhydrous **β-molybdic acid**. These acids have different X-Ray spectra and vapour pressures. **Colloidal molybdic acid** is obtained by dialysing an aqueous solution of (NH₄)₂MoO₄ and HCl, and forms a gum on evaporation. By heating these acids, or the ammonium salts, provide white MoO₃ which on heating further at 500° in the presence of H₂ gives reddish-brown MoO₂, and at 1200° a grey powder of metallic Mo is obtained (see preceding entries). Polymeric forms are known such as **tetramolybdic acid** [H₂Mo₄O₁₆, pK₁ 1.4, pK₂ 1.2, Chauveau et al. *Bull Soc Chim Fr* 1190 1959] and **heptamolybdic acid** [H₆Mo₇O₂₄, pK₅²⁵ ~3.7, pK₆²⁵ 4.33, Sasaki et al *J Inorg Nucl Chem* **9** 93 1959, DOI: 10.1016/0022-1902(59)80016-6]. **Note** that in aqueous alkaline solution the main Mo(VI) acid species are MoO₄²⁻, but in neutral and acidic media polynuclear species exist [cf. Sasaki & Sillén *Arkiv Kemi* **29** 253 1968, and Sasaki et al *J Inorg Nucl Chem* **9** 93 1959, DOI: 10.1016/0022-1902(59)80016-6]. [Herzog et al. in *Handbuch der Preparativen Anorganische Chemie* (Ed. Brauer) Enke Verlag Stuttgart Vol **3** p 1544 1981,

Rosenheim *Z Anorg Chem* **50** 320 1906, DOI: 10.1002/zaac.19060500131; Peters et al. *Z Anorg Allgem Chem* **365** 14 1969, DOI: 10.1002/zaac.19693650103; and for pKs see also Sasaki & Sillén *Acta Chem Scand* **18** 1014 1964, DOI: 10.3891/acta.chem.scand.18-1014; Rohwer & Cruywagen *J S African Chem Inst* **16** 26 1963, for first protonation constant of monomeric molybdic acid see Rohwer & Cruywagen *J S African Chem Inst* **17**(2) 145 1964, ISBN 03794350, http://hdl.handle.net/10520/AJA03794350_1072]

Monocalcium phosphate dihydrate (monobasic) [7789-77-7 ($2\text{H}_2\text{O}$), 7757-93-9 (anhydrous)] **$\text{CaHPO}_4 \cdot 2\text{H}_2\text{O}$, M 154.1, m 200° (dec, loses H_2O at 100°), d 25 2.31g/ml.** Crystallise it from a near-saturated solution in 50% aqueous reagent grade phosphoric acid at 100° by filtering through fritted glass and cooling to room temperature. The crystals are filtered off, and this process is repeated three times using fresh acid. For the final crystallisation the solution is cooled slowly with constant stirring to give thin plate crystals that are filtered off on a fritted glass funnel, washed free of acid with anhydrous acetone and dry in a vacuum desiccator [Egan et al. *J Am Chem Soc* **78** 1811 1956, DOI: 10.1021/ja01590a009].

Neodymium chloride hexahydrate [13477-89-9] **$\text{NdCl}_3 \cdot 6\text{H}_2\text{O}$, M 358.7, 250.6, m 124° , d 25 4.13g/cm³ (anhydr), d 25 2.282g/cm³ (hydrate), pK₁²⁵ 8.43 (for Nd^{3+} hydrolysis).** Neodymium chloride forms large purple-pink prisms from concentrated solutions of dilute HCl. **Hygroscopic.** The crystals are soluble in H_2O (2.46 parts in 1 part of H_2O) and EtOH, and lose H_2O at 160° . [Steele & Wertz 'Solvent effects on the coordination of neodymium(3+) ions in concentrated neodymium trichloride solutions' *Inorg Chem* **16** 1225 1977, DOI: 10.1021/ic50171a050.]

Neodymium(II) iodide [61393-36-0, 61393-39-0] **NdI_2 , M 398.1, 395.7, m 562° , d 25 5.85g/cm³.** This one electron reductant can be prepared in large quantities (~40g) by direct reaction of the metal and iodine at 600° [Evans et al. *Inorg Chem* **42** 3097 2003, DOI: 10.1021/ic0300316]. The black solid NdI_2 can be kept at room temperature for months in the absence of solvent. Solutions can be stored under argon for several hours at -15° but should be used as soon as possible. The stability is considerably reduced if N_2 is used as inert atmosphere, instead of argon, even at -30° . In tetrahydrofuran under argon it promotes a **pseudo-Barbier reaction**, i.e. rapid reductive coupling of primary and secondary (but not tertiary) halides with ketones, e.g. butylchloride and cyclohexanone provide almost quantitative yield of 1-butylhexan-1-ol [Evans et al. *Org Lett* **5** 2041 2003i, DOI: 10.1021/ol030033u]. Thus behaving somewhat like a Grignard reagent. This solution also has **Birch reduction-type reactivity**, e.g. reduces naphthalene to 1,4-dihydronaphthalene.

NdI_2 (THF)_x solution is prepared by transferring pre-cooled THF (45ml, at -15°) via a cannula into a septum-capped flask containing black NdI_2 (1.0g, 2.5mmol) and a magnetic stirrer bar at -15° whereby the colour of the solution becomes purple, and is stirred for 45 minutes. An aliquot should show the concentration to be 0.05M by compleximetric titration [method of Evans & Allen *J Am Chem Soc* **122** 2118 2000, DOI: 10.1021/ja992951m]. This solution is used for reactions that should be performed under argon.

Neodymium(III) iodide (anhydrous) [13813-24-6] **NdI_3 , M 524.7, m 775° , b $1370^\circ/\text{atm}$, d 25 5.85g/ml.** It is an almost black crystalline powder (green when ground) that is soluble in hot and cold H_2O . It can be prepared by the method of Bochkarev [Bochkarev & Fagin *Chem Eur J* **5** 2990 1999, DOI: 10.1002/(SICI)1521-3765(19991001)5:10<2990> where a small quantity of Nd metal is placed in a quartz crucible in a quartz reactor [see Evans et al. *Inorg Chem* **42** 3097 2003, DOI: 10.1021/ic0300316], heated to 600° , and I_2 is added, then small amounts of metal and I_2 are added alternately. Every addition of I_2 results in an orange glow in the mixture. When addition is complete (total ≥ 3 equivalents) the apparatus is cooled, and the crucible containing NdI_3 is transferred to an argon filled glovebox and the salt is ground (with a pestle and mortar) to a green powder. The metal content is analysed by titration (complexometric metal analysis is performed by dissolving the salt in H_2O at $\sim 25^\circ$, evaporating, ashing the residue at 500° , dissolving in HCl and the analysis is carried out in hexamethylenetetramine buffer with xylenol orange as indicator and EDTA as titrant: Schwarzenbach & Flaschka 'Complexometric Titrations' Metheun, London p 194 1969; Evans et al. *J Am Chem Soc* **103** 6672 1981, DOI: 10.1021/ja00412a023). It should contain at least 27.5% Nd (theoretical is 29.5% Nd). [Evans & Workman *Organometallics* **24** 1989 2005, DOI: 10.1021/om050033t.]

Potassium graphite KC_8 with NdI_3 has also been used for reductive coupling and is prepared *in situ*. [Weitz & Rabinovitz *JCS Perkin Trans I* 117 1993, DOI: 10.1039/P19930000117]. KC_8 can be prepared in a Schlenk

line, or in a glove box, by adding K metal (0.541g, 13.8mmol) to a scintillating vial containing a Teflon stirbar then Graphite (1.242g, 12.9mmol), and the mixture is stirred and heated until a bronze-coloured powder results. The *NdI₃/KC₈/THF* is prepared by adding precooled (-15° via a cannula) THF into a septum-capped Schlenk flask (~25ml) containing NdI₃, KC₈ and a Teflon stirbar thus producing a purple solution which is stirred at -15° for 40 minutes and is ready for the reaction. The alkyl halide is injected, stirred for 1 minute, and is followed by the aldehyde or ketone. [Evans & Workman *Organometallics* **24** 1989 2005, DOI: 10.1021/om050033t.]

Neodymium nitrate hexahydrate [16454-60-7 (6H₂O), 13746-96-8 (xH₂O)] **Nd(NO₃)₃ · 6H₂O**, **M 438.4(6H₂O), 330.3(anhydr)**, **m 40°, 69-71°, 70-72°**. It crystallises (red-purple crystals) with 5 and 6 molecules of H₂O from concentrated solutions in dilute HNO₃ by slow evaporation; 1 part is soluble in 10 parts of H₂O. Used for colouring glass.

Neodymium oxide [1313-97-9] **Nd₂O₃**, **M 336.5, m 2320°, d²⁰ 7.24g/cm³**. Dissolve it in HClO₄, precipitate it as the oxalate with doubly recrystallised oxalic acid, wash it free of soluble impurities, dry it at room temperature and ignite it in a platinum crucible at higher than 850° in a stream of oxygen. It is a blue powder. [Tobias & Garrett *J Am Chem Soc* **80** 3532 1958, DOI: 10.1021/ja01547a011.]

Neon [7440-01-9] **Ne**, **M 20.2, m -248.6°, b -246.1°/atm, d 0.9002g/L (at STP: 0°/101.325), d 1.207g/L (at -246.1°)**. Pass the gas through a copper coil packed with 60/80 mesh 13X molecular sieves, which is cooled in liquid N₂, or through a column of *Ascarite* (NaOH-coated silica/asbestos adsorbent).

Nickel bromide [13462-88-9] **NiBr₂**, **M 218.5, m 963°(loses H₂O at ~200°), d²⁵ 5.098g/cm³**. Crystallise it from dilute HBr (0.5ml/g) by partial evaporation in a desiccator. The *anhydrous* salt is yellow, but the *trihydrate* is green.

Nickel chloride hexahydrate [7791-20-0 (6H₂O), 69098-15-3 (xH₂O), 7718-54-9 (anhydrous)] **NiCl₂ · 6H₂O**, **M 237.7(6H₂O), m 140°(6H₂O), 1001°(anhydr), d²⁵ 1.92g/cm³(6H₂O), d²⁵ 3.55g/cm³(anhydr)**. It crystallises from dilute HCl to form the green *hexahydrate*. At 70° this dehydrates to the *tetrahydrate*, and at higher temperatures it forms the *anhydrous* salt. It sublimes in yellow hexagonal scales in a stream of HCl. Store it in a desiccator as it is *deliquescent*. [Hart & Partington *J Chem Soc* 104 1943, DOI: 10.1039/JR9430000104.]

Nickel nitrate hexahydrate [13478-00-7] **Ni(NO₃)₂ · 6H₂O**, **M 290.8, m 56°, 57°, d²⁵ 2.05g/cm³(6H₂O)**, Crystallise it from water (3.3g/ml) by partial evaporation in a desiccator. Store it in a desiccator as it is *deliquescent*.

Nickel(II) perchlorate hexahydrate [13520-61-1] **Ni(ClO₄)₂ · 6H₂O**, **M 365.7, m 140°, 200°, 209° (sealed tube), d⁰ 1.570, d²⁰ 1.583, d⁴⁰ 1.597, d⁵⁰ 1.646, d⁶⁰ 1.597**. The greenish blue hexagonal prisms of this salt are obtained by double decomposition between NiSO₄ and Ca(ClO₄)₂ in aqueous solution, filtering off the insoluble BaSO₄, and concentrating the filtrate. It recrystallises from H₂O, and drying the crystals in a vacuum desiccator over H₂SO₄ or CaCl₂ provides the *hexahydrate*. Alternatively, it is prepared from freshly precipitated NiO or NiCO₃ and aqueous HClO₄, filtering off the excess of NiCO₃ and concentrating the filtrate to crystallisation. The solubility of Ni(ClO₄)₂·6H₂O in H₂O (g/100ml) is 217 (0°), 236 (10°), 245 (20°), 267 (30°), 280 (35°), 273 (40°), 311 (50°), 295 (55°), 280 (60°). The salt form hydrates with 2, 4, 5, 6, 7, and 9 molecules of H₂O. The *hexahydrate* is normally obtained by crystallisation from H₂O and is the more stable hydrate. It forms the *tetrahydrate* at 110°/760mm/36 hours or 70°/1mm/66 hours, which yields the *dihydrate* at 130°/760mm/42 hours or 100°/1mm/72 hours. It should be kept in a desiccator over CaCl₂ as it is quite *hygroscopic*. The UV in EtOH has λ_{max} (logε) at 395 (~0.68), 660 (~0.22), 720 (~0.25) nm. Evaporation of a solution in ether with dry air gives *Ni(ClO₄)₂·Et₂O* as a yellow powder, and with dioxane it forms *Ni(ClO₄)₂·2C₄H₈O₂*. A solution in Me₂CO at ~2 x 10⁻⁴M is green with a characteristic absorption at 410nm [Katzin *Nature* **182** 1013 1958, DOI: 10.1038/1821013a0]; it complexes also with acetylacetone [Izatt et al. *J Phys Chem* **59** 235 1955, DOI: 10.1021/j150525a010], ethylene, trimethylenediamine [Cotton & Harris *J Phys Chem* **59** 1203 1955, DOI: 10.1021/j150534a006], DMF [Pflaum, & Popov *Anal Chim Acta* **13** 165 1955, DOI: 10.1016/S0003-2670(00)87919-2], and with pyridine it forms blue crystals of *[Ni(C₅H₅N)₆](ClO₄)₂·4H₂O* [Weinland et al. *Arch*

Pharm **265** 352 1927, DOI: 10.1002/ardp.19272651026], $Ni(ClO_4)_2 \cdot 4(C_5H_5N)$ and $Ni(ClO_4)_2 \cdot 6(C_5H_5N)$ [Sinha *J Indian Chem Soc* **35** 865 1958, Sinha & Ray *J Indian Chem Soc* **20** 32 1943]. [Salvadori *Gazzetta* **42** I 458 1912, Veeraiah & Qureshi *J Indian Chem Soc* **21** 127 1944, Bernath et al. *J Prakt Chem* **143** 298 1935, DOI: 10.1002/prac.19351431004; Freund & Schneider *J Am Chem Soc* **81** 4780 1959, DOI: 10.1021/ja01527a006; Gmelin's *Handbuch der Anorganischen Chemie, Nickel Teil B-Lieferung 2I*, Verlag Chemie GMBH Weinheim, System 57, pp 596-601 1966.]

Nickel sulfate hexahydrate [10101-97-0] $NiSO_4 \cdot 6H_2O$, **M 262.9, m loses $5H_2O$ at 100° , becomes anhydrous at $\sim 280^\circ$, d^{25} 2.07**. It crystallises from H_2O in bluish-green tetragonal crystals of the α -form which undergo a transition at 53.3° to the β -form that has green transparent monoclinic crystals. The crystals are stable at $\sim 40^\circ$, become blue in colour and then opaque in air due to slow efflorescence. Its solubility in H_2O is 0.6/ml at 0° and 3.4g/ml at 100° . It is sparingly soluble in EtOH, slightly more soluble in MeOH, soluble in aqueous NaOH, and very soluble in aqueous ammonia to give the bluish-green **monoclinic $(NH_4)_2Ni(SO_4)_2 \cdot 6H_2O$ double salt**. The pH of an aqueous solution is about 4.5, i.e. weakly acidic. (See also the heptahydrate below).

Nickel sulfate heptahydrate [10101-98-1, 7786-81-4 (anhydrous)] $NiSO_4 \cdot 7H_2O$, **M 280.9, 154.8(anhydr), m loses $5H_2O$ at 100° , anhydrous m at $\sim 280^\circ$, d^{25} 1.948 g/cm³**. The sulfate crystallises from warm water (4g/ml) or dilute H_2SO_4 as bright green monoclinic crystals on cooling. It is isomorphous with Epsom salt (see $MgSO_4 \cdot 7H_2O$ above). Prolonged exposure to air gives the blue **tetrahydrate**. On heating above 118° , it is converted to the **dihydrate**, which at $>280^\circ$ is converted to the **yellow anhydrous $NiSO_4$** which does not react with HCl.

Nickel(II) sulfide **M 90.8 (NiS, Mellirite, bronze-yellow, d^{25} 5.3-5.6), 240.2 (Ni_3S_2 [12035-72-2] Hazelwoodite, d^{25} 5.8 g/cm³) and (Ni_3S_2/NiS black powder [16812-54-7] d^{25} 5.3-5.6 g/cm³), m 790° , 797°** . The sulfide is generally obtained as a highly insoluble black precipitate by adding NH_4OH and NH_4SH to a nickel salt. It dissolves slowly in excess of $(NH_4)_2S_x$ and re-precipitates on boiling, exposure to air or addition of acid. A black dense sulfide is also obtained by boiling a nickel salt with $Na_2S_2O_3$. It is insoluble in dilute HCl, slowly soluble in concentrated HCl, readily in HNO_3 and **aqua regia**. The sulfides should be washed well with H_2O , dried, and analysed for Ni. Various forms have been prepared [Dunn & Rideal *J Chem Soc* **123** 1242 1923, DOI: 10.1039/CT9232301242.]

Niobium (Colombium) (V) chloride [10026-12-7] $NbCl_5$, **M 270.2, m $204.7-209.5^\circ$, b $\sim 250^\circ/atm$ (begins to sublime at 125°), d^{25} 2.75g/cm³**. It forms yellow, very deliquescent crystals which decompose in moist air to liberate HCl. Keep it in a dry box flushed with N_2 in the presence of P_2O_5 . Wash it with CCl_4 and dry it over P_2O_5 . The yellow crystals can contain a few small, dirty white pellets among the yellow needles. These should be easily picked out. Upon grinding in a dry box, however, they turn yellow. $NbCl_5$ has been sublimed and fractionated in an electric furnace. It forms complexes with Lewis bases. [Epperson et al. *Inorg Synth* **7** 163 1963, DOI: 10.1002/9780470132388.ch45; Alexander & Fairbrother *J Chem Soc* suppl 233 1949, DOI: 10.1039/JR949000S223.]

Nitric acid (Aqua fortis) [7697-37-2] HNO_3 , **M 63.0, m -42° , b 83° , d^{25} 1.5027, [Constant boiling acid has composition w/w of 68% HNO_3 + 32% H_2O , b 120.5° , d_4^{20} 1.41], pK^{25} -1.27 (1.19)**. The acid is obtained colourless (approx. 92%) by direct distillation of fuming HNO_3 under reduced pressure at $40-50^\circ$ with an air leak at the head of the fractionating column. **Concentrated nitric acid** is an aqueous solution containing 70-71% of HNO_3 (d_4^{20} 1.4134). Store it in a desiccator that is kept in a refrigerator away from light which causes the formation of NO_2 . Nitrite-free HNO_3 can be obtained by vacuum distillation from urea. [Ward et al. *Inorg Synth* **3** 11 1950, DOI: 10.1002/9780470132340.ch4; Kaplan et al. *Inorg Synth* **4** 52 1953, DOI: 10.1002/9780470132340.ch4.] If '**fuming nitric acid**' (90%) is yellow in colour (due to the presence of oxides of nitrogen), then treat 100mls with urea (0.5g) and bubble dry air through it until it is colourless (~ 20 minutes) [Freeman & Shepard *Org Synth* **43** 83 1963, DOI: 10.15227/orgsyn.043.0083]. Acid that is free from oxides does not discolour drops of 1N $KMnO_4$. **Anhydrous HNO_3 (100%, d_4^{20} 1.5129)** is obtained on distilling a mixture of equal volumes of '**fuming nitric acid**' and concentrated H_2SO_4 [Liang *Org Synth Coll Vol* **3** 803 1955, DOI: 10.15227/orgsyn.021.0105]. It is an oxidising agent, its vapours irritate and cauterise tissues, and cause the skin to turn yellow in colour. The '**fuming acid**' is particularly dangerous, should be handled in an efficient fume cupboard, and is used more as an oxidant than for acidification — use EYE PROTECTION.

Nitric oxide [10102-43-9] NO , **M 30.0**, **m -163.6°**, **b -151.8°/atm**, **d²⁵ 1.3402g/ml**, **n_D²⁵ 1.0002697**. Bubble the gas through 10M NaOH which removes NO_2 . It can also be freed from NO_2 by passage through a column of *Ascarite* followed by a column of silica gel held at -197°K. The gas is dried with solid NaOH pellets or by passing through silica gel cooled at -78°, followed by fractional distillation from a liquid N_2 trap. This purification does not eliminate nitrous oxide. Other gas scrubbers sometimes used include one containing concentrated H_2SO_4 and another containing mercury. It is freed from traces of NO_2 by the freeze and thaw method. The solubility of the gas in H_2O is 0.0098g/100ml (0°) and 0.0056g/ml (20°). [Blanchard et al. *Inorg Synth* **2** 126 1946, DOI: 10.1002/9780470132333.ch37; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 485-487 1963.] **TOXIC**.

Nitrogen [7727-37-9] N_2 , **M 28.0**, **m -210.00°**, **b -195.8°/atm**, **-195.795°/atm**, **d 1.3402g/L(STP: 0°/101.325kPa)**, **d 0.808g/L(at b -195.8°)**. Cylinder N_2 can be freed from oxygen by passage through *Fieser's solution* [which comprises 2g sodium anthraquinone-2-sulfonate and 15g sodium hydrosulfite dissolved in 100ml of 20% KOH; see Fieser, *J Am Chem Soc* **46** 2639 1924, DOI: 10.1021/ja01677a005] followed by scrubbing with saturated lead acetate solution (to remove any H_2S generated by the Fieser solution), concentrated H_2SO_4 (to remove moisture), then soda-lime (to remove any H_2SO_4 and CO_2). *Alternatively*, after passage through *Fieser's solution*, N_2 can be dried by washing with a solution of the metal ketyl from benzophenone and Na wire in absolute diethyl ether. [If ether vapour in N_2 is undesirable, the ketyl from liquid Na-K alloy under xylene can be used.]

Another method for removing O_2 is to pass the nitrogen through a long, tightly packed column of Cu turnings, the surface of which is constantly renewed by scrubbing it with aqueous ammonia (sg 0.880) solution. The gas is then passed through a column packed with glass beads moistened with concentrated H_2SO_4 (to remove ammonia), through a column of packed KOH pellets (to remove H_2SO_4 and to dry the N_2), and finally through a glass trap packed with chemically clean glass wool immersed in liquid N_2 . Nitrogen has also been purified by passage over Cu wool at 723°K and Cu(II) oxide [prepared by heating $\text{Cu}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ at 903°K for 24 hours] and then into a cold trap at 77°K.

A typical dry purification method consists of a mercury bubbler (as trap), followed by a small column of silver and gold turnings to remove any mercury vapour, towers containing anhydrous CaSO_4 , dry molecular sieves or $\text{Mg}(\text{ClO}_4)_2$, a tube filled with fine Cu turnings and heated to 400° by an electric furnace, a tower containing soda-lime, and finally a plug of glass wool as filter. Variations include tubes of silica gel, traps containing activated charcoal cooled in a Dry-ice bath, copper on Kieselguhr heated to 250°, and Cu and Fe filings at 400°. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 458-460 1963.]

Nitrosyl chloride [2696-92-6] NOCl , **M 65.5**, **m -59.4°**, **b -5.55°/atm**, **d²⁵ 2.872mg/L**. It is an orange gas with a suffocating odour. It has been fractionally distilled at atmospheric pressure in an all-glass, low-temperature still, taking the fraction boiling at -4° and storing it in sealed tubes. *Alternatively*, the gas is dried by CaCl_2 and passed through H_2SO_4 when Cl_2 passes on, but NOCl is absorbed to form *nitrososulfuric acid* ($\text{NO} \cdot \text{HSO}_4$) which on warming with NaCl evolves pure NOCl [Tilden *J Chem Soc* **27** 630 1874, DOI: 10.1039/JS8742700630.] It is decomposed by H_2O and alkali, and forms compounds with metal chlorides e.g. $\text{FeCl}_3 \cdot \text{NOCl}$. [Coleman et al. *Inorg Synth* **1** 55 1939, DOI: 10.1002/9780470132326.ch20; Morton, Wilcox et al. *Inorg Synth* **4** 48 1953, DOI: 10.1002/9780470132357.ch16; Beckham et al. 'Nitrosyl Chloride' *Chem Rev* **48** 319 1951, DOI: 10.1021/cr60151a001]

Nitrous oxide (laughing gas) [10024-97-2] N_2O , **M 44.0**, **m -90.86°**, **b -88.5°/atm**, **d²⁵ 1.977g/L(gas)**, **n 1.000516(STP: 0°/101.325kPa)**. Wash the gas with concentrated alkaline pyrogallol solution, to remove O_2 , CO_2 , and NO_2 , then dry it by passing it through columns of P_2O_5 or Drierite, and collecting in a dry trap cooled in liquid N_2 . It is further purified by freeze-pump-thaw and distillation cycles under vacuum [Ryan & Freeman *J Phys Chem* **81** 1455 1977, DOI: 10.1021/j100530a005; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 484-485 1963].

Osmium(VIII) tetroxide (osmic acid) [20816-12-0] $\text{OsO}_4 \cdot 2\text{H}_2\text{O}$, H_4OsO_6 , **M 524.2**, **m 40.6°**, **b**

59.4°/60mm, 71.5°/100mm, 109.3°/400mm, 130°/760mm, d_D^{20} 5.10, pK_1^{25} 7.2, pK_2^{25} 12.2, pK_3^{25} 13.95, pK_4^{25} 14.17. It is **VERY TOXIC** and should be manipulated in a very efficient fume cupboard. *It attacks the eyes severely (use also eye and face protection)* and is a good oxidising agent. It is volatile and has a high vapour pressure (11mm) at room temperature. It sublimes and volatilises well below its boiling point. It is soluble in $*C_6H_6$, H_2O (7.24% at 25°), CCl_4 (375% at 25°), $EtOH$ and Et_2O . It is estimated by dissolving a sample in a glass-stoppered flask containing 25ml of a solution of KI (previously saturated with CO_2) and acidified with 0.35M HCl . After gentle shaking in the dark for 30 minutes, the solution is diluted to 200ml with distilled H_2O saturated with CO_2 and titrated with standard thiosulfate using starch as indicator. This method is not as good as the gravimetric method. Hydrazine hydrochloride (0.1 to 0.3g) is dissolved in 3M HCl (10ml) in a glass-stoppered bottle. After warming to 55-65°, a weighed sample of OsO_4 solution is introduced, and the mixture is digested on a water bath for 1 hour. The mixture is transferred to a weighed glazed crucible and evaporated to dryness on a hot plate. A stream of H_2 is started through the crucible, and the crucible is heated over a burner for 20-30 minutes. The stream of H_2 is continued until the crucible is cooled to room temperature, and then the H_2 is displaced by CO_2 in order to avoid rapid combustion of H_2 . Finally the crucible is weighed. [Grube in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II pp 1603 1965, Anderson & Yost *J Am Chem Soc* **60** 1822 1938, DOI: 10.1021/ja01275a031; Fieser **1** 759, **10** 290, **12** 258, **13** 222, **14** 235, **15** 240, **16** 249.] § Available commercially on a variety of polymer support.

Oxygen [7782-44-7] O_2 , **M 32.00, m -218.79°, b -182.96°/atm, d^{183} 1.149g/L(liquid), d 1.429(STP: 0°/101.325kPa).** Purify it by passing the gas over finely divided platinum at 673°K and $Cu(II)$ oxide (see under nitrogen) at 973°, then condensed in a liquid N_2 -cooled trap. **HIGHLY EXPLOSIVE with organic matter.**

Palladium (II) chloride [7647-10-1] $PdCl_2$, **M 177.3, m 678-680°, d^{25} 4.0g/cm³.** The *anhydrous salt* is insoluble in H_2O and dissolves in HCl with difficulty. The *dihydrate* forms red *hygroscopic* crystals that are readily reduced to Pd . Dissolve it in concentrated HCl through which dry Cl_2 is bubbled. Filter this solution which contains H_2PdCl_4 and H_2PdCl_6 and on evaporation it yields a residue of pure $PdCl_2$. [Grube in *Handbook of Preparative Inorganic Chemistry* (Ed Brauer) Academic Press Vol II p 1582 1965, Mozingo *Org Synth Coll Vol 3* 685 1955, DOI: 10.15227/orgsyn.026.0077.] Alternatively, (fume cupboard), Pd metal is dissolved in *aqua regia*, and converted to the chloride by repeated evaporation with concentrated HCl . The dark red palladous chloride is dried at 150-180° in a stream of dry chlorine gas to give the *anhydrous* salt. [Kharasch et al. *J Am Chem Soc* **60** 882 1938, DOI: 10.1021/ja01271a035; for use as catalyst see Fieser **1** 782, **2** 303, **4** 369, **5** 500, **6** 447, **7** 277, **8** 384, **9** 352, **10** 300, **11** 393, **12** 371, **13** 234, **15** 248, **16** 268.]

Palladium (II) cyanide [2035-66-7] $Pd(CN)_2$, **M 158.5.** The yellow solid should be washed well with H_2O and dry in air. Its solubility in H_2O is low with a $\log K_{sp}$ of -42. [Bigelow et al. *Inorg Synth* **2** 245 1946, DOI: 10.1002/9780470132333.ch78; Hibble et al. 'Structures of $Pd(CN)_2$ and $Pt(CN)_2$: Intrinsically Nanocrystalline Materials'. *Inorg. Chem.* **50** 104 2011, DOI: 10.1021/ic101358q]. It is an effective catalyst for the stereospecific ring opening of oxiranes in almost quantitative yields [Imi et al. *J Org Chem* **52** 1013 1987, DOI: 10.1021/jo00382a008]. **POISONOUS.**

Perchloric acid [7601-90-3] $HClO_4$, **M 100.5, d^{25} 1.665g/cm³, pK^{25} -2.4 to -3.1 ($HClO_4$).** The 72% acid has been purified by double distillation from silver oxide under vacuum: this frees the acid from metal contamination. Distillation at atmospheric pressure is *dangerous and explosive*. The *anhydrous* acid is obtained by adding gradually 400-500ml of oleum (20% fuming H_2SO_4) to 100-120ml of 72% $HClO_4$ in a reaction flask cooled in an ice-bath. The pressure is reduced to 1mm (or less), with the reaction mixture at 20-25°. The temperature is gradually raised during 2 hours to 85°; the distillate is collected in a receiver cooled in Dry-ice. For further details of the distillation apparatus see Smith [*J Am Chem Soc* **75** 184 1953, DOI: 10.1021/ja01097a048]. **It is HIGHLY EXPLOSIVE; a strong protective screen should be used at all times.** [Schmeisser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 318-320 1963, Fieser **1** 796, **2** 309, **3** 220, **5** 506, **6** 453, **7** 279, **11** 402.]

Perrhenic acid [hydrated rhenium(VII) oxide] [13768-11-1] $HReO_4$ (gas), $H_4Re_2O_9$ (solid), **M 251.2, m sublimes, d^{25} 2.16g/cm³, pK^{25} -1.25.** The acid, which has the formula $[Re_2O_7(OH_2)_2]$, is commercially

available as a 65-70% w/w solution in H₂O, from which it can be obtained by evaporation. It can be prepared freshly from a solution of pure finely powdered **potassium perrhenate** {[10466-65-6] KReO₄ M 289.3, **m** 550° **d**²⁵ 4.887g/cm³} (10.0g, 35.6mmol) in hot H₂O (120ml) at 100° which is placed on the top of a Dowex 50W-X1 cation exchange resin column (40 x 20mm, 50g of 50-100 mesh with a 1cm layer of Pyrex wool). This resin is first washed with 6 M HCl (20ml) and flushed with pure hot H₂O at 100° until the effluent is colourless and does not become turbid on addition of AgNO₃ solution (i.e. chloride free). The hot solution of KReO₄ is applied to the resin in four 30ml portions (to keep the temperature high, boiling if necessary for the salt to remain in solution), and the hot column is rinsed with pure hot H₂O. The rate of flow should be kept at 15-20ml/minute, the first effluent (ca 50ml) is discarded and the acidic fraction (to litmus, ca 150ml) is collected. This solution is concentrated to as small a volume as possible, preferably *in vacuo* over P₂O₅, and an aliquot is diluted and titrated against standard alkali to determine the concentration of HReO₄ in the residue. It is a strong acid (cf. pK_a). It is then diluted with pure H₂O as required. The theoretical amount of acid in the residue should be 8.68g. [Watt et al. *Inorg Synth* **7** 189 1963, DOI: 10.1002/9780470132388.ch51.] It is also prepared by dissolving the **anhydride** rhenium heptoxide {**rhenium(VII) oxide** [1314-68-7] Re₂O₇ M484.4, **m** 220°, **b** 360°/atm, **d**²⁵ 6.103g/cm³} in H₂O. [Smith & Long *J Am Chem Soc* **70** 354 1948, DOI: 10.1021/ja01181a110; Melaven et al. *Inorg Synth* **3** 188 1950, DOI: 10.1002/9780470132340.ch50.] When aqueous solutions of Re₂O₇ [i.e. Re₂O₇ (OH₂)₂ = H₄Re₂O₉] are kept for several months it breaks up, and crystals of **HReO₄·H₂O** separate which contain tetrahedral ReO₄⁻. Generally perrhenic acid and rhenium(VII) oxide have been used interchangeably, from which organorhenium oxides are prepared that catalyse olefin oxidation, metathesis and other transformations [Herrmann et al. *Inorg Chem* **34** 4701 1955, DOI: 10.1021/ic00123a001].

Phosgene [75-44-5] COCl₂, **M** 98.9, **m** -118°, **b** 8.2°/756mm. Dry the gas with Linde 4A molecular sieves, degas it and distil it under vacuum at low temperature. This should be done in a closed system such as a vacuum line. It is hydrolysed slowly by H₂O, but does not fume in moist air. It is available in cylinders and as a ~20% solution in toluene (**d**²⁰ 0.94g/ml); as well as in cartridges (of **triphosgene** [32315-10-9]) for generating phosgene and respective starter kits. **It is HIGHLY TOXIC and should not be inhaled. If it is inhaled, the operator should lie very still, and be made to breathe in ammonia vapour that reacts with phosgene to form urea.** [Atkinson et al. *J Chem Soc* **117** 1410 1920, DOI: 10.1039/CT9201701410; *Beilstein* **3** IV 41.]

Phosphine [7803-51-2] PH₃, **M** 34.0, **m** -133°, **b** -87.7°, **critical temp** 51.3°, **pK**²⁵ -14, **pK_b** 28. PH₃ is best purified in a gas line (in a vacuum) in an efficient fume cupboard. It is spontaneously flammable, has a strong odour of decayed fish and is **POISONOUS**. The gas is distilled through solid KOH towers (two), through a Dry ice-acetone trap (-78°, to remove H₂O, and P₂H₄ which spontaneously ignites with O₂ to form oxides of phosphorus), then through two liquid N₂ traps (-196°), followed by distillation into a -126° trap (Dry-ice/methylcyclohexane slush), allowed to warm in the gas line and then sealed in ampoules preferably under N₂. Its IR as **v_{max}** at 327 (m), 1121 (m) and 900 (m) cm⁻¹. [Klement in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 525-530 1963, Gokhale et al. *Inorg Synth* **9** 56 1967, DOI: 10.1002/9780470132401.ch17.] PH₃ has also been absorbed into a solution of cuprous chloride in hydrochloric acid (when CuCl.PH₃ is formed). PH₃ gas is released when this solution is heated, and the gas is purified by passage through KOH pellets and then over P₂O₅. It ignites spontaneously in air with a luminous flame. Its solubility is 0.26ml/1 ml of H₂O at 20°, forming a crystalline **hydrate** on releasing the pressure on the aqueous solution. [For further detail see 'Miscellaneous As, B, P, Si, S, Se and Te Compounds' in Chapter 3.]

Phosphonitrilic chloride (cyclic tetramer) (octachloro-cyclotetraphosphazene) [1832-07-1] (NPCl₂)₄, **N₄P₄Cl₈**, **M** 463.9. Purify it by zone melting, then recrystallise it from petroleum ether (b 40-60°) or *n*-hexane. [Shaw et al. *Chem Rev* **62** 247 1962, DOI: 10.1021/cr60217a004; Crystal structure, ³¹P, ¹HNMR and reactivity of Cl: van der Huizen et al. *JCS Dalton Trans* 1317 1986, DOI: 10.1039/DT9860001317.]

Phosphonitrilic chloride (cyclic trimer) (hexachloro-cyclotriphosphazene) [940-71-6] (NPCl₂)₃, **N₃P₃Cl₆**, **M** 347.6, **m** 112.8°, **113-114°**, **113-115°**, **128.8°**, **b** 127°/12mm, **d**²⁵ 1.98g/cm³. Purify it by zone melting, by crystallisation from petroleum ether, *n*-hexane or *benzene, and by sublimation. [Nielsen et al. *Inorg Synth* **6** 94 1960, DOI: 10.1002/9780470132371.ch28; Shaw et al. *Chem Rev* **62** 247 1962, DOI: 10.1021/cr60217a004; Paddock *Quart Rev* **18** 168 1964, DOI: 10.1039/QR9641800168; crystal structure, ³¹P, ¹HNMR and reactivity

of Cl see van der Huizen et al. *JCS Dalton Trans* 1317 1986, DOI: 10.1039/DT9860001317; Meirovitch et al. *J Phys Chem* **88** 1522 1984, DOI: 10.1021/j150652a016; Alcock et al. *J Am Chem Soc* **106** 5561 1984, DOI: 10.1021/ja00331a028; ^{31}P , ^1H NMR and reactivity of Cl: Winter & van de Grampel *JCS Dalton Trans* 1269 1986, DOI: 10.1039/DT9860001269; Fieser **2** 206, **4** 386, **5** 322, **6** 469.] Useful for the synthesis of ‘dandelion’ (spherical) dendrimers [Labarre et al. *Synlett* 799 1996, DOI: 10.1055/s-1996-5584].

Phosphonous acid (phosphonic acid) [13598-36-2] H-P=O(OH)_2 , **M 82.0**, **m 73.6°**, **d** 25 **1.651**, **pK₁ 1.43**, **pK₂ 6.67**. Phosphonous acid is best prepared by adding PCl_3 to concentrated HCl , when HCl gas is liberated and the solution is evaporated (fume-cupboard) until the temperature reaches 180° (all HCl gas is driven off). On cooling the acid separates as white hygroscopic (and deliquescent) crystals. It has a garlic taste, melts at 73.6° , and decomposes at 200° to give phosphine and phosphoric acid (*note* that PH_3 ignites in air with bright flashes, see above). It is a dibasic acid that slowly oxidises to phosphoric acid in air. Store it under dry N_2 , or as a 20% aqueous solution under N_2 . [Simon et al. *Z Anorg Allgem Chem* **230** 289 1937, DOI: 10.1002/zaac.19372300312; Voight et al. *Inorg Synth* **4** 55 1953, DOI: 10.1002/9780470132357.ch18.]

Tautomerism of phosphonous acid. This acid is tautomeric with trihydroxyphosphine with the latter tautomer being in extremely small concentrations, e.g. 1 in 10^{12} . The diester [e.g. $(\text{MeO})_2\text{P(H)=O}$] has the H linked to the P atom. The tautomerism is not like that of keto-enol tautomerism in carbon chemistry, and dialkylphosphites $(\text{RO})_2\text{P(H)=O}$ do *not* react with sulfur to form the corresponding P-SH derivatives, but they do form solid salts such as $(\text{RO})_2\text{PONa}$ which *do* add sulfur readily. Also this acid does form triesters such as $(\text{MeO})_3\text{P}$ which is trimethylphosphite or trimethoxyphosphine, and behave more like phosphines. [cf. F.A. Cotton, G. Wilkinson, C.A. Murillo and M. Bochmann, *Advanced Inorganic Chemistry*, 6th Edn, Interscience Publ, 1999, ISBN 0-471-19957-5.]

Phosphoric acid (orthophosphoric acid) [7664-38-2] H_3PO_4 , **M 98.0**, **m 41 to 44°**, **42.3°**, **b 158°/atm**, **d** 25 **1.685–1.830g/cm³**, **pK₁²⁵ 2.15**, **pK₂²⁵ 7.21**, **pK₃²⁵ 12.37**. Pyrophosphate can be removed from phosphoric acid by diluting with distilled H_2O and refluxing overnight. By cooling to 11° and seeding with crystals obtained by cooling a few millilitres in a Dry-ice/acetone bath, 85% orthophosphoric acid crystallises as $\text{H}_3\text{PO}_4 \cdot \text{H}_2\text{O}$. The crystals are collected on a sintered glass filter. Concentrated acid is ~85% $\text{H}_3\text{PO}_4 \cdot x\text{H}_2\text{O}$. [Weber et al. *Inorg Synth* **I** 101 1939, DOI: 10.1002/9780470132326.ch35.]

When orthophosphoric acid (85%) is heated and liquefies (*ca* 350° and above) it is converted into **meta-phosphoric acid** [37267-86-0] $(\text{HPO}_3)_n$ which cools to a **hygroscopic** transparent, glassy solid, or a soft silky mass. It is slowly soluble in H_2O (faster on boiling) to regenerate orthophosphoric acid. It is polymeric, consisting of cyclic trimer, tetramer or linear polymers (with 3 to 7 or more P atoms) depending on the heat treatment. It is soluble in EtOH , can be made into rods and is stabilised with NaPO_3 or sodium phosphate. It has been used as a phosphorylating agent as in the preparation of coenzyme phosphates and polyphosphates. [Viscontini et al. *Helv Chim Acta* **32** 1482 1949, DOI: 10.1002/hlca.19490320515; Viscontini et al. *Helv Chim Acta* **34** 1834, 2198 1951, codecarboxylase DOI: 10.1002/hlca.19510340617, pyridoxal-5'-phosphate DOI: 10.1002/hlca.19510340712.] The sodium phosphate stabilised solid has been used in dentistry for making zinc oxyphosphate cement.

Phosphorus (red) [7723-14-0] **P**, **M 31.0**, **m 590°/43atm**, **ignites at 200°**, **d** 25 **2.34g/ml**. Heat it for 15 minutes in boiling distilled H_2O , allow it to settle and wash it several times with boiling H_2O . Transfer it to a Büchner funnel, wash it with hot H_2O until the washings are neutral, then dry it at 100° and store it in a desiccator.

Phosphorus (white) [7723-14-0] **P**, **M 31.0**, **m 44.1°**, **b 287°**, **d** 25 **1.82**. Purify white phosphorus by melting it under dilute H_2SO_4 —dichromate (possible **carcinogen**) mixture and allow to stand for several days in the dark at room temperature. It remains liquid, and the initial milky appearance due to insoluble, oxidisable material gradually disappears. The phosphorus can then be distilled under vacuum in the dark [Holmes *Trans Faraday Soc* **58** 1916 1962, DOI: 10.1039/TF9625801916]. It sublimes *in vacuo*. Other methods of purification include extraction with dry CS_2 followed by evaporation of the solvent, or washing with 6M HNO_3 , then H_2O , and drying under vacuum. It ignites in air at $\sim 50^\circ$, or by friction if dry. Store and cut it under H_2O . **POISONOUS, use gloves.**

Phosphorus oxychloride (phosphoryl chloride) [10025-87-3] POCl_3 , M 153.3, m 1.25°, b 105.8°/atm, d₄²⁰ 1.675, n_D²⁰ 1.461. Distil the liquid under reduced pressure to separate it from the bulk of the HCl and the phosphoric acid (from hydrolysis); the middle fraction is re-distilled into ampoules containing a little purified mercury. These ampoules are sealed and stored in the dark for 4-6 weeks with occasional shaking to facilitate reaction of any free chloride with the mercury. The POCl_3 is then again fractionally distilled and stored in sealed ampoules in the dark until required [Herber *J Am Chem Soc* **82** 792 1960, DOI: 10.1021/ja01489a007]. Lewis and Sowerby [*J Chem Soc* 336 1957, DOI: 10.1039/JR9570000336] refluxed their distilled POCl_3 with Na wire for 4 hours, then removed the Na and again distilled. *Use Na only with almost pure POCl_3 to avoid explosions.* [Fieser **1** 876, **2** 330, **3** 228, **4** 390, **5** 535, **7** 292, **9** 374, **11** 429, **13** 249, **15** 267, **17** 288.] **HARMFUL VAPOURS; work in an efficient fume cupboard.**

Phosphorus pentabromide [7789-69-7] PBr_5 , M 430.6, m <100°, b 106°/atm(dec). Dissolve it in pure nitrobenzene at 60°, filtering off any insoluble residue on to sintered glass funnel, then allow it to crystallise by cooling. Wash the collected solid with dry Et_2O and remove excess ether in a current of dry N_2 . (All manipulations should be performed in a dry-box.) [Harris & Payne *J Chem Soc* 3715 3732 1958, DOI: 10.1039/JR9580003715.]. It *fumes* in moist air because of hydrolysis. **HARMFUL VAPOURS** (wash burning eyes with aqueous NaHCO_3).

Phosphorus pentachloride [10026-13-8] PCl_5 , M 208.2, m 179-180°(sublimes). [All operations should be carried out in an efficient fume cupboard.] Sublime it at 160-170° in an atmosphere of chlorine. Excess chlorine is then displaced by dry N_2 gas. All subsequent manipulations should be performed in a dry-box [Downs & Johnson *J Am Chem Soc* **77** 2098 1955, DOI: 10.1021/ja01613a019]. It fumes in moist air and attacks the eyes and the mucous membranes of the nose. It should not be breathed in and has very **HARMFUL VAPOURS** (wash burning eyes with aqueous NaHCO_3). A commercial polymer-bound version is available. [Fieser **1** 866, **4** 388, **5** 534, **7** 290.]

Phosphorus pentasulfide [1314-80-3] P_2S_5 , M 444.5, m 277-283°, 290°, b 513-515°/atm, d²⁵ 2.09g/cm³. Purify P_2S_5 by extraction and crystallisation with CS_2 , using a Soxhlet extractor, and is heated in a CO_2 atmosphere at 150° to remove solvent. It liberates H_2S in moist air. **HARMFUL VAPOURS.** [Klements in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 568 1963, Fieser **1** 870, **3** 226, **4** 389, **5** 534, **6** 470, **7** 290, **8** 401, **9** 374, **10** 320, **11** 428.]

Phosphorus pentoxide [1314-56-3] P_2O_5 , M 141.9, m 340°, 562°, b 605°/atm, d²⁵ 2.3g/ml. It has been sublimed at 250° under vacuum into glass ampoules. It fumes in moist air and reacts violently with water. It is an excellent drying agent for use in desiccators. **HARMFUL VAPOURS and attacks skin, use gloves.** [Manley *J Chem Soc* **121** 331 1922, DOI: 10.1039/CT9222100331; Klements in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 541 1963, Fieser **1** 871, **7** 291, **10** 319.]

Phosphorus sesquisulfide [1314-85-8] P_4S_3 , M 220.1, m 172°, b 408°/atm, d²⁵ 2.08g/ml. Extract P_4S_3 with CS_2 , filter it and evaporate it to dryness. *Alternatively*, place it in H_2O , and pass steam through it for an hour. The H_2O is then removed, the solid is dried, and recrystallised from CS_2 (with the usual CS_2 precautions) [Rogers & Gross *J Am Chem Soc* **74** 5294 1952, DOI: 10.1021/ja01141a018].

Phosphorus sulfochloride (phosphorus thiochloride) [3982-91-0] PSCl_3 , M 169.4, m -35°, b 122-124°/atm, 125°(corr)/atm, d₄³⁰ 1.64, n_D³⁰ 1.556. Possible impurities are PCl_5 , H_3PO_4 , HCl and AlCl_3 . Gently mix it with H_2O to avoid a heavy emulsion; the product decolourises immediately and settles to the bottom layer. It is soluble in C_6H_6 and CCl_4 . [Moeller et al. *Inorg Synth* **4** 71 1953, DOI: 10.1002/9780470132357.ch24.] **HARMFUL VAPOURS.**

Phosphorus tribromide [7789-60-8] PBr_3 , M 270.7, m -41.5°, b 168-170°/725mm, 171-173°/atm, 172.9°(corr)/760mm, d₄³⁰ 2.852. It is decomposed by moisture, it should be kept dry and is *corrosive*. Purify it by distillation through an efficient fractionating column [see Whitmore & Lux *J Am Chem Soc* **54** 3451 1932, DOI: 10.1021/ja01347a071] in a slow stream of dry N_2 , i.e. under strictly dry conditions. [Gay et al. *Inorg Synth* **2** 147 1946, DOI: 10.1002/9780470132333.ch43; Noller & Dismore *Org Synth Coll Vol* **2** 358 1943, DOI:

10.15227/orgsyn.013.0020.] Dissolve it in CCl_4 , dry it over CaCl_2 , filter and distil it. Store it in sealed ampoules under N_2 and keep it away from light. It is also commercially available as a 1.0M solution in CH_2Cl_2 (d²⁵ 1.488). [Fieser 1 873, 2 330, 7 292, 8 400.] **HARMFUL VAPOURS.**

Phosphorus trichloride [7719-12-2] PCl_3 , M 137.3, m -112° , b $76^\circ/\text{atm}$, d₄²⁰ 1.575, n_D²⁰ 1.574. Heat it under reflux to expel dissolved HCl, then distil it. It has been further purified by vacuum fractionation several times through a -45° trap into a receiver at -78° . [Forbes et al. *Inorg Synth* 2 145 1946, DOI: 10.1002/9780470132333.ch42.] Also commercially available as 2.0M solution in CH_2Cl_2 . **HARMFUL VAPOURS.**

Phosphorus triiodide [13455-01-1] PI_3 , M 411.7, m 61° , d²⁵ 4.18g/cm³. It decomposes in moist air and must be kept in a desiccator over CaCl_2 . It is crystallised from sulfur-free CS_2 ; otherwise the m decreases to ca 55° . It is best to prepare it freshly. [Germann & Traxler *J Am Chem Soc* 49 307 1927, DOI: 10.1021/ja01401a001; Klement in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 541 1963.] **HARMFUL VAPOURS.**

12-Phosphotungstic acid hydrate [12501-23-4] $\text{H}_3[\text{P}(\text{W}_3\text{O}_{10})_4 \cdot x\text{H}_2\text{O}]$, M 2880.2(anhydr), m 89° (hydrate), 95° , ~ 96° . A few drops of concentrated HNO_3 are added to 100g of phosphotungstic acid dissolved in 75ml of water, in a separating funnel, and the solution is extracted with diethyl ether. The lowest of the three layers, which contains a phosphotungstic acid-ether complex, is separated, washed several times with 2M HCl, then with water and again extracted with ether. Evaporation of the ether under vacuum with mild heating on a water bath gives crystals which are dried under vacuum and ground [Matijevic & Kerker, *J Am Chem Soc* 81 1307 1959, DOI: 10.1021/ja01515a012]. [For gravimetric determination of potassium as phospho-12-tungstate see Rieben & Van Slyke *J Biol Chem* 156 765 1944, <http://www.jbc.org/content/156/2/765.citation>.]

Platinum (IV) ammonium chloride [ammonium hexachloroplatinate (IV)] [16919-58-7] $(\text{NH}_4)_2\text{PtCl}_6$, M 443.9, m 380° , d²⁵ 3.065g/ml. This salt is a useful Pt compound and is a very good source for preparing many amine complexes. It provides *spongy Pt* at high temperature. It is prepared by dissolving the less expensive spongy Pt (9.76g, 50mmol) in aa (200ml, 1HNO₃:4HCl by volume, and allowed to stand for 30 minutes until orange-red in colour) by heating and concentrating in an evaporating dish (*use an efficient fume hood and care as it is a dangerous operation*) to a thick syrup which is then taken almost to dryness on a water bath. Avoid drying at higher temperatures as it becomes incompletely soluble in H_2O . The orange-red crusty residue is dissolved in H_2O (100ml), filtered; and to it is add slowly a solution of NH_4Cl (8g, 150mmol) in H_2O (100ml) with stirring, when a yellow finely crystalline precipitate separates. Set the mixture aside for 30 minutes in an ice bath. Filter off the platinate salt and wash it several times with 1% aqueous NH_4Cl (30ml portions), then with EtOH (which removes any corresponding Pd salt impurity) and Et₂O. Dry the salt first in air, powder the crystals and dry further at 120° for 1 hour to provide an almost quantitative yield. **Note** that although the filtrates may be yellow in colour they only contain minute traces of Pt as the colour can be imparted at as low concentrations as 1 part in 20,000 parts of H_2O . The salt is not hygroscopic. Its solubility (by weight) in H_2O is 0.5% at 20° and 3.37% at 100° , and it is still less soluble in 1M NH_4Cl (0.0028%). It decomposes above 185° and yields *spongy Pt* at higher temperatures. It is reduced to the metal by H_2 at 120° and yields H_2PtCl_6 with Cl_2 . [Kauffman *Inorg Synth* 9 182 1967, DOI: 10.1002/9780470132401.ch51]

Platinum (II) chloride [10025-65-7] PtCl_2 , M 266.0, m 581° (dec), d₄²⁰ 5.87. It is purified by heating at 450° in a stream of Cl_2 for 2 hours. Some sublimation occurs because the PtCl_2 sublimes completely at 560° as red (almost black) needles. This sublimate can be combined to the bulk chloride, and while still at ca 450° it should be transferred to a container and cooled in a desiccator. A probable impurity is PtCl_4 . To test for this add a few drops of H_2O (in which PtCl_4 is soluble) to the salt, filter and add an equal volume of saturated aqueous NH_4Cl to the filtrate. If no precipitate is formed within 1 minute, then the product is pure. If a precipitate appears, then the whole material should be washed with small volumes of H_2O until the soluble PtCl_4 is removed. The purified PtCl_2 is partly dried by suction and then dried in a vacuum desiccator over P_2O_5 . It is insoluble in H_2O , but soluble in HCl to form *chloroplatinic acid* (H_2PtCl_4) by disproportionation. [Cohen & Ferrone *Inorg Synth* 6 209 1960, DOI: 10.1002/9780470132371.ch67.]

Potassium bicarbonate [298-14-6] KHCO_3 , **M 100.1**. It is crystallised from water at 65-70° (1.25ml/g) by filtering and then cooling to 15° (~0.4ml/g). During all operations, CO_2 is passed through the stirred mixture. The crystals are sucked dry at the pump, washed with distilled water, dried in air and then over H_2SO_4 in an atmosphere of CO_2 . It is much less soluble than the carbonate in H_2O (see below).

Potassium biiodate [13455-24-8] KIO_4 , HIO_4 , $\text{KH}(\text{IO}_4)_2$, **M 389.9**. Crystallise the biiodate three times from hot water (3ml/g), and stirring continuously during each cooling. After drying at 100° for several hours, the crystals are suitable for use in volumetric analysis. *Alternatively*, dry it for 2 hours in an oven at 105° and allow to cool to ~25° in a desiccator before use (keep drying temperature below 120° or decomposition occurs).

Potassium bisulfate [7646-93-7] KHSO_4 , **M 136.2**, **m 214°, 218°, d²⁵ 2.32g/cm³**. Crystallise it from H_2O (1ml/g) between 100° and 0°. It is also formed when a warm solution of K_2SO_4 in conc H_2SO_4 is cooled down. It can be freshly fused at 400°, cool and ground before use.

Potassium borohydride [13762-51-1] KBH_4 , **M 53.9**, **m ~500°(dec) d²⁵ 1.18g/cm³**. Crystallise it from liquid ammonia. It is slowly hydrolysed by H_2O . Its solubility at ~20° in H_2O or liquid NH_3 is 20%, in MeOH it is 0.7%, in Me_2NCHO it is 15% and in $\text{MeOH}/\text{H}_2\text{O}$ (1:4) it is 13%. [James & Wallbridge *Progr Inorg Chem* **11** 99-231 1970, DOI: 10.1002/9780470166123.ch3.]

Potassium bromate [7758-01-2] KBrO_3 , **M 167.0**, **m 350°(dec at 370°), d²⁰ 3.27**. Crystallise KBrO_3 from distilled H_2O (2ml/g) between 100° and 0°. To remove bromide contamination, a 5% solution in distilled H_2O , cooled to 10°, is bubbled with gaseous chlorine for 2 hours, then filtered and extracted with reagent grade CCl_4 until colourless and odourless. After evaporating the aqueous phase to about half its volume, it is cooled again slowly to about 10°. The crystalline KBrO_3 that separates, is washed with 95% EtOH and dried in a vacuum [Boyd et al. *J Am Chem Soc* **74** 237 1952, DOI: 10.1021/ja01121a062]. Another way to remove Br^- ions is by stirring several times in MeOH and then drying at 150° [Field & Boyd *J Phys Chem* **89** 3707 1985, DOI: 10.1021/j100263a026]. KBrO_3 is a powerful brominating agent for deactivated aromatic compounds [Kosandal et al. *Org Prep Proc Int* **23** 395, 569 1991, DOI: 10.1080/00304949109458221; amendment: DOI: 10.1080/00304949109458242].

Potassium bromide [7758-02-3] KBr , **M 119.0**, **m 734°, d²⁰ 2.75**. Crystallise the bromide from distilled water (1ml/g) between 100° and 0°. Wash it with 95% EtOH , followed by Et_2O . Dry it in air, then heat it at 115° for 1 hour, pulverise it, then heat it in a vacuum oven at 130° for 4 hours. It has also been crystallised from aqueous 30% EtOH , or EtOH , and dried over P_2O_5 under vacuum before heating in an oven.

Potassium carbonate [584-08-7] K_2CO_3 , **M 138.2**, **m 891°, 898°, d²⁵ 2.43g/cm³**. It crystallises from water between 100° and 0°. The solubility in H_2O is 105% at 0°, 127% at 60° and 205% at 135° (the boiling point of a saturated solution). After two recrystallisations of technical grade material, it had B, Li and Fe at 1.0, 0.04 and 0.01 ppm, respectively. [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 987 1963.]

Potassium chlorate [3811-04-9] KClO_3 , **M 122.6**, **m 356°, 368°**. It has been recrystallised from water (1.8ml/g) between 100° and 0°, and the crystals were filtered onto sintered glass. Keep away from organic material as it oxidises them readily.

Potassium chloride [7447-40-7] KCl , **M 74.6**, **m 770°, 771°, d²⁵ 1.98g/cm³**. Dissolve it in conductivity water, filter it, and saturate it with chlorine (generated from concentrated HCl and KMnO_4). Excess chlorine is boiled off, and the KCl is precipitated by HCl (generated by dropping concentrated HCl into concentrated H_2SO_4). The precipitate is washed with water, dissolved in conductivity water at 90-95°, and crystallised by cooling to about -5°. The crystals are drained at the centrifuge, dried in a vacuum desiccator at room temperature, then fused in a platinum dish under N_2 , cooled and stored in a desiccator. Potassium chloride has also been sublimed in a stream of pre-purified N_2 gas and collected by electrostatic discharge [Craig & McIntosh *Can J Chem* **30** 448 1952, DOI: 10.1139/v52-053].

Potassium chromate [7789-00-6] K_2CrO_4 , M 194.2, m 971°, 975°, d_4^{20} 2.72, pK_1^{25} 0.74, pK_2^{25} 6.49 (for H_2CrO_4). Crystallise it from conductivity water (0.6g/ml at 20°), and dry it between 135° and 170°.

Potassium cobalticyanide [potassium hexacyanocobaltate(III)] [13963-58-1] $\text{K}_3\text{Co}(\text{CN})_6$, M 332.4, m dec on heating, d^{25} 1.878g/cm³. Crystallise it from water to remove traces of HCN. Its solubility in 87-88% EtOH is 1 in 7500 at 20°. [Glemser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II p 1542 1965.]

Potassium cyanate [590-28-3] KOCN , M 81.1, d^{25} 2.05g/cm³, pK^{25} 3.46 (for HCNO). Common impurities include ammonia and bicarbonate ion (from hydrolysis). Purify it by preparing a saturated aqueous solution at 50°, neutralising with acetic acid, filtering, adding two volumes of EtOH and keeping for 3-4 hours in an ice bath. (More EtOH can lead to co-precipitation of KHCO_3 .) Filter, wash it with EtOH and dry it rapidly in a vacuum desiccator (P_2O_5). The process is repeated [Vanderzee & Meyers *J Phys Chem* 65 153 1961, DOI: 10.1021/j100819a044].

Potassium cyanide [151-50-8] KCN , M 65.1, m 634°, d_4^{20} 1.52, pK^{25} 9.216 (for HCN). A saturated solution in H_2O -ethanol (1:3) at 60° is filtered and cooled to room temperature. Absolute EtOH is added, with stirring, until crystallisation ceases. The solution is again allowed to cool to room temperature (during 2-3 hours), then the crystals are filtered off, washed with absolute EtOH, and dried, first at 70-80° for 2-3 hours, then at 105° for 2 hours [Brown et al. *J Phys Chem* 66 2426 1962, DOI: 10.1021/j100818a026]. It has also been purified by melting in a vacuum and by zone refining. Its solubility is 33% (H_2O), 66% (boiling H_2O), 33% (glycerol), 4% (MeOH) and 1% (EtOH). It hydrolyses in H_2O , and a 0.1M aqueous solution of KCN has a pH of 11, i.e. CO_2 in the H_2O makes it acidic enough to liberate a lot of the more poisonous HCN . To keep the HCN in solution it must be made quite alkaline (pH <11). LD₅₀ oral toxicity in rat is ~10mg/Kg. [Fieser 5 553, 7 299, 8 409, 10 324, 11 433, 12 405.] **HIGHLY POISONOUS work in an efficient fume cupboard.**

Potassium dichromate [7778-50-9] $\text{K}_2\text{Cr}_2\text{O}_7$, M 294.2, m 398°(dec), d_4^{20} 2.68. Crystallise it from water (g/ml) between 100° and 0° and dry it under vacuum at 156°. [Fieser 8 410, 9 383, 10 324, 12 405.] Possible **CARCINOGEN**.

Potassium dihydrogen phosphate (moobasic) [7778-77-0] KH_2PO_4 , M 136.1, m 252.6°, d^{25} 2.338g/cm³, pK_1^{25} 2.15, pK_2^{25} 6.82, pK_3^{25} 12.38. Dissolve it in boiling distilled water (2ml/g), keep on a boiling water-bath for several hours, then filter it through paper pulp to remove any turbidity. Cool rapidly with constant stirring, and the crystals are collected on to hardened filter paper, using suction, washed twice with ice-cold water, once with 50% EtOH, and dried at 105°. Alternate recrystallisations are from water, then 50% EtOH, and again water, or from concentrated aqueous solution by addition of EtOH. It is freed from traces of Cu by extracting its aqueous solution with diphenylthiocarbazone in CCl_4 , followed by repeated extraction with CCl_4 to remove traces of diphenylthiocarbazone.

Potassium dithionate [13455-20-4] $\text{K}_2\text{S}_2\text{O}_6$, M 238.3, $\text{pK}_{\text{Est}(1)} -3.4$, pK_2^{25} 0.49 (for dithionic acid). Crystallise it from water (1.5ml/g) between 100° and 0°. [For crystal structure see Stanley *Acta Cryst* 9 897 1956, DOI: 10.1107/S0365110X56002540.]

Potassium ferricyanide (Red prussiate) [13746-66-2] $\text{K}_3\text{Fe}(\text{CN})_6$, M 329.3, pK^{25} <1 (for ferricyanide). It has been recrystallised repeatedly from hot water (1.3ml/g) and dried under vacuum in a desiccator. [For uses see Fieser 1 929, 2 345, 4 406, 5 554, 6 480, 7 300, 8 410, 9 385, 13 255.]

Potassium ferrocyanide trihydrate [14459-95-1] $\text{K}_4\text{Fe}(\text{CN})_6 \cdot 3\text{H}_2\text{O}$, M 422.4, m 70°, pK_3^{25} 2.57, pK_4^{25} 4.35 (for ferrocyanide). It is purified by repeated crystallisation from distilled water, and never heating above 60°. The *anhydrous* salt is prepared by drying at 110° over P_2O_5 in a vacuum desiccator. To obtain the *trihydrate*, it is necessary to equilibrate the salt in a desiccator over a saturated aqueous solution of sucrose and NaCl. It can also be precipitated from a saturated solution at 0° by adding an equal volume of cold 95% EtOH, setting aside for several hours, then centrifuge and wash with cold 95% EtOH. It is finally sucked air dry with water-pump vacuum. The *anhydrous* salt is obtained by drying the hydrate in a platinum boat at 90° in a slow stream of N_2 .

[Loftfield & Swift *J Am Chem Soc* **60** 3083 1938, DOI: 10.1021/ja01279a504]. [*Beilstein* **2** IV 67.]

Potassium fluorosilicate [16871-90-2] K_2SiF_6 , **M 220.3**, $d^{25} 2.665\text{g/cm}^3$, $pK^{25} 1.92$ (for H_2SiF_6). Crystallise it several times from conductivity water (100ml/g) between 100° and 0°.

Potassium hexachloroiridate (III) (K_3IrCl_6) [14024-41-0] K_3IrCl_6 , **M 522.2**. Crystallise it from hot aqueous solution, and the solution should be olive-green in colour. If it has a tinge of red, then some would have oxidised to (K_2IrCl_6) [see following entry]. In this case make a concentrated solution in H_2O , and bubble H_2S through until the solution is clearly olive-green in colour due to Ir(III). Add KCl and $\text{K}_3\text{IrCl}_6 \cdot 3\text{H}_2\text{O}$ deposits on evaporating under N_2 . Filter it off, wash it with a little H_2O , then EtOH and dry it *in vacuo* away from air. [Grube in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** pp 1595 1965.]

Potassium hexachloroiridate (IV) (K_2IrCl_6) [16920-56-2] K_2IrCl_6 , **M 483.1**, $m >385^\circ(\text{dec})$, $d^{25} 3.42\text{g/cm}^3$. Crystallise it from hot aqueous solution containing a few drops of HNO_3 to keep it in the oxidised state. It forms small shiny red-black octahedral crystals which are dried at 100° and give a red powder on grinding [Grube in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** pp 1593-1594 1965].

Potassium hexachloroosmate (IV) [16871-60-6] K_2OsCl_6 , **M 481.1**, $m 600^\circ(\text{dec})$, $d^{25} 3.546\text{g/cm}^3$. Crystallise it from hot dilute aqueous HCl. [Turner et al. *Anal Chem* **30** 1708 1958, DOI: 10.1021/ac60142a602.]

Potassium hexachloroplatinate (IV) [16921-30-5] K_2PtCl_6 , **M 486.0**, $m 250^\circ(\text{dec})$, $d^{25} 3.50\text{g/cm}^3$. It crystallises in orange-yellow crystals from water (20ml/g) between 100° and 0°. Its solubility in H_2O is 0.7% at 0°, 1.12% at 20°, 2.16% at 50° and 5.13% at 100°. [Grube in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1571 1965.]

Potassium hexacyanochromate (III) ($\text{K}_3[\text{Cr}(\text{CN})_6] \cdot 3\text{H}_2\text{O}$) [13601-11-1] $\text{K}_3\text{Cr}(\text{CN})_6$, **M 328.4**, $m >350^\circ$. It forms yellow crystals from water. Recrystallise it two or three times from H_2O and dry it over H_2SO_4 . Its solubility at 20° is 30.96g/100g H_2O , and it is insoluble in EtOH. Aqueous solutions tend to decompose especially in the presence of light or on heating when $\text{Cr}(\text{OH})_3$ separates. [Hein & Herzog in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** pp 1373-1374 1965.]

Potassium hexafluorophosphate [17084-13-8] KPF_6 , **M 184.1**, $m 575^\circ$, $d^{25} 2.75\text{g/cm}^3$. Crystallise it from alkaline aqueous solution, using polyethylene vessels, or from 95% EtOH, and dry it in a vacuum desiccator over KOH. [Kolditz *Z Anorg Allgem Chem* **284** 144 1956, DOI: 10.1002/zaac.19562840116; Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 196 1963.]

Potassium hexafluorozirconate [16923-95-8] K_2ZrF_6 , **M 283.4**, $d_4^{20} 3.48$. Recrystallise it from hot water (solubility is 0.78% at 2° and 25% at 100°).

Potassium hydrogen fluoride [7789-29-9] KHF_2 , **M 78.1**, $m 225^\circ(\text{dec})$, $239^\circ(\text{dec})$, $d^{25} 2.37\text{g/cm}^3$. It crystallises from water. It is very soluble in hot H_2O and 41% at 21°. [Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 237 1963.] A 3.0M aqueous solution ($d^{25} 2.37\text{g/cm}^3$) is also commercially available, and useful for preparing potassium organotrifluoroborates from their respective boronic acids [Molander & Ham *Org Lett* **8** 2031 2006, DOI: 10.1021/ol060375a].

Potassium hydroxide (solution) [1310-58-3] KOH , **M 56.1**, $pK^{25} 16$ (for aquu K^+). Its carbonate content can be reduced by rinsing KOH sticks rapidly with water prior to dissolving them in boiled out distilled water. Alternatively, a slight excess of saturated BaCl_2 or $\text{Ba}(\text{OH})_2$ can be added to the solution which, after shaking well, is set aside so that the BaCO_3 is allowed to separate out. Davies and Nancollas [*Nature* **165** 237 1950, DOI: 10.1038/165237b0] rendered KOH solutions carbonate free by ion exchange using a column of Amberlite IR-100 in the OH^- form. [For applications see Fieser **1** 935, **2** 346, **3** 238, **4** 409, **5** 557, **6** 486, **7** 303, **8** 414, **9** 387, **11** 439, **12** 411, **13** 258.]

Potassium iodate [7758-05-6] KIO_3 , M 214.0, m 560°, d²⁵ 3.93g/cm³, pK²⁵ 0.80 (for HIO_3). It has been crystallised twice from distilled water (3ml/g) between 100° and 0°, dried for 2 hours at 140° and cooled in a desiccator. Analytical reagent grade material dried in this way is suitable for use as an analytical standard.

Potassium iodide [7681-11-0] KI , M 166.0, m 681°, b 1330°/atm, d²⁵ 3.12g/cm³, pK²⁵ -8.56 (for HI). Crystallise it from distilled water (0.5ml/g) by filtering the near-boiling solution and cooling. To minimise oxidation to iodine, the process can be carried out under N_2 and the salt is dried under a vacuum over P_2O_5 at 70-100°. Before drying, the crystals can be washed with EtOH or with acetone followed by petroleum ether. It has also been recrystallised from water/ethanol. After 2 recrystallisations, ACS/USP grade had Li and Sb at <0.02 and <0.01 ppm respectively. [Lingane & Kolthoff *Inorg Synth* **I** 163 1939, DOI: 10.1002/9780470132326.ch55.]

Potassium nickel sulfate hexahydrate [13842-46-1] $\text{K}_2\text{SO}_4 \cdot \text{NiSO}_4 \cdot 7\text{H}_2\text{O}$, M 437.1, m <100°(dec), d₄²⁰ 2.124. Crystallise the blue-green solid from H_2O (1.7ml/g) between 75° and 0°. [For crystal growth see He et al. *J Cryst Growth* **233** 809 2001, DOI: 10.1016/S0022-0248(01)01564-0.]

Potassium nitrate (saltpetre) [7757-79-1] KNO_3 , M 101.1, m 334°, 336°, decomposes at 400°/atm, d¹⁵ 2.109 g/cm³, n_D²⁵ 1.335, 1.5056, 1.5604, pK_b 15.3. It crystallises from hot H_2O (0.5ml/g) on cooling (cf. KNO_2 below). Dry it for 12 hours under vacuum at 70°. The solubility in H_2O is 13.3% at 0°, 110% at 60°, and 246% at 100°. After two recrystallisations, technical grade salt had <0.001 ppm of metals. The fused salt is a powerful oxidising agent. It decomposes between 550° and 790° to KNO_2 and O_2 . Useful for making fireworks, explosives, blasting powders, in candle wicks as it burns evenly, as a flux, tempering steel and as a diuretic. [Spencer *Saltpeter: The Mother of Gunpowder* OUP, UK 2013 pp 256, ISBN 9780199695751.]

Potassium nitrite [7758-09-0] KNO_2 , M 85.1, m 350°(dec at ~441°), d²⁵ 1.915g/ml, pK²⁰ 3.20 (for HNO_2). A saturated solution at 0° is warmed and partially evaporated under vacuum. The crystals so obtained are filtered off from the warm solution. (This procedure is designed to reduce the level of nitrate impurity and is based on the effects of temperature on solubility. The solubility of KNO_3 in water is 13g/100ml at 0°, 247g/100ml at 100°; for KNO_2 the corresponding figures are 280g/100ml and 413g/100ml.) Alternatively, dissolve it in H_2O and precipitate by adding of EtOH. An aqueous solution is alkaline, and it reacts with dilute mineral acid to liberate nitrous acid (blue green colour nitrosating agent) which oxidises to give brown fumes of NO_2 . [Young 'Potassium Nitrite' *J Chem Educ* **85** 779 2008, DOI: 10.1021/ed085p779.]

Potassium nitrosodisulfonate (Fremy's Salt) [14293-70-0] $(\text{K}_2\text{SO}_3)_2\text{NO}$, M 268.3. It forms yellow needles (dimeric) which dissolve in H_2O to give the violet monomeric free radical. It is purified by dissolving (~12g) in 2M KOH (600ml) at 45°, filtering the blue solution and keeping it in a refrigerator overnight. The golden yellow crystals (10g) are filtered off, washed with MeOH (3x), then Et_2O and stored in a glass container in a vacuum over KOH. It is stable indefinitely when dry. [Cram & Reeves *J Am Chem Soc* **80** 3094 1958, DOI: 10.1021/ja01545a048; Schenk *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 505 1963; preparation and use: Wehrli & Pigott *Org Synth Coll Vol* **6** 1010, 1988, DOI: 10.15227/orgsyn.052.0083.] A useful **radical oxidant**, e.g. for the oxidation of anilines and phenols to quinones [Teuber & Benz *Chem Ber* **100** 218 1967, DOI: 10.1002/cber.19671000916; Teuber *Org Synth Coll Vol* **6** 480 1988, DOI: 10.15227/orgsyn.052.0088].

Potassium osmate (VI) dihydrate [19718-36-6, 10022-66-9 ($2\text{H}_2\text{O}$)] $\text{K}_2\text{OsO}_4 \cdot 2\text{H}_2\text{O}$, M 368.4, m 332.4°($2\text{H}_2\text{O}$), 368.4°(anhyr). It forms **hygroscopic POISONOUS** purple crystals which are soluble in H_2O but insoluble in EtOH and Et_2O . It decomposes slowly in H_2O to form the **tetroxide** which **attacks the eyes**. The solid should be kept dry and in this form it is relatively safe. [Lloyd et al. *Synthesis* 610 1972, DOI: 10.1055/s-1972-21948; Grube in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1604 1965.] A reagent for the cleavage of olefins to ketones and carboxylic acids [Whitehead et al. *Tetrahedron Lett* **47** 3797 2006, DOI: 10.1016/j.tetlet.2006.03.087], and the intramolecular amidohydroxylation of carbamoyloxy-tethered olefins to give hydroxyoxazolidinones [Donohoe et al. *J Am Chem Soc* **128** 2514 2006, DOI: 10.1021/ja057389g]

Potassium perchlorate [7778-74-7] KClO_4 , **M 138.6**, **m 400(dec)**, d_4^{20} **2.52**, pK^{25} **-2.4 to -3.1 (for HClO_4)**. It crystallises from boiling water (5ml/g) on cooling. Dry it under vacuum at 105°. **Potentially explosive.**

Potassium periodate (potassium metaperiodate) [7790-21-8] KIO_4 , **M 230.0**, **m 582°**, d_4^{20} **3.62**. Crystallise it from distilled water. Its solubility in H_2O is 0.2% at 0°, 0.4% at 20°, 4.4% at 80°, and 7.9% at 100°. [Willard et al. *Inorg Synth* **I** 168 1939, DOI: 10.1002/9780470132326.ch58; Schmeisser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 325 1963.] I is a powerful oxidant [Fieser **1** 809, **2** 311, **4** 373, **5** 507, **6** 455.]

Potassium permanganate [7722-64-7] KMnO_4 , **M 158.0**, **m 240°(dec)**, d_4^{20} **2.7**, pK^{25} **-2.25 (for HMnO_4)**. Crystallise the purple-bronze solid from hot water (19ml/g at 15°, 4ml/g at 65°), then dry it in a vacuum desiccator over CaSO_4 . Phillips and Taylor [*J Chem Soc* 4242 1962, DOI: 10.1039/JR9620004242] cooled an aqueous solution of KMnO_4 , saturated at 60°, to room temperature in the dark, and filtered it through a No.4 porosity sintered-glass filter funnel. The solution was allowed to evaporate in air in the dark for 12 hours, and the supernatant liquid was decanted from the crystals, which were dried as quickly as possible with filter paper. It is a secondary analytical standard. It is a useful oxidant and is used as a topical antiseptic. [Fieser **1** 942, **2** 348, **4** 412, **5** 562, **8** 416, **9** 388, **10** 330, **11** 440, **12** 413, **13** 258, **14** 267, **15** 273.] Also commercially available on phosphate buffered silica gel (200-400 mesh).

Potassium peroxydisulfate (potassium persulfate) [7727-21-1] $\text{K}_2\text{S}_2\text{O}_8$, **M 270.3**, **m <100°(dec)**, d^{25} **3.477g/cm³**, n_D^{25} **1.467**. Crystallise the persulfate twice from distilled water (10ml/g) and dry it at 50° in a vacuum desiccator. Its solubility in H_2O is 1.6% at 0°, 4.5% at 20°, and 7.2% at 30°. An aqueous solution decomposes on long standing with evolution of O_2 and formation of KHSO_4 . It is a powerful oxidising agent. Store it at ~10°. [Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 390 1963.]

Potassium peroxymonosulfate (Oxone, potassium monopersulfate triple salt; $2\text{KHSO}_5 \cdot \text{KHSO}_4 \cdot \text{K}_2\text{SO}_4$), [37222-66-5, 70693-62-8 (triple salt)] $2\text{KHSO}_5 \cdot \text{KHSO}_4 \cdot \text{K}_2\text{SO}_4$, **M 614.8**. This is a stable form of Caro's acid and should contain >4.7% of active oxygen. It can be used in $\text{EtOH}/\text{H}_2\text{O}$ and $\text{EtOH}/\text{AcOH}/\text{H}_2\text{O}$ solutions. If active oxygen is too low, it is best to prepare it afresh from 1mole of KHSO_5 , 0.5mole of KHSO_4 and 0.5mole of K_2SO_4 . [Kennedy & Stock *J Org Chem* **25** 1901 1960, DOI: 10.1021/jo01081a019; Stephenson US Patent 2,802,722 1957.] A rapid preparation of *Caro's acid* is made by stirring finely powdered potassium persulfate (**M 270.3**) into ice-cold concentrated H_2SO_4 (7ml) and when homogeneous add ice (40-50g). It is stable for several days if kept cold. Keep away from organic matter as it is a **STRONG OXIDANT**. A detailed preparation of *Caro's acid* (*hypersulfuric acid*, H_2SO_5 , [7722-86-3]) in crystalline form **m ~45°** from H_2O_2 and chlorosulfonic acid was described by Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 388 1963. [For applications see Fieser **1** 952, **11** 442, **12** 413, **13** 259, **14** 267.]

Potassium perrhenate [10466-65-6] KReO_4 , **M 289.3**, **m 550°**, **b 1360-1370°/atm**, d^{25} **4.887g/cm³**, n_D^{25} **1.643**, pK^{25} **-1.25 (for HReO_4)**. It forms white tetrahedral crystals from H_2O where its solubility is 12.1g/L at 20° and 140g/L at 100°. The salt is then fused in a platinum crucible in air at 750°. [Smith & Long *J Am Chem Soc* **70** 354 1948, DOI: 10.1021/ja01181a110; Watt et al. *Inorg Synth* **7** 189 1963, DOI: 10.1002/9780470132388.ch51.]

Potassium reineckate [34430-73-4] $\text{K}[\text{Cr}(\text{NH}_3)_2(\text{SCN})_4] \cdot \text{H}_2\text{O}$, **M 357.5**. Crystallise it from KNO_3 solution, then from warm water [Adamson *J Am Chem Soc* **80** 3183 1958, DOI: 10.1021/ja01546a004]. [Use as a chemical actinometer (photochemical reference): *Analyst* **114** 739 1989, DOI: 10.1039/AN9891400739; for charge transfer photochemistry see Mainusch et al. *Inorg Chim Acta* **255** 87 1997, DOI: 10.1016/S0020-1693(96)05343-1]

Potassium ruthenate(VI) [31111-21-4] KRuO_4 , **M 243.3**. Dissolve the ruthenate in H_2O and evaporate until crystals are formed. The crystals are iridescent green prisms which appear red in thin films. A possible impurity is RuO_4 ; in this case wash with CCl_4 (which dissolves RuO_4). The concentration of an aqueous solution of RuO_4^{2-} (orange colour) can be estimated from the absorbance at 385nm (ϵ 1030 $\text{M}^{-1} \text{cm}^{-1}$), or at 460nm (ϵ 1820

$\text{M}^{-1} \text{cm}^{-1}$). [Lee et al. *Can J Chem* **50** 3741 1972, DOI: 10.1139/v72-592; Connick & Hurley *J Am Chem Soc* **74** 5012 1952, DOI: 10.1021/ja01140a007; Grube et al. *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1600 1965]. For the preparation of salts free of chloride ions see Kaziro et al. [*Inorg Chim Acta* **164** 85 1989, DOI: 10.1016/S0020-1693(00)80880-4].

Potassium selenocyanate [3425-46-5] KSeCN , **M 144.1**. Dissolve it in acetone, filter and precipitate it by adding Et_2O . For use in solid state NMR see Bowmaker et al. [*Inorg Chem* **37** 1734 1998, DOI: 10.1021/ic9700112]. A convenient reagent for selenocyanation [Block et al. *J Am Chem Soc* **128** 14949 2006, DOI: 10.1021/ja065037j], cyclocondensation [Köhler et al. *Tetrahedron* **46** 7735 1990, DOI: 10.1016/S0040-4020(01)90069-9] and heterocyclisation reactions [*Chem Commun* 860 1991, DOI: 10.1039/C39910000860].

Potassium sulfate [7778-80-5] K_2SO_4 , **M 174.3**, **m 1069°**, **d₄²⁰ 2.67**. It crystallised from distilled water (4ml/g at 20°; 8ml/g at 100°) between 100° and 0°.

Potassium tetrachloroplatinate(II) [10025-99-7] K_2PtCl_4 , **M 415.1**, **m 500°(dec)**, **d²⁵ 3.38g/cm³**. It forms crystals from aqueous 0.75M HCl (20ml/g) between 100° and 0°. Wash them with ice-cold water and dry. The solubility in HO is 0.93g/100ml (16°) and 5.3g/100ml (100°). [Keller & Moeller *Inorg Synth* **7** 247 1963, DOI:10.1002/9780470132333.]

Potassium tetracyanopalladate (II) trihydrate [10025-98-6] $\text{K}_2\text{Pd}(\text{CN})_4 \cdot 3\text{H}_2\text{O}$, **M 377.4**. All operations should be carried out in an efficient fume cupboard. **Cyanide is very POISONOUS**. Dissolve the complex (ca 5g) in a solution of KCN (4g) in H_2O (75ml) with warming and stirring, and evaporate hot till crystals appear. Cool, filter off the crystals and wash them with a few drops of cold H_2O . Further concentration of the mother liquors provides more crystals. The complex is recrystallised from H_2O as the colourless **trihydrate**. It **effloresces** in dry air and dehydrates at 100° to the **monohydrate**. The **anhydrous salt** is obtained by heating at 200°, but at higher temperatures it decomposes to $(\text{CN})_2$, Pd and KCN. [Bigelow et al. *Inorg Synth* **2** 245 1946, DOI: 10.1002/9780470132333.ch78.]

Potassium tetrafluoroborate (potassium borofluoride) [14075-53-7] KBF_4 , **M 125.9**, **m 530°**, **d₄³⁰ 2.505**, **pK²⁵ -4.9 (for HBF_4)**. Crystallise it from H_2O (solubility% (temperature): 0.3 (3°), 0.45 (20°), 1.4 (40°), 6.27 (100°), and dry it under vacuum. It is a non-hygroscopic salt. A 10% solution is transparent blue at 100°, green at 90° and yellow at 60°. [Vörlander et al. *Chem Ber* **65** 535 1932, DOI: 10.1002/cber.19320650406; Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 223 1963.]

Potassium thiocyanate (potassium rhodanide) [333-20-0] KSCN , **M 97.2**, **m 172°, 173°**, **pK²⁵ -1.85 (for HSCN)**. Crystallise it from H_2O if much chloride ion is present in the salt, otherwise from EtOH or MeOH (optionally by addition of Et_2O). Filter off on a Büchner funnel without paper, and dry it in a desiccator at room temperature before heating for 1 hour at 150°, with a final 10-20 minutes at 200° to remove the last traces of solvent [Kolthoff & Lingane *J Am Chem Soc* **57** 2126 1935, DOI: 10.1021/ja01314a025]. Store it in the dark. [Fieser **1** 954.]

Potassium thiosulfate hydrate [13446-67-8, 10294-66-3 (75% aqueous solution)] $\text{K}_2\text{S}_2\text{O}_3 \cdot x\text{H}_2\text{O}$, **M 190.3**, **pK₁²⁵ 0.6**, **pK₂²⁵ 1.74 (for $\text{H}_2\text{S}_2\text{O}_3$)**. Crystallise it from warm water (0.5ml/g) by cooling in an ice-salt mixture. It is a good reducing agent used in analytical chemistry. [Foerster & Mommsen *Chem Ber* **57** 258 1924, DOI: 10.1002/cber.19240570218.]

Potassium tungstate (ortho dihydrate) [37349-36-3; 7790-60-5] $\text{K}_2\text{WO}_4 \cdot 2\text{H}_2\text{O}$, **M 362.1(anhydr)**, **m 921°**, **d²⁵ 3.12g/cm³**, **pK₁²⁵ 2.20**, **pK₂²⁵ 3.70 (for H_2WO_4)**. Crystallise it from hot water (0.7ml/g).

Praseodymium trichloride hexahydrate [10361-79-2, 19423-77-9 ($6\text{H}_2\text{O}$)] $\text{PrCl}_3 \cdot 6\text{H}_2\text{O}$, **M 247.3(anhydr)**, **m 786°**, **d²⁵ 2.25g/cm³(hydrate)**, **pK₁²⁵ 8.55 (for Pr^{3+} hydrol)**. Its 1M solution in 6M HCl is passed twice through a Dowex-1 anion-exchange column. The eluate is evaporated in a vacuum desiccator to half its volume and allowed to crystallise [Katzin & Gulyas *J Phys Chem* **66** 494 1962, DOI: 10.1021/j100809a029].

Praseodymium (III, IV) oxide [12037-29-5] Pr_6O_{11} , **M 1021.4**. Dissolve the oxide in acid (perchloric acid), precipitate it as the oxalate and the salt is ignited at 650° to give the oxide. The single valence **praseodinium(III) oxide** Pr_2O_3 , **M 329.8**, has **m** 2183°, **b** 2760° and **d**²⁵ 6.9g/ml.

Reineckate salt See ammonium reineckate above.

Rhodium (III) chloride [10049-07-7, 20765-98-4 ($x\text{H}_2\text{O}$)] RhCl_3 , **M 209.3**, **m** >100°(dec), **b** 717°. Probable impurities are KCl and HCl. Wash the chloride well with small volumes of H_2O to remove excess KCl and KOH and dissolve it in the minimum volume of concentrated HCl. Evaporate it to dryness on a steam bath to give wine-red coloured $\text{RhCl}_3 \cdot 3\text{H}_2\text{O}$. Leave it on the steam bath until the odour of HCl is lost—do **not** try to dry further as it begins to decompose above 100° to the oxide and HCl. It is not soluble in H_2O but soluble in alkalis or CN solutions and forms double salts with alkali chlorides. [Anderson et al. *Inorg Synth* **7** 214 1963, DOI: 10.1002/9780470132388.ch57.] Catalyst for converting CH_4 to AcOH [Lin & Sen *Nature* **368** 613 1994, DOI: 10.1038/368613a0], and for conjugate reduction of cinnamaldehydes followed by cross-coupling with arylboronic acids [Wang et al. *Chem Commun* 1192 2004, DOI: 10.1039/B401511H]. [Fieser **6** 504, **7** 313, **16** 292.]

Rubidium bromide [7789-39-1] RbBr_3 , **M 165.4**, **m** 682°, **b** 1340°/atm, **d**²⁵ 3.35g/cm³. The bromide is a white crystalline powder which crystallises from H_2O (solubility: 50% in cold and 67% in boiling H_2O to give a neutral solution). It also crystallises from near-boiling water (0.5ml/g) by cooling to 0°.

Rubidium chlorate [13446-71-4] RbClO_3 , **M 168.9**, **d**²⁰ 3.19. It crystallises from water (1.6ml/g) by cooling from 100°.

Rubidium chloride [7791-11-9] RbCl_3 , **M 120.9**, **m** 715°, **m** 1383°/atm, **d**²⁵ 2.80g/cm³. Crystallise it from water (0.7ml/g) by cooling to 0° from 100°. Its solubility in H_2O is 77.3% at 0.6°, 90.3% at 70° and 147% at boiling point. [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 951-955 1963.]

Rubidium nitrate [13126-12-0] RbNO_3 , **M 147.5**, **m** 305°, **d**²⁵ 2.80g/cm³. Crystallise the nitrate from hot water (0.25ml/g) by cooling to room temperature.

Rubidium perchlorate [13510-42-4] RbClO_4 , **M 184.9**, **m** 281°, **b** 600°/atm(dec), **d**²⁵ 2.88g/cm³, **pK**²⁵ -2.4 to -3.1 (for HClO_4). Crystallise the perchlorate from hot water by cooling to 0° [solubility in g/100ml: 1.09 (0°), 1.30 (25°), 3.44 (50°), 6.72 (70°) and 17.39 (99°)]. Oxidising agent.

Rubidium sulfate [7488-54-2] Rb_2SO_4 , **M 267.0**, **m** 530°, **d**²⁵ 3.61g/ml. Crystallise the sulfate from water (1.2ml/g) between 100° and 0°.

Ruthenium (III) chloride dihydrate (β-form) [10049-08-8 (anhydrous), 13815-94-6 ($3\text{H}_2\text{O}$), 14898-67-0 ($x\text{H}_2\text{O}$)] $\text{RuCl}_3 \cdot x\text{H}_2\text{O}$, **M 207.4 + H_2O** , **m** >500°(dec), **d**²⁵ 3.11g/ml, **pK**₁²⁵ 3.40 (for aquo Rh^{3+} hydrolysis). Dissolve the black salt in H_2O , filter and concentrate to crystallisation in the absence of air to avoid oxidation. Evaporate the solution in a stream of HCl gas while being heated just below its boiling point until a syrup is formed and finally to dryness at 80-100° and dried in a vacuum over H_2SO_4 . When heated at 700° in the presence of Cl_2 the insoluble **α-form** is obtained [Grube in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1598 1965,]. A very useful catalyst particularly when coordinated with organic ligands, e.g. for oxidation reactions with an oxidant [Carlsen et al. *J Org Chem* **46** 3936 1981, DOI: 10.1021/jo00332a045; Hasegawa et al. *Chem Lett (Jpn)* **14** 1385 1985, DOI: 10.1246/cl.1985.1385], the oxidative cyclisation of 1,7-ienes to oxepane diols [Piccialli et al. *Tetrahedron Lett* **48** 5131 2007, DOI: 10.1016/j.tetlet.2007.05.078] and hydroxylation of tertiary carbon-hydrogen bonds of cyclic ethers with periodate or bromate [Lee et al. *J Org Chem* **72** 5820 2007, DOI: 10.1021/jo070382s]. A kit of ruthenium metathesis catalysts kit is available commercially [see Schrodri & Pederson *Aldrichimica Acta* **40** 45 2007].

Ruthenium (IV) oxide [12036-10-1, 32740-79-7 (xH_2O)] RuO_2 , **M 133.1, b 1200°/atm(sublimes), d²⁵ 6.97g/cm³**. Free the black oxide from nitrates by boiling in distilled water and filtering. A more complete purification is based on fusion in a KOH/KNO₃ mix to form the soluble ruthenate and perruthenate salts. The melt is dissolved in water, and filtered, then acetone is added to reduce the ruthenates to the insoluble hydrated oxide which, after making a slurry with paper pulp, is filtered and ignited in air to form the *anhydrous* oxide [Campbell et al. *Anal Chem* **33** 58 1961, DOI: 10.1021/ac60169a016]. It is an electrocatalyst for producing chlorine, its oxides and oxygen [Mills 'Heterogeneous redox catalysts for oxygen and chlorine evolution' *Chem Soc Rev* **18** 285 1989, DOI: 10.1039/CS9891800285], and hydration of the oxide strongly affects its catalytic activity [Mills & Davies, *Inorg Chim Acta*, **189** 149 1991, DOI: a10.1016/S0020-1693(00)80183-8].

Samarium (II) iodide [32248-43-4] SmI_2 , **M 404.2, m 520°, b 1580, d²⁵ 6.97g/cm³**. A possible impurity is SmI_3 , [13813-25-7] **M 531.1**, from which it is made. If present, grind the solid to a powder and heat it in a stream of pure H_2 . The temperature (~ 500-600°) should be below the **m** (~ 628°) of SmI_3 , since the molten compounds react very slowly. [Wetzel in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** pp 1149, 1150 1965.] It promotes a *pseudo-Barbier reaction* between aldehydes and ketones with aliphatic or alicyclic halides (see NdI_2) [Namy et al. *J Organomet Chem* **328** 81 1987, DOI: 10.1016/S0022-328X(00)99769-9.] It is a *one-electron reducing agent* [Brendt et al. *Aldrichimica Acta* **24** 15 1991.]

SmI_2 in THF (0.1M, d²⁵ 0.99g/cm³) is available commercially for reactions can also be prepared in a dry box, or dry gloves, filled with N_2 (not generally necessary for synthetic purposes, but should be used for Grignard-type conditions, i.e. exclusion of moisture) from samarium powder (3g, 20mmol, or distilled metal or ingot), placed in a flask fitted with a dropping funnel containing 1,2-diiodoethane (22.82g, 10mmol) in THF (250ml) which is added dropwise. The reaction has no induction period if the solvent is pure. [Excess metal can be re-used later.] An intense blue-green solution results with a SmI_2 concentration of 4×10^{-2} M in THF that can be diluted to 10^{-1} M. These solutions can be stored for a long time without loss of Sm^{2+} concentration if kept in an inert atmosphere in the presence of a small amount of metal. However, it is best to use them within a few days to ensure reproducible results. In addition to promoting coupling reactions, it is also a powerful and selective reducing agent for olefins [Girard et al. *J Am Chem Soc* **102** 2693 1980, DOI: 10.1021/ja00528a029DOI: 10.1021/ja00528a029; see also Asano et al. *Synthesis* 1309 2007, DOI: 10.1055/s-2007-966006], promotes the reaction of chlorofluoroacetonitrile [359-05-7] with aldehydes to form cyanofluorohydrins in the presence of HMPA [Asano et al. *Synthesis* 1309 2007, DOI: 10.1055/s-2007-966006], intramolecular cyclisation reactions [Molander & McKie *J Org Chem* **59** 3186 1994, DOI: 10.1021/jo00090a041], and intermolecular ketone-olefin coupling reactions [Kawatsura et al. *J Org Chem* **59** 6900 1994, DOI: 10.1021/jo00102a010]. [See also Fieser **10** 344, **11** 464, **12** 429, **13** 270, **14** 276, **15** 282, **16** 294, **17** 307.]

Selenous acid [7783-00-8] H_2SeO_3 , **M 129.0, m 70°(dec), d²⁵ 3.004g/cm³, pK₁²⁵ 2.62, pK₂²⁵ 8.32 (H_2SeO_3)**. Recrystallise the acid from water. On heating it loses water and SeO_2 sublimes. [Waitkins & Clark *Chem Rev* **36** 235 1945, DOI: 10.1021/cr60115a001; for use see Fieser **1** 992.]

Selenium [7782-49-2] **Se, M 79.0, m 217.4°, 220.85°, b 684.1°/atm, d²⁵ 4.81g/cm³**. Dissolve selenium in small portions in hot concentrated HNO_3 (2ml/g), filter and evaporate to dryness to give selenious acid which is then dissolved in concentrated HCl. Pass SO_2 gas through the solution whereby selenium (but not tellurium) precipitates. It is filtered off and washed with concentrated HCl. This purification process is repeated. The selenium is then converted twice to the selenocyanate by treating with a 10% excess of 3M aqueous KCN (CARE), heated for half an hour on a sand-bath and filtered. Add an equal weight of crushed ice to the cold solution, followed by an excess of cold, concentrated HCl, with stirring (in an efficient fume cupboard as HCN is evolved) which precipitates selenium powder. This is washed with water until colourless, and then with MeOH and is heated in an oven at 105°. Finally it is fused for 2 hours *in vacuo*. It is cooled, crushed and stored in a desiccator [Tideswell & McCullough *J Am Chem Soc* **78** 3026 1956, DOI: 10.1021/ja01594a025].

Selenium dioxide [7446-08-4] SeO_2 , **M 111.0, m 315°(sublimes), 340°, d²⁵ 3.95g/cm³**. Purify it by sublimation at 315°, or by solution in HNO_3 , precipitation of selenium which, after standing for several hours or

boiling, is filtered off, then re-oxidised by HNO_3 and cautiously evaporated to dryness below 200° . The dioxide is dissolved in H_2O and again evaporated to dryness. In H_2O it forms selenious acid (see selenious acid above). Its solubility in H_2O in g/100ml is 38.4 (20°), 39.5 (25°) and 82.5 (65°), and it is soluble in EtOH (6.7g/100ml at 15°), in Me_2CO (4.4g/100ml at 15°) in MeOH (10.16/100ml at 12°) and in AcOH (1.11/100ml at 14°). It is a useful and selective oxidant. [Waitkins & Clark *Chem Rev* **36** 235 1945, DOI: 10.1021/cr60115a001; Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed Brauer) Academic Press Vol **I** p 421 1963, Rabjohn *Org Reactions* **5** 331 1949.]

Silica [7631-86-9 (colloidal), 60676-86-0 (Quartz, Cristobalite, sand), 112945-52-5 (fumed)] SiO_2 , **M 60.1, m >1600°, b 2230°**. Purification of silica for high technology applications uses isopiestic vapour distillation from concentrated volatile acids and is absorbed in high purity water. The impurities remain behind. Preliminary cleaning to remove surface contaminants uses dip etching in HF or a mixture of HCl, H_2O_2 and deionised water [Phelan & Powell *Analyst* **109** 1269 1984, DOI: 10.1039/AN9840901269].

Silica gel [63231-67-4, 112926-00-8]. Before use as a drying agent, silica gel is heated in an oven, then cooled in a desiccator. Conditions in the literature range from heating at 110° for 15 hours to 250° for 2-3 hours. Silica gel has been purified by washing with hot acid (in one case successively with *aqua regia*, concentrated HNO_3 , then concentrated HCl; in another case it was digested overnight with hot concentrated H_2SO_4), followed by exhaustive washing with distilled water (one week in a Soxhlet apparatus has also been used), and prolonged oven drying. Alternatively, silica gel has been extracted with acetone until all soluble material was removed, then dried in a current of air, washed with distilled water and oven dried. Silica gel has also been washed successively with water, M HCl, water, and acetone, then activated at 110° for 15 hours. Commercially available in various grades. [For applications see also Fieser **9** 410, **10** 346, **11** 466, **12** 431.]

Silicon monoxide [10097-28-6] SiO , **M 44.1, m > 1700°, d²⁵ 2.18g/cm³**. Purify the monoxide by sublimation in a porcelain tube in a furnace at 1250° (4 hours) in a high vacuum (10^{-4} mm) in a stream of N_2 . It is obtained as brownish black scales. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 696 1963.]

Silicon tetrachloride [10026-04-7] SiCl_4 , **M 169.9, m -70° , b $57.6^\circ/\text{atm}$, d²⁵ 1.483g/cm³**. Distil it under vacuum and store it in sealed ampoules under N_2 . It fumes in moist air and is very sensitive to moisture. It is soluble in organic solvents. It is a **strong irritant**. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 682-683 1963.] Also available as a 1.0M solution in CH_2Cl_2 (d²⁵ 1.34g/cm³). Useful for preparing silicon compounds.

12-Silicotungstic acid (tungstosilicic acid;) [12027-43-9, 11130-20-4] $\text{H}_4\text{SiW}_{12}\text{O}_{40}$, **M 2878.2**. Extract the acid with diethyl ether from a solution acidified with HCl. The diethyl ether is evaporated under vacuum, and the free acid is crystallised twice [Matijevic & Kerker *J Phys Chem* **62** 1271 1958, DOI: 10.1021/j150568a026].

Silver (metal) [7440-22-4] **Ag, M 107.9, m 961.9° , b $2212^\circ/\text{atm}$, d²⁵ 10.49g/cm³**. For purification by electrolysis, see Craig et al. [*J Res Nat Bur Stand* **64A** 381 1960]. For purification of crude, or silver residues to pure silver see Glemser & Sauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 1028-1030 1963, and for the preparation of *colloidal silver* see *ibid* (Ed. Brauer) p 1034.

Silver acetate [563-63-3] **AgOAc, M 166.9, dec on heating, d²⁵ 3.259**. The salt is a white (or slightly grey if exposed to light) crystalline powder or lustrous needles, which should not be exposed to light for long periods. It can be purified by recrystallisation from H_2O (preferably containing a little AcOH) as its solubility (w/v) in cold H_2O is 1% and in boiling H_2O it is 35%. Alternatively, it is boiled with Ac_2O , the insoluble salt is collected and recrystallised from glacial AcOH. It is freely soluble in dilute HNO_3 . Store in dark bottles away from light. [Beilstein **2** H 110, **2** I 48, **2** II 116, **2** III 189, **2** IV 112.]

It oxidises a green solution of U(IV) acetate in liquid NH_3 to the yellow U(VI) in a few hours with the formation of the tris complex $(\text{NH}_4)\text{U}(\text{VI})\text{O}_2(\text{AcO})_3$ [isolated], acetamide and Ag metal [Kline & Kershner *Inorg Chem* **5** 932 1966, DOI: 10.1021/ic50039a047].

At a 0.2-2 mole%, AgOAc catalyses efficient cycloaddition reactions of methyl isocynoacetate with olefin

Michael-acceptors at ambient temperature to produce Δ^1 - or Δ^2 -pyrrolines in good yields. Isocyanoacetates undergo AgOAc catalysed cyclodimerisation to imidazoles in excellent yields in the absence of a suitable olefin. Also, when combined with azomethine ylide, AgOAc catalyses a 1,3-dipolar cycloaddition reaction in a one-pot sequential cascade process to yield the 7-azabicyclo[2.2.1]heptane ring system which is characteristic of the naturally occurring potent non-opioid analgesic epibatidine [originally isolated from the skin of the Ecuadoran poison frog *Epipedobates tricolor*, Spande et al. *J Am Chem Soc* **114** 3475 1992, DOI: 10.1021/ja00035a048]. [Grigg et al. *Tetrahedron* **55** 2025 1999, DOI: 10.1016/S0040-4020(98)01216-2.]

When combined with I_2 , AgOAc in aqueous or anhydrous AcOH reacts with olefins to form *vicinal diols* (glycols). The hydroxylations are not always *cis* [Woodward & Brucher *J Am Chem Soc* **80** 209 1958, DOI: 10.1021/ja01534a053; Ginsberg *J Am Chem Soc* **75** 5746 1953, DOI: 10.1021/ja01118a515; Barkley et al. *J Am Chem Soc* **76** 5014 1954, DOI: 10.1021/ja01649a003; Klass et al. *J Am Chem Soc* **77** 3829 1955, DOI: 10.1021/ja01619a04; Slates & Wendler *J Am Chem Soc* **78** 3749 1956, DOI: 10.1021/ja01596a054; Jefferies & Milligan *J Chem Soc* 2363 1956, DOI: 10.1039/JR9560002363; Gunstone & Morris *J Chem Soc* 487 1957, DOI: 10.1039/JR9570000487; Ellington et al. *J Chem Soc* 1327 1966, DOI: 10.1039/J39660001327], but can produce *cis/trans* diol mixtures depending on the olefin [Bunton & Carr *J Chem Soc* 770 1963, DOI: 10.1039/JR9630000770]. It was used in a novel preparation of highly reflective and conductive silvered polymeric films [Southward et al. *Chem Mater* **11** 501 1999, DOI: 10.1021/cm981014v].

Silver bromate [7783-89-3] $AgBrO_3$, **M 235.8, m dec on heating, $d^{25} 5.28g/cm^3$** . It crystallises from hot water (80ml/g). It reacts with bromine water to form bromic acid ($HBrO_3$) which is a strong oxidising agent. Store it in the dark.

Silver bromide [7785-23-1] $AgBr$, **M 187.8, m 432°, $d^{25} 6.473g/cm^3$, $n_D^{25} 2.253$** . Purify it from Fe, Mn, Ni and Zn by zone melting in a quartz vessel under vacuum. It is insoluble in dilute HNO_3 or dilute NH_3 but is soluble in concentrated NH_3 . Its solubility product in H_2O at 20° is 3.5×10^{-13} . Store it in the dark as it is affected by light. Used in photography and as antiseptic.

Silver chlorate [7783-92-8] $AgClO_3$, **M 191.3, m 230°, b 250°/atm(dec), $d^{25} 4.43g/cm^3$** . Recrystallise the chlorate three times from water (10ml/g at 15°; 2ml/g at 80°). Also soluble in EtOH. Store it in the dark. [Nicholson et al. *Inorg Synth* **2** 4 1946, DOI: 10.1002/9780470132333.ch2; Glemser & Sauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 1037 1963.]

Silver chloride [7783-90-6] $AgCl$, **M 143.3, m 455°, b 1547°/atm, $d^{25} 5.56g/cm^3$, $n_D^{25} 2.071$** . Recrystallise it from conc NH_3 solution by acidifying with HCl, filtering off the solid, washing it with H_2O and drying it in a vacuum. Its solubility product in H_2O at 20° is 1.2×10^{-10} . It is soluble in NH_3 and should be kept in the dark as light decomposes it.

Silver chromate [7784-01-2] Ag_2CrO_4 , **M 331.8, $d^{25} 5.625g/cm^3$, $pK_1^{25} 0.74$, $pK_2^{25} 6.49$ (for H_2CrO_4), $K_{sp} 1.1 \times 10^{-12}$** . Wash the red-brown powder with H_2O , dry it in a vacuum, then powder well and dry again in a vacuum at 90°/5 hours. Its solubility in H_2O is 0.0014% at 10°, and its solubility product in H_2O at 20° is 1.7×10^{-12} . Store it in the dark. [Cardillo & Shimizu *J Org Chem* **42** 4268 1977, DOI: 10.1021/jo00862a023.]

Silver cyanide [506-64-9] $AgCN$, **M 133.9, m dec at 320°, $d^{25} 3.95g/cm^3$** . It is a **POISONOUS** white or greyish white powder. Stir it thoroughly with H_2O , filter, wash well with EtOH and dry it in air in the dark. It is very insoluble in H_2O (0.00023g in 100ml H_2O), but is soluble in HCN or aqueous KCN to form the soluble $Ag(CN)_2^-$ complex. [Schnitz-Dumont *Chem Ber* **72** 298 1939, DOI: 10.1002/cber.19390720210; Randall & Halford *J Am Chem Soc* **52** 178 1930, DOI: 10.1021/ja01364a026.]

Silver (II) difluoride [7783-95-1] AgF_2 , **M 145.9, m 690°, b 700°/atm(dec) $d^{25} 4.58g/cm^3$** . It is highly **TOXIC** because it liberates HF and F_2 . It is very **hygroscopic** and reacts **violently** with H_2O . It is a powerful oxidising agent and liberates O_3 from dilute acids, and I_2 from I^- solution. Store it in quartz or iron ampoules. It is white when pure; otherwise it is brown-tinged. It is thermally stable up to 700°. [Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 241 1963.] [For surface characterization see Wolan & Hoflund *Appl Surf Sci* **125** 251 1998, DOI: 10.1016/S0169-4332(97)00498-4].

Silver (I) fluoride [7775-41-9] AgF , M 126.9, m $>300^\circ$, 435° , b *ca* 1150° , d^{25} 5.855g/cm^3 . The fluoride is a *hygroscopic* solid with a solubility in H_2O (g/100ml) of 65.78 (0°), 179.1 (25°) and 213.4 (50°); in MeOH of 1.5 (25°) and 83 (12°), and forms an insoluble basic fluoride in moist air. Purify it by washing with AcOH and dry $^*\text{C}_6\text{H}_6$, then keep it in a vacuum desiccator at room temperature to remove $^*\text{C}_6\text{H}_6$, and store it in opaque glass bottles. The flaky *hygroscopic* crystals darken on exposure to light. It *attacks* bone and teeth. [Sharpe *J Chem Soc* 4518 4538 1952, DOI: 10.1039/JR9520004518; Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 240 1963.]

Silver iodate [7783-97-3] AgIO_3 , M 282.8, m $>200^\circ$, b 1150° , d^{25} 5.525g/cm^3 . Wash the iodate with warm dilute HNO_3 , then H_2O and dry it at 100° , or recrystallise it from NH_3 solution by adding HNO_3 , filtering, washing with H_2O and drying at 100° . It is poorly soluble in H_2O : 0.003g/100ml at 10° and 0.019g/100ml at 50° ; and soluble in ammonia.

Silver nitrate [7761-88-8] AgNO_3 , M 169.9, m 212° , b $444^\circ/\text{atm}(\text{dec})$, d^{25} 4.35g/cm^3 , n_D^{25} 1.744. Purify it by recrystallisation from hot water (solubility of AgNO_3 in water is 912g/100ml at 100° and 122g/100ml at 0°). It has also been purified by crystallisation from hot conductivity water by slow addition of freshly distilled EtOH.

CAUTION: avoid using EtOH for washing the precipitate; and avoid concentrating the filtrate to obtain further crops of AgNO_3 owing to the risk of EXPLOSION (as has been reported to us) caused by the presence of silver fulminate. When using EtOH in the purification, the apparatus should be enveloped in a strong protective shield. [Tully, *News Ed (Am Chem Soc)* 19 3092 1941; Henderson & Garin *J Chem Educ* 47 741 1970, DOI: 10.1021/ed047p741; Bretherick, *Handbook of Reactive Chemical Hazards* 4th edn, Butterworths, London, 1985, pp 13-14.] Before being used as a standard in volumetric analysis, analytical reagent grade AgNO_3 should be finely powdered, dried at 120° for 2 hours, then cooled in a desiccator. Its solubility in AcOH is 0.776g/Kg (30°), 1.244g/Kg (40°), 5.503g/Kg (93°); in EtOH is 1.3g/100g (19°); in EtOAc is 2.7g/100g (20°); in Me_2CO is 0.35g/100g (14°), 0.44g/100g (18°); and in $^*\text{C}_6\text{H}_6$ is 0.22g/Kg (35°) and 0.44g/Kg (40.5°). [For applications see Fieser 1 1008, 2 366, 3 252, 4 427, 5 528, 7 321, 9 411, 10 350, 12 433.] Recovery of silver residues as AgNO_3 [use protective shield during the whole of this procedure] can be achieved by washing with hot water and adding 16M HNO_3 to dissolve the solid. Filter this through glass wool and concentrate the filtrate on a steam bath until precipitation commences. Cool the solution in an ice-bath and filter the precipitated AgNO_3 . Dry it at 120° for 2 hours, then cool it in a desiccator in a vacuum. Store it over P_2O_5 in a vacuum in the dark. **AVOID contact with hands due to formation of black stains.** It is a standard in volumetric analysis, and is used as a precursor for preparing silver-containing nanomaterials [Zhou et al. *Ind Eng Chem Res* 45(10) 3503 2006, DOI: 10.1021/ie051098z] and complexes [Konno et al. *Chem Lett (Jpn)* 35(3) 316 2006, DOI: 10.1246/cl.2006.316]. Also available as a loading of ~ 10 wt% on silica gel (+230 mesh).

Silver nitrite [7783-99-5] AgNO_2 , M 153.9, m $141^\circ(\text{dec})$, d^{25} 4.45g/cm^3 . Crystallise the salt from hot conductivity water (0.0142g/ml) in the dark. Its solubility in H_2O (g/100ml) is 0.155 (0°), 0.275 (15°) and 1.363 (60°). Dry it in the dark under vacuum. [Glemser & Sauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 1048 1963.] Used in Meyer-type nucleophilic substitution with organic bromides and iodides to form nitro compounds [Kornblum & Ungnade *Org Synth* 38 75 1958, DOI: 10.15227/orgsyn.038.0075], and nitroalkene synthesis with nitril iodide [Sy & By *Tetrahedron Lett* 26 1193 1985, DOI: 10.1016/S0040-4039(00)98431-4] generated *in situ* from AgNO_3 and I_2 [Waldman et al. *Tetrahedron Lett* 37 7889 1996, DOI: 10.1016/0040-4039(96)01807-2].

Silver(I) oxide [20667-12-3] Ag_2O , M 231.7, m $>200^\circ(\text{dec})$, d^{25} 7.13g/cm^3 , K_{sp} 1.52×10^{-8} (20° AgOH). Leach the oxide with hot water in a Soxhlet apparatus for several hours to remove any entrained electrolytes. Its solubility in H_2O is 0.013g/L (20°), 0.025g/L (40°) and 0.053g/L. [Glemser & Sauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II p 1037 1965, for applications see Fieser 1 1011, 2 368, 3 252, 4 430, 5 583, 6 515, 7 321, 8 442, 10 350, 11 468.]

Silver (I, III) oxide [1301-96-8] AgO , M 123.9, m $>100^\circ(\text{dec})$, d^{25} 7.48g/cm^3 . It is soluble in 40,000 parts of H_2O , and should be protected from light. Stir it with an alkaline solution of potassium peroxysulfate ($\text{K}_2\text{S}_2\text{O}_8$) at 85 – 90° . The black AgO is collected, washed free from sulfate with H_2O made slightly alkaline and dried in air in the dark. [Hammer et al. *Inorg Synth* 4 12 1953, DOI: 10.1002/9780470132357.ch3.]

Silver perchlorate monohydrate [14242-05-8 (H_2O), 331717-44-3 (xH_2O), 7783-93-9 (anhydrous)] **AgClO₄, M 207.3, pK²⁵ -2.4 to -3.1 (for HClO₄)**. Reflux it with *benzene (6ml/g) in a flask fitted with a Dean and Stark trap until all the water is removed azeotropically (*ca* 4 hours). The solution is cooled and diluted with dry pentane (4ml/g of AgClO₄). The precipitated AgClO₄ is filtered off and dried in a desiccator over P₂O₅ at 1mm for 24 hours [Radell et al. *J Am Chem Soc* **83** 3958 1961, DOI: 10.1021/ja01480a007]. It has also been recrystallised from perchloric acid. [**Caution due to its EXPLOSIVE nature in the presence of organic matter.**] Store it in the dark. [For applications see Fieser **2** 369, **3** 121, **4** 432, **5** 585, **6** 69, **6** 518, **7** 142, **9** 413, **10** 354, **11** 469, **15** 121, **16** 300.]

Silver permanganate [7783-98-4] **AgMnO₄, M 226.8, m >160°(dec), d²⁵ 4.27/cm³**. The salt forms violet crystals which can be recrystallised from hot H₂O (solubility is 5.5g/L at 0° and 16.9g/L at 30°. Store it in the dark. This oxidising agent is decomposed by light. [Lux in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 1463 1963, for crystal structure see Boonstra *Acta Cryst Section B* **24** (8) 1053 1968, DOI: 10.1107/S0567740868003699]

Silver perrhenate (silver tetraoxorhenate) [20654-56-2] **Ag(ReO₄), M 358.1, m 430°, d²⁵ 7.05/cm³, pK_a -1.25**. It is prepared by adding slowly, with ice cooling, H₂O₂ (35%, 100ml) to a stirred suspension of Re powder (10g, 54mmol) in H₂O (20ml) during 4 hours, stirring at ~25° for 30 minutes, and then at 80° for 3 hours. The mixture is filtered from a small amount of insoluble material, and the clear solution is treated with Ag(NO₃) (10g, 59mmol), when Ag(ReO₄) immediately separates as a white precipitate which is filtered off, washed with Et₂O (3 x 25ml) to remove H₂O and H₂O₂, and dried *in vacuo* to give the silver salt (18.7g, 97%). [Herrmann & Kratzer *Inorg Synth* **33** 111 2002, DOI: 10.1002/0471224502.ch2; Herrmann et al. *Angew Chem Int Ed* **36** 2652 1997, DOI: 10.1002/anie.199726521.]

Silver sulfate [10294-26-5] **Ag₂SO₄, M 311.8, m 652°, b 1085°/atm(dec), d²⁵ 5.45/cm³**. Crystallise the sulfate from hot concentrated H₂SO₄ containing a trace of HNO₃, and dilute with H₂O while being strongly cooled. The precipitate is filtered off, washed with H₂O and dried at 120°. Its solubility in H₂O is 0.8% at 17°, and 1.46% at 100°. Store it in the dark. [Glemser & Sauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1042 1965.] [Fieser **1** 1015, **3** 254, **4** 435.]

Silver thiocyanate [1701-93-5] **AgSCN, M 165.9, m 265°(dec), d₄²⁰ 3.746, pK²⁵ -1.85 (for HSCN)**. Digest the solid salt with dilute aqueous NH₄NCS, filter, wash it thoroughly with H₂O and dry it at 110° in the dark. It is soluble in dilute aqueous NH₃. *Alternatively*, dissolve it in strong aqueous NH₄NCS solution, filter and dilute with large volumes of H₂O when the Ag salt separates. The solid is washed with H₂O by decantation until free from NCS⁻ ions, collected, washed with H₂O, EtOH and dried in an air oven at 120°. It has also been purified by dissolving in dilute aqueous NH₃ when single crystals are formed by free evaporation of the solution in air. Store it in the dark. [Garrick & Wilson *J Chem Soc* 835 1932, DOI: 10.1039/JR9320000835; Occleshaw *J Chem Soc* 2404 1932, DOI: 10.1039/JR9320002404; IR and Raman: Kinell & Strandberg *Acta Chem Scand* **13** 1607 1957, DOI: 10.3891/acta.chem.scand.13-1607; Lindqvist *Acta Cryst* **10** 29 1957, DOI: 10.1107/S0365110X57000067; Beilstein **3** IV 303.]

Sodium (metal) [7440-23-5] **Na, M 23.0, m 97.5°, 97.794°, 97.8°, b 882.94°/atm, d²⁵ 0.968/cm³**. The metal is placed on a coarse grade of sintered-glass filter, melted under vacuum and forced through the filter using argon. The Pyrex apparatus is then re-evacuated and sealed off below the filter, so that the sodium could be distilled at 460° through a side arm and condenser into a receiver bulb which is then sealed off [Gunn & Green *J Am Chem Soc* **80** 4782 1958, DOI: 10.1021/ja01551a008]. [Fieser **1** 1022, **4** 437, **5** 589, **7** 324, **13** 277.] **EXPLODES and IGNITES in water.**

Sodium amide [7782-92-5] **NaNH₂, M 39.0, m 210°, b 400°/atm, d²⁵ 1.39/cm³, pK_a 38 (conjugate acid)**. It reacts *violently* with H₂O and is soluble in liquid NH₃ (1% at 20°). It should be stored in wax-sealed containers in small batches. It is very *hygroscopic* and absorbs CO₂ and H₂O. If the solid is discoloured by being yellow or brown in colour, then it should be destroyed as it can be highly **EXPLOSIVE**. It should be replaced if discoloured. It is best destroyed by covering it with much toluene and slowly adding dilute EtOH with stirring until all the ammonia is liberated (FUME CUPBOARD). [Dennis et al. *Inorg Synth* **1** 74 1939, DOI: 10.1002/

9780470132326.ch25; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 465 1963, Bergstrom *Org Synth Coll Vol 3* 778 1955, DOI: 10.15227/orgsyn.020.0086; Fieser **1** 1034, **2** 373, **4** 439, **5** 591, **6** 525, **8** 446, **13** 278, **14** 284, **15** 288.]

Sodium ammonium hydrogen phosphate [13011-54-6, 7783-13-3 ($4\text{H}_2\text{O}$)] $\text{NaNH}_4\text{HPO}_4$, **M 209.1(anhydr)**, **m 79°(dec)**, **80°(dec)**, **d²⁵ 1.544/cm³**. Crystallise it from hot water (1ml/g).

Sodium arsenate heptahydrate [10048-95-0] $\text{Na}_2\text{HAsO}_4 \cdot 7\text{H}_2\text{O}$, **M 312.0(hydrate)**, **m 50° (loses 5H₂O)**, **m 130°**, **d²⁰ 1.88**, **pK₁²⁵ 2.22**, **pK₂²⁵ 6.98 (for H₃AsO₄)**. Crystallise it from water (2ml/g). [For arsenic and arsenic compounds see Grund et al. *Ullmann's Encyclopedia of Industrial Chemistry* 2005, Wiley-VCH, Weinheim, DOI: 10.1002/14356007.a03_113].

Sodium azide [26628-22-8] NaN_3 , **M 65.0**, **m 300°(dec, explosive)**, **pK²⁵ 4.72 (for HN₃)**. Crystallise sodium azide from hot water or from water by adding absolute EtOH or acetone. Also purify it by repeated crystallisation from an aqueous solution saturated at 90° by cooling it to 10°, and adding an equal volume of EtOH. The crystals are washed with acetone, and the azide is dried at room temperature under vacuum for several hours in an Abderhalden pistol. Its solubility in H_2O is 42% at 18°, and in EtOH it is 0.22% at 0°. [Das et al. *JCS Faraday Trans 1* **78** 3485 1982, DOI: 10.1039/F19827803485; Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 474-475 1963, Browne et al. *Inorg Synth* **1** 79 1939, DOI: 10.1002/9780470132326.ch27; Miller et al. *Inorg Synth* **2** 139 1946, DOI: 10.1002/9780470132333.ch40.] Used also for the efficient 1,4-addition of azide to a α,β -unsaturated aldehydes in the presence of tertiary amines [Kim & Park *Synth Commun* **37** 1027 2007, DOI: 10.1080/00397910601168306]. **HIGHLY POISONOUS and potentially explosive.**

Sodium bicarbonate [144-55-8] NaHCO_3 , **M 84.0**, **m ~50°(dec, -CO₂)**, **b 851°**, **d²⁵ 2.20/cm³**, **n_D²⁵ 1.3344**, **pK₁²⁵ 6.35**, **pK₂²⁵ 10.33 (for carbonic acid)**. Crystallise it from hot water. Its solubility in H_2O is 69g/L (0°), 165g/L (60°) and 236g/L (100°); and 0.02 %wt in Me_2CO ; 2.13 %wt in MeOH, and insoluble in EtOH. The solid should not be heated above 40° due to the formation of carbonate. [For the solubilities of sodium carbonate and sodium bicarbonate in acetone-water and methanol-water mixtures see Ellingboe & Runnels *J Chem Eng Data* **11**(3) 323 1966, DOI: 10.1021/jc60030a009.]

Sodium bisulfite [7631-90-5] NaHSO_3 , **M 104.1**, **d²⁵ 1.48/cm³**. Crystallise it from hot H_2O (1ml/g). Dry it at 100° under vacuum for 4 hours. It dehydrates on warming to give a solid which is sodium metabisulfite $\text{Na}_2\text{S}_2\text{O}_5$ ($2\text{NaHSO}_3 - 2\text{H}_2\text{O}$). An aqueous solution of *sodium metabisulfite* is in fact a solution of *sodium bisulfite*.

Sodium borate (borax) [1330-43-4] $\text{Na}_2\text{B}_4\text{O}_7$, **M 201.2**, **m 741°**, **d²⁵ 2.367/cm³**. Most of the water of hydration is removed from the *decahydrate* (see below) by evacuation at 25° for three days, followed by heating to 100° and evacuation with a high-speed diffusion pump. The dried sample is then heated gradually to fusion (above 966°), allowed to cool gradually to 200°, then transferred to a desiccator containing P_2O_5 [Grenier & Westrum *J Am Chem Soc* **78** 6226 1956, DOI: 10.1021/ja01605a004]. [Becher in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 794-795 1963, Fieser **4** 461, **15** 295.]

Sodium borate (decahydrate, hydrated borax) [1303-96-4] $\text{Na}_2\text{B}_4\text{O}_7 \cdot 10\text{H}_2\text{O}$, **M 381.2**, **m 75°(loses 5H₂O at 60°)**, **d²⁵ 1.73/cm³**. Crystallise the borate from water (3.3ml/g), keeping below 55° to avoid formation of the pentahydrate. Filter it off at the pump, wash it with water and equilibrate it for several days in a desiccator containing an aqueous solution saturated with respect to sucrose and NaCl. Borax can be prepared more quickly (but its water content is somewhat variable) by washing the recrystallised material at the pump with water, followed by 95% EtOH, then Et_2O , and dried in air at room temperature for 12-18 hours on a clock glass. [Becher in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 794-795 1963.]

Sodium borohydride [16940-66-2] NaBH_4 , **M 37.8**, **m ~400°(dec)**, **b 500°/atm(dec)**, **d²⁵ 1.0740/cm³**. After adding NaBH_4 (10g) to freshly distilled diglyme (120ml) in a dry three-necked flask fitted with a stirrer, nitrogen inlet and outlet, the mixture is stirred for 30 minutes at 50° until almost all of the solid has dissolved. Stirring is

stopped, and, after the solid has settled, the supernatant liquid is forced under N_2 pressure through a sintered-glass filter into a dry flask. [The residue is centrifuged to obtain more of the solution which is added to the bulk.] The solution is cooled slowly to 0° and then decanted from the white needles that separated. The crystals are dried by evacuating for 4 hours to give anhydrous $NaBH_4$. *Alternatively*, after the filtration at 50° the solution is heated at 80° for 2 hours to give a white precipitate of substantially anhydrous $NaBH_4$ which is collected on a sintered-glass filter under N_2 , then evacuated at 60° for 2 hours [Brown et al. *J Am Chem Soc* **77** 6209 1955, DOI: 10.1021/ja01628a044].

$NaBH_4$ has also been crystallised from isopropylamine by dissolving it in the solvent at reflux, cooling, filtering and allowing the solution to stand in a filter flask connected to a Dry-ice/acetone trap. After most of the solvent has passed over into the cold trap, crystals are removed with forceps, washed with dry diethyl ether and dried under vacuum. [Kim & Itoh *J Phys Chem* **91** 126 1987, DOI: 10.1021/j100285a029.] Somewhat less pure crystals were obtained more rapidly by using Soxhlet extraction with only a small amount of solvent and extracting for about 8 hours. The crystals that formed in the flask are filtered off, then washed and dried as before. [Stockmayer et al. *J Am Chem Soc* **77** 1980 1955, DOI: 10.1021/ja01612a082.] Other solvents in which it is soluble include H_2O (but reacts vigorously with it, but less if alkaline, stable at $\sim pH$ 14), MeOH (13g/100ml, but reacts slowly with it), EtOH (3.16g/100ml, reacts very slowly with it), diglyme (5.15g/100ml), liquid NH_3 , amines and pyridine, but is almost insoluble in Et_2O . It is destroyed by acids and liberates H_2 . [Banfi et al. Sodium Borohydride in *Encyclopedia of Reagents for Organic Synthesis* (L. Paquette ed) 2004, J. Wiley & Sons, NY, DOI:10.1002/047084289X.rs052.] Also commercially available in 2-methoxyethyl ether (0.5M), ~ 12 wt% in 14M NaOH, in tetraglyme (3.0M), in triethylene glycol dimethyl ether (2.0M) and on alumina (~ 10 wt% loading).

Sodium bromate [7789-38-0] $NaBrO_3$, M 150.9, m 381° , d^{25}_4 3.339/cm³. It is crystallised from hot water (1.1ml/g) to decrease contamination by NaBr, bromine and hypobromite. [Noszticzius et al. *J Am Chem Soc* **107** 2314 1985, DOI: 10.1021/ja00294a019; Fieser **1** 1055.]

Sodium bromide [7647-15-6] $NaBr$, M 102.9, m 747° , 755° , b $1390^\circ/atm$, d^{20}_4 3.2. Crystallise the bromide from water (0.86ml/g) between 50° and 0° , and dry it at 140° under vacuum (this purification may not eliminate chloride ion). [Fieser **12** 445.]

Sodium carbonate (soda ash) [497-19-8 (anhydrous), 5968-11-6 (H_2O), 6132-02-1 ($10H_2O$)] Na_2CO_3 , M 106.0, m 851° (anhydr, dec), 858° (anhydr, dec), 100° (H_2O , dec), 33.5° ($7H_2O$, dec), 34° ($10H_2O$, dec), d^{25}_4 2.54/cm³(anhydr), n^{25}_D 1.487 (anhydrous), n^{25}_D 1.420 (H_2O), n^{25}_D 1.405 ($10H_2O$), pK_1^{25} 6.35, pK_2^{25} 10.33 (for carbonic acid). It crystallises from water as the *decahydrate* which is redissolved in water to give a near-saturated solution. By bubbling CO_2 , $NaHCO_3$ is precipitated. It is filtered off, washed and ignited for 2 hours at 280° [MacLaren & Swinehart *J Am Chem Soc* **73** 1822 1951, DOI: 10.1021/ja01148a121]. Before being used as a volumetric standard, analytical grade material should be dried by heating at 260 – 270° for 0.5 hour and allowed to cool in a desiccator. It has a transition point at 450° , and its solubility in water is 21.58% at 20° (*decahydrate* in solid phase), 49.25% at 35° (*heptahydrate* in solid phase) and 44.88% at 75° (*monohydrate* in solid phase) [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 987-988 1963]. After three recrystallisations, technical grade Na_2CO_3 had Cr, Mg, K, P, Al, W, Sc and Ti at 32, 9.4, 6.6, 3.6, 2.4, 0.6, 0.2 and 0.2 ppm respectively; another technical source had Cr, Mg, Mo, P, Si, Sn and Ti at 2.6, 0.4, 4.2, 13.4, 32, 0.6, 0.8 ppm respectively.

Sodium chlorate [7775-09-9] $NaClO_3$, M 106.5, m 248° , 248 – 261° , b $>300^\circ$ (dec), d^{25}_4 2.50g/cm³, n^{20}_D 1.515. It is crystallised from hot water [solubility: 0.79g/ml (0°), 1.06g/ml (25°), 1.25g/ml (40°) and 2.20g/ml (100°)]. A strong oxidising agent, and should be kept clear from organic matter. [Preparation: Scott & Allen *Org Synth Coll Vol* **2** 128 1943, DOI: 10.15227/orgsyn.018.0015; Fieser **1** 1056.] It is used for tanning leather, as a herbicide and a sulfation biosynthesis inhibitor.

Sodium chloride (Rock salt) [7647-14-5] $NaCl$, M 58.4, m 800.7° , b $1413^\circ/atm$, d^{25}_4 2.165g/cm³, n^{25}_{589} 1.5442. Recrystallise from a saturated aqueous solution (2.7ml/g) by passing in HCl gas, or by adding EtOH or acetone. It can be freed from bromide and iodide impurities by adding chlorine water to an aqueous solution and boiling it for some time to expel free bromine and iodine. Traces of iron can be removed by prolonged boiling

of solid NaCl in 6M HCl; the crystals are then washed with EtOH and dried at *ca* 100°. Sodium chloride has been purified by sublimation in a stream of pre-purified N₂ and collected by electrostatic discharge [Ross & Winkler *J Am Chem Soc* **76** 2637 1954, DOI: 10.1021/ja01639a010]. For use as a primary analytical standard, analytical reagent grade NaCl should be finely ground, dried in an electric furnace at 500-600° in a platinum crucible, and allowed to cool in a desiccator. For most purposes, however, drying at 110-120° is satisfactory. Its solubility (g/kg at 25°) in H₂O is 360, in formamide is 94.0, in glycerine is 83.0, in propylene glycol is 71.0, in formic acid is 52.0, in liq NH₃ is 30.2, in MeOH is 14.0, in EtOH is 0.65, in DMF is 0.4, in *n*-PrOH is 0.124, in sulfolane is 0.05, in *n*-BuOH is 0.05, in 2-PrOH is 0.03, in 1-pentanol is 0.018, in MeCN is 0.003 and in Me₂CO is 0.00042.

Sodium chlorite [7758-19-2, 49658-21-1 (3H₂O)] NaClO₂, **M 90.4(anhydr)**, **m 180-200°(dec)**, **d²⁵ 2.468g/cm³**, **pK_a 10.11 (HClO₂)**. Crystallise the chlorite from hot water and store it in a cool place. It has also been crystallised from MeOH by counter-current extraction with liquid ammonia [Curti & Locchi *Anal Chem* **29** 534 1957, DOI: 10.1021/ac50162a033]. A major impurity is chloride ion which can be removed by recrystallisation from 0.001M NaOH. [Schmeisser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 312 1963, for applications see Fieser **9** 423, **11** 481, **13** 280.]

Sodium chromate tetrahydrate [10034-82-9 (4H₂O), 7775-11-3] Na₂CrO₄, **M 234.0(4H₂O)**, **162.0(anhydr)**, **m ~20°(for 10H₂O)**, **792°(anhydr)**, **d²⁵ 2.698g/cm³**, **pK₁²⁵ 0.74**, **pK₂²⁵ 6.49 (for H₂CrO₄)**. Crystallise the yellow chromate from hot water [solubility is 0.32g/ml (0°), 0.85g/ml (25°) and 1.27g/ml (100°)]. It is *deliquescent*. [Fieser **5** 605.] Useful anti-rust in paints.

Sodium cyanate [917-61-3] NaOCN, **M 65.0**, **m 550°**, **d²⁵ 1.893g/cm³**, **pK₂₅²⁵ 3.47 (for HCNO)**. It forms colourless needles from EtOH (0.22g/100ml at 0°). Its solubility in EtOH is 0.22g/100g at 0°. It is soluble in H₂O (11.6g/100ml at 25°), but can be recrystallised from small volumes of it. [Beilstein **3** IV 80.]

Sodium cyanide [143-33-9] NaCN, **M 49.0**, **m 563.7°**, **b 1496°/atm**, **d²⁵ 1.5955g/cm³**, **n_D²⁵ 1.452**, **pK₂₅²⁵ 9.216 (for HCN)**. NaCN can be purified in the same way as KCN ([151-50-8] above). It has also been prepared by bubbling HCN gas through an alcoholic solution of NaOH. This white solid is slightly *more soluble* in H₂O *than KCN*, with solubilities of 48.15g/100ml at 10° and 63.7g/100ml at 25°. It has some solubility in MeOH, EtOH and NH₃, is slightly soluble in DMF and SO₂ but is insoluble in DMSO. Because it is the salt of a weak acid it hydrolyses in aqueous solution generating HCN (*prussic acid* see [74-90-8]) in the equilibrium, which is liberated into the air, consequently it has the odour of HCN (bitter almonds). **Note** that some individuals (genetic variants) cannot smell this odour (the HCN molecule does not span across their olfactory sensors) so should take great care when working with compounds that liberate HCN. Earlier chemists with this olfactory deficiency smoked tobacco pipes near the equipment because the HCN vapours sucked through the pipes produced a foul taste in the mouth. Cyanides, and particularly HCN are extremely poisonous, readily absorbed in the mouth, the lethal dose is of the order of 150-300mg, depending on the individual. It is a respiratory inhibitor, blocks electron transport by acting on mitochondrial cytochrome oxidase, leading to anaerobic metabolism and lactic acidosis. This is evidenced by the blue colouration of the face and body. NaCN (and KCN) is used in gold mining as it forms water soluble *Na[Au(CN)₂]* **complex**, in fumigation for insects and vermin by releasing HCN, in electroplating baths and in case hardening of steel. [For applications see Fieser **4** 446, **5** 606, **6** 535, **7** 333, **8** 453, **13** 185.]

Sodium cyanoborohydride [25895-60-7] NaBH₃CN, **M 62.8**, **m 240-242°(dec)**, **d²⁸ 1.199 g/cm³**. It is a very *hygroscopic* solid, soluble in H₂O (212g/100ml at 29°, 121g/100ml at 88°), tetrahydrofuran (37g/100ml at 28°, 42.2g/100ml at 62°), very soluble in MeOH, slightly soluble in EtOH, but insoluble in Et₂O, *C₆H₆ and hexane. It is stable to acid up to pH 3 but is hydrolysed in 12N HCl. The rate of hydrolysis at pH 3 is 10⁻⁸ times that of NaBH₄. The fresh commercially available material is usually sufficiently pure. If very pure material is required, one of the following procedures can be used [Lane *Synthesis* 135 1975, DOI: 10.1055/s-1975-23685]: **(a)** The NaBH₃CN is dissolved in tetrahydrofuran (20% w/v), filtered and the filtrate is treated with a fourfold volume of CH₂Cl₂. The solid is collected and dried in a vacuum [Wade et al. *Inorg Chem* **9** 2146 1970, DOI: 10.1021/ic50091a036]. **(b)** NaBH₃CN is dissolved in dry MeNO₂, filtered, and the filtrate is poured into a 10-fold volume of CCl₄ with vigorous stirring; the white precipitate is collected, washed several times with CCl₄

and dried in a vacuum (yield 75%) [Berschied & Purcell *Inorg Chem* **9** 624 1970, DOI: 10.1021/ic50085a039].

(c) When the above procedures fail to give a clean product, then dissolve NaBH_3CN (10g) in tetrahydrofuran (80ml) and add N MeOH/HCl until the pH is 9. Pour the solution with stirring into dioxane (250ml). The solution is filtered and heated to reflux. A further volume of dioxane (150ml) is added slowly with swirling. The solution is cooled slowly to room temperature, then chilled in ice and the crystalline dioxane complex is collected, dried in a vacuum for 4 hours at 25° , then 4 hours at 80° to yield the amorphous dioxane-free powder (6.7g) with purity >98% [Borch et al. *J Am Chem Soc* **93** 2897 1971, DOI: 10.1021/ja00741a013]. The purity can be checked by iodometric titration [Lyttle et al. *Anal Chem* **24** 1843 1952, DOI: 10.1021/ac60071a041].

NaBH_3CN is a selective reducing agent [Boesch et al. *J Am Chem Soc* **93** 2897 1971, DOI: 10.1021/ja00741a013; Lane *Aldichimica Acta* **8** 3 1975], albeit **less powerful than NaBH_4** due to the electron withdrawing CN group, but is more stable in aqueous acidic solutions up to pH~3.0. In 12N HCl however, it decomposes very rapidly, but at a rate of $ca\ 10^{-8}$ that of NaBH_4 . It has been used for the reductive amination of aldehydes [Manescalchi et al. *Tetrahedron Lett* **35** 2775 1994, DOI: 10.1016/S0040-4039(00)77030-4], ketones [Mattson et al. *J Org Chem* **55** 2552 1990, DOI: 10.1021/jo00295a060], reductive alkylation of amines [Balogh et al. *Synth Commun* **24** 701 1994, DOI: 10.1080/00397919408012649], and used in the synthesis of a new phenolate-bridged dilanthanum (III) complex [Wang et al. *Inorg Chem* **36** 629 1997, DOI: 10.1021/ic960665v]. [See also Fieser, **4** 448, **5** 607, **6** 537, **7** 334, **8** 454, **9** 424, **10** 360, **11** 481, **12** 445, **13** 287, **14** 286, **16** 305.] **POISONOUS, FATAL if swallowed due to cyanide.**

Sodium dichromate [7789-12-0] $\text{Na}_2\text{Cr}_2\text{O}_7$, M **298.0**, m **84.6°**, **91°(2H₂O)**, **356° (anhydrous)**; b **400°/atm(dec)**, d_4^{25} **2.348**. Crystallise the dichromate from small volumes of H_2O by evaporation to crystallisation. Its solubility in H_2O is 238% at 0° and 508% at boiling. The red *dihydrate* is slowly dehydrated by heating at 100° for long periods. It is *deliquescent* and is a powerful oxidising agent—*do not place it in contact with skin— wash immediately as it is caustic.* (Possible **carcinogen**.)

Sodium dihydrogen orthophosphate dihydrate [13472-35-0 ($2\text{H}_2\text{O}$), 10049-21-5 (H_2O), 7558-80-7 (anhydrous)] NaH_2PO_4 , M **120.0**, m **60°(dec)**, d_4^{20} **1.91**. Crystallise it from warm water (0.5ml/g) by chilling.

Sodium dithionite dihydrate (Reductone, sodium hydrosulfite, hydrolin) [7631-94-9 ($2\text{H}_2\text{O}$), 7775-14-6 (anhydrous)] $\text{Na}_2\text{S}_2\text{O}_4$, M **174.1(nhydr)**, **210.1 ($2\text{H}_2\text{O}$)**, m **110°(loses $2\text{H}_2\text{O}$)**, **267°(dec)**, d^{25} **2.38g/cm³(anhydr)**, $\text{pK}_{\text{Est}(1)}^{25}$ **-3.4**, pK_2^{25} **0.49 (for dithionic acid)**. Crystallise it from hot water (1.1ml/g) by cooling (solubility at 20° :18.2g/100ml for anhydrous salt and 21.9g/100ml for dihydrate). [Fieser **3** 266, **11** 487.]

Sodium ferricyanide monohydrate [14217-21-1, 13601-19-9 (anhydrous)] $\text{Na}_3\text{Fe}(\text{CN})_6 \cdot \text{H}_2\text{O}$, M **298.9**, pK^{25} **<1 (for ferricyanide)**. Crystallise the ferricyanide from hot water (1.5ml/g) or by precipitation from 95% EtOH. Solubility in H_2O is 18.9g/100g at 0° and 67g/100g at 100° , and is insoluble in EtOH.

Sodium ferrocyanide decahydrate [13601-19-9] $\text{Na}_4\text{Fe}(\text{CN})_6 \cdot 10\text{H}_2\text{O}$, M **484.1**, m **50-80° (loses $10\text{H}_2\text{O}$)**, **435°(anhydr, dec)**, d^{25} **1.458g/cm³**, n_D^{25} **1.530**, pK_3^{25} **2.57**, pK_4^{25} **4.35 (for ferrocyanide)**. Crystallise the pale yellow solid from hot water until free of ferricyanide as shown by the absence of formation of Prussian Blue colour with ferrous sulfate solution. Its solubility in H_2O is 10.2g/100ml at (10°), 17.6g/100ml at (20°), and 39.7g/100ml at (96.6°).

Sodium fluoride [7681-49-4] NaF, M **42.0**, m **993°**, **996°**, b **1695°/atm**, d^{25} **2.558g/cm³**, n_D^{25} **1.3252**. Crystallise NaF from water by partial evaporation in a vacuum desiccator, or dissolve it in water, and precipitate *ca* half of it by adding EtOH. The precipitate is dried in an air oven at 130° for one day, and then stored in a desiccator over KOH. Its solubility in H_2O is 4% at 15° and 4.3% at 25° . [Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 235 1963, Fieser **1** 1073, **2** 382.]

Sodium fluoroborate [13755-29-8] NaBF_4 , M **109.8**, m **384°**, d^{25} **2.47g/cm³**, pK^{25} **-4.9 (for fluoroboric acid $\text{H}_3\text{O}^+\text{BF}_4^-$)**. Crystallise the fluoroborate from hot water (50ml/g) by cooling to 0° . *Alternatively*, free it from insoluble material by dissolving it in a minimum amount of water; then fluoride ions are removed by adding concentrated lanthanum nitrate in excess. After removing lanthanum fluoride by centrifugation, the supernatant is passed through a cation-exchange column (Dowex 50, Na^+ -form) to remove any remaining lanthanum [Anbar

& Guttman *J Phys Chem* **64** 1896 1960, DOI: 10.1021/j100841a021]. It has also been recrystallised from anhydrous MeOH and dried in a vacuum at 70° for 16 hours. Keep it dry as it is *hygroscopic*. [Delville et al. *J Am Chem Soc* **109** 7293 1987, DOI: 10.1021/ja00258a008.]

Sodium hexafluorophosphate [21324-39-0] NaPF_6 , **M 167.9**, **m >200°**, **d**²⁵ **2.369g/cm³**, **pK₁²⁵ ~0.5**, **pK₂²⁵ 5.12** (for fluoro-phosphoric acid $\text{H}_2\text{PO}_3\text{F}$). Recrystallise it from acetonitrile and dry it in a vacuum for 2 days at room temperature. It is an **irritant** and is *hygroscopic*. [Delville et al. *J Am Chem Soc* **109** 7293 1987, DOI: 10.1021/ja00258a008.].

Sodium hexafluorosilicate [16893-85-9] Na_2SiF_6 , **M 188.1**, **d**²⁵ **2.7g/cm³**, **n_D²⁵ 1.312**. Crystallise it from hot water by cooling. Its solubility in H_2O is 0.64g/100ml at (20°), 1.27g/100ml at (50°), and 2.45g/100ml at (100°).

Sodium hexanitrocobaltate III [13600-98-1] $\text{Na}_3[\text{Co}(\text{NO}_2)_6]$, **M 403.9**, **m >220°(dec)**. Dissolve the salt (ca 60g) in H_2O (300ml), filter to obtain a clear solution, add 96% EtOH (250ml) with vigorous stirring. Allow the precipitate to settle for 2 hours, filter it off, wash it with EtOH (4 x 25ml), twice with Et_2O and dry it in air [Glemser in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1541 1965]. It forms yellow to brown crystals which are very soluble in H_2O , are decomposed by acid and form an insoluble K salt. Used for estimating potassium.

Sodium hydride [7646-69-7] NaH , **M 24.00**, **m 300°(dec)**, **~425°(dec)**, **800°(dec)**, **d**²⁵ **1.396g/cm³**, **n_D²⁵ 1.470**. NaH is a granular grey powder which **ignites** in moist air and should be stored in an inert atmosphere. It is prepared in the laboratory by passing H_2 through a dispersion of Na in an industrial white oil at ca 250°. [Mattson et al. *Inorg Synth* **5** 10 1957, DOI: 10.1002/9780470132364.ch3]. Like sodium it is very reactive with H_2O liberating hydrogen and NaOH . Fortunately it is commercially available as a 50% to 60% (w/w) dispersion in oil, which protects the NaH from the atmosphere, moderates its reaction with moisture, and can be handled routinely without recourse to using dry box techniques. In this form its reaction with H_2O (and alcohols) can be controlled safely, unlike the dry solid. The oil is freely soluble in petroleum ethers, hexanes, Et_2O , $^*\text{C}_6\text{H}_6$, toluene and THF without affecting NaH reactivity. Removal of the oil is achieved by adding any of these solvents, decanting them off, repeating the process, then adding the desired reaction solvent to cover the NaH . NaH is insoluble in inert organic solvents and in liquid NH_3 . When the dispersion is kept for long periods, or if the lid of the container in which it is kept is removed a considerable number of times, the NaH may have deteriorated and the mixture will contain a high percentage of NaOH and/or NaHCO_3 or Na_2CO_3 . In this case a new batch should be used as laboratory purification is not practicable. The NaH content can, however, be determined by carefully adding H_2O to a known weight of the dispersion oil in a flask and measuring the volume of H_2 released using a manometer. The aqueous solution can then be titrated with standard acid (phenolphthalein indicator) to provide the total Na content. [For many examples of the use of NaH in organic synthesis see Fieser **1** 1075, **2** 382, **4** 452, **5** 610, **6** 541, **7** 335, **8** 458, **9** 427, **12** 447, **14** 288, **16** 307.]

Sodium hydroxide (anhydrous, 'caustic soda') [1310-73-2] NaOH , **M 40.0**, **m 318°, 323°**, **b 1390°/atm**, **d**²⁵ **2.13g/cm³**, **n_D²⁵ 1.3576**, **pK_b 0.2**. Common impurities are water and sodium carbonate. Sodium hydroxide can be purified by dissolving 100g in 1L of pure EtOH, filtering the solution under vacuum through a fine sintered-glass disc to remove insoluble carbonates and halides. (This and subsequent operations should be performed in a dry, CO_2 -free box.) The solution is concentrated under vacuum, using mild heating, to give a thick slurry of the mono-alcoholate which is transferred to a coarse sintered-glass disc and evacuated free of mother liquor. After washing the crystals several times with purified alcohol to remove traces of water, they are dried in a vacuum, with mild heating, for about 30 hours to decompose the alcoholate, leaving a fine white crystalline powder [Kelly & Snyder *J Am Chem Soc* **73** 4114 1951, DOI: 10.1021/ja01153a019]. **CAUSTIC**.

Sodium hydroxide solutions (caustic), NaOH aq, pK²⁵ 14.77. Carbonate ion can be removed by passage through an anion-exchange column (such as Amberlite IRA-400; OH^- -form). The column should be freshly prepared from the chloride form by slow prior passage of sodium hydroxide solution until the effluent gives no test for chloride ions. After use, the column can be regenerated by washing with dilute HCl , then water. Similarly, other metal ions are removed when a 1M (or more dilute) NaOH solution is passed through a column

of Dowex ion-exchange A-1 resin in its Na^+ -form.

Alternatively, carbonate contamination can be reduced by rinsing sticks of NaOH (analytical reagent quality) rapidly with H_2O , then dissolving in distilled H_2O , or by preparing a concentrated aqueous solution of NaOH and drawing off the clear supernatant liquid. (Insoluble Na_2CO_3 is left behind.) Carbonate contamination can be reduced by adding a slight excess of concentrated BaCl_2 or $\text{Ba}(\text{OH})_2$ to a NaOH solution, shaking well and allowing the BaCO_3 precipitate to settle. If the presence of Ba in the solution is unacceptable, an electrolytic purification can be used. For example, sodium amalgam is prepared by the electrolysis of 3L of 30% NaOH with 500ml of pure mercury for cathode, and a platinum anode, passing 15 Faradays at 4Amps, in a thick-walled polyethylene bottle. The bottle is then fitted with inlet and outlet tubes, the spent solution being flushed out by CO_2 -free N_2 . The amalgam is then washed thoroughly with a large volume of deionised water (with the electrolysis current switched on to minimise loss of Na). Finally, a clean steel rod is placed in contact in the solution with the amalgam (to facilitate hydrogen evolution), reaction being allowed to proceed until a suitable concentration is reached, before being transferred to a storage vessel and diluted as required [Marsh & Stokes *Aust J Chem* **17** 740 1964, DOI: 10.1071/CH9640740].

Sodium hypophosphite monohydrate [10039-56-2] $\text{NaH}_2\text{PO}_2 \cdot \text{H}_2\text{O}$, M 106.0 (see pK of hypophosphorous acid). Dissolve it in boiling EtOH, cool and add dry Et_2O till all the salt separates. Collect and dry it in vacuum. It is soluble in 1 part of H_2O . It liberates PH_3 on heating and can *ignite* spontaneously when heated. The anhydrous salt is soluble in ethylene glycol (33% w/w) and propylene glycol (9.7%) at 25°.

Sodium iodate [7681-55-2] NaIO_3 , M 197.9, m dec on heating $>425^\circ$, d^{25}_{40} 4.28g/cm³. Crystallise sodium iodate from water by cooling. Its solubility in H_2O is 2.5g/100ml at (0°), 9.47g/100ml at (25°), and 32.59g/100ml at (100°); and it is soluble in AcOH, poorly soluble in DMF (0.5g/kg) and insoluble in EtOH.

Sodium iodide [7681-82-5] NaI , M 149.9, m 660°, b 1304°/atm, d^{20}_4 3.67. Crystallise NaI from water/ethanol solution and dry it for 12 hours under vacuum, at 70°. *Alternatively*, dissolve it in acetone, filter it and cool it to -20°; the resulting yellow crystals are filtered off and heated in a vacuum oven at 70° for 6 hours to remove acetone. The NaI is then crystallised from very dilute NaOH, dried under vacuum, and stored in a vacuum desiccator [Verdin *Trans Faraday Soc* **57** 484 1961, DOI: 10.1039/TF9615700484]. [For applications see Fieser **1** 1087, **2** 384, **3** 267, **4** 456, **6** 543, **7** 338, **10** 365, **12** 449, **13** 282, **14** 289.] **Sodium iodide dihydrate** [13517-06-1] $\text{NaI} \cdot 2\text{H}_2\text{O}$, M 185.9, crystallises from concentrated H_2O solution by evaporation and has d^{25} 2.448g/ml.

Sodium-mercury amalgam (sodium amalgam) [11110-52-4] Na/Hg , M depends on Na%, m (Na-Hg/m): 0.5%/0°, 1.0%/50°, 1.5%/100°, 2.0%/130°, 2.5%/156°, 3.3%/250°, 4.0%/320°. The composition of the amalgam depends on the amount of Na added to the Hg. The reaction is exothermic and can be violent on first mixing. Generally 1-3% of Na in Hg is used, but higher amalgams can be obtained or prepared. Amalgams with more than 1% are solids with melting points that increase with the percentage of Na reaching $>360^\circ$ at $>5\%$ Na.

Unless it has been prepared recently, it is always best to prepare the amalgam afresh. Three general procedures can be adopted. Use a fume cupboard as Hg vapour is **TOXIC**.

Method 1: For a ~1.2% amalgam, clean dry Na (9g) is placed in a 500ml flask with a reflux condenser, covered with dry toluene (20ml), and warmed gently until the Na has melted. Hg is added dropwise (separating funnel) with swirling. The *exothermic reaction* during the addition of the first few mls of Hg will cause the toluene to reflux, and Hg is added at such a rate as to keep the toluene refluxing gently. When addition is complete, allow the solvent to evaporate and the amalgam will solidify. **Note** that it is low melting (~50°) but will solidify on cooling. The solid amalgam is pulverised in a mortar (under N_2 ; to avoid spluttering of small quantities of solid Na-Hg, the mortar is covered with a thick filter paper, or cardboard, with a hole in the centre to allow the pestle to go through), and stored away from moisture in a tightly stoppered container. Up to a ~2%Na amalgam can be prepared by this method and using 15.2g Na and 750ml of Hg.

Method 2: To a flask with dry N_2 or argon flushing continuously through it, clean freshly cut Na (22.8g) is added followed by Hg (10ml) from a separating funnel. The flask is warmed gently until the reaction commences then more Hg (total 750g) is added, with shaking, at such a rate as to keep the amalgam molten. After half the Hg is added, the amalgam begins to solidify and should be kept liquid by external heating. When addition is complete,

the molten amalgam is poured (under N₂ or Argon) into a cold mortar and powdered (see Method 1 above). This provides a 3% amalgam. It should be stored as before. A 5%Na amalgam corresponds to a composition of NaHg₂, and a 20% amalgam corresponds to a composition of Na₂Hg.

Method 3: This method is convenient and commonly used. To the required volume of Hg in a wide-necked flask continuously flushed with N₂ or Argon, is added Na which is cut into small cubes, *ca* 5mm, and each cube is speared with a pointed glass rod and pushed into the liquid Hg. If traces of metal are not a concern, then it is better to use a stainless steel spike or a long tweezers to pick the Na cubes. A vigorous reaction occurs (perhaps with a flash of light occasionally) as each piece of Na dissolves into the Hg or amalgam. The temperature rises and the amalgam is kept in liquid form by the heat released on adding the Na. Do not allow the amalgam to solidify before all the Na is added. If it does, then heat the flask with a burner to keep the amalgam in a liquid state. To avoid damage from flying pieces of Na resulting from the vigorous reaction, the glass rod or metal pick spearing the Na cubes is inserted through a hole in the centre of a thick cardboard sheet which is placed over the mouth of the flask. The amalgam is then powdered and stored as above.

The amalgams are good reducing agents, and react with H₂O releasing 'nascent H' and H₂, but much less vigorously than Na metal. [Brauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II p 1802 1965, Audrieth *Inorg Synth* Vol I 5 1939, DOI: 10.1002/9780470132326.ch3; Babcock et al. *Inorg Synth* 1 10 1939, DOI: 10.1002/9780470132326.ch4; Holleman *Org Synth Coll* Vol 1 554 1941, DOI: 10.15227/orgsyn.007.0088; Renfrow & Hauser *Org Synth Coll* Vol 2 607 1943, DOI: 10.15227/orgsyn.019.0083; Deulofeu & Guerrero *Org Synth Coll* Vol 3 586 1955, DOI: 10.15227/orgsyn.022.0089.]

Sodium metaperiodate (sodium periodate) [7790-28-5] NaIO₄, M 213.9, m ~300°(anhydr, dec), d²⁵ 3.865g/cm³(anhydr). Recrystallise it from hot water. [Willard et al. *Inorg Synth* I 168 1939, DOI: 10.1002/9780470132326.ch58; Bernays et al. *Inorg Synth* 2 212 1946, DOI: 10.1002/9780470132333.ch66.] Useful oxidant; generates quinones *via* glycol cleavage and oxidation of hydroquinones [Daumas et al. *Synthesis* 64 1989, DOI: 10.1055/s-1989-27155].

Sodium metasilicate pentahydrate [10213-79-3 (5H₂O), 6834-92-0 (anhydrous)] Na₂SiO₃ · 5H₂O, M 122.1 (anhydr), 212.1(5H₂O), m 1088°, d₄²⁰ 2.4. Crystallise it from aqueous 5% NaOH solution. [Schwartz & Mathis *Z Anorg Allgem Chem* 126 55 1923, DOI: 10.1002/zaac.19231260105.]

Sodium molybdate dihydrate [10102-40-6] Na₂MoO₄ · 2H₂O, M 241.9, m 100°(loses 2H₂O), 687°(anhydr), d²⁵ 3.78g/cm³, n_D²⁵ 1.714, pK²⁵ 4.08 (for H₂MoO₄). Crystallise the white powder from hot water (solubility is 84g/100ml at 100°) by cooling to 0°. It is incompatible with alkali metals and oxidising agents, e.g. will *explode* when in contact with molten Mg, and its reaction with hot Na, K or Li is *incandescent*. Used in agriculture as a necessary Mo trace element, and in the metal industry as a corrosion inhibitor. [Braithwaite & Haber *Molybdenum: An outline of its Chemistry and Uses*. 1994, Elsevier Science Amsterdam, The Netherlands.]

Sodium nitrate (Chile saltpeter) [7631-99-4] NaNO₃, M 85.0, m 307°, b 380°/atm, d²⁵ 2.257g/cm³, n_D²⁵ 1.587(trigonal). Crystallise NaNO₃ from hot water (0.6ml/g) by cooling to 0°, or from a concentrated aqueous solution by adding MeOH. Its solubility in H₂O is 73g/100ml at (0°), 91.2g/100ml at (25°), and 180g/100ml at (100°). Dry it under a vacuum at 140°. After two recrystallisations, technical grade sodium nitrate had K, Mg, B, Fe Al, and Li at 100, 29, 0.6, 0.4, 0.2 and 0.2 ppm respectively. (See KNO₃.)

Sodium nitrite [7632-00-0] NaNO₂, M 69.0, m 271°, b 320°/atm, d²⁵ 2.168g/cm³, n_D²⁵ 1.65. Crystallise NaNO₂ from hot water by cooling to 0°, or from its own melt. Its solubility in H₂O is 71.4g/100ml at (0°), 84.8g/100ml at (25°), and 160g/100ml at (100°); in MeOH it is 4.4g/100ml (25°), in Et₂O it is 0.3g/100ml (25°). Dry it over P₂O₅. (See KNO₂.)

Sodium perborate monohydrate and tetrahydrate (PBS, hydrogen peroxide sodium borate adduct) [7632-04-4 and 90568-23-3 (anhydrous), 10332-33-9 (H₂O), 10486-00-7 (4H₂O)] NaBO₃ · nH₂O, M 99.81 (PBS-1, H₂O), 153.9 (PBS-4, 4H₂O), m dec (anhydr), 60-65°, 63° (4H₂O, melts in its own H₂O then dec at 130-150°), pK²⁵ 7.91 [for hypothetical (H₃BO₃, H₂O)⁻ H⁺]. The *perborate* is prepared by reaction of borax (Na₂B₄O₇ · 5H₂O), H₂O₂ and NaOH; and upon recrystallisation from H₂O provides the *tetrahydrate* (solubility of 2.3w/v% at 20°) which on gentle heating dehydrates to the *monohydrate* (solubility of 1.6w/v% at 20°). They

hydrolyse slowly in H_2O producing H_2O_2 and borate, liberating ~10% and ~15% w/w of oxygen respectively. Oxygen release is rapid $>60^\circ$, but they are active at lower temperatures ($40\text{--}60^\circ$) in the presence of additives, e.g. acid or tetraacetylenediamine. X-ray crystallography showed that the structural unit consists of a cyclic **dimer anion** $\text{B}_2\text{O}_4(\text{OH})_4^{2-}$ where the two boron atoms are joined by two oxo bridges in a chair-shaped 6-membered ring [Carrondo & Skapski *Acta Crystallogr* **34B** 3551 1978, DOI: 10.1107/S0567740878011565]. The perborate has oxidising properties and is used in industry as a disinfectant, a bleach in detergents (laundry), cleaning products, dental products, eye drops and is less aggressive than hypochloride in degrading dyes and textiles. [Greenwood & Earnshaw *Chemistry of the Elements* (2nd ed.) 1997, Butterworth-Heinemann. ISBN 0080379419; JRC European Commission, Institute for Health and Consumer Protection (IHPC), Toxicology and Chemical Substance (TCS), European Chemicals Bureau (ECB), 1-21027 Ispra (VA) Italy, 2007 EUR 22973 EN/2]. [For applications see Fieser **1** 1102, **2** 387, **4** 59, **12** 452, **14** 290, **16** 310.]

Also used as a chemical oxidising agent, e.g. in AcOH as solvent it oxidises *p*-substituted anilines to the respective 4,4'-disubstituted azobenzenes [Mehta & Vakilwala *J Am Chem Soc* **74** 563 1952, DOI: 10.1021/ja01122a514; Santurri et al. *Org Synth* **40** 18 1960, DOI: 10.15227/orgsyn.040.0018], and shown to be a useful epoxidising agent for quinones including (\pm) terric acid in high yields [Rashid & Read *J Chem Soc C* 1323 1967, DOI: 10.1039/J39670001323]. **IRRITANT** on skin and if inhaled.

Sodium percarbonate ($\text{Na}_2\text{CO}_3 \cdot 1.5\text{H}_2\text{O}_2$, hydrogen peroxide sodium carbonate adduct) [15630-89-4] $\text{Na}_2\text{CO}_3 \cdot 1.5\text{H}_2\text{O}_2$, M 157.0, dec $>120^\circ$. This is not a per-salt, but is crystalline Na_2CO_3 with H_2O_2 of crystallisation with a specific composition. The crystals are consistent in having 1.5 molecules of H_2O_2 for each molecule of Na_2CO_3 . It is a stable, inexpensive, a non-toxic source of H_2O_2 that is easily handled in the laboratory. It is prepared by the addition of excess of H_2O_2 (~3 to 15%) to a saturated aqueous solution of analytically pure Na_2CO_3 or NaHCO_3 at room temperature, stirring for 30 minutes then absolute EtOH is added. The percarbonate separates as white needle-shaped crystals. These are filtered off, washed with EtOH then Et_2O and dried in air. Its available oxygen content is determined by titration with standard KMnO_4 after acidification, and is ~13-15%. [Galwey & Hood *J Phys Chem* **83** 1810 1979, DOI: 10.1021/j100477a003.] Its solubility in H_2O at $\sim 25^\circ$ is ~1 mol/L and the pH is ~10.5. The **kinetics of thermal decomposition** have been studied in detail and are autocatalytic, and best represented by the equation $\text{Na}_2\text{CO}_3 \cdot 1.5\text{H}_2\text{O}_2 = \text{Na}_2\text{CO}_3 + 1.5 \text{H}_2\text{O} + 0.75\text{O}_2$. [Galwey & Hood *JCS Faraday Trans 1* **78** 2815 1982, DOI: 10.1039/F19827802815.] It is a useful oxidising agent; converting nitriles to amides [Kabalka et al. *Synth Commun* **20** 1445 1990, DOI: 10.1080/00397919008052860], oxidises α -diketones to dicarboxylic acids [Yang et al. *Synth Commun* **23** 1183 1993, DOI: 10.1080/00397919308018595], it promotes the Baeyer-Villiger reaction in TFA at 0° [Olah et al. *Synthesis* 739 1991], forms epoxides with olefins, hydroxylates arenes, forms *N*-oxides, and converts sulfides into sulfones [Review: Muzart *Synthesis* 1325 1995, DOI: 10.1055/s-1995-4128], and converts selectively primary and secondary alcohols to their respective carbonyl compounds in the presence of catalytic amounts of $\text{K}_2\text{Cr}_2\text{O}_7$ and Adogen 464 [see 72749-59-8] [Muzart et al. *Tetrahedron Lett* **35** 1989 1994, DOI: 10.1016/S0040-4039(00)73029-2; Mohand & Muzart *Synth Commun* **25** 2373 1995, DOI: 10.1080/00397919508015440]. It is a useful disinfectant and antiseptic, and is best stored below 10° .

For X-ray crystallography, the crystals were prepared by slowly evaporating in air a solution of Na_2CO_3 in 10% H_2O_2 , or in an excess of 15% H_2O_2 or D_2O_2 by standing for 1 hour at -5° before precipitating with EtOH, washing with EtOH and Et_2O , and drying in air. $\text{Na}_2^{13}\text{CO}_3$ was similarly used. The X-ray structure revealed the presence of H_2O_2 molecules hydrogen bonded to the CO_3 ions in the crystal, with two types of H_2O_2 molecules. The IR and Raman spectra of the normal, ^{13}C and ^2H isotopically substituted 'percarbonates' are consistent with the X-ray data where the two H_2O_2 sites are different. The IR (nujol, 25°) of $\text{Na}_2\text{CO}_3 \cdot 1.5\text{H}_2\text{O}_2$ has ν_{max} at 3050w, 2900m, 2720sh, 2490s, 2350m, 1550m, 985m, 960s, 973vw, 868w cm^{-1} , and $\text{Na}_2\text{CO}_3 \cdot 1.5^2\text{H}_2\text{O}_2$ has ν_{max} at 2470w, 2210m, 1940sh, 1890s, 1820m, 1130m, 1040s, 1035m, 873vw, 869w, 660m, 635s cm^{-1} [Carrondo et al. *JCS Dalton Trans* 2323 1977, DOI: 10.1039/DT9770002323.]

Sodium perchlorate (anhydrous) [7601-89-0 (anhydrous), 7791-07-3 (H_2O)] NaClO_4 , M 122.4, m 130° (for monohydrate), 468° (anhydrous), $d^{25} 2.4994\text{g/cm}^3$, $n_D^{25} 1.4618$, $\text{pK}^{25} -2.4$ to -3.1 (for HClO_4). Because its solubility in water is high (2.1g/ml at 15°) and it has a rather low-temperature coefficient of solubility, sodium perchlorate is usually crystallised from acetone, MeOH, water/ethanol or dioxane/water (33g dissolved in 36ml of water and 200ml of dioxane is added). After filtering and recrystallising, the solid is dried under vacuum at

140-150° to remove solvent of crystallisation. Basic impurities can be removed by crystallisation from hot acetic acid, followed by heating at 150°. If NaClO₄ is precipitated from distilled water by adding HClO₄ to the chilled solution, the precipitate contains some free acid. **EXPLOSIVE.**

Sodium phosphoamidate [3076-34-4] $\text{H}_2\text{N-P(ONa)}_2\text{O}$, **M 121.0**. Dissolve it in water below 10°, and acetic acid is added dropwise to pH 4.0 so that the monosodium salt is precipitated. The precipitate is washed with water and Et₂O, then dried in air. Addition of the stoichiometric amount of NaOH to the solution gives the disodium salt, the solution being adjusted to pH 6.0 before use [Rose & Heald *Biochem J* **81** 339 1961, DOI: 10.1042/bj0810339].

Sodium-potassium alloy [11135-81-2] **Na/K, M depends on composition, liquid**. This is generally in the form of a liquid globule from which an aliquot can be pipetted under dry N₂. It is available commercially in two ratios, viz K (78wt%), Na (22 wt%) and K (56 wt%), Na (44 wt%), in ampoules. A fresh *ca* ~5:1 NaK_{2.8} liquid alloy can be prepared from clean K metal (77g, 2.0mol) and clean Na (16g, 0.7mol) by heating gently in xylene (100ml, freshly distilled from Na) until the metals coalesce, then dry oxygen-free diglyme (50ml) is added in order to keep the alloy in one globule (stir with a glass rod). Small globules of the alloy can be united by adding a few drops of *iso*-PrOH to the mixture. Cool to room temperature, and this can be stored under dry N₂ indefinitely. An aliquot can be safely pipetted out from the central globule by syringe equipped with a metal stopcock. The alloy can be more safely disposed of than the Na/Hg alloy, but must be handled carefully as it ignites in moist air. Residues are readily destroyed by a 1:1 mixture of *iso*-PrOH/petroleum ether (b 60-100°). [Ellis & Flom *J Organomet Chem* **99** 263 1975, DOI: 10.1016/S0022-328X(00)88455-7; Gilman & Young *J Org Chem* **1**, 315 1936, DOI: 10.1021/jo01233a001.] When it reacts, e.g. in THF, to form metal carbonyl anions, and with some hydrocarbons, e.g. cyclopentadienes, diphenyl ether, it always forms the K-metallide, i.e. cyclopentadienylpotassium and C₆H₅K, and not the Na-metallide [Bryce-Smith & Turner *J Chem Soc* 861 1953, DOI: 10.1039/JR9530000861; Ziegler & Schnell *Justus Liebigs Ann Chem* **437** 227 1924, DOI: 10.1002/jlac.19244370114; Müller & Bunge *Chem Ber* **69** 2164 1936, DOI: 10.1002/cber.19360690929].

Sodium pyrophosphate decahydrate (tetrasodium pyrophosphate) [13472-36-1 (10H₂O), 7722-88-5 (anhydrous)] $\text{Na}_4\text{P}_2\text{O}_7 \cdot 10\text{H}_2\text{O}$, **M 446.1(10H₂O), 265.9(anhydr)**, **d²⁵ 2.534g/cm³, n_D²⁵ 1.425, m 988°(anhydr), 79.5°(10H₂O), pK₁²⁵ 1.52, pK₂²⁵ 2.36, pK₃²⁵ 6.60, pK₄²⁵ 9.25 (for pyrophosphoric acid, H₄P₂O₇)**. Crystallise the salt from hot H₂O and dry it in air at room temperature. Its solubility in H₂O is 2.61g/100ml at (0°), 6.7g/100ml at (25°), and 42.2g/100ml at (100°). Useful buffer, and a laxative.

Sodium selenate [13410-01-0, 10102-23-5 (10H₂O)] Na_2SeO_4 , **M 188.9(anhydr)**, **d²⁵ 3.098g/cm³, pK₁²⁵ ~0, pK₂²⁵ 1.66 (for selenic acid, H₂SeO₄)**. Crystallise sodium selenate from hot water. [Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 433 1963]. Used in the glass industry to produce a red hue. Also used as a fungicide and insecticide. **TOXIC.**

Sodium selenite [10102-18-8] Na_2SeO_3 , **M 172.9, m >350°, 710°(dec), d²⁵ 3.1g/cm³, pK₁²⁵ 2.62, pK₂²⁵ 8.32 (for H₂SeO₃)**. Crystallise sodium selenite from a saturated aqueous solution where its solubility is 68% at 20° to give the **pentahydrate** salt. The solubility at 20° is 85g/100ml of H₂O. This yields the **anhydrous** salt on heating at 40°. [Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 431 1963]. It imparts a pink colour to glass which cancels the green colour from Fe impurities. Used as a trace element supplement. **TOXIC.**

Sodium silicate solution (water glass, sodium trisilicate) [1344-09-8, 6834-92-0] $\text{Na}_2\text{Si}_3\text{O}_7$, **M 242.2, d²⁵ 3.1g/cm³, n_D²⁵ 1.52, pK₁²⁵ 9.51, pK₂²⁵ 11.77 (for silicic acid, H₄SiO₄)**. Purify by contact filtration through activated charcoal. Its solubility in H₂O is 2.2g/100ml at (25°) and 160.6g/100ml at (80°). Used to preserve food and eggs; as an adhesive in concrete, stucco, caulking and plasters, automotive repair leaks, e.g. exhaust pipes, as an abrasive in drilling, in detergents and waterproofing.

Sodium sulfate decahydrate (Glauber's salt) [7727-73-3 (10H₂O), 7757-82-6 (anhydrous)] $\text{Na}_2\text{SO}_4 \cdot 10\text{H}_2\text{O}$, **M 140.0(anhydr), m 32.3°(hydrate, dec), 884° (anhydrous), d²⁵ 2.664g/cm³ (anhydrous), d²⁵ 1.464 g/cm³, (10H₂O), n_D²⁵ 1.468(anhydr), n_D²⁵ 1.394(10H₂O)**. Crystallise sodium sulfate from water at 30° (1.1ml/g) by

cooling to 0°. It becomes anhydrous at 32°. The solubility of the **anhydrous salt** in H₂O is 4.76g/100ml at (0°), 13.9g/100ml at (20°), and 42.7g/100ml at (100°); and the **heptahydrate** is 2.19.5g/100ml at (0°) and 44g/100ml at (20°). Useful for the manufacture of wood pulp, glass industry as a fining agent (removes air bubbles), in textiles, as an inert drying agent for solvents when anhydrous, and for heat storage in solar panels. The **decahydrate** was historically used as a laxative. [Fieser 1 1103.]

Sodium sulfide nonahydrate [1313-84-4 (9H₂O), 1313-82-2 (anhydrous)] Na₂S 10H₂O, M 240.2(9H₂O), 78.0(anhydr), m ~50(loses H₂O), 950(anhydrous), d²⁵ 1.856g/cm³ (anhydrous), d²⁵ 1.58g/cm³ (anhydrous), (5H₂O), d²⁵ 1.43g/cm³ (9H₂O), m 50°(9H₂O, dec), 100°(5H₂O, dec), 1176° (anhydrous), pK₁²⁵ 7.04, pK₂²⁵ 11.96 (for H₂S). Some purification of the pale yellow hydrated salt can be achieved by selecting large crystals and removing the surface layer (contaminated with oxidation products) by washing with distilled water. Other metal ions can be removed from Na₂S solutions by passage through a column of Dowex ion-exchange A-1 resin, Na⁺-form. The hydrated salt can be rendered **anhydrous** by heating it in a stream of H₂ or N₂ until water is no longer evolved. (The resulting cake should not be heated to fusion because it is readily oxidised.) Recrystallise it from distilled water [Andersson & Azoulay *JCS Dalton Trans* 469 1986, DOI: 10.1039/DT9860000469]. **Note** that sodium sulfide hydrolyses in H₂O to form NaHS + H₂O, and is therefore alkaline and the solution has the odour of rotten eggs, as does the solid. A 0.1N solution in H₂O is 86% hydrolysed at room temperature. Its solubility in H₂O is 12.4% at 0°, 18.6% at 20° and 39% at 50°. The **anhydrous** salt is obtained by allowing it to stand in a vacuum over concentrated H₂SO₄ or P₂O₅ at 45° to start with, then at 30-35° when the salt contains 4% of water. The last traces of water are removed by heating to 700° in a glass or porcelain tube in a stream of H₂ to give pure H₂S. [Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 358-360 1963, for applications see Fieser 1 1104, 11 492, 12 543.] It has found use in the pulp and paper industry, in water treatment to scavenge O₂ and precipitate heavy metals, and used in the leather rubber industry.

Sodium sulfite [7757-83-7] Na₂SO₃, M 126.0, m 33.4°(7H₂O, dec), 500°(anhydr, dec), d²⁵ 2.633g/cm³, n_D²⁵ 1.565, pK₁²⁵ 1.82, pK₂²⁵ 7.205 (for H₂SO₃). Crystallise the sulfite from warm water (solubility at 20° is 27%) by cooling to 0°. Also purify it by repeated crystallisation from deoxygenated water inside a glove-box, and finally drying it under vacuum. [Rhee & Dasgupta *J Phys Chem* 89 1799 1985, DOI: 10.1021/j100255a052.] It is a useful reagent for particular photochemical reductions [see Fieser 2 387]. Used in the pulp and paper industry, as a preservative; also useful in photography to protect developer solutions and for the production of SO₂ by addition of dilute acid.

Sodium tetrametaphosphate [13396-41-3, 77489-25-9] Na₄P₄O₁₂, M 429.9, pK₁²⁵ 2.60, pK₂²⁵ 6.4, pK₃²⁵ 8.22, pK₄²⁵ 11.4 (tetrametaphosphoric acid, H₄P₄O₁₂). Crystallise it twice from water at room temperature by adding EtOH (300g of Na₄P₄O₁₂·H₂O, 2L of water, and 1L of EtOH), wash it first with 20% EtOH then with 50% EtOH and dry it in air [Quimby *J Phys Chem* 58 603 1954, DOI: 10.1021/j150518a006].

Sodium thioantimonate (Schlippe's salt) [13776-84-6] Na₃SbS₄·9H₂O, M 481.1, m 87°, b 234°/atm, d²⁵ 1.806g/cm³. Crystallise the pale yellow solid from warm water (2ml/g made weakly alkaline with a few drops of dilute aqueous NaOH) by cooling to 0°. It forms a yellow **nonahydrate** which readily **effloresces** in air. The salt was used for preparing antimony quinsulfide Sb₂S₅ (gold sulfur) on acidification which is used to vulcanise rubber. The antimony (V) sulfide was used as an **amplifier** in Ag-based photography, as a flammable component of matches and for the electrodeposition of zinc. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 619 1963, and for the crystal structure see Mereiter et al. *Acta Cryst* 35B 19 1979, DOI: 10.1107/S0567740879002442].

Sodium thiocyanate (sodium rhodanate) [540-72-7] NaSCN, M 81.1, m 300°, pK²⁵ -1.85 (for HSCN). It is recrystallised from EtOH or Me₂CO, and the mother liquor is removed from the crystals by centrifugation. It is very **deliquescent** and should be kept in an oven at 130° before use. It can be dried in a vacuum at 120°/P₂O₅ [Partington & Winterton *Trans Faraday Soc* 30 1104 1934, DOI: 10.1039/TF9343001104]. Its solubility in H₂O is 113% at 10°, 178% at 46°, 225.6% at 101.4°; in MeOH 35% at 15.8°, 51% at 48°, 53.5% at 52.3°; in EtOH 18.4% at 18.8°, 24.4% at 70.9°; and in Me₂CO 6.85% at 18.8° and 21.4% at 56° [Hughes & Mead *J Chem Soc* 2282 1929, DOI: 10.1039/JR9290002282].

Sodium thiocyanate has also been recrystallised from water, acetonitrile or from MeOH using Et₂O for washing,

then dried at 130°, or dried under vacuum at 60° for 2 days. [Strasser et al. *J Am Chem Soc* **107** 789 1985, DOI: 10.1021/ja00290a009.] (The latter purification removes material reacting with iodine.) Sodium thiocyanate solutions can be freed from traces of iron by repeated batch extractions with Et₂O. [For preparation of *isopropylthiocyanate* [625-59-2] **b** 152-153°/atm, see Shriner *Org Synth Coll Vol* **2** 366 1943, DOI: 10.1002/0471264180.os011.27; and for the synthesis of thiaza heterocycles see Allen and van Allan *Org Synth Coll Vol* **3** 76 1955, DOI: 10.15227/orgsyn.022.0016.]

Sodium thiosulfate pentahydrate (hypo, hydrosulfite) [10102-17-7 (5H₂O), 7772-98-7 (anhydrous)] **Na₂S₂O₃·5H₂O**, **M 248.2(H₂O)**, **158.1°(anhydrous)**, **m 48°(rapid heat)**, **d²⁵ 1.667g/cm³ (anhydrous)**, **n_D²⁵ 1.489**, **pK₁²⁵ 0.6**, **pK₂²⁵ 1.74 (for H₂S₂O₃)**. Crystallise it from EtOH/H₂O solutions or from water (solubility is 70.1g/100ml at 20° and 231g/100ml at 100°) below 60° by cooling to 0°, and dry it at 35° over P₂O₅ under vacuum. [Foerster & Mommsen *Chem Ber* **57** 258 1924, DOI: 10.1002/cber.19240570218.] This salt is used as a secondary standard in volumetric analysis (iodometry) [Kilpatrick *J Am Chem Soc* **45** 2132 1923, DOI: 10.1021/ja01662a014], and is used as '**Hypo**' in photography [Hargreaves & Dunningham *J Soc Chem Ind* **42** 147T 1923, DOI: 10.1002/jctb.5000421517]. The reaction of Na₂S₂O₃ with HCl which produces NaCl + S + SO₂ + H₂O is known as the **Clock Reaction** because when the S reaches a certain concentration, the solution changes from colourless to pale yellow.

Sodium trimetaphosphate hexahydrate [7785-84-4] **Na₃P₃O₉·6H₂O**, **M 305.9(anhydr)**, **m 53°(6H₂O)**, **d²⁵ 2.49g/cm³ (anhydrous)**, **d²⁵ 1.786g/cm³ (6H₂O)**, **n_D²⁵ 1.489 (6H₂O)**, **pK₂²⁵ 1.64**, **pK₃²⁵ 2.07 (for trimetaphosphoric acid, H₃P₃O₉)**. It is precipitated from an aqueous solution at 40° by adding EtOH. Its solubility in HO is 22% at ~25°. It is dried in air.

Sodium tripolyphosphate (STPP, tripolyphosphate TPP, sodium triphosphate STP) [7758-29-4] **Na₅P₃O₁₀**, **M 367.9**, **m 622°**, **pK₁²⁵ ~1**, **pK₂²⁵ 2.0**, **pK₃²⁵ 2.13**, **pK₄²⁵ 5.78**, **pK₅²⁵ 8.56 (for tripolyphosphoric acid, H₅P₃O₁₀)**. Purify it by repeated precipitation from aqueous solution by slow addition of MeOH and dried in air. Also a solution of anhydrous sodium tripolyphosphate (840g) in water (3.8L) is filtered, MeOH (1.4L) is added with vigorous stirring to precipitate Na₅P₃O₁₀·6H₂O. The precipitate is collected on a filter, air dried by suction, then left to dry in air overnight. It is crystallised twice more in this way, using a 13% aqueous solution (w/w), and leaching the crystals with 200ml portions of water [Watters et al. *J Am Chem Soc* **78** 4855 1956, DOI: 10.1021/ja01600a011]. Similarly, EtOH can be added to precipitate the salt from a filtered 12-15% aqueous solution, the final solution containing *ca* 25% EtOH (v/v). Air drying should be at a relative humidity of 40-60%. Heat and vacuum drying should be avoided. Its solubility in H₂O is 14.5g/100ml at (25°). [Quimby *J Phys Chem* **58** 603 1954, DOI: 10.1021/j150518a006; Klement in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 547 1963.] This phosphate is used in detergents, ceramics, leather tanning, anti-caking, anticorrosion, antifreeze, textiles, and fermentation.

Sodium tungstate dihydrate [10213-10-2 (2H₂O), 13472-45-2] **Na₂WO₄·2H₂O**, **M 329.9**, **m 698°**, **d²⁵ 4.172g/cm³ (anhydrous)**, **d²⁵ 3.25g/cm³ (2H₂O)**, **pK₁²⁵ 2.20**, **pK₂²⁵ 3.70 (for tungstic acid, H₂WO₄)**. The salt crystallises from hot water (0.8ml/g) on cooling to 0°. Its solubility in H₂O is 57.5g/100ml at (0°), 74.2g/100ml at (25°), 96.9g/100ml at (100°). [Grüttner & Jender in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1727 1965; for applications see Fieser **6** 455, **13** 145, **15** 295.] The oxidation of alkenes and alcohols are catalysed by this salt. Sodium tungstate is known also for its anti-diabetic properties [Dominiguez et al. *J Biol Chem* **278** 42785 2003, DOI: 10.1074/jbc.M308334200].

Stannic chloride (tin IV chloride, stannic tetrachloride) [7646-78-8, 10026-06-9 (5H₂O)] **SnCl₄**, **M 260.5**, **m -33°, -30.2°, b 114°/760mm**, **d²⁵ 2.226g/cm³ (anhydrous)**, **d²⁵ 2.04g/cm³ (5H₂O)**, **n_D²⁵ 1.512**, **pK²⁵ 14.15 (for aquo Sn⁴⁺ hydrolysis)**. SnCl₄ fumes in moist air due to formation of a *hydrate*. Fractionate it in a ground glass still and store it in the absence of air. Possible impurities are SO₂ and HCl [Baudler in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 729 1963]. It forms a solid **pentahydrate** [10026-06-9] which smells of HCl and is obtained when the **anhydrous salt** is dissolved in a small volume of H₂O. Also reflux it with clean mercury or P₂O₅ for several hours, then distil it under (reduced) N₂ pressure into a receiver containing P₂O₅. Finally redistil it. *Alternatively*, distil it from Sn metal under vacuum in an all-glass system and seal off in large ampoules. SnCl₄ is available commercially as 1M solutions in CH₂Cl₂ or hexane. It

is a Lewis acid catalyst [Hirabayashi et al. *Chem Lett (Jpn)* **36** 826 2007, DOI: 10.1246/cl.2007.826], and stabilises polyaniline (by doping) allowing it to be processed in MeNO₂ solution [Kulszewicz-Bajer et al. *Chem Mater* **11** 552 1999, DOI: 10.1021/cm980727a]. [Also see Fieser **1** 1111, **3** 269, **5** 627, **6** 553, **7** 342, **9** 436, **10** 370, **11** 522, **12** 486, **13** 300, **14** 304, **15** 311.] **HARMFUL VAPOURS.**

Stannic iodide [7790-47-8] **SnI₄**, **M 626.3**, **m 144°**, **b 340°/atm**, **d²⁵ 4.47g/cm³**. It is recrystallised from anhydrous CHCl₃, dry it under vacuum and store it in a vacuum desiccator. It sublimes at 180°.

Stannic oxide (SnO₂) [18282-10-5] **SnO₂**, **M 150.7**, **m 1630°**, **d²⁵ 6.95g/cm³**. Reflux it repeatedly with fresh HCl until the acid shows no tinge of yellow. The oxide is then dried at 110°. It is abrasive and used for polishing glass, ceramic and metal, and as a mordant in dyeing and textile printing.

Stannous chloride (anhydrous) [7772-99-8, 10025-69-1 (2H₂O)] **SnCl₂**, **M 189.6**, **m 247°(anhydr)**, **37.7°(2H₂O)**, **b 606°/atm**, **b 652°/atm(dec)**, **d²⁵ 3.94g/cm³(anhydrous)**, **d²⁵ 2.71g/cm³(2H₂O)**, **pK₁²⁵ 1.7**, **pK₂²⁵ 3.7 (for aquo Sn²⁺ hydrolysis)**. Analytical reagent grade stannous chloride *dihydrate* is dehydrated by adding it slowly to vigorously stirred, redistilled acetic anhydride (120g salt per 100g of anhydride) in a fume cupboard. After *ca* an hour, the *anhydrous* SnCl₂ is filtered on to a sintered-glass or Büchner funnel, washed free from acetic acid with dry Et₂O (2 x 30ml), and dried under vacuum. It is stored in a sealed container. [Stephen *J Chem Soc* 2783 2786 1930, DOI: 10.1039/JR9300002783; Williams *Org Synth Coll Vol* **3** 626 1955, DOI: 10.15227/orgsyn.023.0063; Fieser **1** 1113, **3** 389, **5** 631, **6** 554, **11** 521, **13** 298, **15** 309, **16** 329.] Used in tin-plating, dyeing, as a reducing agent, and for many chemical reactions.

Strontium bromide [10476-81-0, 7789-53-9 (6H₂O)] **SrBr₂**, **M 247.4**, **m 643°/atm**, **d²⁵ 4.216g/cm³(anhydrous)**, **d²⁵ 2.386g/cm³(6H₂O)**. Crystallise the bromide from water (0.5ml/g). It is soluble in EtOH and is *deliquescent*. [Ehrlich in *Handbook of Preparative Inorganic Chemistry (Ed. Brauer)* Academic Press Vol **I** p 930 1963.] It is a white crystalline powder that burns bright red in air and is used in flares. [For crystal structure see Sass; et al. *J Phys Chem* **67** 2862 1963, DOI: 10.1021/j100806a516].

Strontium chloride hexahydrate [10025-70-4 (6H₂O), 10476-85-4] **SrBr₂ · 6H₂O**, **M 266.6(6H₂O)**, **158.5(anhydr)**, **m 61°(rapid heating)**, **114-150°(loses 5H₂O)**, **868°(anhydrous)**, **b 1250°/atm**, **d²⁵ 3.052g/cm³(anhydrous)**, **d²⁵ 1.930g/cm³(6H₂O)**, **n_D²⁵ 1.650(anhydrous)**, **n_D²⁵ 1.536(6H₂O)**. It crystallises from warm water (solubility of *hexahydrate* is 106g/100ml at 0° and 206g/100ml at 40°, and for *anhydrous salt* it is 53.8g/100ml at 20°) and cooling to 0°. It *dehydrates* at 150-160° in a vacuum. [Ehrlich in *Handbook of Preparative Inorganic Chemistry (Ed. Brauer)* Academic Press Vol **I** p 930 1963.] It is used in dental care.

Strontium chromate [7789-06-2] **SrCrO₄**, **M 203.6**, **d²⁵ 3.353g/cm³**, **pK₁²⁵ 0.74**, **pK₂²⁵ 6.49 (for H₂CrO₄)**. Crystallise yellow strontium chromate from water (solubility is 0.12g/100ml at 15° and 3.0g/100ml at 100°) by cooling. The salt is used as a colorant, an inhibitor in pigments, anti-corrosion primer for Zn, Mg, Al and alloys in aircrafts, and in pyrotechnics.

Strontium hydroxide octahydrate [1311-10-0 (8H₂O), 18480-07-4 (anhydrous)] **Sr(OH)₂ · 8H₂O**, **M 121.6(anhydr)**, **m >100°(loses H₂O)**, **375°(anhydr)**, **b 710°/atm(anhydr, dec)**, **d²⁵ 3.625g/cm³(anhydr)**, **d²⁵ 1.90g/cm³(8H₂O)**, **pK_b -2.19**. Crystallise the hydroxide from hot water (solubility is 0.41g/100ml at 0°, and 1.77g/100ml at 20°, and 21.8g/100ml at 100°) by cooling to 0°. Readily absorbs CO₂ to form SrCO₃. *Deliquescent*. [Georg in *Handbook of Preparative Inorganic Chemistry (Ed. Brauer)* Academic Press Vol **I** p 935 1963.] Used in refining beet sugar and as a plastic stabiliser. **IRRITANT.**

Strontium nitrate [10042-76-9] **Sr(NO₃)₂**, **M 211.6**, **m 570°**, **b 645°/atm (dec)**, **d²⁵ 2.986g/cm³**. Crystallise it from hot water (solubility of *tetrahydrate* is 60.43g/100ml at 0° and 206.5g/100ml at 100°; and of the *anhydrous salt* is 71.0g/100ml at 18° and 66.0g/100ml at 20°) by cooling to 0°. Its physiological effects are similar to those of Ca²⁺, and lessens skin irritations. It burns with a rich red flame in fireworks and road flares.

Sulfamic acid [5329-14-6] **NH₂SO₃H**, **M 97.1**, **m 205°(dec)**, **pK²⁵ 0.99 (NH₂SO₃H)**. Crystallise the white

solid from water at 70° (300ml per 25g), after filtering, by cooling a little and discarding the first batch of crystals (about 2.5g) before standing in an ice-salt mixture for 20 minutes. The crystals are filtered off by suction, washed with a small quantity of ice cold water, then twice with cold EtOH and finally with Et₂O. Dry it in air for 1 hour, then store it in a desiccator over Mg(ClO₄)₂ [Sisler et al. *Inorg Synth* **2** 176 1946, DOI: 10.1002/9780470132333.ch52; for use as standard of reference in acidimetry see Butler et al. *Ind Eng Chem (Anal Ed)* **10** 690 1938, DOI: 10.1021/ac50128a011]. For the preparation of primary standard material see *Pure Appl Chem* **25** 459 1969. Aqueous solutions are unstable and hydrolyse to ammonium bisulfate, but the crystals are indefinitely stable on normal storage. It is moderately soluble in DMF, slightly soluble in MeOH but insoluble in hydrocarbon solvents. Used for cleaning metals, removing rust and limescale, and household cleaning. [For sulfamic acid and its *N*-substituted derivatives see Benson & Spillane *Chem Rev* **80** 151 1980, DOI: 10.1021/cr60324a002.]

Sulfamide [7803-58-9] (NH₂)₂SO₂, **M 96.1, m 91.5°, 93°, b 250°/atm (dec), d²⁵ 1.611g/cm³**. Crystallise sulfamide from absolute EtOH. It decomposes at 250°. [Schenk in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 482-483 1963, for the role of sulfamide derivatives in Medicinal Chemistry see Reitz et al. A Patent Review (2006 – 2008) *Expert Opinion on Therapeutic Patents* **19** 1449–1453 2009, DOI: 10.1517/13543770903185920.]

Sulfur [7704-34-9] **S, M 32.1, m between 112.8° and 120°, depending on form.** Murphy, Clabaugh & Gilchrist [*J Res Nat Bur Stand* **64A** 355 1960] have obtained sulfur of about 99.999% purity by the following procedure: Roll sulfur was melted and filtered through a coarse-porosity glass filter funnel into a 2L round-bottomed Pyrex flask with two necks. Conc H₂SO₄ (300ml) was added to the sulfur (2.5kg), and the mixture was heated to 150°, stirring continuously for 2 hours. Over the next 6 hours, conc HNO₃ was added in about 2ml portions at 10-15 minutes intervals to the heated mixture. It was then allowed to cool to room temperature and the acid was poured off. The sulfur was rinsed several times with distilled water, then remelted, cooled, and rinsed several times with distilled water again, this process being repeated four or five times to remove most of the acid entrapped in the sulfur. An air-cooled reflux tube (*ca* 40cm long) was attached to one of the necks of the flask, and a gas delivery tube (the lower end about 2.5cm above the bottom of the flask) was inserted into the other. While the sulfur was boiled under reflux, a stream of helium or N₂ was passed through to remove any water, HNO₃ or H₂SO₄, as vapours. After 4 hours, the sulfur was cooled so that the reflux tube could be replaced by a bent air-cooled condenser. The sulfur was then distilled, rejecting the first and the final 100ml portions, and transferred in 200ml portions to 400ml glass cylinder ampoules (which were placed on their sides during solidification). After adding about 80ml of water, displacing the air with N₂, the ampoule was cooled, and the water was titrated with 0.02M NaOH, the process being repeated until the acid content was negligible. Finally, entrapped water was removed by alternate evacuation to 10mm Hg and refilling with N₂ while the sulfur was kept molten. The ampoules were then sealed. Other purifications include crystallisation from CS₂ (which is less satisfactory because the sulfur retains appreciable amounts of organic material), *benzene or *benzene/acetone, followed by melting and degassing. It has also been boiled with 1% MgO, then decanted, and dried under a vacuum at 40° for 2days over P₂O₅. [For the purification of **S₆**, '**recrystallised S₈**' and '**Bacon-Fanelli sulfur**' see Bartlett et al. *J Am Chem Soc* **83** 103, 109 1961, DOI: 10.1021/ja01462a020.] [For applications see Fieser **1** 1118, **3** 273, **5** 632, **6** 556, **15** 297.]

Sulfur dichloride (sulfur chloride) [10545-99-0] SCl₂, **M 103.0, m -121°, -78°, b 59°/760mm(dec), d²⁵ 1.621g/cm³, n_D²⁵ 1.5570**. Distil the red sulfur chloride twice in the presence of a small amount of PCl₃ through a 12in Vigreux column, the fraction boiling between 55-61° being redistilled (in the presence of PCl₃), and the fraction distilling between 58-61° retained. (The PCl₃ is added to inhibit the decomposition of SCl₂ into S₂Cl₂ and Cl₂). The SCl₂ must be used as quickly as possible after distillation — within 1 hour at room temperature. The sample contains 4% of S₂Cl₂. On long standing this reaches 16-18%. [Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** pp 370-372 1963.] It hydrolyses with release of HCl, and old samples contain Cl₂. **HARMFUL VAPOURS.**

Sulfur dioxide [7446-09-5] SO₂, **M 64.1, m -73°, -72°, b -10°/atm, d²⁵ 2.6288g/cm³**. Dry the pungent irritating gas by bubbling through conc H₂SO₄ and by passage over P₂O₅, then through a glass-wool plug. Freeze it with liquid air and pump it to a high vacuum to remove dissolved gases. It is easily liquefied by

compression (2.5 atmospheres at 15°), or by passing it through a glass spiral column in a freezing mixture of ice and salt. It is a colourless liquid with a density of 1.434 at 0°, which on rapid evaporation forms a snow white solid. It has been used as a solvent in certain reactions, a reducing agent, a refrigerant, in wine making, and is a physiologically active compound and an air pollutant. **HARMFUL SUFFOCATING VAPOURS.**

Sulfurous acid [7782-99-2] H_2SO_3 , **M 82.0**, $d^{25} 1.03\text{g/cm}^3$, $\text{pK}_1^{25} 1.857$, $\text{pK}_2^{25} 7.172$, is a weak acid obtained by bubbling SO_2 through H_2O , and a *ca* 6% solution is available commercially. It is a reducing agent (e.g. reducing I_2 to iodide), used for making sulfites and bisulfites, a disinfectant and a mild bleaching agent. [For 'Generation and Characterization of Sulfurous Acid (H_2SO_3) and of Its Radical Cation as Stable Species in the Gas Phase' see Sülzle et al. *Angew Chem Int Ed* **27** 1533 1988, DOI: 10.1002/anie.198815331.]

Sulfuric acid (oil of vitriol) [7664-93-9] H_2SO_4 , **M 98.08**, $d_4^{20} 1.836$ (96-98%), **1.805** (100%), **m 3.0°** (98%), **10.36°** (100%), **b 330.0 ± 0.5°/atm** (100%), **dec > 340.0°**, *dihydrate* **M 134.11**, **m -38.9°**, **b 167°/atm**, $d^{25} 1.84\text{g/cm}^3$, $\text{pK}_1^{25} \sim -8.3$, $\text{pK}_2^{25} 1.99$. Sulfuric acid, and also 30% fuming H_2SO_4 , can be distilled in an all-Pyrex system, optionally from potassium persulfate. It has been purified by fractional crystallisation of the *monohydrate* from the liquid. It has a very strong dehydrating action and attacks skin—wash immediately with cold H_2O *and protect eyes*; otherwise the skin can be scarred for life *and could be blinding*. It causes tissues and paper to char. It is very *hygroscopic* and has been used as a desiccant in desiccators. *Dilution with H_2O is highly exothermic, and because the concentrated acid is much more dense than H_2O it is diluted by running the concentrated acid down the side of the container of H_2O with slow stirring while cooling the outside of the container in an ice bath. If these precautions are not taken, the H_2O is likely to splatter and boil vigorously over.* **CORROSIVE.**

Sulfuric acid (fuming) (oleum) [8014-95-7] H_2SO_4 , **M 98.08 (+ x SO_3)**. This is sulfuric acid with various amounts of SO_3 dissolved in it up to 80% free SO_3 . The specific gravities, **Sp gr** d_4^{20} , of oleums (% free SO_3) are: 1.899 (15%), 1.915 (20%), 1.952 (30%), 2.001 (50%), 2.102 (60% maximum density) and 1.949 (80%). They are an almost colourless viscous liquids which emit choking fumes of SO_3 , and should be handled with great care: **use eye and body protection**. It reacts explosively with H_2O to form strong sulfuric acid, and is a strong sulfonating agent.

Sulfur monochloride (sulfur chloride) [10025-67-9, 12771-08-3] S_2Cl_2 , **M 135.0**, **m -80°**, **-77°**, **b 19.1°**, **29-30°/12mm**, **72°/100mm**, **138°/760mm**, $d^{25} 1.688\text{g/cm}^3$, $n_D^{20} 1.658$. It is a *pungent, irritating golden yellow liquid*. When impure its colour is orange to red due to SCl_2 formed. It fumes in moist air and liberates HCl , SO_2 and H_2S in the presence of H_2O . Distil it and collect the fraction boiling above 137° at atmospheric pressure. Fractionate this fraction over sulfur at *ca* 12mm using a ground glass apparatus (**b 29-30°**). *Alternatively*, purify it by distillation below 60° from a mixture containing sulfur (2%) and activated charcoal (1%), under reduced pressure (e.g. 50mm). It is soluble in EtOH , $^*\text{C}_6\text{H}_6$, Et_2O , CS_2 and CCl_4 . Store it in a closed container in the dark in a refrigerator. [Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 371 1963, see also Fieser **1** 1122, **3** 275, **6** 560, **9** 442, **11** 495.] **VERY HARMFUL VAPOURS.**

Sulfuryl chloride [7791-25-5] SO_2Cl_2 , **M 135.0**, **m -54.1°**, **b 69.3°/760mm**, $d^{25} 1.67\text{g/cm}^3$, $n_D^{20} 1.4473$. It is a *pungent, irritating colourless liquid*. It becomes yellow with time due to decomposition to SO_2 and HCl . Distil it and collect fraction boiling below 75°/atm which is mainly SO_2Cl_2 . To remove HSO_3Cl and H_2SO_4 impurities, the distillate is poured into a separating funnel filled with crushed ice and *briefly* shaken. The lower cloudy layer is removed, dried for some time in a desiccator over P_2O_5 and finally fractionate it at atmospheric pressure. The middle fraction boils at 69-70° and is pure SO_2Cl_2 . It decomposes gradually in H_2O to H_2SO_4 and HCl . It reacts **violently** with EtOH and MeOH and is soluble in $^*\text{C}_6\text{H}_6$, toluene, Et_2O and acetic acid. [Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed Brauer) Academic Press Vol I p 383 1963, Allen et al. *Inorg Synth* **1** 114 1939, DOI: 10.1002/9780470132326.ch42; Maynard *Encyclopedia of Reagents for Organic Synthesis* John Wiley 2001 DOI: 10.1002/047084289X.rs140; for a versatile alternative to chlorine see Moussa *Aust J Chem* **65** 95 2012, DOI: 10.1071/CH11367.] **HARMFUL VAPOURS, LACHRYMATOR.**

Tantalum (V) chloride (tantalum pentachloride) [7721-01-9] TaCl_5 , **M 358.2**, **m 216.2°**, **216.5-220°**,

b 239°/atm, d²⁵ 3.68g/cm³. Purify it by sublimation in a stream of Cl₂. It forms colourless needles when pure (yellow when contaminated with even less than 1% of NbCl₅). It is sensitive to H₂O forming TaOCl₃; even in conc HCl it decomposes to tantalum acid. It is soluble in EtOH, Et₂O and CCl₄. [Rolsten *J Am Chem Soc* **80** 2952 1958, DOI: 10.1021/ja01545a011; Brauer in *Handbook of Preparative Inorganic Chemistry* (Ed Brauer) Academic Press Vol II p 1302 1965.]

Telluric acid [7803-68-1] H₆TeO₆, M 229.6, m 136°, d²⁵ 3.07g/cm³, pK₁²⁵ 7.68, pK₂²⁵ 11.04 (H₆TaO₆). Crystallise it once from nitric acid, then repeatedly from hot water (0.4ml/g). [Fehér in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I pp 451-453 1963.] It has a strong tendency to polymerise, e.g. dehydrates to *polymetatelluric acid* (H₂TeO₄)₁₀ on heating >100°.

Tellurium [13494-80-9] Te, M 127.6, m 450°, 722.6°, b 990°/atm, 1261°/atm, d²⁵ 6.24g/cm³. Purify the silver white tellurium by zone refining and repeated sublimation to an impurity of less than 1 part in 10⁸ (except for surface contamination by TeO₂). [Machol & Westrum *J Am Chem Soc* **80** 2950 1958, DOI: 10.1021/ja01545a010.] Tellurium is volatile at 500°/0.2mm. It has also been purified by electrode deposition [Mathers & Turner *Trans Amer Electrochem Soc* **54** 293 1928]. [Fieser **11** 498.] With KOH, it promotes the pinacolisation of aromatic carbonyl compounds quantitatively [Khan et al. *Synth Commun* **27** 2193 1997, DOI: 10.1080/00397919708003371].

Tellurium dioxide [7446-07-3] TeO₂, M 159.6, m 733°, b 1245°/atm, d²⁵ 5.67g/cm³, n_D²⁰ 1.24. Dissolve the cream coloured solid in 5M NaOH, filter it and precipitate it by adding 10M HNO₃ (CARE) to the filtrate until the solution is acid to phenolphthalein. After decanting the supernatant, the precipitate is washed five times with distilled water, then dried for 24 hours at 110° [Horner & Leonhard *J Am Chem Soc* **74** 3694 1952, DOI: 10.1021/ja01134a508]. It is soluble in acid and alkali.

Terbium (III, IV) oxide [12037-01-3] Tb₄O₇, M 747.7, d²⁵ 7.3g/cm³, pK²⁵ 8.16 (for Tb³⁺ hydrolysis). Dissolve the dark brown-black hygroscopic solid in acid (e.g. perchloric acid), precipitate it as its oxalate and ignite the oxalate at 650°.

Thallium (III) nitrate trihydrate [TTN, thallic nitrate trihydrate, Tl(NO₃)₃·3H₂O] [13453-38-8] Tl(NO₃)₃·3H₂O, M 444.4, m 102-105°, 103°. It is prepared from Tl(III)oxide (45g, [1314-32-5]) by stirring it in hot concentrated HNO₃ (120ml) at ~50° for 30 minutes when all the oxide dissolves to give a clear colourless solution. On cooling to 0°, a white solid crystallises out. It is filtered off, washed with a little cold H₂O, dilute HNO₃, dried in a vacuum desiccator over P₂O₅ to give TTN (75g, 85%) as hard, colourless crystals which should be stored in a cold tightly sealed container. It has reducing properties. Treat carefully as it is **POISONOUS**. [McKillop et al. *J Am Chem Soc* **95** 3635 1973, DOI: 10.1021/ja00792a028.] [For applications see Fieser **4** 492, **5** 656, **6** 578, **7** 362, **8** 476, **9** 460, **10** 395, **12** 481, **14** 302, **16** 326.]

Thallous bromide [7789-40-4] TlBr, M 284.3, m 460°, 480°, b 815°/atm, d²⁵ 7.557g/cm³. Thallous bromide (20g) is purified by refluxing for 2-3 hours with water (200ml) containing 3ml of 47% HBr. It is then washed until acid-free, heated to 300° for 2-3 hours and stored in brown bottles. Its solubility in H₂O (w/w) is 0.034% at 0°, 0.048% at 20°, and 0.204% at 100°. [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 870 1963, for applications see Fieser **2** 405, **3** 286, **5** 655, **7** 361.] **POISONOUS**.

Thallous carbonate [6533-73-9] TlCO₃, M 468.7, m 268-270°, 272°, d²⁵ 7.11g/cm³. It crystallises from hot water (solubility is 5.2g/100ml at 25° and 27.2g/100ml at 100°) on cooling. It is soluble in EtOH, Et₂O and Me₂CO. **POISONOUS**.

Thallous chlorate [13453-30-0] TlClO₃, M 287.8, d₄²⁰ 5.05. It crystallises from hot water (2ml/g) on cooling. Its solubility in H₂O (w/w) is 0.17% at 0°, 0.32% at 20°, and 2.4% at 100°. [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 870 1963]. **POISONOUS**.

Thallous chloride [7791-12-0] TlCl, M 239.8, m 429.9°, b 806°/atm, d²⁵ 7.004g/cm³, n_D²⁰ 1.2.247. Recrystallise it from 1% HCl and wash it until acid-free, or crystallise it from hot water (50ml/g), then dry it at

140° and store it in brown bottles. Also purify it by subliming it in a vacuum, followed by treatment with dry HCl gas and filtering while molten. (It is soluble in 260 parts of cold water and 70 parts of boiling water). It is soluble in EtOH, Me₂CO and NH₄OH. **POISONOUS.**

Thallous hydroxide [12026-06-1] TlOH, M 221.4, m 139°(dec), d²⁵ 7.44g/cm³, pK²⁵ 13.2 (for Tl⁺). Crystallises the yellow solid from hot water (0.6ml/g) on cooling. It is a strong base. **POISONOUS.**

Thallous iodide [7790-30-9] TlI, M 331.3, m 440°, 441.8°, b 824°/atm, d²⁵ 7.29g/cm³. Reflux the yellow solid (red above 170°) for 2-3 hours with water containing HI, then wash it until acid-free, and dry it at 120°. Store it in brown bottles. Almost insoluble in H₂O. [Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 870 1963]. **POISONOUS.**

Thallous nitrate [10102-45-1] TlNO₃, M 266.4, m 206°, b 430°/atm, 450°/atm (dec), d₄²⁰ 5.55. The nitrate crystallises from warm water (1ml/g) on cooling to 0°. [Fieser 12 481.] **POISONOUS.**

Thallous perchlorate [13453-40-2] M 303.8, d²⁰ 4.8g/cm³, pK²⁵ -2.4 to -3.1 (for HClO₄). It crystallises from hot water (0.6ml/g) on cooling. Dry it under vacuum for 12 hours at 100° (protect from possible **EXPLOSION**). [For thermal behavior see Udupa *Thermochim Acta* 16 128 1976, DOI: 10.1016/0040-6031(76)85051-4.]

Thallous sulfate [7446-18-6] Tl₂SO₄, M 504.8, m 633°, d²⁵ 6.77g/cm³, n_D²⁵ 1.860. The sulfate crystallises from hot water (solubility is 2.70g/100ml at 0°, 4.87g/100ml at 20°, and 18.45g/100ml at 100°) by cooling; then dry it under vacuum over P₂O₅. The salt inhibits plant growth. **POISONOUS.** It is the precursor of Tl₂S [1314-97-2] M 440.8, (precipitated by bubbling H₂S through an aqueous solution of the sulfate) which exhibits high electrical conductivity on exposure to infrared light. [For the 'Thalofide Cell—a New Photo-Electric Substance' see Case *Phys Rev* 15 289 1920, DOI: 10.1103/PhysRev.15.289; and for the Case thallous sulfide cell see Kaplan *J Opt Soc Am* 14(3) 251 1927, DOI: 10.1364/JOSA.14.000251.]

Thionyl chloride [7719-09-7] SOCl₂, M 119.0, m -105°, b 74.6°/atm, 77°/atm, d²⁵ 1.638g/cm³, n_D²⁰ 1.517. Crude SOCl₂ can be freed from sulfuryl chloride, sulfur monochloride and sulfur dichloride by refluxing it with sulfur and then fractionally distilling twice. [The SOCl₂ is converted to SO₂ and sulfur chlorides. The S₂Cl₂ (b 135.6°) is left in the residue, whereas SCl₂ (b 59°) passes over in the forerun.] The usual purification is to distil it from quinoline (50g SOCl₂ to 10g quinoline) to remove acid impurities, followed by distillation from boiled linseed oil (50g SOCl₂ to 20g of oil). Precautions must be taken to exclude moisture. Thionyl chloride is used extensively in organic syntheses and can be prepared by distillation of technical SOCl₂ in the presence of *diterpene* (12g/250ml SOCl₂), and avoiding overheating. Further purification is achieved by redistillation from *linseed oil* (1-2%) [Rigby *Chem Ind (London)* 1508 1969]. Gas chromatographically pure material is obtained by distillation from 10% (w/w) triphenyl phosphite [Friedman & Wetter *J Chem Soc (A)* 36 1967, DOI: 10.1039/J19670000036; Larsen et al. *J Am Chem Soc* 108 6950 1986, DOI: 10.1021/ja00282a020]. It is soluble in most aprotic solvents, e.g. toluene, CHCl₃, Et₂O, but reacts with protic solvents, e.g. H₂O, MeOH etc. **HARMFUL VAPOURS.**

Thorium (IV) chloride [10026-08-1] ThCl₄, M 373.8, m 770°, b 921°/atm, d²⁵ 4.59g/cm³, pK₁²⁵ 10.45, pK₂²⁵ 10.62, pK₃²⁵ 10.80, pK₄²⁵ 11.64 (for aquo Th⁴⁺). It is freed from anionic impurities by passing a 2M solution of ThCl₄ in 3M HCl through a Dowex-1 anion-resin column. The eluate is partially evaporated to give crystals which are filtered off, washed with Et₂O and stored in a desiccator over H₂SO₄ to dry. *Alternatively*, a saturated solution of ThCl₄ in 6M HCl is filtered through quartz wool and extracted twice with ethyl, or isopropyl ether (to remove iron), then evaporated to a small volume on a hot plate. (Excess silica precipitates and is filtered off. The filtrate is cooled to 0° and saturated with dry HCl gas.) It is shaken with an equal volume of Et₂O, shaken with HCl gas, until the mixture becomes homogeneous. On standing, ThCl₄·8H₂O precipitates out and is filtered off, washed with Et₂O and dried [Kremer *J Am Chem Soc* 64 1009 1942, DOI: 10.1021/ja01256a505].

Thorium sulfate tetrahydrate [33088-16-3 (4H₂O), 10381-37-0 (9H₂O)] Th(NO₃)₄·4H₂O, M 496.2(4H₂O),

M 424.2 (anhydr), m 42°(loses H₂O), 500°(anhydr, dec), d²⁰ 2.80g/cm³. Crystallise it from water. The solubility of the *decahydrate* increases with increase in temperature, whereas the solubility of the *tetrahydrate* decreases with increase of temperature.

Tin (powder) [7440-31-5] Sn, M 118.7, m 231.9°, b 2603°/atm, d²⁵ 5.769g/cm³(grey, α form, diamagnetic), d²⁵ 7.365g/cm³(white, β form, paramagnetic). The melting point varies with the form of the metal. Tin powder is purified by adding it to about twice its weight of 10% aqueous NaOH and shaking vigorously for 10 minutes. (This removes oxide film and stearic acid or similar material that is sometimes added for pulverisation.) It is then filtered, washed with water until the washings are no longer alkaline to litmus, rinsed with MeOH and dried in air. [Sisido et al. *J Am Chem Soc* **83** 538 1961, DOI: 10.1021/ja01464a008; for applications see Fieser **1** 1168, **2** 414, **13** 298, **17** 333.]

Titanium tetrabromide [7789-68-6] TiBr₄, M 367.5, m 28.3°, 39°, b 233.5°/atm, d²⁵ 3.25g/cm³. Purify it by distillation. The distillate forms light orange hygroscopic crystals. Store it in the dark under N₂ preferably in sealed brown glass ampoules. It is soluble in chlorocarbons and *C₆H₆ but reacts *violently* with H₂O to liberate HBr. [Olsen & Ryan *J Am Chem Soc* **54** 2215 1932, DOI: 10.1021/ja01345a008.] It has been used as a Lewis acid catalyst in organic synthesis.

Titanium tetrachloride [7550-45-0] TiCl₄, M 189.7, m -24°, b 136.4°/atm 154°, d²⁵ 1.726g/cm³, n_D^{10.5} 1.61, pK₁²⁵ 0.3, pK₂²⁵ 1.8, pK₃²⁵ 2.1, pK₄²⁵ 2.4 (for aquo Ti⁴⁺ hydrolysis). Reflux it with mercury or a small amount of pure copper turnings to remove the last traces of colour [due to FeCl₃ and VCl₄], then distil it under N₂ in an all-glass system, taking precautions to exclude moisture. Clabaugh et al. [*J Res Nat Bur Stand* **55** 261 1955] removed organic material by adding aluminium chloride hexahydrate as a slurry with an equal amount of water (the slurry being *ca* one-fiftieth the weight of TiCl₄), refluxed it for 2-6 hours while bubbling in chlorine, the excess of which is subsequently removed by passing a stream of clean dry air. The TiCl₄ is then distilled, refluxed with copper and again distilled, taking precautions to exclude moisture. Volatile impurities are then removed using a technique of freezing, pumping and melting. The *titanium tetrachloride 2-tetrahydrofuran complex*, **TiCl₄ (THF)₂ [31011-57-1] M 333.9**, has **m 126-128°** and is easier to handle than TiCl₄ [Hamilton et al. *J Am Chem Soc* **75** 2881 1953, DOI: 10.1021/ja01108a027; Abrahamson et al. *Organometallics* **3** 1379 1984, DOI: 10.1021/om00087a010; Beilstein **17/1** V 33]. [Baxter & Fertig *J Am Chem Soc* **45** 1228 1923, DOI: 10.1021/ja01658a017; Baxter & Butler *J Am Chem Soc* **48** 3117 1926, DOI: 10.1021/ja01691a014.] It is useful as a Lewis acid in organic synthesis, in the formation of organometallic complexes with useful properties, reacts with H₂O to form a white smoke screen of TiO₂, and with THF it forms yellow crystals of TiCl₄(THF)₂. [For applications see Fieser **1** 1169, **2** 414, **3** 291, **4** 507 **5** 671, **6** 590, **7** 370, **8** 483, **9** 468, **10** 401, **11** 529, **11** 261, **12** 494, **13** 304, **14** 309, **15** 317, **16** 332, **17** 344.] It is also available as a 0.09M solution in 10% HCl; a 1.0M solution in CH₂Cl₂; 1.0M in toluene; and a TiCl₄.2THF complex [31011-57-1] M 333.9, m 126-128°. A **low valent titanium reagent** which can be prepared *in situ* by adding Mg powder (25mmol, 50 mesh) to TiCl₄ (15mmol) in THF (60ml) and activated by addition of 1,2-dibromomethane (10mmol) during 5 minutes at 0° under N₂. This reagent was shown to reduced Schiff's bases as well as dimerise them to provide imidazolines in moderate to good yields with some control in the stereochemistry [Periasamy et al. *Tetrahedron Lett* **37** 4767 1996, DOI: 10.1016/0040-4039(96)00930-6], and low valent titanium for carbonyl coupling [McMurry *Chem Rev* **89** 1513 1989, DOI: 10.1021/cr00097a007]. TiCl₄ vapours are **HARMFUL** and head and body protection should be used.

Titanium trichloride [7705-07-9] TiCl₃, M 154.3, m 425°(dec), 440°(dec), b 960°/atm (dec), d²⁵ 2.64g/cm³, n_D²⁵ 1.4854, pK₁²⁵ 2.55 (for hydrolysis of Ti³⁺ to TiOH²⁺). It is a brown purple powder that is very reactive to H₂O and pyrophoric when dry. It should be manipulated in a dry box. It is soluble in CH₂Cl₂ and tetrahydrofuran, and is used as a M solution in these solvents in the ratio of 2:1, and stored under N₂. It is a powerful reducing agent. [Ingraham et al. *Inorg Synth* **6** 52 1960, DOI: 10.1002/9780470132371.ch16.] In alcoholic solvent, with *tert*-butylhydroperoxide it mediates the reductive coupling of aryl aldehydes to form 1,2-diols [Clerici et al. *Eur J Org Chem* 4050 2007, DOI: 10.1002/ejoc.200700252]. [Fieser **2** 415, **4** 506, **5** 669, **6** 587, **7** 369, **8** 482, **9** 467, **10** 400, **11** 529, **11** 529, **12** 492, **13** 302, **15** 316, **16** 330.] Also commercially available

is a ~10wt% in 20-30wt% in hydrochloric acid for the quantitative titration of NO₂ groups to NH₂ groups.

Titanium trichloride—aluminium chloride [12003-13-3] **3:1 (TiCl₃)₃.AlCl₃ complex, M 596.0, m >250°**, which is a useful reducing agent for catalysis. A 1:3 complex **TiCl₃/3THF** [18039-90-2] **M 370.5, >250°**, for making bis(β-diketone) ligands of bimetallic Ti(III) complexes with a triple-helix structures is commercially available [Grillo et al. *JCS Chem Commun* 1561 1997, DOI: 10.1039/A702971C].

Titanyl (IV) sulfate [13825-74-6] **TiOSO₄. 2H₂O, M 160.0, d²⁵ 1.361g/cm³(~15 wt% in dil H₂SO₄)**. Dissolve it in water, filter and crystallise it three times from boiling 45% H₂SO₄, washing with EtOH to remove excess acid, then with Et₂O. Dry it in air for several hours, then in an oven at 105-110°. [Hixson & Fredrickson *Ind Eng Chem* **37** 678 1945, DOI: 10.1021/ie50427a021.]

Trisodium orthophosphate dodecahydrate [10101-89-0] **Na₃PO₄. 12H₂O, M 380.1, m 75°(dec), b 100°/atm, (12H₂O, dec), d²⁵ 1.62g/cm³ (12H₂O), pK₁²⁵ 2.15, pK₂²⁵ 7.21, pK₃²⁵ 12.33 (for H₃PO₄)**. It crystallises from warm dilute aqueous NaOH (1ml/g) on cooling to 0°. The solubility of the *dodecahydrate* in H₂O is 25.8g/100ml at 25°, and the *anhydrous* salt is 5.4g/100ml at 0°, 14.5g/100ml at 25°, 23.3g/100ml at 40°, and 94.6g/100ml at 100°.

Tritium [10028-17-8] **³H₂, M 6.0**. Purify tritium from hydrocarbons and ³H by diffusion through the wall of a hot nickel tube, as well as diffusion of H₂ and D₂ [Landecker & Gray *Rev Sci Instrum* **25** 1151 1954, <http://dx.doi.org/10.1063/1.1770969>]. Its half life is 12.32 years. **RADIOACTIVE**.

Tungsten (rod) [7440-33-7] **W, M 183.8, m 3410°, 3695°, b 5900°/atm, 6203°/atm, d²⁵ 19.25g/cm³**. Clean the solid with concentrated NaOH solution, rub it with very fine emery paper until its surface is bright, wash it with previously boiled and cooled conductivity water and dry it with filter paper. [Hein & Herzog in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1417 1965.] It has many radioactive isotopes.

Tungsten (VI) trichloride (tungsten hexachloride) [13283-01-7] **WCl₆, M 396.6, m 265°(dec), 275°, b 346.7°/atm, d²⁵ 3.52g/cm³, pK₁²⁵ 2.20, pK₂²⁵ 3.70 (for tungstic acid, H₂WO₄)**. Sublime the dark blue solid in a stream of Cl₂ in a high temperature furnace and collect it in a receiver cooled in a Dry-ice/acetone bath in an inert atmosphere because it is sensitive to moisture. At low temperatures its colour becomes red-wine. The red form can be produced also by condensing the vapour, but reverts to the blue-black form on gentle heating. It is soluble in CS₂, CCl₄, CHCl₃, POCl₃, *C₆H₆, petroleum ether and Me₂CO. Its solutions decompose on standing. Good crystals can be obtained by heating WCl₆ in CCl₄ to 100° in a sealed tube, followed by slow cooling (tablets of four-sided prisms). Store it in a desiccator over H₂SO₄ in the dark as it [Leitzke et al. *Inorg Synth* **3** 163 1950, DOI: 10.1002/9780470132340.ch44; Porterfield et al. *Inorg Synth* **9** 133 1967, DOI: 10.1002/9780470132401.ch36; Hein & Herzog in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1420 1965.] The trichloride mediates the conversion of 1,3-dithiolanes and 1,3-dithianes to chlorinated dihydro-1,4-dithiins and dihydro-1,4-dithiepinines [Firouzbadi et al. *Synlett* 413 1999, DOI: 10.1055/s-1999-2628], and is a precursor to tungsten-based carbenes which catalyse the olefin metathesis reactions of 1- and 2-octene [Düz et al. *Tetrahedron Lett* **47** 5167 2006, DOI: 10.1016/j.tetlet.2006.05.054].

Uranium hexafluoride [7783-81-5] **UF₆, M 352.0, m 64.0°, 64.8° (sealed tube??), b 0°/17.4mm, 56.2°/765mm(sublimes), d²⁵ 5.02g/cm³, pK²⁵ 1.68 (for hydrolysis of U⁴⁺ to UOH³⁺)**. Purify uranium hexafluoride by fractional distillation to remove HF. Also purify it by low-temperature trap-to-trap distillation over pre-dried NaF, and it sublimes at 56.5°/atm. [Anderson & Winfield *JCS Dalton Trans* 337 1986, DOI: 10.1039/DT9860000337].

Uranium trioxide [1344-58-7] **UO₃, M 286.0, m ~200-650°(dec), d²⁵ 5.5—8.7g/cm³**. The yellow-orange oxide is dissolved in HClO₄ (to give a uranium content of 5%), and the solution is adjusted to pH 2 by addition of dilute ammonia. Dropwise addition of 30% H₂O₂, with rapid stirring, precipitated U(VI) peroxide, the pH being held constant during the precipitation, by addition of small amounts of the ammonia solution. Then H₂O₂

is added until further quantities caused no change in pH. After stirring for 1 hour, the slurry is filtered through coarse filter paper in a Büchner funnel, washed with 1% H₂O₂ acidified to pH 2 with HClO₄, then heated at 350° for three days in a large platinum dish [Baes *J Phys Chem* **60** 878 1956, DOI: 10.1021/j150541a011]. It is **POISONOUS** and **RADIOACTIVE** and should not be inhaled, ingested or be in contact with the skin — take extreme precautions.

Uranyl nitrate (UO₂(NO₃)₂ · 6H₂O) [13520-83-7, 10102-06-4] UO₂(UO₃)₂ · 6H₂O, M 394.0, 502.1, m 60.2°, b 118°/atm (dec), d²⁵ 2.81g/cm³, pK²⁵ 5.82 (for aquo UO₂²⁺). Crystallise the nitrate from water by cooling to -5°, taking only the middle fraction of the solid which separates. Its solubility in H₂O is 98g/100g at 0° and 122g/100g at 20°, and 474g/100g at 100°, and is soluble in tributyl phosphate. Dry the *deliquescent* rhombic yellow crystals of the *hexahydrate* over 35-40% H₂SO₄ in a vacuum desiccator. The crystals reflect a greenish lustre. They are remarkable because on crushing, rubbing or shaking they show *triboluminescence* with occasional detonation. They are very soluble in EtOH, and solutions of the nitrate in Et₂O can explode in the presence of sunlight. **HIGHLY TOXIC** see previous entry.

Vanadium (metal) [7440-62-2] V, M 50.9, m 1910°, b 3407°/atm, d²⁵ 6.0g/cm³(solid), 5.5g/cm³(melt). Clean the blue-silver-grey metal by rapid exposure consecutively to HNO₃, HCl, HF, de-ionised water and reagent grade acetone, then dry it in a vacuum desiccator. [Brauer in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol II pp 1252-1255 1965.] Used in hardening steel, trace metal in some enzymes, e.g. vanadium nitrogenases and haloperoxidases contain vanadium.

Vanadyl trichloride (VOCl₃) [7727-18-6] VOCl₃, M 173.3, m -79.5°, b 124.5-125.5°/744mm, 127.16°/760mm, d⁰ 1.854g/cm³, d³² 1.826 g/cm³, d³² 1.811 g/cm³. VOCl₃ should be lemon yellow in colour. If it is red, it may contain VCl₄ and Cl₂. Fractionally distil it, and then redistil it over metallic Na, but be careful to leave some residue because the residue can become **EXPLOSIVE** in the presence of the metal. **USE A SAFETY SHIELD** and avoid contact with moisture. It readily hydrolyses to vanadic acid and HCl. Store it in a tightly closed container or in sealed ampoules under N₂. [Brown et al. *Inorg Synth* **1** 106 1939, DOI: 10.1002/9780470132326.ch38; Brown et al. *Inorg Synth* **4** 80 1953; DOI: 10.1002/9780470132357.ch28; Oppermann 'Untersuchungen an Vanadinoxidchloriden und Vanadinchloriden. I. Gleichgewichte mit VOCl₃, VO₂Cl und VOCl₂'. *Z Anorg Allgem Chem* **351** 113 1967, DOI: 10.1002/zaac.19673510302.]

Water [7732-18-5] H₂O, M 18.0, m 0°, b 100°/atm, d^{3.98} 1.000000g/ml (0.999972g/cm³, n_D²⁰ 1.33335, pK²⁵ 14.00. Conductivity water (specific conductance ca 10⁻⁷ mho) can be obtained by distilling water in a steam-heated tin-lined still, then, after adding 0.25% of solid NaOH and 0.05% of KMnO₄, distilling once more from an electrically heated Barnstead-type still, taking the middle fraction into a Jena glass bottle. During these operations suitable traps must be used to protect against entry of CO₂ and NH₃. Water, only a little less satisfactory for conductivity measurements (but containing traces of organic material) can be obtained by passing ordinary distilled water through a mixed bed ion-exchange column containing, for example, Amberlite resins IR 120 (cation exchange) and IRA 400 (anion exchange), or Amberlite MB-1. This treatment is also a convenient one for removing traces of heavy metals. (The metals Cu, Zn, Pb, Cd and Hg can be tested for by adding pure concentrated ammonia to 10ml of sample and shaking vigorously with 1.2ml of 0.001% dithizone in CCl₄. Less than 0.1μg of metal ion will impart a faint colour to the CCl₄ layer.) For almost all laboratory purposes, simple distillation yields water of adequate purity, and most of the volatile contaminants such as ammonia and CO₂ are removed if the first fraction of distillate is discarded. Most laboratories have glass stills that 'doubly' or 'triple' distil water for chemical and biological purposes. [See 'water' in Chapter 1.] Water also has d²⁵ 0.997, d₄⁰ 0.999868 (liq) and d⁰ (ice) 0/917g/cm³.

Wood's Metal [76093-98-6] Bi,Pb,Sn,Cd, M ~1486, m ~75°, 73-77° (low melting granular) It is a fusible alloy of Bi,Pb,Sn,Cd with the atomic composition of 4:2:1:1 (m 71°) It is commonly used as a metal bath for heating reaction flasks as it melts at a convenient temperature. Its temperature rises rather rapidly when heated with a Bunsen burner in a crucible (e.g. steel or nickel), so care should be taken when heating a reaction or distilling flask containing chemical reactants immersed into it. Heating should be applied slowly, and the temp-

erature should be monitored continuously with a thermometer immersed in the molten metal. The temperature of the metal can be taken up to $\sim 350^\circ$, and it is better to work in an efficient fume cupboard as the vapours are toxic. At higher temperature some oxidation of the metal takes place. A bath that has been heated at high temperatures and then the metal cooled back to the solid, and melted again has a solid scum of the oxides on its surface. These can be easily removed with a nickel spatula to leave a nice clean surface. The melting point of the re-used alloy is not seriously affected, and the alloy can be used over and over again without considerable loss in weight. It is commercially available in a *low melting granular* form, or as *sticks* with the composition of Bi(50 wt%), Pb (25 wt%), Sn (12.5 wt%) and Cd (12.5 wt%). Other such alloys are **Rose's metal** (**m** 93.75°) with the composition of Bi (50%), Pb (25-28%), Sn (22-25%), i.e. in the ratio of *ca* 2:1:1; **Lipowitz alloy** [**60° (soft) to 70° (melt)**] with the composition of Bi, Pb, Sn, Cd in the ratio 15:8:4:3; **Cerrolow alloy** (**m** 47.2°) with the composition Bi (44.7%), Pb (22.6%), Sn (8.3%) In (19.1%) and Cd (5.3%) but they are not very commonly used in the laboratory probably because they contain Cd. Alloys of Bi, Pb and Sn which melt slightly above 100° have been used as melting plugs in sprinkler alarms.

Zinc (dust) [7440-66-6] **Zn**, **M 65.4**. **m** 419.5° , **m** 420° , **b** $907^\circ/\text{atm}$, **d** $^{25} 7.14\text{g}/\text{cm}^3$, **d** $^{420} 6.57\text{g}/\text{cm}^3$ (**melt**), **n** $^{25}_{\text{D}}$ **1.4854**. Commercial zinc dust (1.2kg) is stirred with 2% HCl (3L) for 1 minute, then the acid is removed by filtration, and washed in a 4L beaker with a 3L portion of 2% HCl, three 1L portions of distilled water, two 2L portions of 95% EtOH, and finally with 2L of absolute Et₂O. (Wash solutions were removed by filtration.) The material is dried thoroughly, and break lumps in a mortar. [Wagenknecht & Juza *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1067 1965.] Or, fuse it *in vacuo*, cool, and wash it with acid to remove any oxide. Used in galvanising, in alloys, batteries, brass and bronze production, mineral supplements, zinc ointments; as antibacterial, and is an efficient Lewis acid. **Zinc nano metallic powder** is available in 75-125nm p.size with high surface area. It is very reactive and forms organo-Zn compounds with halides and used in azo couplings. Sonicate to de-agglomerate to 35nm size. ZnNP is available in <50nm p.size.

Zinc bromide [7699-45-8, 18921-13-6 ($2\text{H}_2\text{O}$)] **ZnBr₂**, **M 225.2(anhydr)**, **m** 384° , 394° (**anhydr**), **b** $697^\circ/\text{atm}$, **d** $^{25} 4.22\text{g}/\text{cm}^3$, **n** $^{25}_{\text{D}}$ **1.5452**. Heat ZnBr₂ to 300° under vacuum (2×10^{-2} mm) for 1 hour, then sublime it. Its solubility in H₂O is 311g/100ml at 0° , 447g/100ml at 20° , and 538g/100ml at 100° ; and it is soluble in EtOH, Et₂O, Me₂CO and THF. [Wagenknecht & Juza *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1072 1965; Fieser **2** 463, **5** 762, **8** 535, **9** 520, **10** 461, **11** 600, **13** 349, **15** 368, **16** 389.] The **anhydrous salt** is a Lewis acid which catalyses stereo and region selective reactions.

Zinc chloride [7646-85-7] **ZnCl₂**, **M 136.3(anhydr)**, **m** 283° , 290° , 293° , **b** $732^\circ/\text{atm}$, **d** $^{25} 2.907\text{g}/\text{cm}^3$, **n** $^{25}_{\text{D}}$ **1.5452**. The anhydrous white material can be sublimed under a stream of dry HCl, followed by heating to 400° in a stream of dry N₂. It sublimes at high vacuum. Also purify it by refluxing (50g) in dioxane (400ml) with 5g zinc dust, filtering hot and cooling to precipitate ZnCl₂. Crystallise it from dioxane and store it in a desiccator over P₂O₅. It has also been dried by refluxing in thionyl chloride. [Weberg et al. *J Am Chem Soc* **108** 6242 1986, DOI: 10.1021/ja00280a022.] *Hygroscopic: minimal exposure to the atmosphere is necessary.* However, its solubility in H₂O is 43.2g/100ml at 25° , and is soluble in EtOH (43g/100ml), Me₂CO and glycerol. [Wagenknecht & Juza *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1070 1965, Fieser **1** 1289, **2** 464, **3** 338, **5** 763, **6** 676, **8** 536, **9** 522, **10** 461, **11** 601, **12** 574, **13** 349, **15** 368, **16** 391.] Also available commercially as a 1M solution in Et₂O (**d** $^{25} 0.835\text{g}/\text{ml}$).

Zinc-Copper Couple [53801-63-1] **Zn/Cu**, **m** $419-420^\circ$. It is available in the form of a powder made of Zn and Cu. If the Zn contains some ZnO, then it should be treated with dilute HCl for a very short period, washed H₂O, a little EtOH then Et₂O, and dried *in vacuo* (see also zinc dust above). The **couple** can be prepared in either of the following ways. (1) Zinc dust [(120g, MW of Zn is 65.4) and powdered copper oxide (10g, MW of Cu is 63.6) in a round-bottomed flask ($\sim 200\text{ml}$) are heated gently over a flame in a current of H₂ and N₂ with stirring or rotating the flask until the CuO is reduced, and a uniformly grey mixture is obtained. The temperature should be maintained below the point of fusion during heating ($\sim 500^\circ$) and the mixture is stored in a desiccator. (2) Zinc powder (24.6g) is stirred with 3% aqueous HCl (20ml) for 1 minute, washed by decantation with 3% aqueous HCl (3 x 20ml), with H₂O (5 x 20ml), 2% aqueous CuSO₄ (2 x 40ml), distilled H₂O (5 x 20ml), with EtOH (4 x 20ml), with Et₂O (2 x 20ml), filtered off on a Büchner funnel, sucked dry then stored in a desiccator over P₂O₅.

overnight. [Corbin et al. *Org Synth Coll Vol* **5** 328 1973; DOI: 10.15227/orgsyn.044.0030 Smith & Simmons *Org Synth Coll Vol* **5** 855 1973, DOI: 10.15227/orgsyn.041.0072.] (3) A suitable zinc/copper alloy with 5–8% of Cu can be prepared by melting zinc (m 419.5°) with clean brass turnings (*ca* 2Cu/1Zn) and casting into bars, which can be turned into fine shavings prior to use. [Noller *Org Synth Coll Vol* **2** 184 1943, DOI: 10.15227/orgsyn.012.0086.] Note that the couple loses activity in moist air, so it should be kept dry, and used as soon as possible after it is prepared for best results. When it is required to reduce organic halides to the respective hydrocarbon or deuterated derivative, it is best to prepare the Zn-Cu couple in an oxygen-free system. Thus Zn dust (6.5g, 100mmol) suspended in H₂O (10ml) is vigorously stirred with acidic cupric chloride solution (15ml of 0.15M in 5% hydrochloric acid) until evolution of gas ceases, the black solid is filtered off, and washed with H₂O until the filtrate gives a negative chloride test with 6% AgNO₃ solution. The **couple** is washed twice with Me₂CO, and in order to obtain the highest deuterium incorporation, the Me₂CO wash should be followed by a D₂O wash, two Me₂CO washes, two Et₂O washes and dried *in vacuo* at ~25°. This preparation of the **couple** is exceptional for replacing the halogen in a variety of organic halides by H or D in water-containing ether solvents, under mild conditions, and in reproducibly moderate to high yields (e.g. 1,4-dideuterobutadiene from 1,4-dichloro-1,3-butadiene, or 2-butanone-3-*d*₁ from 3-bromo-2-butanone). The deuterium derivatives provide a convenient ¹³C NMR method for deuterium analysis. [Stephenson et al. *J Org Chem* **42** 212 1977, DOI: 10.1021/jo00422a006.]

Zinc cyanide [557-21-1] Zn(CN)₂, **M 117.4, m 800°(dec), d²⁵ 1.852g/ml**. It is a **POISONOUS** white powder which becomes black on standing if Mg(OH)₂ and carbonate are not removed in the preparation. Thus, wash it well with H₂O, then well with EtOH, Et₂O and dry it in air at 50°. Analyse it by titrating the cyanide with standard AgNO₃. Other likely impurities are ZnCl₂, MgCl₂ and traces of basic zinc cyanide; the first two salts can be washed out. It is soluble in aqueous KCN solutions. However, if purified in this way Zn(CN)₂ is not reactive in the Gattermann synthesis. For this, the salt should contain at least 0.33 mols of KCl or NaCl which will allow the reaction to proceed faster. Its solubility in H₂O is 0.00005g/100ml at 20°, and it reacts with alkalis, KCN and NH₄OH. [Adams & Levine *J Am Chem Soc* **45** 2373 1923, DOI: 10.1021/ja01663a020; Arnold & Sprung *J Am Chem Soc* **60** 1699 1938, DOI: 10.1021/ja01274a501; Fuson et al. *Org Synth Coll Vol* **3** 549 1955, DOI: 10.15227/orgsyn.023.0057; Fieser **1** 1293, *Beilstein* **2** III 86.]

Zinc fluoride [7783-49-5, 13986-18-0 (4H₂O)] ZnF₂, **M 103.4(anhydr), m 872°(anhydr), 100°(anhydr, dec), b 1500°/atm, d²⁵ 4.95g/ml**. A possible impurity is H₂O which can be removed by heating at 100° or by heating to 800° in a dry atmosphere. Heating in the presence of NH₄F produces larger crystals. It is sparingly soluble in H₂O (*anhydrous* 0.00005g/100ml at 20°; *tetrahydrate* 1.52g/100ml at 20°), but is slightly more soluble in HCl, HNO₃ and NH₄OH. It can be stored in glass bottles. [Kwasnik in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 242 1963, for application see Fieser **6** 514.]

Zinc iodide [1314-13-2] ZnI₂, **M 319.2, m 445°, 446°, b 1150°(dec), d²⁵ 4.74g/ml**. Heat the iodide to 300°/2 x 10⁻²mm) for 1 hour, then sublime it. Its solubility in H₂O is 450g/100ml at 20°, and it is soluble in EtOH. Store it in the dark. It is an X-ray opaque material in radiology, electrolyte and with OsO₄ is a stain in electron microscopy. [For applications see Fieser **1** 1294, **5** 765, **10** 462, **11** 604, **12** 576, **13** 350.]

Zinc oxide [10139-47-6] ZnO, **M 81.4 m 1975°(dec), b 1975°(dec), d²⁵ 5.606g/ml, n_D²⁵ 2.0041**. Zinc oxide, obtained by burning zinc vapour (at > 907°) in air with emission of bright luminescence and lowering temperature, is called **zinc white** and used as a pigment. For pharmaceutical purposes it is prepared by treating an aqueous solution of ZnSO₄ with Na₂CO₃, and heating the basic zinc carbonate which is precipitated. When pure salts are used then pure ZnO thus obtained is used. The oxide is white and is **thermochromic** as it changes colour to yellow on heating reversibly, i.e. changes back to white on cooling. This is apparently due to a very small loss of some oxygen at *ca* 800° which is re-absorbed on cooling. The oxide is **amphoteric**: dissolving in acids to form **salts** containing Zn⁺ or Zn²⁺ ions and in alkalis to form **zincates** containing [ZnO₂]²⁻ ions. The oxide has numerous applications which include pigments (paints), ceramic industry, rubber manufacture, food additives, pharmaceutical industry, UV absorbers, coatings, corrosion prevention electronics, and sensors among many applications. [De Liedekerke '2.3. Zinc Oxide (Zinc White): Pigments, Inorganic, 1' in *Ullmann's Encyclopedia of Industrial Chemistry* 2006 Wiley-VCH, Weinheim, DOI:10.1002/14356007.a20_243.pub2; Özgür et al. A comprehensive review of ZnO materials and devices *J Appl Phys* **98** 041301 2005, DOI: 10.1063/

1.1992666; Klingshirn 'ZnO: Material, Physics and Applications' *Chem Phys Chem* **8** 782 2007, DOI: 10.1002/cphc.200700002.] [For applications see Fieser **1** 1294, **5** 622.] The oxide is also used to prepare *NaZnSiO₃OH* which is a novel chiral zincosilicate framework material with potential application in ion exchange for adsorption or catalysis [Healey et al. *Inorg Chem* **38** 455 1999, DOI: 10.1021/ic9810135].

Zinc oxide hydrate [55204-38-1] **ZnO. xH₂O, M 81.4 (anhyd), m >390°, d²⁵ 3.05g/ml**, a *hydrated form* of the oxide, is also available commercially.

Zinc perchlorate hexahydrate [13637-61-1 (anhydrous), 10025-64-6 (6H₂O)] **Zn(ClO₄)₂. 6H₂O, M 372.4, m 105-107°, 106°, b 200°/atm, d²⁵ 2.252g/cm³, n_D²⁵ 1.50, pK²⁵ -2.4 to -3.1 (for HClO₄)**. Crystallise the white solid from a small volume of H₂O. It is soluble in EtOH. Useful catalyst for epoxide ring opening by amines [Shivani & Chakraborti *J Org Chem* **72** 3713 2007, DOI: 10.1021/jo062674j] **Corrosive and Potentially EXPLOSIVE**.

Zinc-Silver Couple [37350-66-6] **Zn/Ag**. It is best to prepare the couple fresh when used to replace a vinylic bromine atom in organic compounds by a hydrogen atom, deuterium atom or a tritium atom. The zinc dust to be used should first be cleaned from any zinc oxide or greasy material. Zinc dust (22g) is stirred under N₂ at ~25° as a suspension in 10% aqueous HCl (110ml) for 20 minutes. Acetone (50ml) is added to the mixture, the liquid is decanted from the metal, and the grey slurry is washed thoroughly with Me₂CO (6 x 50ml) by decantation. The residual Zn is treated with a boiling suspension of AgOAc (~6.6g) in AcOH (125ml), stirred rapidly without further heating, the liquid is decanted off and the black granular solid that is formed is washed with AcOH (2 x 60ml), Me₂CO (2 x 20ml), and finally dry THF (2 x 50ml). Store it under N₂ in the dark. If it is used for deuterium replacement of bromine it is best to stir the couple with D₂O (5ml) in dry THF (50ml) for 15 minutes, decant, repeat the process a second time and then wash the solid with dry THF (50ml). This reduces the amount of proton incorporation while maximising deuterium incorporation. [Fryzuk & Bosnich *J Am Chem Soc* **101** 3043 1979, DOI: 10.1021/ja00505a035; Clark & Heathcock *J Org Chem* **41** 636 1976, DOI: 10.1021/jo00866a012.] It is a mild and convenient reagent for reductive dehalogenation of aryl halides to arenes [Chung et al. *Synth Commun* **18**(5) 507 1988, DOI: 10.1080/00397918808060744].

Zinc sulfate heptahydrate (white vitriol) [7446-20-0 (7H₂O), 13786-24-8 (6H₂O), 7446-19-7 (H₂O), 7733-02-0 (anhydrous)] **ZnSO₄. 7H₂O, M 287.5(7H₂O), 161.5(anhydr), m 100°(7H₂O, dec), 280°(loses all 7H₂O), >680°(anhydrous), d²⁵ 3.54g/cm³ (anhydr), d²⁵ 2.072g/cm³ (7H₂O), n_D²⁵ 1.658(anhydr), n_D²⁵ 1.4357(7H₂O)**. Crystallise it from aqueous EtOH or dilute H₂SO₄ below 39° when it forms the *heptahydrate*, and between 39° and 70° it forms the *hexahydrate*, and above 70° the *monohydrate* is stable. The *anhydrous salt* is obtained from the hydrates by heating at 280° or lower temperatures in a current of dry air. It decomposes to ZnO and SO₂ at 767°. The solubility of the *heptahydrate* in H₂O is 5.88% at 0°, 61.92% at 30°, 66.61% at 35° and 70.05% at 39°. [Rohe & Wolf 'Zinc Compounds' in *Ullmann's Encyclopedia of Industrial Chemistry* 2005, Wiley-VCH, Weinheim. DOI: 10.1002/14356007.a28 537].

Zirconium tetrachloride [10026-11-6] **ZrCl₄, M 233.0, m >300°(sublimes), d²⁵ 2.80g/cm³, pK₁²⁵ -0.32, pK₂²⁵ 0.06, pK₃²⁵ 0.35, pK₄²⁵ 0.46 (for hydrolysis of aquo Zr⁴⁺)**. Crystallise the white solid repeatedly from concentrated HCl. It is hydrolysed by H₂O to form white ZrOCl₂ (see below) and HCl. [Krebs *Z Anorg Allgem Chem* **378** 263 1970, DOI: 10.1002/zaac.19703780306; Bora 'Zirconium Tetrachloride' *Synlett* (7) 1073 2003, DOI: 10.1055/s-2003-39323.]

Zirconyl chloride hexahydrate [7699-43-6 (30% in HCl), 15461-27-5 (xH₂O)] **ZrOCl₂. 6H₂O, M 286.2, m 150°(loses 6H₂O)**. Crystallise it repeatedly from 8M HCl to give ZrOCl₂.8H₂O (see below). On drying, ZrOCl₂.6H₂O, **m 150°**, is formed. The product is not free from hafnium. [Blumenthal *J Chem Ed* **39** 604 1962, DOI: 10.1021/ed039p604.] **Zirconyl chloride octahydrate** [13520-92-8] **ZrOCl₂. 8H₂O, M 322.3, m 150°(loses 6H₂O), 210°(loses all H₂O), 400°(anhydrous dec), d²⁵ 1.91g/cm³**. Recrystallise the chloride several times from water [Ferragina et al. *JCS Dalton Trans* 265 1986, DOI: 10.1039/DT9860000265]. Recrystallisation from 8M HCl gives the *octahydrate* as white needles on concentrating. It is also formed by hydrolysing ZrCl₄ with water. After one recrystallisation from H₂O, 99+% grade zirconyl chloride had Ag, Al, As, Cd, Cu, Hf, Mg, Na, Sc and V at 20, 1.8, 0.6, 0.6, 0.4, 8.4, 0.4, 2.4, 80 and 3 ppm, respectively. (See above.) [For structure see Mak 'Refinement of the crystal structure of zirconyl chloride octahydrate' *Can J Chem* **46**, 3491 1968, DOI: 10.1139/v68-579.]

METAL-ORGANIC COMPOUNDS

This section contains metal-organic compounds, ammonium and metal derivatives of organic alcohols, amines and carboxylic acids (salts), as well as ionophores that form complexes with metal ions. Note that there are a large number of metal-organic catalysts, and reagents for preparing some of these catalysts, in Chapter 5, that can be considered as an extension of this section. (For Introduction to this section see p 635.)

Acetylenedicarboxylic acid monopotassium salt [928-04-1] $\text{C}_4\text{HO}_4\text{K}$, **M 152.2**, **m 133°**. The white powder is very soluble in H_2O , but can be crystallised from a small volume of H_2O in small crystals. These are washed with EtOH and dried over H_2SO_4 at 125°. [Bandrowski *Chem Ber* **10** 838 1877, DOI: 10.1002/cber.187701001231; Lossen *Justus Liebigs Ann Chem* **272** 127 1893, DOI: 10.1002/jlac.18932720202; *Beilstein* **2** H 801, **2** I 317, **2** II 670, **2** III 1991, **2** IV 2290.]

Acetylferrocene (ferrocenyl methylketone) [1271-55-2] $\text{C}_{12}\text{H}_{12}\text{OFe}$, **M 228.1**, **m 86°, 86-87°, b 161-163°/4mm, 369.2-373.5°/760mm**. Orange-red (red-brown) crystals are obtained when it is recrystallised from isooctane or $^*\text{C}_6\text{H}_6$, and then sublimed at 100°/1mm. Insoluble in H_2O , but soluble in most organic solvents. The *oxime* has **m 167-170°** (from Et_2O or aqueous EtOH). The *semicarbazone* has **m 198-201°** (from EtOH). [Richmond & Freiser *J Am Chem Soc* **77** 2022 1955, DOI: 10.1021/ja01612a100; Weinmayr *J Am Chem Soc* **77** 3009 1955, DOI: 10.1021/ja01616a026; Broadhead et al. *J Chem Soc* 650 1958, DOI: 10.1039/JR9580000650; Donahue & Donahue 'Beyond Acetylferrocene: The Synthesis and NMR Spectra of a Series of Alkanoylferrocene Derivatives' *J Chem Edu* **90** 1688 2013, DOI: 10.1021/ed300544n; *Beilstein* **16** IV 1798.]

Allylpalladium(II) chloride dimer [12012-95-2] $\text{C}_6\text{H}_{10}\text{Cl}_2\text{Pd}_2$, **M 365.9**, **m 120°(dec), ~160°, 155-156°(dec)**. Recrystallise the pale yellow solid from benzene. It is soluble in MeOH, Et_2O and CHCl_3 . [Hüttel et al. *Chem Ber* **94** 766 1961, DOI: 10.1002/cber.19610940329; Dent et al. *J Chem Soc* 1585 1964, DOI: 10.1039/JR9640001585; Armstrong *J Org Chem* **31** 618 1966, DOI: 10.1021/jo01340a522; Tatsuno et al. '(η^3 -allyl)palladium(II) Complexes' *Inorg Synth* **28** 342 1990, DOI: 10.1002/9780470132593.ch88; for applications see Feiser **2** 109, **5** 189, **7** 5, **9** 132.] Useful catalyst for efficient and accelerated Suzuki cross-coupling reactions [Wallow & Novak *J Org Chem* **59** 5034 1994, DOI: 10.1021/jo00096a056], Heck coupling of aryl halides with simple olefins [Fall et al. *Synthesis* 1683 2007, DOI: 10.1055/s-2007-966063], and enol ethers with aryl bromides [Battace et al. *Eur J Org Chem* 3122 2007, DOI: 10.1002/ejoc.200700152].

Allyl tri-*n*-butylstannane (allyl tributyl tin) [24850-33-7] $\text{C}_{15}\text{H}_{33}\text{Sn}$, **M 331.1**, **b 88-92°/0.2mm, 115°/17mm, d²⁰ 1.068g/ml, n_D²⁰ 1.487**. A possible impurity is tributylchlorostannane — test for Cl as Cl ion after hydrolysing. Dissolve it in $^*\text{C}_6\text{H}_6$ (or toluene), shake this with dilute aqueous NaOH, dry (CaCl_2), filter, evaporate and distil the residue in a vacuum [Jones et al. *J Chem Soc* 1446 1947, DOI: 10.1039/JR9470001446; Bristow *Aldrichimica Acta* **17** 75 1984, Yamamoto *Aldrichimica Acta* **20** 45 1987]. [*Beilstein* **4** IV 4317; for applications see Fieser **9** 8, **11** 15, **12** 21, **13** 10, **14** 14, **16** 7, **17** 12.] Used in an efficient synthesis of homoallylic alcohols and amines from 2,4,6-trichloro-1,3,5-triazine [Das et al. *Tetrahedron Lett* **47** 9103 2006, DOI: 10.1016/j.tetlet.2006.10.068].

Aluminum acetylacetonate (tris[2,4-pentandionate]aluminium, Al(acac)₃) [13963-57-0] $\text{C}_{15}\text{H}_{21}\text{O}_6\text{Al}$, **M 324.3**, **m 192-194°, 195°, b 315°/atm, d²⁰ 1.42g/ml**. Recrystallise the white solid several times from $^*\text{benzene}$ or aqueous MeOH, λ_{max} 216 and 286nm. [Charles & Pawlikowski *J Phys Chem* **62** 440 1958, DOI: 10.1021/j150562a017.] It can be purified by sublimation and has the following solubilities in g percent: $^*\text{C}_6\text{H}_6$ 35.9 (20°), 47.6 (40°), toluene 15.9 (20°), 22.0 (40°) and acetylacetone 6.6 (20°), 10.4 (40°). [Fernelius & Bryant *Inorg Synth* **5** 105 1957, DOI: 10.1002/9780470132364.ch29; *Beilstein* **1** IV 3668.]

Aluminum ethoxide [555-75-9] $\text{C}_6\text{H}_{15}\text{O}_3\text{Al}$, **M 162.2**, **m 154-159°, 146-151°, b 187-190°/7mm, 210-214°/13mm, d²⁰ 1.142g/ml**. Crystallise the white powder from CS_2 [**m** 139°, CS_2 complex] and distil it in a vacuum. It reacts violently with H_2O , but is slightly soluble in xylene and PhCl. The molecular weight

corresponds to $[\text{Al}(\text{OEt})_3]_4$ [Robinson & Peak *J Phys Chem* **39** 1125 1935, DOI: 10.1021/j150368a009; Vilani & Nord *J Am Chem Soc* **69** 2605 1947, DOI: 10.1021/ja01203a011]. [*Beilstein* **1** III 1284, **1** IV 1289.]

Aluminium isopropoxide [555-31-7] $\text{C}_9\text{H}_{21}\text{O}_3\text{Al}$, **M 204.3**, **m 119°(98%+purity)**, **138-142°(99.99%+purity)**, **b 94°/0.5mm**, **135°/10mm**, **d²⁰ 1.035g/ml**. Redistil the white solid under vacuum. *Hygroscopic*, decomposes in H_2O , but is soluble in *iso*-PrOH. [Robinson & Peak *J Phys Chem* **39** 1125 1935, DOI: 10.1021/j150368a009; *Beilstein* **1** IV 1468; Fieser **1** 35, **3** 10, **4** 15, **5** 14, **6** 19, **8** 15, **9** 14, **11** 29.] Commonly used in the preparation of Al_2O_3 [Sim *J Korean Ceram Soc* **52**(1) 56 2015, DOI: 10.4191/kcers.2015.52.1.56].

Aluminum triethyl (triethyl aluminum) [97-93-8] $\text{C}_6\text{H}_{15}\text{Al}$, **M 114.2**, **m -50°**, **b 69°/1.5mm**, **76°/2.5mm**, **129-131°/55mm**, **d²⁰ 0.695**, **n_D²⁰ 1.394**. Purify $\text{Al}(\text{Et})_3$ by fractionation in an inert atmosphere under a vacuum in a 50cm column containing a heated nichrome spiral, taking the fraction **b 112-114°/27mm**. It is very sensitive to H_2O and should be stored under N_2 . It should not contain chloride ions which can be shown by hydrolysis and testing with AgNO_3 . [Baker & Sisler *J Am Chem Soc* **75** 4828 5193 1953, DOI: 10.1021/ja01115a502, DOI: 10.1021/ja01117a013; NMR: Brownstein et al. *J Am Chem Soc* **81** 3826 1959, DOI: 10.1021/ja01524a006; *Beilstein* **4** IV 4398; for applications see Fieser **1** 1197, **2** 427, **3** 299, **4** 526, **5** 688, **10** 415, **17** 367.] Also available commercially as solutions in heptane (1M), in hexanes (1M) and in toluene (25 wt%).

Aluminium tri-tert-butoxide [556-91-2] $\text{C}_{12}\text{H}_{27}\text{O}_3\text{Al}$, **M 246.3**, **m 208-210°(dec)**, **>300°**. Crystallise pale yellow $\text{Al}(\text{O}-t\text{Bu})_3$ from $^*\text{C}_6\text{H}_6$ and sublime it at 180°. [McElvain & Davie *J Am Chem Soc* **73** 1400 1951, DOI: 10.1021/ja01148a002; *Beilstein* **1** IV 1612; for applications see Fieser **1** 23, **2** 21.]

Aluminium trimethanide (trimethyl aluminium, TMA) [75-24-1] $\text{C}_3\text{H}_9\text{Al}$, **M 72.1**, **m 15.2°**, **b 111.5°/488.2mm**, **124.5°/atm**, **d²⁰ 0.752g/ml**. Distil $\text{Al}(\text{Me})_3$ through a 10-20 theoretical plates column under 1 atmosphere pressure of N_2 (better with very slow take-off). It attacks grease (use glass joints). It has been distilled over Al in absence of grease, into small glass vials and sealed under N_2 . The purity is measured by its freezing point. It reacts with H_2O , is non-conducting in $^*\text{C}_6\text{H}_6$ and is **HIGHLY FLAMMABLE**. [Bamford et al. *J Chem Soc* 468 1946, DOI: 10.1039/JR9460000468; Pitzer & Gutowsky *J Am Chem Soc* **68** 2204 1946, DOI: 10.1021/ja01215a027; *Beilstein* **4** IV 4397; for applications see Fieser **8** 506, **15** 341, **17** 371.]

4-Aminophenylmercuric acetate (APMA) [6283-24-5] $\text{C}_8\text{H}_9\text{NO}_2\text{Hg}$, **M 371.8**, **m 163-165°**, **168°**, **175°(dec)**, **180°(dec)**. Recrystallise it from hot dilute AcOH and dry it in air. Highly **TOXIC**. [Mahapatra et al. *J Indian Chem Soc* **32** 613 1955, Albert & Schneider *Justus Liebigs Ann Chem* **465** 257 1928, DOI: 10.1002/jlac.19284650113; *Beilstein* **16** III 1411, **16** IV 1754.]

Ammonium acetate [631-61-8] $\text{C}_2\text{H}_3\text{NO}_2$, **M 77.1**, **m 112-114°**, **114°**, **d²⁰ 1.17g/ml**. Crystallise it twice from anhydrous acetic acid, and dry under vacuum for 24 hours at 100° [Proll & Sutcliffe *Trans Faraday Soc* **57** 1078 1961, DOI: 10.1039/TF9615701078]. Its solubility in H_2O in g/100ml is 102 (0°), 148 (4°), 143 (20°) and 533 (80°); in MeOH it is 7.89 (15°) and 131.2 (94.2°); and 0.1 (25°) in DMF. [*Beilstein* **2** IV 121; Fieser **1** 38, **7** 11.]

Ammonium benzoate [1863-63-4] $\text{C}_7\text{H}_9\text{NO}_2$, **M 139.2**, **m 198°**, **200°(dec)**, **d²⁰ 1.26g/cm⁻³**. Crystallise it from EtOH. Its solubility in H_2O in w/v% is 21.3 (2°) and 83 (100°); and soluble in MeOH [*Beilstein* **9** IV 273.]

Ammonium dodecylsulfate (ammonium laurylsulfate) [2235-54-3] $\text{C}_{12}\text{H}_{29}\text{NO}_4\text{S}$, **M 283.4**, **b 418°/atm**, **d²⁰ 1.02g/cm⁻³** and **n_D²⁰ 1.374** (for a ~30 wt% in H_2O). Recrystallise it first from 90% EtOH and then twice from absolute EtOH, and finally dry it in a vacuum. [*Beilstein* **1** III 1786.] Useful anionic surfactant. **IRRITANT**.

Ammonium ferric oxalate trihydrate [13268-42-3] $\text{C}_6\text{H}_{12}\text{N}_3\text{O}_{12}\text{Fe} \cdot 3\text{H}_2\text{O}$, **M 428.1**, **m ~160°(dec)**, **d²⁰ 1.77g/ml**. Crystallise it from hot water (0.5ml/g). [*Beilstein* **3** III 1103.]

Ammonium formate [540-69-2] CH_5NO_2 , **M 63.1**, **m 116°**, **117.3°**, **b 180°/atm (dec)**, **d²⁵ 1.26g/cm³**, **d₄⁴⁵ 1.280**. Heat the white solid in NH_3 vapour and dry it in a vacuum till the NH_3 odour is faint (*note* that it can evaporate completely in a vacuum). Recrystallise it from absolute EtOH and then keep it in a desiccator over 99% H_2SO_4 *in vacuo*. It is very *hygroscopic*. It exists in two forms, stable needles and less stable plates. It also

forms acid salts, i.e. $\text{HCO}_2\text{NH}_4 \cdot 3\text{HCO}_2\text{H}$ and $\text{HCO}_2\text{NH}_4 \cdot \text{HCO}_2\text{H}$. [Kendall & Adler *J Am Chem Soc* **43** 1470 1921, DOI: 10.1021/ja01440a010; *Beilstein* **2** IV 18.] Its solubility in H_2O in g/100ml is 102 (0°), 142.7 (20°), 202.4 (40°) and 516 (80°); and is soluble in liquid NH_3 . HCO_2NH_4 , in the presence of microencapsulated $\text{Pd}(0)$ under microwave irradiation, removes phenolic benzyl protecting groups rapidly [Quai et al. *Tetrahedron Lett* **48** 1241 2007, DOI: 10.1016/j.tetlet.2006.12.041].

Ammonium ionophore I (Nonactin) [6833-84-7] $\text{C}_{40}\text{H}_{64}\text{O}_{12}$, **M 736.9**, **m 147-148°**, $[\alpha]_{\text{D}}^{20}$ **0** (c 1.2, CHCl_3). Crystallise it from MeOH (colourless needles), and dry at 20° in high a vacuum. It is a selectophore with high sensitivity for transporting NH_4^+ ions through biological membranes. [Corbaz et al. *Helv Chim Acta* **38** 1445 1955, DOI: 10.1002/hlca.19550380617; Dominguez et al. *Helv Chim Acta* **45** 129 1962, DOI: 10.1002/hlca.19620450117; Dobler *Helv Chim Acta* **55** 1371 1972, DOI: 10.1002/hlca.19720550504; *Beilstein* **19/12** V 751.] Used for preparing potentiometric electrodes for determination of K^+ and NH_4^+ ions.

Ammonium oxalate dihydrate [6009-70-7 (H_2O), 1113-38-8 (anhydrous)] $\text{C}_2\text{H}_8\text{N}_2\text{O}_4 \cdot \text{H}_2\text{O}$, **M 142.1**, **m 70° (H_2O)**, **133° (anhydr, dec)**, **b 365.1°/760mm**, **d²⁰ 1.50g/cm³**. Crystallise it from water (10ml/g) at 50°. [*Beilstein* **2** IV 1846.] It is a constituent of kidney stones.

Ammonium picrate (Dunnite explosive) [131-74-8] $\text{C}_6\text{H}_6\text{N}_4\text{O}_7$, **M 246.1**, **m EXPLODES above 200°**, **d²⁵ 1.719g/cm³**. Crystallise it from EtOH and acetone. Its solubility in H_2O at ~20° is 1g/100ml. [Mitchell & Bryant *J Am Chem Soc* **65** 128 1943, DOI: 10.1021/ja01242a001; *Beilstein* **6** II 262, **16** III 879, **16** IV 1392.]

n-Amylmercuric chloride (pentylmercuric chloride) [544-15-0] $\text{C}_5\text{H}_{11}\text{HgCl}$, **M 307.2**, **m 110°**. Crystallise it from EtOH. The **bromide** has **m 122°**. [Larock & Brown *J Am Chem Soc* **92** 2467 1970, DOI: 10.1021/ja00711a043; Marvel et al. *J Am Chem Soc* **47** 3009 1925, DOI: 10.1021/ja01689a026; *Beilstein* **14** H 706, 725.]

Aurothioglucose (gold thioglucose, Solganal) [12192-57-3] $\text{C}_6\text{H}_{11}\text{AuO}_5\text{S}$, **M 392.2**. Purify it by dissolving it in H_2O (0.05g in 1ml) and precipitating it by adding EtOH. It yields yellow crystals with a slight mercaptan odour. It decomposes slowly in H_2O , and is soluble in propylene glycol but insoluble in EtOH and other common organic solvents. [Caterson & Taylor *FEBS Lett* **98** 351 1979; DOI: 10.1016/0014-5793(79)80215-X; Cooney et al. *Biochem J* **259** 651 1989; DOI: 10.1042/bj2590651; Shaw 'Gold-Based Therapeutic Agents' *Chem Rev* **99** 2589 1999, DOI: 10.1021/cr980431o.]

Barium acetate [543-80-6] $\text{C}_4\text{H}_6\text{O}_4\text{Ba}$, **M 255.4**, **m 450°**, **d²⁵ 2.468g/cm³ (anhydr)**. Crystallise the white salt twice from anhydrous acetic acid and dry it under vacuum for 24 hours at 100°. Its solubility in H_2O in w/v% is 55.8 (0°) and 72 (20°), but it is only slightly soluble in EtOH. It decomposes to BaCO_3 upon heating in air. [*Beilstein* **2** III 192, **2** IV 114.] Used as a mordant for printing fabrics, drying paints and varnishes, and in lubricating oil.

Barium ionophore I [*N,N,N',N'*-tetracyclohexyloxy-bis-(*o*-phenyleneoxy)diacetamide] [96476-01-6] $\text{C}_{40}\text{H}_{56}\text{N}_2\text{O}_5$, **M644.9**, **m 156-158°**. Purify the ionophore by chromatography on a Kieselgel column, elute it with $\text{CH}_2\text{Cl}_2/\text{EtOAc}$ (5:1), and recrystallise the residue from the evaporated effluent from EtOH/ Me_2CO to give colourless crystals. It is an electrically neutral ionophore with high selectivity for Ba^{2+} ions and with high lipophilicity. [Kleiner et al. *Chem Ber* **118** 1071 1985, DOI: 10.1002/cber.19851180324; Läubli et al. *Anal Chem* **57** 2756 1985, DOI: 10.1021/ac00291a005; Peshkova et al. *Electroanalysis* **22(19)** 2147 2010, DOI: 10.1002/elan.201000222.]

Barium propionate (H_2O) [5908-77-0] $\text{C}_6\text{H}_{10}\text{O}_4\text{Ba} \cdot \text{H}_2\text{O}$, **M 301.5**, **283.5 (anhydr)**, **m 300°**, **d₄²⁷ 1.44**, **pK²⁴ 4.88 (for propionic acid)**. Crystallise the white powder from warm water (50ml/g) by adding EtOH and cooling. [*Beilstein* **2** III 517, **2** IV 702.]

Benzenechromium(0) tricarbonyl [12082-08-5] $(\text{C}_6\text{H}_6)(\text{CO})_3\text{Cr}$ **M 214.1**, **m 162-163°**, **163-166°**. Purify the yellow complex by sublimation *in vacuo*. A possible impurity is 2-picoline which can be removed by washing

with pentane and drying. It is then purified further by sublimation at 80-85°/10⁻³mm, or by recrystallisation from Et₂O to give yellow crystals. ¹H NMR in CDCl₃ should give a single peak at τ 4.68. Soluble in THF, Et₂O and *C₆H₆, but insoluble in H₂O. [Rausch *J Org Chem* **39** 1787 1974, DOI: 10.1021/jo00925a052; Fischer & Ófele *Chem Ber* **90** 2532 1957, DOI: 10.1002/cber.19570901117; Pauson in Houben-Weyl *Meth Org Chem* V, E 18 Pt I p226 Theme Verlag, Stuttgart 1986, *Beilstein* **5** IV 625; for applications see Fieser **6** 27, **13** 19.]

Beryllium acetate [Be(OAc)₂] [543-81-7] C₄H₆O₄Be, M 127.1, m 65-100° (slow heating), 155-180° (rapid heating). It is obtained by dissolving the *basic acetate* (4g, see below) in boiling glacial AcOH containing acetyl chloride (4.5g) and refluxing for ~15 minutes. The Be(OAc)₂ which separates during this time, is filtered off, washed with AcOH, cold CHCl₃ and dried *in vacuo* to give 90-94% yield of the salt. It is stable for several weeks in a tightly stoppered container at ~ 25°. It slowly loses Ac₂O, and more rapidly on heating, to give the basic acetate which sublimes out. Strong heating partially decomposes it to give Ac₂O and BeO. It is not readily attacked by cold H₂O but forms a hydrate on warming, and is insoluble in most solvents. Used in the ceramic industry. [Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 901 1963, Besson & Hardt *Z Anorg Allgem Chem* **277** 188 1954, DOI: 10.1002/zaac.19542770310.] Beryllium compounds are potentially **carcinogenic**.

Beryllium acetate (basic) [Be₄O(OAc)₆] [1332-52-1, 19049-40-2] C₁₂H₁₈O₁₃Be₄, M 406.3, m 283°, 285-286°, b 330-331°/atm. On evaporating Be(OH)₂ with AcOH this covalent basic acetate is produced. Alternatively, BeCO₃ (40g, see [744998-97-8]) and AcOH (80ml) are heated with stirring until evolution of CO₂ ceases. When the reaction is complete, white semi-translucent crystals separate. After cooling to ~ 25°, the basic acetate crystals are filtered off and dried in air. These are treated with warm CHCl₃ (60-80ml), any insoluble material is filtered off, and the colourless octahedral crystals of the basic acetate which separate on cooling (with partial evaporation) are collected and freed from any CHCl₃ *in vacuo* to give pure basic salt (28g, ~47%, m 284°). It is readily volatile, it can be distilled, and it sublimes in a vacuum leaving a small residue of 0.3-0.5% of BeO. It is very soluble in CHCl₃, soluble in non-polar organic solvents such as boiling *C₆H₆, toluene, xylene, tetralin, and in AcOH; less so in CCl₄, Ac₂O and AcCl, but is sparingly soluble (~0.3%) in Et₂O. It dissolves in anhydrous boiling MeOH but soon liberates Ac₂O to form highly aggregated basic acetates. It is quite stable in cold H₂O, but is rapidly hydrolysed in hot H₂O. [Moeller et al. *Inorg Synth* **3** 9 1950, DOI: 10.1002/9780470132340.ch3; Ehrlich in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol I p 901 1963, and for the basic formate or propionate salts see Besson & Hardt *Z Anorg Allgem Chem* **277** 188 1954, DOI: 10.1002/zaac.19542770310; Hendus & Hardt *Z Anorg Allgem Chem* **277** 127 1954, DOI: 10.1002/zaac.19542770303; Hendus & Hardt *Z Anorg Allgem Chem* **286** 265 1956, DOI: 10.1002/zaac.19562860511; and Hardt *Z Anorg Allgem Chem* **286** 254 1956, DOI: 10.1002/zaac.19562860510; *Beilstein* **2** H 111, **2** I 48, **2** II 116, **2** III 190, **2** IV 112.] The X-ray crystal structure shows that the four beryllium atoms are arranged at the corners of a regular tetrahedron with oxygen at the centre, and the six edges are occupied by acetate groups. Thus the four Be atoms at the apexes of the tetrahedron are coordinated to the carboxyl oxygen atoms of the six acetate groups, i.e. each Be atom is coordinated to three oxygen atoms. [Bragg & Morgan *Proc Roy Soc* **104** 437 1923, DOI: 10.1098/rspa.1923.0119; for partial structure see also F.A. Cotton and G. Wilkinson *Advanced Inorganic Chemistry (A Comprehensive Text)* p. 175, Interscience Publ. 1962, Library of Congress Catalog Number 62-14818.] Beryllium compounds are potentially **carcinogenic**.

Beryllium acetylacetonate [Be(acac)₂] [10210-64-7] C₁₀H₁₄O₄Be, M 207.2, m 108°, 108.5-109°, b 270°/atm, d²⁵ 1.168g/ml. This complex has been prepared by adding 3M aqueous NaOH (100ml) to a solution of BeSO₄ 4H₂O (10g, see [7787-50-0]) in distilled H₂O (100ml) to give a strong basic solution to which is added freshly distilled acetylacetone (30ml) dropwise with stirring, and a white precipitate separates. This is collected, washed with H₂O and dried *in vacuo* to give the complex (5.1g 43%) which is recrystallised by dissolving in a small volume of *C₆H₆ and adding gradually petroleum ether until crystallisation is complete. The crystals are collected, washed with petroleum ether and dried *in vacuo*. It can be recrystallised further from a small volume of *C₆H₆ or a large volume of petroleum ether. It has also been prepared in 70% yield from basic beryllium carbonate (3g) in H₂O (45ml) by conversion to the chloride with 6N aqueous HCl (ca 20ml) until slightly acidic. To this is added dropwise a clear solution of acetylacetone (10g suspended in 45ml of H₂O to which is added 6N ammonia until completely clear) and stirred until precipitation of the complex is complete (pH should be near neutral). The solid is collected washed with H₂O, dried, and crystallised as before. It is also sublimed through a

plug of glass wool at a vacuum of 0.1mm by heating in a bath at 80 to 100°. It is very soluble in most organic solvents such as EtOH, Et₂O, Me₂CO, *C₆H₆ and in CS₂ but considerably less in petroleum ethers. However, it is practically insoluble in H₂O which hydrolyses it on boiling; and it is also hydrolysed by acids and alkalis. [Jones *J Am Chem Soc* **81** 3188 1959, DOI: 10.1021/ja01522a005; Arch et al. *Inorg Synth* **2** 17 1946, DOI: 10.1002/9780470132333.ch5.] Beryllium compounds are potentially **carcinogenic**.

Bicyclo[2.2.1]hepta-2,5-diene rhodium (I) chloride dimer (norbornadiene rhodium chloride complex dimer) [12257-42-0] C₁₄H₁₆Cl₂Rh₂, **M 461, m 239-241°(dec), 240°(dec)**. It recrystallises from hot CHCl₃/petroleum ether as fine crystals soluble in CHCl₃ and *C₆H₆ but is almost insoluble in Et₂O or petroleum ether. [Abel et al. *J Chem Soc* 3178 1959, DOI: 10.1039/JR9590003178.] It is a **pre-catalyst** for asymmetric and cross-coupling reactions, and is a catalyst for hydrosilylation. [cf. Ngai et al. *J Am Chem Soc* **129** 280 2007, DOI: 10.1021/ja0670815.]

2,2'-Bipyridinium chlorochromate [76899-34-8] C₁₀H₉N₂CrO₃Cl, **M 292.6**. Wash it with cold conc HCl, then H₂O (sintered glass funnel) and dry it in a vacuum (CaCl₂) to give a free-flowing yellow-brown powder. Store it in the dark. [Guziec & Luzzio *Synthesis* 691 1980, DOI: 10.1055/s-1980-29172; Chakraborty & Chandrasekaran *Synth Commun* **10** 951 1980, DOI: 10.1080/00397918008061857; Fieser **10** 30, **11** 44.] Useful oxidant. Together with *m*-chloroperbenzoic acid it mediates the cleavage of benzyldene acetals [Luzzio & Bobb *Tetrahedron Lett* **38** 1733 1997, DOI: 10.1016/S0040-4039(97)00203-7]. **SUSPECTED CARCINOGEN**.

2,2'-Biquinolin-4,4'-dicarboxylic acid dipotassium salt [63451-34-3, 207124-63-8 (3H₂O)] C₂₀H₁₀N₂O₄K₂, **M 420.51(anhydr), m >300°**. Recrystallise it from H₂O. The **Cu salt** has UV with λ_{max} at 562nm. [Mopper & Gindler *Anal Biochem* **56** 440 1973, DOI: 10.1016/0003-2697(73)90210-8; Beilstein **25** IV 1148.] Useful charged ligand for catalyst immobilisation and analysis [Review: Chisholm & McIndoe *JCS Dalton Trans* **30** 3933 2008, DOI: 10.1039/B800371H].

Bis(1,2,3,4,5-η-cyclooctadienyl)ruthenium [Ru(COD)₂] [63395-36-8] C₁₆H₂₂Ru, **M 315.4, m 89-90°, 113-114°**. The bis-complex is prepared by adding a solution of RuCl₃·3H₂O (624mg, 1.33mol) in EtOH (20ml) under argon (or N₂) slowly over 30 minutes to a stirred suspension of Zn powder (7g), EtOH (5ml) and redistilled 1,2-5,6-cyclooctadiene (11ml, 80mol, COD see [111-78-4, 1552-12-1], then stirring at room temperature for 2 hours, and filtering the mixture under N₂. The filtrate is concentrated on a vacuum line to remove volatiles, and the residual brown oil is placed on top of an Al₂O₃ column in hexane after flushing with N₂. The yellow band is eluted with hexane, concentrated *in vacuo*, the eluate is evaporated, pentane is added to dissolve the residue and the solution is cooled at -78° for several hours, to provide very pale yellow (almost colourless) crystals of the analytically **pure complex** (283mg, 35%). The ¹H NMR (60MHz, C₆D₆, TMS) has τ at 4.42 (t, 2H), 6.15 (dd, 4H), 6.45 (m, 4H) and 7.49-8.95 (m, 12H); and the ¹³C NMR (22.6MHz, C₆D₆, TMS) has δ at 22.5, 27.3, 36.7, 35.3, 38.9, 62.4, 76.7, 87.6. [Preparation: Pertici et al. *JCS Dalton Trans* 1961 1980 (gave m 89-90°), DOI: 10.1039/DT9800001961; Itoh et al. *J Organomet Chem* **272** 179 1984 (gave m 113-114°), DOI: 10.1016/0022-328X(84)80465-9.]

Bis(η⁵-2,4-cyclopentadien-1-yl)samarium II [SmCp₂, samarocene] [80695-16-5] C₁₀H₁₀Sm, **M 280.5, decomposes at high temperatures**. Samarium dicyclopentadienyl can be prepared in Schlenk equipment under argon by adding SmI₂ (0.1M in THF, 60ml, 6mmol) to a 0.4M solution of sodium pentadienide in THF (30ml, 12mmol) when the complex separates immediately and is decanted within 1 hour. The red powder of SmCp₂ is collected, washed twice with THF to remove NaI, dried *in vacuo* and stored under argon. The red powder and the purple solid THF complex (**M 352.6**) are **pyrophoric** in air. SmCp₂ can also be stored for a few days in a Schlenk tube under THF without decomposition. Freshly prepared suspensions of SmCp₂ have been used in **pseudo-Barbier couplings** between carbonyl compounds and aliphatic or allylic halides and are more efficient than SmI₂ [Namy et al. *J Organometal Chem* **328** 81 1986, DOI: 10.1016/S0022-328X(00)99769-9].

SmCp₂-THF has been prepared in one arm of a U-tube by stirring in an anhydrous oxygen-free helium atmosphere) tris(cyclopentadienyl)samarium III (SmCp₃, 0.499g), K metal (0.527g, naphthalene (0.128g, 25% molar deficiency) and THF (25ml, freshly distilled under reduced pressure from Na-benzophenone) for 48 hours. Purple SmCp₂-furanate was separated by decantation, washed by successive back-distillation with THF and the solvent is removed at 24°/10⁻³mm during 18 hours. It is **pyrophoric** and should be handled in an anhydrous

oxygen-free atmosphere. Its IR has ν_{\max} (Nujol) at 3080 (C-H stretch), 1475, 1347, 1308, 1263, 1163, 1070, 1008 (C-H bend parallel), 775, 740 (C-H bend perpendicular) and 350 (antisym metal-ring vibration) cm^{-1} , and other bands at 2980, 2880, 1375, 725 and 565 cm^{-1} due to coordinated THF with intensities similar to the weaker metallocene bands. On exposure to traces of air this complex immediately changes in colour from deep purple to yellow-grey with drastic reduction in the paramagnetism. [Watt & Gillow *J Am Chem Soc* **91** 775 1969, DOI: 10.1021/ja01031a061.]

Bis(2,9-dimethyl-1,10-phenanthroline) copper(I) perchlorate (Cuproine) [54816-44-5] $\text{C}_{28}\text{H}_{24}\text{N}_4\text{Cu} \cdot \text{ClO}_4$, **M 579.6**, pK^{25} -2.4 to -3.1 (for HClO_4) and 6.15 (for dimethylphenanthroline). Crystallise it from acetone. It has UV with λ_{\max} in isopentanol or hexan-1-ol at 454nm. [Smith & McCurdy *Anal Chem* **24** 371 1952, DOI: 10.1021/ac60062a029; Gahler *Anal Chem* **26** 577 1954, DOI: 10.1021/ac60087a052; Beilstein **23** III/IV 1737.]

1,1'-Bis(diphenylphosphino)ferrocene (DPPF) [12150-46-8] $\text{C}_{34}\text{H}_{28}\text{FeP}_2$, **M 554.4**, **m 181-182°(dec), 184-194°**. Wash the ferrocene with distilled H_2O and dry it in a vacuum. Dissolve it in *ca* 5 parts of hot dioxane and cool to give orange crystals **m 181-183°**. Recrystallisation from $^*\text{C}_6\text{H}_6$ /heptane (1:2) gives a product with **m 183-184°**. [Bishop et al. *J Organomet Chem* **27** 241 1971, DOI: 10.1016/S0022-328X(00)80571-9.] This ligand, *via* Pd-phosphine, is used to prepare new functionalised furan derivatives by sequential C-C and C-O bond formation [Tanimori et al. *Synthesis* **5** 865 2006, DOI: 10.1055/s-2006-926321], and when complexed with Pd(II), catalyses cross-coupling synthesis of oxazepine ring systems [Ma et al. *Tetrahedron* **62** 9002 2006, DOI: 10.1016/j.tet.2006.07.009].

Bis(ethyl)titanium(IV) chloride [2247-00-9] $\text{C}_4\text{H}_{10}\text{Cl}_2\text{Ti}$, **M 176.9**. Recrystallise it from boiling toluene. [See Marek (ed) *Titanium and Zirconium in Organic Synthesis* Wiley-VCH 2002 ISBN 3-527-304-28-2, Wailes et al. *Organometallic Chemistry of Titanium, Zirconium and Hafnium* Academic Press 1974 ISBN 0-127303502.] **Bis(ethyl)zirconium(IV) chloride** [92212-70-9] $\text{C}_4\text{H}_{10}\text{Cl}_2\text{Zr}$, **M 220.3**. Recrystallise it from boiling toluene. [See Marek (ed) *Titanium and Zirconium in Organic Synthesis* Wiley-VCH 2002 ISBN 3-527-304-28-2, Wailes et al. *Organometallic Chemistry of Titanium, Zirconium and Hafnium* Academic Press 1874 ISBN 0-127303502.]

***N,N'*-Bis(salicylidene)ethylenediamine cobalt (II) [Co(SALEN)₂, salcomine]** [14167-18-1] $\text{C}_{16}\text{H}_{14}\text{N}_2\text{O}_2\text{Co}$, **M 325.2**. The powder should have an oxygen capacity of 4.7-4.8% as measured by the increase in weight under O_2 at 100 pounds pressure at *ca* 20°. The O_2 is expelled on heating the material to 65°. It crystallises from pyridine, CHCl_3 or $^*\text{C}_6\text{H}_6$, and the solvent may be removed by heating at 120° in a vacuum. However, this heating may mean reduced O_2 capacity. [Diehl & Hack *Inorg Synth* **3** 196 1950, DOI: 10.1002/9780470132340.ch53; Fieser **2** 360, **6** 507, **12** 429.] In the dry state and in solvent it absorbs O_2 reversibly, turning from a maroon colour to black. [Vogt et al. *Chem Rev* **63** 269 1963, DOI: 10.1021/cr60223a004; for metal complexes in catalysis Cozzi *Chem Soc Rev* **33**(7) 410 2004, DOI: 10.1039/B307853C]. The uncomplexed ligand, ***N,N'*-bis(salicylidene)ethylenediamine (SALEN ligand)** [94-93-9] $\text{C}_{16}\text{H}_{16}\text{N}_2\text{O}_2$, **M 268.3**, has **m 125-126°** (from EtOH), **127-128°** and **125-129°** were also reported. [Original paper: Pfeiffer et al. *Justus Liebig's Ann Chem* **503** 84 1933, DOI: 10.1002/jlac.19335030106] **VERY TOXIC**.

Bis(tetrabutylammonium) dichromate [56660-19-6] $\text{C}_{32}\text{H}_{72}\text{N}_2\text{O}_7\text{Cr}_2$, **M 700.9**, **m 139-142°, 139-143°**. Wash the dichromate with water and dry it in a vacuum. Recrystallise it from hexane (**m 79-80°**). Useful oxidant. [Santaniello & Ferraboschi *Synth Commun* **10** 75 1980, DOI: 10.1080/00397918008080055; Beilstein **4** IV 556.] (Possible **CARCINOGEN**.)

***n*-Butylmercuric chloride** [543-63-5] $\text{C}_4\text{H}_9\text{N}_2\text{ClHg}$, **M 293.1**, **m 130°**. Crystallise it from 95% EtOH (white needles or leaflets). Its solubility (g/100ml) in H_2O is 1×10^{-4} at 18° and 3.3×10^{-4} at 100°; in EtOH is 1.5 at 18° and 9.0 at 70°; and in CHCl_3 is 5.6 at 18°. [Larock & Brown *J Am Chem Soc* **92** 2467 1970, DOI: 10.1021/ja00711a043; Marvel et al. *J Am Chem Soc* **47** 3009 1925, DOI: 10.1021/ja01689a026.]

***n*-Butylstannoic acid [*n*-BuSnO(OH)_n, *n*-butyltinhydroxy oxide hydrate]** [2273-43-0, 22719-01-3] $\text{C}_4\text{H}_{10}\text{O}_2\text{Sn}$, **M 208.8(anhydr)**, **d²⁵ 1.46g/cm³**. Purify it by adding excess KOH in CHCl_3 to remove small

amounts of $n\text{-BuSn(OH)Cl}_2$ and $n\text{-BuSn(OH)}_2\text{Cl}$, and isolate it by acidification. It is soluble in acetone and dries as an infusible white powder and is **polymeric**. Its Mössbauer parameters are δ .68 and 0.70mm/sec for C, and its $H\Delta E_Q$ are 1.52 and 1.65mm/sec (see Davies et al. below). It forms a **hexameric** (by Xray, see Holmes et al. below) ***n*-butyloxy *o*-nitrobenzoate ester** [$n\text{-BuSn(O)O}_2\text{CC}_6\text{H}_4\text{NO}_2\text{-2}]_6\cdot 3\text{C}_6\text{H}_6$, thus: $n\text{-BuSn(O)(OH)}$ (2.17g, 10.4mmol) is added to a stirred solution of $\text{C}_2\text{H}_2\text{Cl}_2$ (50ml) and $^*\text{C}_6\text{H}_6$ (80ml) followed by *o*-nitrobenzoic acid (1.74, 11.4mmol), and refluxed for 18 hours under a Dean-Stark trap to remove H_2O azeotropically. The original clear yellow solution changes to a cream-white suspension, and after 24 hours standing a white powder containing small clear crystals result, which is filtered off and washed with $^*\text{C}_6\text{H}_6$. On standing larger crystals separate in the filtrate. Both the white powder and the crystals have the same **m 295-305°** (total yield is 3.65g, 98%). They have IR (Nujol) with ν at 1530, 1550 (COO), 563 and 530 (Sn-O) cm^{-1} , and analyse correctly after drying. It forms **hexameric esters** with other acids, e.g. with cyclopentane carboxylic acid [see Chandrasekhar et al. *Inorg Chem* **26** 1050 1987, DOI: 10.1021/ic00254a017]. [Holmes et al. *J Am Chem Soc* **109** 1408 1987, DOI: 10.1021/ja00239a022; Luijten *Recl Trav Chim Pays-Bas* **85** 873 1966, DOI: 10.1002/recl.19660850904; for Mössbauer and NMR spectra see Davies et al. *J Organomet Chem* **39** 279 1792, DOI: 10.1016/S0022-328X(00)80452-0; Pfeiffer *Z Anorg Chem* **68** 102 1910, DOI: 10.1002/zaac.19100680111].

Cadion [1-(4-nitrophenyl)-3-(4-phenylazophenyl)-triazene] [5392-67-6] $\text{C}_{18}\text{H}_{14}\text{N}_6\text{O}_2$, **M 346.3, m 189°(dec), d²⁵ 1.30g/cm³**. Commercial cadion is purified by recrystallisation from 95% EtOH and is dried *in vacuo*. It is stable in 0.2 N KOH (in 20% aqueous EtOH) at 25°. It is a sensitive reagent for **Cd**, and the **Cd complex** has UV with λ_{max} (EtOH) at 475nm. [Chavanne & Geronimi *Anal Chim Acta* **19** 377 1958, DOI: 10.1016/S0003-2670(00)88180-5; *Beilstein* **16** III 664.] The method uses a dual-wavelength spectrophotometric assay and the difference value in absorbance in the λ_{max} (at 475nm) of the Cd chelate and that of Cadion (at 566nm) [Watanabi & Ohmori *Talanta* **26** 959 1979, DOI: 10.1016/0039-9140(79)80133-2].

Cadmium acetate dihydrate [543-90-8(anhydr), 5743-04-4($2\text{H}_2\text{O}$), 89759-80-8($x\text{H}_2\text{O}$),] $\text{C}_4\text{H}_6\text{O}_4\text{Cd}$, **M 230.5(anhydr), m 130° (hydrate, dec), 255°(anhydr), d²⁵ 2.01g/cm³ (hydrate), 2.341g/cm³ (anhydr), pK₁²⁵ 9.7, pK₂²⁵ ~11.0 (for Cd^{2+})**. Recrystallise it twice from anhydrous acetic acid and dry it under vacuum for 24 hours at 100°. The salts are soluble in MeOH and EtOH, and very soluble in H_2O . [*Beilstein* **2** IV 114.] **Cd compounds are considered Group 1 carcinogens by the IARC.**

Cadmium ionophore I [N,N,N',N' -tetramethyl-3,6-dioxooctanedi-(thioamide)] [73487-00-0] $\text{C}_{22}\text{H}_{44}\text{N}_2\text{O}_2\text{S}_2$, **M 432.7, m 35-36°**. Wash it well with petroleum ether, then several times with 2N HCl (if it has a slight odour of pyridine), then H_2O and dry it in a vacuum over H_2SO_4 . It is a **polar selectrophore** for Cd. [Schneider et al. *Helv Chim Acta* **63** 217 1980, DOI: 10.1002/hlca.19800630121; Simon & Carafoli *Methods Enzymol* **56** 439 1979, DOI: 10.1016/0076-6879(79)56043-1.]

Cadmium lactate [16039-55-7] $\text{C}_6\text{H}_{10}\text{O}_6\text{Cd}$, **M 290.6, b 227.6°/760mm**. Recrystallise the lactate from water (10ml/g) by partial evaporation in a desiccator. [*Beilstein* **3** H 277, **3** III 465, **2** IV 637.] **Cd compounds are considered Group 1 carcinogens by the IARC.**

Cadmium salicylate monohydrate [19010-79-8] $\text{C}_{14}\text{H}_{10}\text{O}_6\text{Cd}$, H_2O , **M 386.6(anhydr), m 242°(dec)**. Recrystallise the white solid from distilled Prasad et al. *J Indian Chem Soc* **35** 267 1958, H_2O by evaporation in a desiccator. It is readily soluble in acids and NH_4OH , slightly soluble in cold H_2O , MeOH and EtOH, but freely soluble in hot H_2O . It has been used as a pesticide, and as an antiseptic. [*Beilstein* **10** H 60, **10** I 25, **10** II 33, **10** III 94, **10** IV 128.] **Cd compounds are considered Group 1 carcinogens by the IARC.**

Calcein sodium salt [2',7'-bis-{ N,N -di(carboxymethyl)aminomethyl}fluorescein Na salt, Fluorexon, Fluorescein Complexon] [108750-13-6 diNa salt, 1461-15-0 free acid] $\text{C}_{30}\text{H}_{24}\text{N}_2\text{O}_{13}\text{Na}_2$, **M 666.5, pK_{Est(1)} ~ 1.9, pK_{Est(2)} ~ 2.5, pK_{Est(3)} ~ 8.0, pK_{Est(4)} ~ 10.5 (all for $\text{N-CH}_2\text{COOH}$), and pK_{Est(5)} ~ 3.5 (for benzoic COOH)**. Dissolve it in distilled H_2O and acidify with dilute HCl to pH 3.5. Filter off the solid acid and wash it well with H_2O . Redissolve *ca* 10g in 300ml H_2O containing 12g of NaOAc. Precipitate it again by adding HCl, filtering and washing with H_2O . Add the solid to 200ml of EtOH stir for 1 hour and filter. Repeat the EtOH

wash and dry the bright yellow solid in a vacuum. This acid decomposes on heating at *ca* 180°. See below for the preparation of the Na salt. [Diehl & Ellingboe *Anal Chem* **28** 882 1956, DOI: 10.1021/ac60113a030].

Alternatively, dissolve it in H₂O and acidify with 3N HCl to pH 3.5. Collect the solid and wash it with H₂O. The air-dried precipitate is extracted with 70% aqueous EtOH, filtered hot and cooled slowly. Fine yellow needles of the acid crystallise out; they are filtered off and dissolved in the minimum quantity of 0.01N NaOH and re-precipitated by adding N HCl to pH 3.5. It is then recrystallised from 70% aqueous EtOH (3x). The final product (**acid**) is dried at 80° in a vacuum for 24 hours, **m** >300°(dec). It contains one molecule of water per molecule of **acid** (C₃₀H₃₆N₄O₁₃·H₂O). The product is pure **acid** as revealed by electrophoresis at pH 5.6 and 8.6, and by TLC in *i*-BuOH/*i*-PrOH/AcOH/H₂O (60:60:5:5 by volume) or *i*-PrOH or pH 8.0 borate buffer. [Wallach et al. *Anal Chem* **31** 456 1959, DOI: 10.1021/ac60147a041.] The **Na salt** is prepared by dissolving the pure acid in H₂O containing 2 mols of NaOH per mol of acid reagent and lyophilising. It complexes with Ca and Mg ions [Pribil *Analyst* **83** 188 1958, DOI: 10.1039/AN9588300188]. [*Beilstein* **19** III/IV 4338.]

Calcium acetate monohydrate [5743-26-0 (H₂O), 62-54-4 (xH₂O)] C₄H₆O₄Ca, **M 176.2 (H₂O), m 150° (loses H₂O), 150°(dec to CaCO₃ + Me₂CO), d²⁵ 2.01g/cm³, n_D²⁵ 1.55, pK²⁵ 12.7 (for Ca₂⁺)**. Recrystallise the acetate from water (3ml/g) by partial evaporation in a desiccator. Its solubility in H₂O in g/100ml is 37.4 (0°), 34.7 (20°), and 29.7 (100°); it is slightly soluble in MeOH and hydrazine but insoluble in Me₂CO, EtOH and hydrocarbon solvents. [*Beilstein* **2** IV 113, for applications see Fieser **13** 60.]

Calcium benzoate trihydrate [2090-05-3, 5743-30-6] C₁₄H₁₆O₇Ca·3H₂O, **M 336.4(3H₂O)**. Recrystallise the benzoate from water (10ml/g) between 90° and 0°. Dry it in a crucible for 2 hours at 105° and cool in a dessicator. [*Beilstein* **9** I 60, **9** H 85, **9** III 377, **9** IV 280.] Antimicrobial preservative.

Calcium butyrate [5743-36-2] C₈H₁₄O₄Ca, **M 214.3, d₄³⁰ 1.271**. Recrystallise the butyrate from water (5ml/g) by partial evaporation in a desiccator and dry it in a vacuum to constant weight. [Pathak & Bhide *J Indian Chem Soc* **30** 47, 48 1953.] Its **dissociation constant** at 25° is 0.29 [Colman-Porter & Monk *J Chem Soc* 4363 1952, DOI: 10.1039/JR9520004363; *Beilstein* **2** IV 785].

Calcium carbamate [543-88-4] [H₂NCO₂] (H₂NCO₂)₂ Ca, **M 160.1**. Recrystallise calcium carbamate from aqueous ethanol. [*Beilstein* **4** H 75, **4** I 336, **4** II 557, **4** III 149, **4** IV 234.]

Calcium formate [544-17-2] [HCO₂]₂ Ca, **M 130.1, m 300°(dec on heating), d²⁵ 2.009g/cm³**. Recrystallise white food additive from water (5ml/g) by partial evaporation in a desiccator. The solubility of this white solid in H₂O in g/100ml is 16.1 (0°), 16.6 (20°), and 18.4 (100°); and is insoluble in EtOH. [*Beilstein* **2** IV 16.]

Calcium D-gluconate monohydrate [299-28-5, 18016-24-5, 66905-23-5 (H₂O)] C₁₂H₂₂O₁₄Ca·H₂O, **M 430.4(anhydr), m dec on heating, [α]₅₄₆²⁰ +11.0, [α]_D²⁰ +9.0 (c 1.2, H₂O)**. Calcium gluconate is soluble in H₂O (3.5g in 100g at 25°). Dissolve it in H₂O, filter and precipitate it by adding MeOH. Filter off the solid and dry it in a vacuum at 85°. **Alternatively**, dissolve it in H₂O, filter (from insoluble inorganic Ca) and evaporate it to dryness under vacuum at 85°. [March et al. *J Am Pharm Assoc* **41** 366 1952, DOI: 10.1002/jps.3030410709; *Beilstein* **3** IV 1255.] Used as a Ca supplement, protects against osteoarthritis.

Calcium D-heptagluconate dihydrate [17140-60-2] C₁₄H₂₆O₁₆Ca·2H₂O, **M 526.4 (2H₂O), [α]₅₄₆²⁰ +5.2, [α]_D²⁰ +4.4 (c 5, H₂O)**. Purify it in the same way as for calcium D-gluconate. [*Beilstein* **3** III 1112.]

Calcium ionophore I {ETH 1001, [(*-*)-4*R*,5*R*)-4,5-dimethyl-1,8-dioxo-3,6-dioxaoctamethylene]bis(12-methylaminododecanoate)} [58801-34-6] C₃₈H₇₂N₂O₈, **M 685.0**. This is a neutral Ca **selectophore**. It can be purified by thick layer (2mm) chromatography (Kieselgel F₂₄₅) and eluted with Me₂CO/CHCl₃ (2:1). [Ammann et al. *Helv Chim Acta* **56** 1780 1973, DOI: 10.1002/hlca.19730560538; Simon & Carafoli *Methods Enzymol* **56** 439 1979, DOI: 10.1016/0076-6879(79)56043-1.]

Calcium ionophore II (ETH 129, *N,N,N',N'*-tetracyclohexyl-3-oxapentanediamide, *N,N,N',N'*-tetra(cyclohexyl)diglycolic acid diamide) [74267-27-9] C₂₈H₄₈N₂O₃, **M 460.7, m 153-154°**. Recrystallise it from Me₂CO. It forms 1:2 and 1:3 metal/ligand complexes with Mg²⁺ and Ca²⁺ ions, respectively, and induces

selectivity in membranes for Ca^{2+} over Mg^{2+} by a factor of *ca* 10^4 . [Pretsch et al. *Helv Chim Acta* **63** 191 1980, DOI: 10.1002/hlca.19800630117; Simon & Carafoli *Methods Enzymol* **56** 439 1977, DOI: 10.1016/0076-6879(79)56043-1.]

Calcium ionophore III [A23187 **calcimycin antibiotic**] [52665-69-7] $\text{C}_{29}\text{H}_{37}\text{N}_3\text{O}_6$, **M 523.6, m 181-182°**, $[\alpha]_{\text{D}}^{25}$ **-56.0 (c 1, CHCl_3)**. It recrystallises from Me_2CO as colourless needles. Protect it from light and moisture, store in a refrigerator. It is soluble in Me_2SO (50mg/ml by heating or sonication), EtOH, EtOAc, MeOH and CHCl_3 (20mg/ml) and can be stored for 3 months without loss of activity. Aqueous solutions can be made by dissolving in Me_2SO , and diluting with H_2O . If stored at 2-8°, in a dessicated state and protected from light, it is stable for ~5years. The Mg and Ca salts are soluble in organic solvents and cross biological membranes. It has a pKa of 6.9 in 90% Me_2SO . The Ca complex crystallises from 50% EtOH as colourless prisms. It is **highly TOXIC**. [Pressman *Ann Rev Biochem* **45** 501 1976, DOI: 10.1146/annurev.bi.45.070176.002441; Chaney et al. *J Am Chem Soc* **96** 1932 1974, DOI: 10.1021/ja00813a047; Suzuki et al. *Anal Chem* **61** 382 1989, DOI: 10.1021/ac00179a018; Simon & Carafoli *Methods Enzymol* **56** 439 1977, DOI: 10.1016/0076-6879(79)56043-1; for a new role see Verma et al. *Cell Calcium* **50** 510 2011, DOI: 10.1016/j.ceca.2011.08.007.]

Calcium isobutyrate [533-90-4] $\text{C}_8\text{H}_{14}\text{O}_4\text{Ca}$, **M 248.2, 214.3**. Crystallise it from water (3ml/g) by partial evaporation in a desiccator. It forms a **pentahydrate** at low temperatures, but the crystals filtered from a saturated solution at 80° are the **monohydrate**; the transition temperature is 62.5°. [Lumsden *J Chem Soc* **81** 350 1902, DOI: 10.1039/CT9028100350.] It has a **dissociation constant** of 0.31 [Colman-Porter & Monk *J Chem Soc* 4363 1952, DOI: 10.1039/JR9520004363]. [*Beilstein* **2** H 290, **2** II 290, **2** IV 845.]

Calcium lactate ($5\text{H}_2\text{O}$) [814-80-2, 15743-47-5] $\text{C}_6\text{H}_{10}\text{O}_6\text{Ca}$, **M 308.3($5\text{H}_2\text{O}$), m 120°($5\text{H}_2\text{O}$), 240°(anhydr), d_{D}^{25} 1.494g/cm³, n_{D}^{25} 1.470**. Crystallise calcium lactate from warm water (10ml/g; solubility at 30° is 7.9g/100ml) by cooling to 0°. It is very soluble in EtOH. [*Beilstein* **3** IV 636.] The *S*(+)-**calcium lactate** (**calcium 2-hydroxypropionate**) [41372-22-9] has $[\alpha]_{\text{D}}^{20}$ **-4.2 (c 5, H_2O)**. Used as anti-acid, in foods, e.g. baking powder, added to sugar-free foods to prevent tooth decay, and prolongs fresh-cut fruit to shelf life.

Calcium propionate [4075-81-4] $\text{C}_6\text{H}_{10}\text{O}_4\text{Ca}$, **M 186.2, m dec on heating**. Crystallise this antifungal salt from water (solubility in g/100ml is 49 at 0° and 55.8 at 100°) by partial evaporation in a desiccator. It is slightly soluble in MeOH and EtOH, but insoluble in Me_2CO and hydrocarbon solvents. [*Beilstein* **2** H 238, **2** II 218, **2** III 516, **2** IV 702.] It is used as a food additive.

Calcium salicylate ($2\text{H}_2\text{O}$) [824-35-1, 59788-56-6] $\text{C}_{14}\text{H}_{14}\text{O}_8\text{Ca}$, **M 314.0(anhydr), pK₁²⁰ 3.08, pK₂²⁰ 13.43 (for acid)**. Recrystallise calcium salicylate from water (3ml/g) between 90° and 0°. [*Beilstein* **10** H 60, **10** II 33, **10** III 94, **10** IV 128.] Used in overbased greases [Patent US7407920 B2, applied US 10/911,132 aug 5, 2008.]

Carbonate ionophore I [ETH 6010] (**heptyl 4-trifluoroacetylbenzoate**) [129476-47-7] $\text{C}_{16}\text{H}_{19}\text{F}_3\text{O}_3$, **M 316.3, b 170°/0.02mm, d_4^{20} 0.909**. Purify the ionophore by flash chromatography (2g of reagent with 30g of Silica Gel 60) and elute with EtOAc/hexane (1:19). The fractions that absorb light at 260nm are pooled, evaporated and dried at room temperature (10.3 Torr). The oily residue is distilled in a bubble-tube apparatus (170°/0.02 Torr). Its IR (CHCl_3) has peaks at 1720, 1280, 940cm⁻¹, and its solubility in tetrahydrofuran is 50mg/0.5ml. It is a lipophilic **neutral ionophore selective** for carbonate as well as being an optical humidity sensor. [Behringer et al. *Anal Chim Acta* **233** 41 1990, DOI: 10.1016/S0003-2670(00)83459-5.]

Cerium (III) acetate (cerium triacetate) [537-00-8, 206996-60-3 ($x\text{H}_2\text{O}$)] $\text{C}_6\text{H}_9\text{O}_6\text{Ce}$, **M 317.3(anhydr), pK₁²⁵ 8.1 (9.29), pK₂²⁵ 16.3, pK₃²⁵ 26.0 (for Ce^{3+})**. Recrystallise it twice from anhydrous acetic acid, then pump dry under a vacuum at 100° for 8 hours. [*Beilstein* **2** I 50, **2** II 119, **2** III 196, **2** IV 115.]

Cesium oleate [31642-12-3] $\text{C}_{6}\text{H}_9\text{O}_6\text{Cs}$, **M 414.4**. Recrystallise cesium oleate from EtOAc, dry it in an oven at 40° and store it over P_2O_5 . It forms stable emulsions [Finkle et al. *J Am Chem Soc* **45** 2780 1923, DOI: 10.1021/ja01665a002.] [*Beilstein* **2** II 437, **2** III 1405.]

Cesium perfluoro-octanoate (cesium pentadecafluorooctanoate, CsPFO) [17125-60-9] ($\text{C}_8\text{F}_{15}\text{O}_2$)₃Cs, **M 546.0**. Recrystallise it from a butanol/petroleum ether mixture, dry it in an oven at 40° and store it over P₂O₅ under vacuum. It is prepared by neutralising an aqueous solution of pentadecafluorooctanoic acid {PFO [335-67-1], $\text{C}_8\text{HF}_{15}\text{O}$, M 414.1, m 40-50°, b 189-192°/atm, solubility in H₂O ~9.5g/100ml at ~25°} with Cs₂(CO₃)₃ {[54451-25-1] M 460.3(anhydr)}, evaporating to dryness in an oven at 100°, then recrystallising twice from *n*-BuOH. [For preparation and phase diagram by ¹³³Cs NMR see Boden et al. *J Phys Chem* **97**(29) 7678 1993, DOI: 10.1021/j100131a043.] [*Beilstein* **2** IV 994.] It is a water surfactant.

Chloro-tri-isopropoxy titanium (IV) [20717-86-6] $\text{C}_9\text{H}_{21}\text{ClO}_3\text{Ti}$, **M 260.6, m 45-50°, b 61-65°/0.1mm, d²⁵ 1.091g/ml**. When distilled under vacuum, the distillate sets slowly to a solid on standing. Stock reagents are made by dissolving the warm liquid in pentane, toluene, Et₂O, THF, CH₂Cl₂, and can be stored in a pure state or in solution under dry N₂ for several months. The reagent is **HIGHLY FLAMMABLE** (wear protective clothing), and is **hygroscopic** being readily hydrolysed by H₂O. Also commercially available in hexanes (1M, d²⁵ 1.091g/ml), and use syringes techniques and closed systems under N₂. [Reetz et al. *Chem Ber* **118** 1421 1985, DOI: 10.1002/cber.19851180412; Fieser **14** 87.] This reagent increases the region- and stereo- control in aldol reactions [Kanemasa et al. *Tetrahedron Lett* **34** 677 1993, DOI: 10.1016/S0040-4039(00)61650-7].

Chromium (0) hexacarbonyl [13007-92-6] $\text{Cr}(\text{CO})_6$, **M 220.1, m 130°(dec), >150°(dec), b 210°/atm(dec), d²⁰ 1.77**. Wash the complex with cold EtOH, then Et₂O, and allow it to dry in air. *Alternatively*, recrystallise it from dry Et₂O. This is best accomplished by placing the hexacarbonyl in a Soxhlet extractor and extracting exhaustively with dry Et₂O. Pure $\text{Cr}(\text{CO})_6$ is filtered off and dried in air. Completely colourless refracting crystals are obtained by sublimation at 40-50°/0.5mm in an apparatus where the collecting finger is cooled by Dry-ice and in which there is a wide short bore between the hot and cold sections to prevent clogging by the crystals. Loss of product in the crystallisation and sublimation is slight. It is important not to overdo the drying as the solid is appreciably volatile and **TOXIC** [vapour pressure is 0.04(8°), 1.0(48°) and 66.5(100°) mm]. Also, do not allow the Et₂O solutions to stand too long as a brown deposit is formed which is sensitive to light, and to avoid the possibility of violent decomposition. It sinters at 90°, decomposes at 130°, and **EXPLODES** at 210°. It is also soluble in CHCl₃, CH₂Cl₂ and THF. [Owen et al. *Inorg Synth* **3** 156 1950, DOI: 10.1002/9780470132340.ch42; Podall et al. *J Am Chem Soc* **83** 2057 1961, DOI: 10.1021/ja01470a008.] Useful precursor which mediates [5+5] cycloaddition reactions to provide phenanthrenes [Menon et al. *Tetrahedron* **63** 8788 2007, DOI: 10.1016/j.tet.2007.06.032]. **POISONOUS, possible carcinogen**.

Chromocene [bis(cyclopentadienyl) chromium II] [1271-24-5] $\text{C}_{10}\text{H}_{10}\text{Cr}$, **M 182.2, m 168-170°, 173°, d²⁵ 1.43g/cm³**. Chromocene forms dark red (scarlet) **pyrophoric** crystals on sublimation at 50°/0.1mm followed by resublimation at 75-90°/0.1mm. Although it is stable at least to 300°, it is readily oxidised in air, and effervesces slowly in H₂O to give cyclopentadiene. All operations should be carried out in a dry box. It decomposes in CCl₄ or CS₂, and for IR even grinding with nujol, KBr or KI causes some decomposition. [Wilkinson *J Am Chem Soc* **76** 209 1954, DOI: 10.1021/ja01630a053; Wilkinson et al. *J Inorg Nucl Chem* **3** 104 1956, DOI: 10.1016/0022-1902(56)80073-0; *Beilstein* **16** IV 1774, for some reactions see Beneš et al. *J Organomet Chem* **290** 147 1985, DOI: 10.1016/0022-328X(85)87428-3.]

Chromoionophore I [ETH 5294] [9-diethylamino-5-octadecanoyl-imino-5-*H*-benzo[a]-phenoxazine] [125829-24-5] $\text{C}_{38}\text{H}_{53}\text{N}_3\text{O}_2$, **M 583.9, m 91-93°, b ~680°/atm, d²⁵ 1.1g/ml, n_D²⁰ 1.57**. Purify it by flash chromatography (Silica Gel) and elute with EtOAc. The coloured fractions are pooled, evaporated and recrystallised from EtOAc. It is fluorescent with λ_{ex} 614nm; λ_{em} 663 (protonated). It is a **lipophilic fluorescent chromoionophore** and is a selectophore for K⁺ and Ca²⁺ ions. [Morf et al. *Anal Chem* **62** 738 1990, DOI: 10.1021/ac00206a018; Lapresta-Fernandez & Capitan-Vallvey *Anal Chim Acta* **706** 328 2011, DOI: 10.1016/j.aca.2011.08.042.]

Cobalt (II) meso-5,10,15,20-tetraphenylporphine complex (Co TTP) [14172-90-8] $\text{C}_{44}\text{H}_{28}\text{N}_4\text{Co}$, **M 671.7, m >300°**. It yields brown crystals from Et₂O or CHCl₃/MeOH (cf. *iron chloride complex*). It crystallises on extraction (Soxhlet) with *C₆H₆. It is soluble in most organic solvents except MeOH and petroleum ether. It has λ_{max} at 409nm and 524nm. [UV, IR: Rothmund & Menotti *J Am Chem Soc* **70** 1808 1948, DOI: 10.1021/ja01185a047; Thomas & Martell *J Am Chem Soc* **81** 5111 1959, DOI: 10.1021/ja01528a024; *Beilstein*

26 III 1960.] Used in a new synthetic approach for substituted dihydrooxepines [Sengül & Balci *JCS Perkin Trans 1* 2071 1997, DOI: 10.1039/A701105I.]

Cobaltic (III) acetylacetonate $[\text{Co}(\text{acac})_3]$ [21679-46-9] $(\text{C}_5\text{H}_7\text{O}_2)_3\text{Co}$, **M 356.3**, **m 211°**, **210-213°**, **211.5°**, Recrystallise it from C_6H_6 /petroleum ether and dry it in a vacuum. [Charles & Pawlikowski *J Phys Chem* **62** 440 1938, DOI: 10.1021/j150562a017; *Beilstein 1* H 783.] Used for the preparation of a new family of monoporphyrinates [Spyroulias et al. *JCS Chem Commun* 783 1997, DOI: 10.1039/A608013H].

Cobaltous acetate tetrahydrate [6147-53-1($4\text{H}_2\text{O}$), 71-48-7 (anhydr)] $(\text{C}_2\text{H}_3\text{O}_2)_2\text{Co} \cdot 4\text{H}_2\text{O}$, **M 249.1**, **177.0(anhydr)**, **m 140°($4\text{H}_2\text{O}$)**, **d²⁵ 1.705g/ml($4\text{H}_2\text{O}$)**, **n_D²⁰ 1.452($4\text{H}_2\text{O}$)**, **pK₁²⁵ 9.85 (for Co^{2+})**. Several recrystallisations from 50% aqueous acetic acid give the **tetrahydrate** (intense red crystals). It is converted to the **anhydrous** salt (pink crystals) by drying at 80°/1mm for 60 hours. It is soluble in EtOH, dilute acids, PhCO_2Et . [*Beilstein 2* IV 120, *Fieser 4* 99, **16** 95.]

Cobaltous acetylacetonate [14024-48-7, 123334-29-2 ($x\text{H}_2\text{O}$)] $(\text{C}_5\text{H}_7\text{O}_2)_2\text{Co}$, **M 257.2**, **m 165-170°**, **172°**. Recrystallise it from Me_2CO or MeOH and dry it in a vacuum. [*Beilstein 1* H 783; *Fieser 16* 26, **17** 87.]

12-Crown-4 (lithium ionophore V, 1,4,7,10-tetraoxacyclododecane) [294-93-9] $\text{C}_8\text{H}_{16}\text{O}_4$, **M 176.2**, **m 16°**, **17°**, **b 61-70°/0.5mm**, **d²⁵ 1.089g/cm³**, **n_D²⁰ 1.463**. The distilled crude ionophore has to be recrystallised from pentane at -20° to remove acyclic material. It is then dried over P_2O_5 . It complexes Li^+ . [Anet et al. *Acta Chem Scand* **27** 3395 1973, DOI: 10.3891/acta.chem.scand.27-3395; *Beilstein 19/11* V 334; *Fieser 6* 133, **9** 126.]

Cupric acetate [142-71-2 (anhydrous), 6046-93-1 (H_2O), 66923-66-8 ($x\text{H}_2\text{O}$)] $(\text{C}_2\text{H}_3\text{O}_2)_2\text{Cu}$, **M 181.6(anhydr)**, **m 115°(hydrate)**, **b 240°/atm(dec)**, **d²⁵ 1.882g/cm³ (H_2O)**, **n_D²⁰ 1.450(H_2O)**, **pK₁²⁵ 8.0**, **pK₂²⁵ 13.1 (for Cu^{2+})**. Recrystallise the blue $\text{Cu}(\text{OAc})_2$ twice from warm dilute acetic acid solutions (5ml/g) by cooling. Its solubility (g/100ml) in H_2O is 7.2 at ~10° and 20 at ~90°; and is soluble in EtOH, but slightly in Et_2O and glycerol. [Richardson 'Copper Compounds' in *Ullmann's Encyclopedia of Industrial Chemistry* 2005, Wiley-VCH, Weinheim, DOI: 10.1002/14356007.a07_567; *Beilstein 2* IV 94, 111; for applications see *Fieser 1* 159, **2** 84, **3** 65, **4** 105, **6** 138, **10** 103, **12** 140, **13** 85, **15** 99.]

Cupric benzoate [533-01-7] $\text{C}_{14}\text{H}_{10}\text{O}_4\text{Cu}$, **M 305.8**, **b 249.3°/760mm**, **d²⁵ 1.197g/cm³**. Recrystallise the blue solid from hot water. Dry it at ~90° for 3 hours. Its solubility in EtOH/ C_6H_6 (90%) at 25° is 0.1%. [Crawford & Stewart *J Chem Soc* 288 1953, DOI: 10.1039/JR9530000288; *Beilstein 9* I 60, **9** III 376, **9** IV 280.]

Cupric lactate monohydrate [814-81-3] $\text{C}_6\text{H}_{10}\text{O}_6\text{Cu} \cdot \text{H}_2\text{O}$, **M 259.7**. The **monohydrate** crystallises from hot H_2O (3ml/g) on cooling. [*Beilstein 3* II 203, **3** III 465, **3** IV 636.]

Cupric oleate [1120-44-1] $\text{C}_{36}\text{H}_{66}\text{O}_6\text{Cu}$, **M 626.5**. Crystallise cupric oleate from diethyl ether. Slightly soluble in EtOH and insoluble in H_2O . [*Beilstein 2* H 465, **2** I 196, **2** I202, **2** II 436, **2** III 1404, **2** IV 1646.] Also sold as a green-blue liquid fungicide for fruits and vegetables; and used as a dispersant, lubricant antioxidant and polyamide stabiliser.

Cupric phthalocyanine (phthalocyanine blue, Pigment Blue 15) [147-14-8] $\text{C}_{32}\text{H}_{16}\text{N}_8\text{Cu}$, **M 576.1**, **m 600°(dec)**, **CI 74160 ICI**. Precipitate it twice from concentrated H_2SO_4 by slow dilution with water. It has also been purified by two or three sublimations at 580° in an argon flow at 300-400Pa. Its solubility in H_2O is 0.1g/100ml at 20°, and has it λ_{max} at 678nm in H_2O . [*Beilstein 26* III/IV 4256.]

(1,5-Cyclooctadiene)(1,3,5-cyclooctatetraene)ruthenium $[\text{Ru}(\text{COD})(\text{COT})]$ [127382-91-6] $\text{C}_{16}\text{H}_{22}\text{Ru}$, **M 315.4**, **m 88-94°**, **92-94°**. It can be prepared directly under argon and pressure equalisation, by adding freshly distilled cycloocta-1,5-diene (12.5ml, COD, see [111-78-4, 1552-12-1]) in MeOH (5ml) to Zn dust (6g) in a flask which is placed in an ultrasound bath (Branson, Bransonic Model B220) thermostated at 70°. A solution of $\text{RuCl}_3 \cdot 3\text{H}_2\text{O}$ (530mg) in MeOH (12ml) is slowly added during 20 minutes to the refluxing COD mixture under N_2 flow and ultrasound. After addition is complete the mixture is sonicated for a further 2 hours under

reflux at 70°. The solids are filtered off and the filtrate is evaporated with a vacuum line. The residual brown oil is extracted with the minimum volumes of hexane (3 x, total ~70ml), placed on an Al₂O₃ (Merck 1079, or Brockman Activity II-III) column under N₂ and eluted with hexane. The yellow band is collected, the volatiles are removed *in vacuo* to give orange crystals (590mg, 93%). Analytically pure **Ru(COD)(COT)** [C₁₆H₂₂Ru] is obtained as yellow crystals upon recrystallisation (70-90% recovery) from hexane at ~ -78°/~6 hours preferably under N₂ or argon. [Itoh et al. *J Organomet Chem* **272** 179 1984, DOI: 10.1016/0022-328X(84)80465-9; Pertici et al. *JCS Dalton Trans* 1961 1980, DOI: 10.1039/DT9800001961.] The ¹H NMR (60MHz, *C₆D₆, TMS) has τ at 4.78 (dd, 2H), 5.22 (m, 2H), 6.21 (m, 2H), 7.08 (m, 4H), 7.78 (m, 8H), 8.36 (m, 2H) and 9.10 (m, 2H); compare with the ¹H NMR (60MHz, *C₆D₆, TMS) of **Ru(COD)₂** (see below) which has τ at 4.42 (t, 2H), 6.15 (dd, 4H), 6.45 (m, 4H), and 7.49–8.95 (m, 12H).

Alternatively, thermal isomerisation of **Ru(COD)₂** (75mg, see [63395-36-8]) in CDCl₃ (0.5ml) in an NMR tube filled with argon occurs when the sealed tube is heated at 70° for 12 minutes as observed by the spectrum. Evaporation of the solution and purification as above gave yellow-orange crystals of **Ru(COD)(COT)** in >85% yield with no deuterium incorporation as observed by NMR. Isomerisation in *C₆D₆ is very slow at 70° giving a mixture of three isomers after 5 hours [Itoh et al. *J Organomet Chem* **272** 179 1984, DOI: 10.1016/0022-328X(84)80465-9; see also Petrici et al. *JCS Dalton Trans* 1961 1980, DOI: 10.1039/DT9800001961]. A synthesis *via* a Grignard reduction of [RuCl₂(COD)]_n has also been described [Pelzer et al. *Chem Mater* **16** 4937 2004, DOI: 10.1021/cm049086b]. For its catalytic activity see Chapter 5, Catalysts-Part 1 [Mistudo et al. *J Am Chem Soc* **121** 1839 1999, DOI: 10.1021/ja9835741]. It is a key precursor for the preparation of stabilised Ru nanoparticles [Pan et al. *J Am Chem Soc* **123** 7584 2001, DOI: 10.1021/ja003961m; cf. Chapter 7].

Cyclopentadienyl iron(II) dicarbonyl dimer [di(cyclopentadienyl)tetracarbonyl-diiron], (C₅H₅Fe)₂(CO)₄ [38117-54-3] **C₁₄H₁₀O₄Fe₂, M 353.9, m 194° (dec) (sealed tube)**. This key precursor to a variety of cyclopentadienyl-carbonyl complexes is prepared from Fe(CO)₅ (10ml, [13463-40-6], **CARE**, poisonous, work in an efficient fume cupboard) by refluxing with an excess of dicyclopentadiene (10ml, cyclopentadiene dimer see 77-73-6) for *ca* 40 hours under a N₂ or argon atmosphere in the absence of light, removing the solvent, adding CHCl₃ (50ml), and removing any Fe₂(CO)₉ and insoluble material by centrifugation. The decanted supernatant is cooled in a Dry-ice/Me₂CO bath, the crystals are collected by centrifugation and recrystallised from CHCl₃ to give a ~30% yield (based on CO) of analytically pure dark reddish-purple crystals of **(C₅H₅Fe)₂(CO)₄, m 194° (dec) (sealed tube)**. The solid is stable to air and light, but solutions deteriorate under these conditions. It forms ferrocene (in 75% yield based on C₅H₅) when heated at 210°. It is readily soluble in EtOH, CHCl₃, pyridine, less soluble in CCl₄ and CS₂, and sparingly soluble in light petroleum giving red solutions in each case. It is insoluble in H₂O and is unaffected by it. The **measured molecular weight** by isothermal distillation is **368**. The IR has strong bands at λ_{max} at 4.95μ, 5.1 μ and 5.6 μ (in CCl₄) and 12.1 μ (CS₂). In the presence of O₂ a solution of the complex in EtOH/CHCl₃/concHCl/3hours provides red crystals of **C₅H₅Fe(CO)₂Cl** (from CHCl₃/petroleum ether 85:15, decomposing > 87°). This chloride is not ionic but gives the orange cation **C₅H₅Fe(CO)₂⁺** with AgNO₃/HNO₃, and yellow crystals of **C₅H₅Fe(CO)₂CN** (from CHCl₃, decomposing > 120°) by reaction with NaCN/MeOH/24 hours in 46% yield. [Piper, Cotton and Wilkinson *J Inorg Nucl Chem* **1** 165 1955, DOI: 10.1016/0022-1902(55)80053-X; *Beilstein* **16** IV 1826.] When **(C₅H₅Fe)₂(CO)₄** is treated with NaHg it provides the **anion C₅H₅Fe(CO)₂⁻** that can be alkylated with MeI to furnish **C₅H₅Fe(CO)₂Me** and related alkyl derivatives. [Aktogu et al. *J Organomet Chem* **262** 49 1984, DOI: 10.1016/S0022-328X(00)99122-8; R.B. King *Organometallic Synthesis Vol 1*, p 45, Academic Press, NY, 1965.] [*Beilstein* **16** IV 1826; for further applications see Fieser **8** 41.]

2,4-Cyclopentadien-1-yl lithium (lithium cyclopentadienide, CpLi) [16733-97-4] **C₅H₅Li, M 72.0**. CpLi is a white air and moisture sensitive solid which should be stored under N₂ preferably at low temperature. Like CpNa [4984-82-1] and CpK, it is handled in a dry box or Schlenk equipment in an inert atmosphere (compare with CpTi below). To avoid excessive manipulation, the reagents are sometimes prepared *in situ* in the reactions without being isolated. The solid is prepared in Schlenk equipment under a N₂ atmosphere by adding dropwise redistilled cyclopentadiene (2.34ml, 28.5mmol) to a solution of EtLi (28.5mmol) in *benzene (50ml) and stirred magnetically until precipitation of a voluminous white solid is complete. This is filtered, or centrifuged, off under N₂, washed with dry Et₂O and dried *in vacuo*. It is sparingly soluble in Et₂O but is soluble in THF. [Wagner & Ebel *Tetrahedron* **26** 5155 1970, DOI: 10.1016/S0040-4020(01)93167-9.] For the

preparation of a standard solution (use drybox, or Schlenk equipment under N₂) cyclopentadiene (~10.0mmol) in dry THF (7ml) is refluxed with excess of lithium metal (see Ford *J Am Chem Soc* **92** 2857 1970, DOI: 10.1021/ja00712a042) for 4 hours, filtered through a sintered glass funnel and diluted to a total volume of 10ml with THF which provides a 1.00M solution for further use. [Ford *J Organomet Chem* **32** 27 1971, DOI: 10.1016/S0022-328X(00)80156-4.] Its UV in THF has no maxima above 210nm, and its IR has ν_{\max} (solid in Nujol) at 3048 (m), 2906 (w), 1742 (m), 1629 (m), 1548 (w), 1513 (w), 1426 (m), 1364 (sw), 1304 (w), 1258 (w), 1166 (w), 1110 (w), 1003 (s), 935 (sw), 887 (sw) and 746 (ss) cm⁻¹; the ¹H NMR [100MHz, shift relative to β -CH₂ of THF (1.767ppm of 2% TMS in THF) at 27°] has a peak at δ 5.59 (at 1.0M) or 5.70 (at 0.01M) [Ford *J Organomet Chem* **32** 27 1971, DOI: 10.1016/S0022-328X(00)80156-4; the ¹³C NMR (22.6MHz, MeOCH₂CH₂OMe/*C₆D₆ 3:1, under N₂ at 30°, TMS) has a multiplet with δ centered at 103.59 (m, ¹J_{13C-H} 159.4 and 6.9Hz). [Fischer et al. *J Organomet Chem* **116** 65 1976, DOI: 10.1016/S0022-328X(00)87196-X.]

Cyclopentadienyl thallium [(η^5 -2,4-cyclopentadien-1-yl)thallium, thallium(I) cyclopentadienide, CpTl] [34822-90-7] C₅H₅Tl, **M 269.5**, blackens at ~60° in air and at ~230° in a sealed capillary, but does not melt below 270°. CpTl is purified by repeated sublimation through a glass plug at ~80°/0.005mm, 75°/0.1mm or 100-110°/10mm to give pale yellow acicular (small pointed needles) crystals. It is moderately stable in air but darkens superficially after several months. As it is somewhat light sensitive, it should be stored in dark brown screw cap containers, and thus keeps for prolonged periods of time. It reacts with some solvents, e.g. CHCl₃ and CCl₄ (forming TlCl), and CS₂ forming black and red materials), or is very poorly soluble, e.g. in THF, Et₂O and *C₆H₆. However, it is soluble enough in the latter solvents, even at low temperatures (e.g. -20°), to react almost completely to give high yields of products. ***It is a more convenient reagent for cyclopentadienylation than CpNa, CpK or CpLi because it is less basic, can be prepared in aqueous medium, can be prepared on a large scale, and is stable in air at room temperature for long periods.*** The cyclopentadiene ring of CpTl has IR in accord with C_{5v} symmetry, exhibiting one C—H stretching band at ν_{\max} 3101 cm⁻¹ in the gas phase at 150° to 249° which compares with ferrocene (3109 cm⁻¹), nickelocene (3107 cm⁻¹), di-cyclopentadienylmanganese (3101 cm⁻¹), di-cyclopentadienyl-magnesium(II) (3095 cm⁻¹), and benzene (3099 cm⁻¹), and is supported by theoretical considerations. The five membered ring is flat and ionic with the positive charge delocalised within it, and the Tl bears the counter negative charge and is centered on top of it. [Cotton & Reynolds *J Am Chem Soc* **80** 269 1958, DOI: 10.1021/ja01535a004; Roberts et al. *J Mol Spectr* **35** 476 1970, DOI: 10.1016/0022-2852(70)90188-8; Fritz *Chem Ber* **92** 780 1959, DOI: 10.1002/cber.19590920405.]

CpTl is conveniently prepared by dissolving or suspending a thallium(I) salt [TlCl 3.6g, TlBr 5.26g, TlI does not react, TlSCN 3.94g, TlNO₃ 4.00g or Tl(I) acetylacetonate 4.55g] in a solution of KOH (10g) in H₂O (100ml) in a Waring Blender (900ml jar capacity), adding cyclopentadiene (2.0ml, freshly distilled [542-92-7]) and stirring for 30 seconds. Longer stirring gives a fine suspension that is difficult to collect. The white product is filtered off, washed with EtOH (2 x 10ml), dried in a desiccator over anhydrous CaSO₄ to give CpTl in 91%, 93%, 0%, 99%, 97% and 87% yields respectively which is usually used without purification. The preparation can be scaled up to give ~50g of CpTl, and the preferred salt is TlBr. If desired, further purification can be performed by high vacuum sublimation as described above. [Hunt & Doyle *Inorg Nucl Chem Lett* **2** 283 1966, DOI: 10.1016/0020-1650(66)80043-0; Cotton & Reynolds *J Am Chem Soc* **80** 269 1958, DOI: 10.1021/ja01535a004; Corey et al. *J Am Chem Soc* **93** 1489 1971, DOI: 10.1021/ja00735a031.] **Note that all thallium compounds are POISONOUS and due precautions have to be exercised.**

CpTl is a source of the C₅H₅ ligand, and reactions are performed under N₂ with highly purified solvents. The following are a few of the many reported reactions of CpTl. It has been used to prepare ferrocene, cobaltocene, nickelocene, also complexes such as Cp₂TiCl₂ with TiCl₄ in *C₆H₆, or of Cp(MeOCp₂)Pd(II) from [CpPd(OMe)Cl]₂ with CpTl in *C₆H₆ [Hunt & Doyle *Inorg Nucl Chem Lett* **2** 283 1966, DOI: 10.1016/0020-1650(66)80043-0.], as well as compounds like the 1-methoxymethylcyclopenta-2,4-diene key intermediate for the synthesis of prostaglandins [Corey et al. *J Am Chem Soc* **93** 1489 1971, DOI: 10.1021/ja00735a031]; it reacts with [CpMo(NO)I₂]₂ in THF to give Cp₂Mo(NO)I where the iodine can be replaced [King *Inorg Chem* **7** 90 1968, DOI: 10.1021/ic50059a018]; CpTl reacted with electrophilic olefins, e.g. tetracyanoethylene, in MeCN or THF to form compounds of the type Tl⁺[Cp(CN)C(CN)₂]⁻ [Freeman & Sneddon *Inorg Chem* **19** 1125 1980, DOI: 10.1021/ic50207a005], and the reaction of CpTl (10-80% molar excess) with 7-chloronorbornadiene in dry diglyme at 150°/3-4 hours gave, after filtration of Tl salts, the hydrocarbon **hexahydro-3,4,7-methenocyclopenta[a]pentalene** in 8-12% yield from a one-step synthesis (a hydrocarbon which would have required several synthetic steps to prepare) [Battiste & Timberlake *J Org Chem* **42** 176 1977, DOI: 10.1021/

jo00421a043]. [Meister *Angew Chem* **69** 533 1957 DOI: 10.1002/ange.19570691606; *Beilstein* **16** IV 1690, Fieser **8** 138.]

6,6-Dibenzyl-14-crown-4 (lithium ionophore VI, 6,6-dibenzyl-1,4,8,11-tetra-oxa-cyclo-tetradecane) [106868-21-7] $C_{24}H_{32}O_4$, **M 384.5, m 102-103°**. Dissolve it in $CHCl_3$, wash this with saturated aqueous NaCl, dry ($MgSO_4$), evaporate and purify it by chromatography on silica gel and gradient elution with $*C_6H_6/MeOH$ followed by preparative reverse phase HPLC on an octadecyl silanised silica (ODS) column and eluting with MeOH. It can be recrystallised from MeOH (IR has ν_{max} at 1120 cm^{-1} , C-O-C in KBr). Its solubility in H_2O is 50mg/ml at $\sim 25^\circ$. [Kimura et al. *Anal Chem* **59** 2331 1987, DOI: 10.1021/ac00146a004; see also Maruyama et al. *JCS Perkin Trans 1* 2069 1986, DOI: 10.1039/P19860002069; and Tsukube et al. *JCS Perkin Trans 1* 1033 1986, DOI: 10.1039/P19860001033.] It complexes selectively with, and is a neutral carrier for, Li^+ ions; used for making ion selective electrodes with good selectivity over K^+ and Na^+ ions.

Di-*n*-butyltin (IV) oxide [818-08-6] $C_8H_{18}OSn$, **M 248.9, m >300° (starts dec at $\sim 210^\circ$), d $^{25} 1.6\text{g/cm}^3$** . The oxide is prepared by hydrolysis of di-*n*-butyltin dichloride with KOH. Hence wash it with a little aqueous M KOH, then H_2O and dry at $\sim 80^\circ/10\text{mm}$ until the IR is free from OH bands. [Cummins *Aust J Chem* **18** 98 1965, DOI: 10.1071/CH9650098; *Beilstein* **4** I 588; Fieser **12** 160, **13** 95, **15** 116, **16** 112; Cervantes et al. 'Organotin catalysts in Organosilicon Chemistry' *Appl Organometal Chem* **26** 157 2012, DOI: 10.1002/aoc.2832.]

Dicarbonyl(cyclopentadienyl)Co (I) [12078-25-0] $C_7H_5O_2Co$, **M 180.1, m -22°, b 7537-38.5°/2mm, 75°/22mm, b 139-140°(dec)/710mm, d $^{25} 1.35\text{g/cm}^3$** . Best distilled in an atmosphere of CO in a vacuum. The red brown liquid decomposes slightly on distillation even in a vacuum to liberate some CO. Operations should be performed in an efficient fume cupboard. It is soluble in organic solvents and stable in air but decomposes slowly in sunlight and rapidly under UV. [Piper et al. *J Inorg Nucl Chem* **1** 165 1955, DOI: 10.1016/0022-1902(55)80053-X; King & Stone *Inorg Synth* **7** 99 1963, DOI: 10.1002/9780470132388.ch31; *Beilstein* **16** IV 1827; for applications see Fieser **5** 172, **6** 153, **7** 84, **12** 160, **13** 96, **14** 116, **16** 112, **17** 102.] It catalyses the cyclotrimerisation of alkynes [Cammak et al. *J Org Chem* **61** 4798 1996, DOI: 10.1021/jo960143x], and promotes [2+2+2]-cyclisation of allenediynes to tricyclic steroid skeletons [Petit et al. *Tetrahedron* **62** 10582 2006, DOI: 10.1016/j.tet.2006.05.091]. **TOXIC**.

Dichloro(2,2':6', 2''-terpyridine)platinum(II) dihydrate [151120-25-1] $C_{15}H_{11}Cl_2N_3Pt \cdot 2H_2O$, **M 535.3, decomposes at 240-260°**. The aqueous filtrate from the reaction between terpyridyl (2.3g) and potassium platinochloride (4.0g) in H_2O at 90° for 6 hours, is evaporated, cooled, and treated with HCl whereby the red chloride separates as the *dihydrate*, whereas the black *trihydrate* slowly crystallises from a cold aqueous solution and is air-dried. It is converted to the *dihydrate* in a desiccator over H_2SO_4 , by washing with EtOH, heating in H_2O (slowly), or by precipitating from a warm aqueous solution with hydrochloric acid. This salt is not decomposed by boiling HCl, and is insoluble in most organic solvents. It *intercalates* with helical and double stranded DNA [Lippard *Acc Chem Res* **11** 211 1987, DOI: 10.1021/ar50125a006]. [Morgan & Burstall *J Chem Soc* 1498 1934, DOI: 10.1039/JR9340001498; *Beilstein* **26** IV 260.]

Diethyl aluminium chloride [96-10-6] $C_4H_{10}ClAl$, **M 120.6, m -75.5°, -50°, b 106.5-108°/24.5mm, 125-126°/50mm, d $^{25} 0.961\text{g/ml}$** . Distil it from excess dry NaCl (to remove ethyl aluminium dichloride) in a 50-cm column containing a heated nichrome spiral. **HIGHLY FLAMMABLE AND TOXIC**. [*Beilstein* **4** IV 4403; for applications see Fieser **4** 144, **15** 2, **16** 1, **17** 204.]

***N,N'*-Diheptyl-*N,N'*-5,5-tetramethyl-3,7-dioxanonanediarnide** [lithium ionophore I (ETH 149)] [58821-96-8] $C_{25}H_{50}N_2O_4$, **M 442.7**. Purify it by chromatography on Kieselgel using $CHCl_3$ as eluent (IR has ν_{max} at 1640cm^{-1}). [Kirsch et al. *Helv Chim Acta* **60** 2326 1977, DOI: 10.1002/hlca.19770600723; Simon & Carafoli *Methods Enzymol* **56** 439 1977, DOI: 10.1016/0076-6879(79)56043-1.]

Diphenylmercury (II) [587-85-9] $C_{12}H_{10}Hg$, **M 354.8, m 121-124°, 125.5-126°, 128-129°, b 204°/atm, d $^{25} 2.32\text{g/ml}$** . Sublime Ph_2Hg , then crystallise it from nitromethane or ethanol. If phenylmercuric halides are present, they can be converted to phenylmercuric hydroxide which, being much more soluble, remain in the

alcohol or *benzene used for crystallisation. Thus, crude material (10g) is dissolved in warm ethanol (*ca* 150ml) and shaken with moist Ag₂O (*ca* 10g) for 30 minutes, then heated under reflux for 30 minutes and filtered hot. Concentrating the filtrate by evaporation gives diphenylmercury, which is then recrystallised from *benzene [Blair et al. *J Chem Soc* 3174 1959, DOI: 10.1039/JR9590003174]. [Calvery *Org Synth Coll Vol* 1 228 1941, DOI: 10.15227/orgsyn.009.0054; *Beilstein* 16 IV 1702.] **POISONOUS.**

Disodium calcium ethylenediaminetetraacetate [662-33-9, 39208-14-5, 23411-34-9 ($2\text{H}_2\text{O}$)] $\text{C}_{10}\text{H}_{12}\text{N}_2\text{O}_8\text{CaNa}_2$, M 374.3, (see pKs for EDTA in entry below). Dissolve it in a small amount of water, filter it and precipitate it with excess EtOH. Dry it at 80°. [*Beilstein* 4 IV 2451.] It is a useful buffer.

Disodium dihydrogen ethylenediaminetetraacetic acid ($2\text{H}_2\text{O}$) [6381-92-6] $\text{C}_{10}\text{H}_{14}\text{N}_2\text{O}_8\text{Na}_2 \cdot 2\text{H}_2\text{O}$, M 372.2, m 248°(dec), pK₁²⁵ 0.26, pK₂²⁵ 0.96, pK₃²⁵ 2.60, pK₄²⁵ 2.67, pK₅²⁵ 6.16, pK₆²⁵ 10.26 (see EDTA). Analytical reagent grade material can be used as primary standard after drying at 80°. Commercial grade material can be purified by crystallisation from water or by preparing a 10% aqueous solution at room temperature, then adding ethanol slowly until a slight permanent precipitate is formed, filtering, and adding an equal volume of ethanol. The precipitate is filtered off onto a sintered-glass funnel, is washed with acetone, followed by diethyl ether, and dried in air overnight to give the *dihydrate*. Drying at 80° for at least 24 hours converts it to the *anhydrous* form. [*Beilstein* 4 IV 2451.] It is a useful buffer.

Disodium magnesium ethylenediaminetetraacetate [14402-88-1] $\text{C}_{10}\text{H}_{12}\text{N}_2\text{O}_8\text{Na}_2\text{Mg} \cdot x\text{H}_2\text{O}$, M 358.5, pK₁²⁵ 0.26, pK₂²⁵ 0.96, pK₃²⁵ 2.60, pK₄²⁵ 2.67, pK₅²⁵ 6.16, pK₆²⁵ 10.26 (see EDTA). Dissolve it in a small amount of water, filter and precipitate it with an excess of MeOH. Dry it at 80°. [*Beilstein* 4 IV 2450.] It is a useful buffer.

Disodium succinate [150-90-3] $\text{C}_4\text{H}_4\text{O}_4\text{Na}_2$, M 162.1, m 120°. Crystallise it twice from water (1.2ml/g, its solubility in HO is 30g/100ml at 25°) and dry it at 125°. It has been freed from other metal ions by passage of a 0.1M solution through a column of Dowex resin A-1 (Na form). It is *hygroscopic*. [*Beilstein* 2 H 606, 2 IV 1908.]

Di-*p*-tolylmercury [50696-65-6, 537-64-4] $\text{C}_{14}\text{H}_{14}\text{Hg}$, M 382.8, m 244-246°. Crystallise it from xylene. [Whitmore et al. *Org Synth Coll Vol* 1 231 1941, DOI: 10.1002/0471264180.os003.18; *Beilstein* 16 H 947, 16 I 667, 16 III 1329, 16 IV 1705.]

Di-*p*-tolyl phenylphosphonate [94548-75-1] $\text{C}_{20}\text{H}_{19}\text{O}_3\text{P}$, M 388.3, n_D²⁵ 1.5758. Purify as described under diisooctyl phenylphosphonate.

Eosin B (Eosine bluish, Eosin Scarlet, Acid Red 91, 4',5'-dibromo-2',7'-dinitrofluorescein disodium salt) [548-24-3] $\text{C}_{20}\text{H}_6\text{Br}_2\text{N}_2\text{O}_9\text{Na}_2$, M 624.1, λ_{max} 395nm and 514nm, CI 45400. Free it from inorganic halides by repeated crystallisation from butan-1-ol. [See as below.] [*Beilstein* 19/6 V 469.]

Eosin Y (as di-Na salt) (2',4',5',7'-tetrabromofluorescein di-Na salt) [17372-87-1] $\text{C}_{20}\text{H}_6\text{Br}_4\text{O}_5\text{Na}_2$, M 691.9, λ_{max} 518nm. Dissolve it in water and precipitate it by adding dilute HCl. The precipitate is washed with water, crystallised from ethanol, then dissolved in the calculated amount of dilute NaOH solution and evaporated to dryness on a water-bath. The purified disodium salt is then recrystallised twice from ethanol [Parker & Hatchard *Trans Faraday Soc* 57 1894 1961, DOI: 10.1039/TF9615701894]. [*Beilstein* 19 III/IV 2917.]

Ethylmercuric chloride [107-27-7] $\text{C}_2\text{H}_5\text{ClHg}$, M 265.1, m 192.5°, 193-194°. Mercuric chloride can be removed by suspending ethylmercuric chloride in hot distilled water, filtering with suction onto a sintered-glass crucible and drying it. Then crystallise it from ethanol and sublime it under reduced pressure. It can also be crystallised from water. [Marvel et al. *J Am Chem Soc* 47 3009 1925, DOI: 10.1021/ja01689a026.] **Ethylmercuric iodide** [2440-42-8] $\text{C}_2\text{H}_5\text{IHg}$, M 356.6, m 182°, 186°. Crystallise it once from water (50ml/g). [See previous entry, Marvel et al. *J Am Chem Soc* 4 3009 1925, DOI: 10.1021/ja01689a026.] **POISONOUS.**

Ethylzinc (zinc diethyl) [557-20-0] $\text{C}_4\text{H}_{10}\text{Zn}$, M 123.5, m -28°, -33.8°, b 116.8°/761mm, d₂₅²⁰ 1.205/ml, n_D²⁰ 1.498. The presence of EtI in the liquid can be detected by its absorption spectrum. This can be removed by 2 or 3 passages over a Zn/Cu couple at ~150°. Any hydrocarbon impurities are removed by distillation at ~760mm in an inert atmosphere as it is flammable. It is moisture sensitive, hydrolyses to give ethane so store it in sealed

ampoules under N₂ or argon. It is commercially available also as a ~15% solution in toluene (d_4^{20} 0.915) or ~1M *n*-hexane (d_4^{20} 0.726). Solutions should be stored under dry N₂ or argon, and if they contain some precipitate, the solutions should be filtered through a sintered frit under inert gas pressure. *Alternatively*, siphon carefully the required volume of supernatant. [Noller *Org Synth Coll Vol* **2** 184 1943, DOI: 10.15227/orgsyn.012.0086; Bamford et al. *J Chem Soc* 468 1946, DOI: 10.1039/JR9460000468; *Beilstein* **4** III 1999, **4** IV 4423; for applications see Fieser **1** 253, **2** 134, **3** 257, **4** 153, **5** 219, **6** 194, **11** 182, **17** 115.]

Ethynyl tri-*n*-butylstannane [994-89-8] C₁₄H₂₈Sn, **M 315.1**, **b 76°/0.2mm**, **130-135°/0.7mm**, **200°/2mm**, **d**²⁵ **1.089g/ml**, **n_D²⁰ 1.4770**. Purify the stannane by dissolving the reagent (*ca* 50g) in heptane (250ml), washing it with H₂O (100ml), drying (MgSO₄), evaporating and distilling in a vacuum. It has IR with ν_{\max} at 3280 (= C-H), 2950, 2850, 2005 (C≡C), 1455, 1065 and 865cm⁻¹. [Bottaro et al. *J Org Chem* **46** 5221 1981, DOI: 10.1021/jo00338a035; Stille & Simpson *J Am Chem Soc* **109** 2138 1987, DOI: 10.1021/ja00241a035; for interaction with di-*n*-butylstannane see Nesmeyanov & Borisov *Bull Acad Sci USSR, Div Chem Sci* **16** 227 1967, DOI: 10.1007/BF00907149 (translated in *January*, 1967).]

Europium (III) acetate dihydrate [62667-64-5] C₆H₉O₆Eu. 2H₂O, **M 329.1(anhydr)**, **383.1**, **pK_i²⁵ 8.31** (for **aquo Eu³⁺**). Recrystallise it several times from water [Ganapathy et al. *J Am Chem Soc* **108** 3159 1986, DOI: 10.1021/ja00272a001]. [*Beilstein* **2** II 119.] See lanthanide shift reagents in 'Aliphatic Compounds', Chapter 3

Ferrocene [102-54-5] C₁₀H₁₀Fe, **M 186.0**, **m 173-174°**, **b 249°/atm**, **λ_{\max} 358nm**. Purify ferrocene by crystallisation from pentane or cyclohexane (also *C₆H₆ or MeOH can be used). It is moderately soluble in Et₂O and sublimes readily above 100°. Crystallisation from EtOH gave material **m 172.5-173°**. [Wilkinson *Org Synth Coll Vol* **4** 473 1963, DOI: 10.15227/orgsyn.036.0031; Miller et al. *J Chem Soc* 632 1952, DOI: 10.1039/JR9520000632.] It has also been crystallised from methanol and sublimed *in vacuo*. [Saltiel et al. *J Am Chem Soc* **109** 1209 1987, DOI: 10.1021/ja00238a034; *Beilstein* **16** IV 1783.]

Ferrocene carboxaldehyde [12093-10-6] C₁₁H₁₀OFe, **M 214.1**, **m 117-120°, 118-120°, 121°, 124.5°**. The aldehyde forms red crystals from heptane/CH₂Cl₂, EtOH or petroleum ether and sublimes at 70°/1mm. The **cyanohydrin** has **m 104°** (from *C₆H₆/EtOH). The **semicarbazone** has **m 217-219°(dec)** after recrystallisation from aqueous EtOH. The **oxime** provides two isomers from petroleum ether *viz* **m 96-99°** and **m 155°**. The **O-acetyloxime** has **m 80-81°** after recrystallisation from hexane [Lindsay & Hauser *J Org Chem* **22** 355 1957, DOI: 10.1021/jo01355a001]. The **2,4-dinitrophenylhydrazone** has **m 248°(dec)**. [*Beilstein* **16** IV 1798, Graham et al. *J Am Chem Soc* **79** 3416 1957, DOI: 10.1021/ja01570a027; Broadhead et al. *J Chem Soc* 650 1958, DOI: 10.1039/JR9580000650.] **Chiral** ferrocene aziridinylmethanols were prepared for selective azomethine ylide cycloaddition reactions [Dogan et al. *Org Lett* **8** 4687 2006, DOI: 10.1021/ol061521f].

Ferrocene carboxylic acid [1271-42-7] C₁₁H₁₀O₂Fe, **M 230.1**, **m 210°(dec)**, **225-230°(dec)**, **pK_a²⁰ 4.4 (H₂O)**, **6.29 (68% aqueous MeOH)**. The acid crystallises as yellow crystals from petroleum ether (**m 225-230°dec**), CHCl₃ (**m 208.5°dec**), toluene/petroleum ether (**m 195-205°dec**), or aqueous ethanol. [Matsue et al. *J Am Chem Soc* **107** 3411 1985, DOI: 10.1021/ja00298a003.] The **acid chloride** **m 49°** crystallises from pentane, and has UV with λ_{\max} at 458nm [Lau & Hart *J Org Chem* **24** 280 1959, DOI: 10.1021/jo01084a647]. The **methyl ester** crystallises from aqueous MeOH with **m 70-71°**. The **anhydride** has **m 143-145°** when recrystallised from petroleum ether [Acton & Silverstein *J Org Chem* **24** 1487 1959, DOI: 10.1021/jo01092a024]. The **amide** has **m 168-170°** when crystallised from CHCl₃/Et₂O or **m 167-169°** when crystallised from *C₆H₆/MeOH. [Reeves *Org Synth* **56** 28 1977, DOI: 10.15227/orgsyn.056.0028; Arimoto & Haven *J Am Chem Soc* **77** 6295 1955, DOI: 10.1021/ja01628a068; Benkeser et al. *J Am Chem Soc* **76** 4025 1954, DOI: 10.1021/ja01644a047.] [Hall et al. *J Am Chem Soc* **90** 4972 1968, DOI: 10.1021/ja01020a034; *Beilstein* **16** IV 1807.] Esterification of alcohols and phenols with this acid gave esters used for GCMS analysis [Wasinski & Andersson *J Chromatogr A* **1157** 376 2007, DOI: 10.1016/j.chroma.2007.04.060].

Ferrocene-1,1'-dicarboxylic acid [1293-87-4] C₁₂H₁₀O₄Fe, **M 274.1**, **m >250°(dec)**, **>300°**, **pK_i²⁵ 3.9**, **pK₂²⁵ 5.3**. The dicarboxylic acid crystallises in orange-yellow crystals from AcOH and sublimes above 230°. The **monomethyl ester** has **m 147-149°** [Nesmeyanov & Reutov *Dokl Acad Nauk USSS* **115** 518 1957, *Chem Abstr*

26 469 1960]. The *dimethyl ester* has **m 114-115°** [Woodward et al. *J Am Chem Soc* **74**, 3458 1953, DOI: 10.1021/ja01133a543]. The *diacid chloride* has **m 92-93°** when recrystallised from petroleum ether. [Nesmeyanov & Reutov *Dokl Acad Nauk SSSR* **120** 1267 1958, Kazitsyna et al. *Dokl Acad Nauk SSSR* **127** 333 1959, *Beilstein* **16** IV 1811.]

Ferrocene-1,1,-dimethanol [1291-48-1] $C_{12}H_{14}O_2Fe$, **M 246.1**, **m 107-108°**. The diol is obtained from the diacid by $LiAlH_4$ reduction and recrystallised from Et_2O /petroleum ether. [Rinehart et al. *J Am Chem Soc* **82** 4111 1960, DOI: 10.1021/ja01500a074; *Beilstein* **16** IV 1795.]

1-(Ferrocenyl)ethanol (α -methylferrocenemethanol) [1277-49-2] $C_{12}H_{14}OFe$, **M 230.1**, **m 73-75°, 76-77°, 76-79°**. This versatile reagent is obtained by reduction of acetylferrocene (22.8g, see [1271-55-2]) in dry Et_2O (500ml) with $LiAlH_4$ (1.9g) solution in anhydrous Et_2O and refluxing for 2 hours. Excess of hydride is destroyed with $EtOAc$, the mixture is treated with a solution of NH_4Cl (28g) in H_2O , stirred for 0.5 hours at 0°, filtered, the organic layer is washed twice with H_2O , dried ($MgSO_4$), and evaporated to dryness. The residue (20.5g, 89%) is recrystallised from Et_2O /petroleum ether to give the *carbinol* as orange rods. It is free from a $C=O$ band but has a strong broad OH band in the IR spectrum. The *acetyl derivative* (Ac_2O /pyridine/0°/15 hours), after sublimation at 60°/0.2mm, has **m 67-68°** and an IR band at 5.78μ (s, $C=O$), but no band in the OH region. [Arimoto & Haven *J Am Chem Soc* **77** 6295, DOI: 10.1021/ja01628a068; *Beilstein* **16** IV 1975.] Ligand in $Ti(NEt_2)_4$ -catalysed intramolecular hydroamination of terminal alkynes (Markownikov vs anti-Markownikov addition) [Tillack et al. *Eur J Org Chem* 5001 2005, DOI: 10.1002/ejoc.200500423].

Germanium tetraethoxide [14165-55-0] $C_8H_{20}O_4Ge$, **M 252.8**, **m -72°, b 54.5°/5mm, 71-72°/11mm, 188-190°/722mm, d^{25}_D 1.1288g/ml, n^{20}_D 1.407**. Distil $Ge(OEt)_4$ through a 10cm Vigreux column under reduced pressure. Alternatively, distil it through a Fenske glass helices column fitted with a total condensation variable take-off stillhead. Fractionate it under reduced pressure using a reflux ratio of 10:1. [Johnson & Fritz *J Am Chem Soc* **75** 718 1953, DOI: 10.1021/ja01099a061; Bradley et al. *J Chem Soc* 4916 1956, DOI: 10.1039/JR9560004916; *Beilstein* **1** IV 1308.] It is readily hydrolysed so use strictly anhydrous conditions.

Hexabutyldistannane {hexabutylditin, bis[(tributyl)tin]} [813-19-4] $C_{24}H_{54}Sn_2$, **M 580.4**, **b 160-162°/0.3mm, 198°/10mm, d^{25}_D 1.1520g/cm³, n^{20}_D 1.5120**. Purify bis[(tributyl)tin] by distilling it in a vacuum and store it in the dark. Alternatively, purify by reverse-phase flash chromatography using a C-18 column. ¹¹⁹Sn NMR recommended δ -83ppm, $^1J(^{119}Sn-^{119}Sn)$ 2748Hz [Mitchell e-EUROS *Encyclopedia of Reagents for Organic Synthesis* 15 April 2001, DOI: 10.1002/047084289X.rh002]. [Shirai et al. *Yakugaku Zasshi* **90** 59 1970, *Chem Abstr* **72** 90593 1970, *Beilstein* **4** I 590; for applications see Fieser **14** 173, **16** 174.] This reagent was used in the radioiodination of styrylpyridines [Qu et al. *J Med Chem* **50** 2157 2007, DOI: 10.1021/jm070025+], and to stannylate aryl halides for **Stille coupling** [Bourderieux & Routier *Tetrahedron* **63** 9465 2007, DOI: 10.1016/j.tet.2007.06.118]. **TOXIC substance.**

Hexamethylditin {hexamethyldistannane, bis[(trimethyl)tin]} [661-69-8] $C_6H_{18}Sn_2$, **M 327.6**, **m 23-24 °, 23.5°, b 85-88°/45mm, 182°/756mm, d^{25}_D 1.58g/ml**. Wash bis[(trimethyl)tin] with H_2O and extract with $*C_6H_6$, dry by filtering through powdered Na_2SO_4 , remove $*C_6H_6$ on a rotary evaporator and fractionally distil the oily residue under vacuum (**b 85-88°/45mm**). *It boils at ca 182° at atmospheric pressure, but it cannot be distilled in air because the hot vapours 'flash' in the condenser.* [Kraus & Sessions *J Am Chem Soc* **47** 2361 1925, DOI: 10.1021/ja01686a015; Morris & Selwood *J Am Chem Soc* **63** 2509 1941, DOI: 10.1021/ja01854a056; Pedley et al. *Trans Faraday Soc* **53** 1612 1957, DOI: 10.1039/TF9575301612; *Beilstein* **4** IV 4346; for applications see Fieser **6** 273, **13** 142, **17** 143.] Aryl tin compounds were prepared with this reagent for microwave-assisted **Stille cross-coupling** with halogenated pyridines [Dehlinger et al. *Tetrahedron Lett* **47** 8973 2006, DOI: 10.1016/j.tetlet.2006.10.015], and for copper-promoted *O*-arylation of phenols [Vakalopoulos et al. *Tetrahedron Lett* **47** 8607 2006, DOI: 10.1016/j.tetlet.2006.09.095].

Hexarhodium(0) hexadecacarbonyl [28407-51-4] $Rh_6(CO)_{16}$, **M 1065.6**, **m 220° (dec, in air), 235°, d^{20}_4**

2.87. It slowly loses CO when heated in air, but may be regenerated by heating at 80-200° in the presence of CO at 200 atmospheres pressure for 15 hours, preferably in the presence of Cu. It forms **black crystals** which are insoluble in hexane. It has bands at 2073, 2026 and 1800 cm⁻¹ in the IR. [Hieber & Lagally *Z Anorg Allgem Chem* **251** 96 1963, DOI: 10.1002/zaac.19432510110; Corey et al. *J Am Chem Soc* **85** 1202 1963, DOI: 10.1021/ja00891a040; DOI: 10.1021/ja00891a040; James et al. *Inorg Synth* **16** 49 1976, DOI:10.1002/9780470132470.ch15; Doyle et al. *Tetrahedron Lett* **22** 1783 1981, DOI: 10.1016/S0040-4039(01)90438-1]. Useful catalyst. **POISONOUS.**

Iron(II) acetylacetonate [iron(II) bis(2,4-pentanedionato-κO2,κO4), Fe(2⁺)(acac)₂] [14024-17-0] C₁₀H₁₄O₄Fe, M 254.1, m 175°(dec). The preparation and handling of this complex should be done in a dry box or Schlenk equipment under dry pre-purified N₂; and solvents should be de-gassed prior to use. Fe(acac)₂ is prepared by adding FeCl₂·4H₂O (25g) in degassed H₂O (50ml) containing a small amount of sodium dithionite to reduce any Fe(III) impurities, to a degassed aqueous solution (300ml) containing piperidine (27.3ml) and redistilled acetylacetone (28.1ml, [123-54-6]). After 15 minutes the resulting precipitate is filtered off and washed with degassed H₂O, EtOH then Et₂O and the yellow-brown powder [presumed to be Fe(acac)₂·2H₂O Emmert & Jarczyński *Chem Ber* **64** 1072 1931, DOI: 10.1002/cber.19310640521] is transferred into a sublimation apparatus with a large cold finger and pumped dry at 10⁻³ mm for 6 hours; and the hydrated complex is then heated at 90°/10⁻³mm for at least 12 hours to dehydrate it and the orange-brown solid is sublimed twice at 165-175°/10⁻³mm. The small orange-brown crystals that sublime at first become darker as the crystals grow, giving finally larger black clusters which are analytically pure anhydrous solid. **Note** that the complex is very sensitive to O₂ in air, darkens in colour upon oxidation, and great care should be exercised to avoid oxidation when using it. It may be **pyrophoric** when dry. Its **molecular weight** (cryoscopic and ebullioscopic) in *C₆H₆ increases as the concentration increases reaching a maximum for a hexamer, and the data are supported by the UV absorption spectra. [Buckingham et al. *Aust J Chem* **20** 281 1967, DOI: 10.1071/CH9670281; Dwyer & Sargeson *Pr J Soc NSWales* **90**, 141, 142 1956; for the formation constant see Izatt et al. *J Phys Chem* **59** 80 1955, DOI: 10.1021/j150523a022; [Murray Jr e-EUROPS *Encyclopedia of Reagents for Organic Synthesis* 15 October 2010, DOI: 10.1002/047084289X.rn01225]. *Beilstein* **1** III 3122, **1** IV 3675.] **Note: iron(1+) bis(2,4-pentanedionato-κO2,κO4)** has [20149-10-4], see Chapter 5, Catalysts-Part 1 for Fe(III)(acac)₃, [14024-18-1].

Iron(II) bis(1,1,1,5,5,5-hexafluoropentan-2,4-dionate) [iron(II) bis(1,1,1,5,5,5-hexafluoro-2,4-pentanedionato-κO2,κO4), Fe(2⁺)(facac)₂, Fe(hfacac)₂] [28736-68-7] C₁₀H₂F₁₂O₄Fe, M 470.1, mp 40°. The preparation and handling of this complex [cf. Fe(acac)₂, 14024-17-0] should be carried out in a dry box or Schlenk equipment under dry pre-purified N₂, and solvents should be de-gassed prior to use. Fe(hfacac)₂ is prepared by adding FeSO₄·7H₂O (11.8g) in H₂O (50ml) slowly to a mixture of redistilled hexafluoroacetylacetone (12.2ml, see [1522-22-1]) and piperidine (8.5ml), whereby a dark violet precipitate separates immediately. This solid is filtered off, washed with a large volume of H₂O (1000 ml), placed in a sublimation apparatus and evacuated at 10⁻³mm at ~25° for hours. The temperature is slowly raised to 40° and kept at 45-50° and 10⁻³mm for 3 days, and the dark violet crystals that formed on the cold finger are collected, and if necessary are ground to a fine powder and re-sublimed to give analytically **pure Fe(hfacac)₂** (IR is free of bands at 3300-3500 cm⁻³, i.e. absence of H₂O). It requires a much longer period of evacuation than Fe(acac)₂ for dehydration. The strong electron withdrawing effects of the CF₃ groups make this complex relatively more stable to aerial oxidation than Fe(acac)₂. It can be recrystallised from very dry *C₆H₆ without alteration, where its molecular weight (cryoscopic and ebullioscopic) increases with increase in concentration up to a value of 1.7 (compare with 6 for Fe(acac)₂ above). [Buckingham et al. *Aust J Chem* **20** 281 1967, DOI: 10.1071/CH9670281; for crystal structure see Dickman *Acta Cryst C* **56**(1) 58 2000, DOI: 10.1107/S0108270199013487.]

Iron(II) dicyanobis(2,2'-dipyridine) trihydrate [Fe(bpy)₂(CN)₂·3H₂O] [15603-10-8] C₂₂H₁₆N₆Fe·3H₂O, M 474.3, dec. on heating. The iron complex is prepared from 2,2'-dipyridine (4.7g, 30mmol) and ferrous ammonium sulfate hexahydrate (3.9g, 10mmol) in H₂O (400ml) which are heated to just below boiling, and a freshly prepared solution of KCN (10g) in H₂O (20ml) is added all at once to it, stirred, and the hot mixture is allowed to cool to room temperature. The complex separates as very dark violet (almost black) fine crystals rapidly, but is allowed to stand for 1 hour. The solid is collected and washed liberally with H₂O. Unused bipyridine is recovered from the filtrate by extraction with Et₂O. The iron complex can be purified by dissolving

it in concentrated H_2SO_4 (25ml) and slowly diluting it with H_2O (800ml, CARE as it will warm up) with stirring. The crystalline solid is collected washed free from H_2SO_4 with small volumes of H_2O , sucked dry, rinsed with Me_2CO and dried *in vacuo* to give analytically **pure trihydrated complex** (3.5g, 75%). It is soluble in H_2O to give a pale red solution, in EtOH to give a deep red-violet liquid, and in CHCl_3 to give a deep blue colour; but is only very slightly soluble in basic solution. Large amounts can be dissolved in concentrated H_2SO_4 to give the yellow di-protonated species which on gradual dilution with H_2O the solution changes its hue to orange then to red and finally to the dark violet colour of the original crystals. It is a diamagnetic Fe(II) complex which is readily oxidised to the corresponding Fe(III) complex and can be titrated with Ce(IV) sulfate in H_2SO_4 . On titration with HClO_4 in anhydrous AcOH (**Care** due to possible **explosion**; Schilt *J Am Chem Soc* **82** 5779 1960, DOI: 10.1021/ja01507a004), the first equivalence point leads to a mono-protonated orange species, and at the second equivalence point (not quite as sharp as the first) it provides the yellow di-protonated species. The electronic (Schilt *J Am Chem Soc* **82** 3000 1960, DOI: 10.1021/ja01497a007; Schilt *J Am Chem Soc* **82** 5779 1960, DOI: 10.1021/ja01507a004; Madeja & König *J Inorg Nucl Chem* **25** 377 1963, DOI: 10.1016/0022-1902(63)80188-8; and vibrational spectra (Schilt *Inorg Chem* **3** 1323 1964, DOI: 10.1021/ic50019a029) have been reported, and the **cis-configuration** has been deduced from IR (Hamer & Orgel *Nature* **190** 439 1961, DOI: 10.1038/190439a0; Schilt *Inorg Chem* **3** 1323 1964, DOI: 10.1021/ic50019a029) and stereochemical (Madeja *Chem Zvesti* **19** 186 1965) considerations. [Schilt et al. *Inorg Synth* **12** 247 1970, DOI: 10.1002/9780470132432.ch43.]

Iron enneacarbonyl (di-iron nonacarbonyl) [15321-51-4] $\text{Fe}_2(\text{CO})_9$, M 363.9, m $100^\circ(\text{dec})$ d $^{25} 2.08\text{g}/\text{cm}^3$. Wash the orange crystals with EtOH and Et_2O , then dry it in air. It sublimes at 35° in a high vacuum. It forms dark yellow plates which are stable for several days when kept in small amounts. Large amounts, especially when placed in a desiccator, spontaneously **ignite** in a period of one day. It decomposes in moist air. It is insoluble in hydrocarbon solvents but forms complexes with several organic compounds. [Sheline & Pitzer *J Am Chem Soc* **72** 1107 1950, DOI: 10.1021/ja01159a012; Speyer & Wolf *Chem Ber* **60** 1424 1927, DOI: 10.1002/cber.19270600626; for applications see Fieser **1** 259, **2** 139, **3** 101, **4** 157, **5** 221, **6** 195, **7** 110, **8** 498, **13** 320, **15** 334, **16** 351.] **TOXIC**.

Iron(III) ethoxide $[\text{Fe}(\text{OEt})_3]$ [5058-42-4] $\text{C}_6\text{H}_{15}\text{O}_3\text{Fe}$, M 191.0, m 120° , volatilities at $155^\circ/0.1\text{mm}$ in a molecular still, b $72.6^\circ/760\text{mm}$, d $^{25} 0.87\text{g}/\text{cm}^3$. The ethoxide is prepared by adding excess of ammonia to a solution of anhydrous FeCl_3 (15g) in $^*\text{C}_6\text{H}_6$ (170ml) and EtOH (76ml), which results in an exothermic reaction at the end of which the mixture is evaporated to dryness *in vacuo*. The residue is extracted with $^*\text{C}_6\text{H}_6$ (150ml), the NH_4Cl is filtered off, the filtrate is evaporated to dryness *in vacuo* to leave a viscous brown residue. This is dissolved in hot EtOH (30ml) and brown crystals of analytically **pure ferric ethoxide** (4.2g) deposit slowly. It is **trimeric** in $^*\text{C}_6\text{H}_6$ solution (molecular weight by ebullioscopy) and is used for preparing other useful ferric trialkoxides by alcohol exchange in benzene solution, e.g. with *n*-PrOH, *n*-BuOH, *iso*-BuOH or *n*-amyl alcohol, *via* the azeotropic removal of EtOH. It is soluble in anhydrous hexane and alcohols, and is commercially available as a 1M solution in EtOH. [Bradley et al. *J Chem Soc* 126 1958, DOI: 10.1039/JR9580000126; see also synthesis revisited Seisenbaeva et al. *Inorg Chim Acta* **358(12)** 3506 2005, DOI: 10.1016/j.ica.2005.03.048; for synthesis, magnetic and electronic spectra see Adams et al. *Aust J Chem* **19** 363 1966, DOI: 10.1071/CH9660363; Beilstein **1** IV 1243.]

Iron pentacarbonyl (pentacarbonyl iron) [13463-40-6] $\text{Fe}(\text{CO})_5$, M 195.9, m -20° , b $102.8^\circ/749\text{mm}$, $103^\circ/760\text{mm}$, n_D²⁰ 1.5196, d $^{25} 1.490\text{g}/\text{cm}^3$. It is a pale yellow viscous liquid that is **PYROPHORIC** and readily absorbed by the skin. **HIGHLY TOXIC (protect from light and air)**. It should be purified in a vacuum line by distilling and collecting in a trap at -96° (toluene-Dry-ice slush). It has been distilled at atmospheric pressure (use a very efficient fume cupboard). At $180^\circ/\text{atmospheric pressure}$ it decomposes to give Fe and CO. In UV light in petroleum ether it forms $\text{Fe}_2(\text{CO})_9$ (see previous entry). [Hagen et al. *Inorg Chem* **17** 1369 1978, DOI: 10.1021/ic50183a055; Ewens et al. *Trans Faraday Soc* **35** 681 1939, DOI: 10.1039/TF9393500681; for applications see Fieser **1** 519, **2** 229, **3** 167, **4** 268, **5** 357, **6** 304, **10** 221, **13** 152, **17** 270.]

Iron(III) meso-5,10,15,20-tetraphenylporphine chloride complex [5,10,15,20-tetraphenyl-21H, 23H-porphine iron(III) chloride] [16456-81-8] $\text{C}_{44}\text{H}_{28}\text{ClN}_4\text{Fe}$, M 704.0, λ_{max} 418nm. Purify the complex by extraction from a thimble (Soxhlet) with CHCl_3 . Concentrate the extract to *ca* 10ml and add *ca* 80ml of hot

MeOH. Dark blue crystals separate on cooling. It can be recrystallised several times from $\text{CHCl}_3/\text{MeOH}$. Avoid prolonged heating. It is quite soluble in organic solvents but insoluble in petroleum ether. [Rothmund & Menotti *J Am Chem Soc* **70** 1808 1948, DOI: 10.1021/ja01185a047; UV: Dorrough et al. *J Am Chem Soc* **73** 4315 1951, DOI: 10.1021/ja01153a085; *Beilstein* **26** III 1960.] [As catalyst for silylation of OH groups see Firouzabadi et al. *Synth Commun* **27** 2709 1997, DOI: 10.1080/00397919708004140.]

Lanthanide shift reagents See in 'Aliphatic Compounds', Chapter 3, europium (III) acetate above and $\text{Eu}(\text{tmc})_3$ and $\text{Eu}(\text{tfc})_3$ below.

Lead II acetate (sugar of lead) [301-04-2 (anhydrous), 6080-56-4 ($3\text{H}_2\text{O}$)] $\text{C}_4\text{H}_6\text{O}_4\text{Pb} \cdot 3\text{H}_2\text{O}$, **M 325.3**(anhydr), **m** 75° ($3\text{H}_2\text{O}$) 280° (anhydr), **d** 20 **3.25g/cm³**(anhydr), **d** 20 **2.55g/cm³**($3\text{H}_2\text{O}$), **d** 20 **1.69g/cm³**($10\text{H}_2\text{O}$), **n** $^{25}_{\text{D}}$ **1.567**($3\text{H}_2\text{O}$), **pK** $^{25}_1$ **7.1** (for Pb^{2+}), **pK** $^{25}_2$ **10.1** (HPbO_2^-), **pK** $^{25}_3$ **10.8** (PbO_2^{2-}). Crystallise $\text{Pb}(\text{OAc})_2$ twice from anhydrous acetic acid and dry it under vacuum for 24 hours at 100° . The solubility of anhydrous salt in H_2O (g/100ml) is 19.8 (0°), 44.31 (20°), 69.5 (30°) and 218.3 (50°); and in MeOH it is 102.75 (66.1°) for **anhydrous salt**, and for the **trihydrate** it is 74.75 (15°) and 214.95 (66.1°); and in glycerol it is 20 (15°) for the anhydrous salt and 143 (20°) for the trihydrate. [*Beilstein* **2** IV 118; Fieser **1** 532, **2** 233, **4** 276.] **POISONOUS**.

Lead (bis-cyclopentadienyl) (plumbocene, Cp_2Pb) [1294-74-2] $\text{C}_{10}\text{H}_{10}\text{Pb}$, **M 337.4**, **sublimes at $150^\circ/10^{-7}\text{mm}$** . Purify it by vacuum sublimation. Handle and store it under N_2 . It was prepared by reaction of sodium cyclopentadienide with a Pb salt, e.g. $\text{Pb}(\text{NO}_3)_2$ (in DMF) or PbI_2 (in THF) [Fischer & Grubert *Z Anorg Allgem Chem* **286**(5-6) 237 1956, DOI: 10.1002/zaac.19562860507; Izod et al. *Organometallics* **27** 4386 2008, DOI: 10.1021/om800598b]. With the PbI_2 (in THF) method the plumbocene was less **pyrophoric** on sublimation. [Dave, et al. *J Chem Soc* 3684 1959, DOI: 10.1039/JR9590003684; *Beilstein* **16** IV 1614.] [For photoelectron spectrum of Cp_2Pb see Cradock & Duncan *JCS Faraday Trans 2* 194 1978, DOI: 10.1039/F29787400194].

Lead tetraacetate [546-67-6] $\text{C}_8\text{H}_{12}\text{O}_8\text{Pb}$, **M 443.2**, **m** 175° , **175-180 $^\circ$** , **d** 20 **2.228g/cm³**. Colourless prisms or needles purified by dissolving in hot glacial acetic acid containing a little acetic anhydride, treated with decolorising charcoal, collected on a hot water funnel or preheated Büchner funnel with minimum contact with moist air, and dried in a vacuum desiccator over KOH pellets. Store it in a well-stoppered vessel as it is decomposed readily by moisture to form brown PbO_2 . It attacks skin, is soluble in hot AcOH , C_6H_6 , CHCl_3 , tetrachloroethane, decomposes by alcohols, and is used as a **powerful oxidising agent**. [Bailar et al. *Inorg Synth* **I** 47 1939, DOI: 10.1002/9780470132326.ch17; Baudler in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 767 1963, *Beilstein* **2** IV 118; Fieser **1** 537, **2** 234, **3** 168, **4** 278, **5** 365, **6** 313, **8** 269, **9** 265, **10** 228, **12** 270, **13** 155, **14** 188, **16** 193.] The tetraacetate induces the addition of CF_2I_2 to alkenes and alkynes at 60° [Li & Chen *Synthesis* 1481 1997, DOI: 10.1055/s-1997-1378]. **TOXIC**.

Lithium benzoate [553-54-8] $\text{C}_7\text{H}_5\text{O}_2\text{Li}$, **M 128.1**, **m** **>300 $^\circ$** , **370 $^\circ$** , **375-441 $^\circ$ (decomposition)**, **d** 25 **1.19g/cm³**. Crystallise the salt from EtOH (13ml/g) by partial evaporation. Its solubility in H_2O (wt %) is 25.68 (-11°), 26.76 (-5°), 27.03 (0°), 27.22 (5.7°), 29.54 (22°), and 29.87 (25°); and the presence of benzoic acid increases its solubility in H_2O . [*Beilstein* **9** I 107, **9** IV 279.]

Lithium diisopropylamide (LDA) [4111-54-0] $\text{C}_6\text{H}_{14}\text{NLi}$, **M 107.1**, **b** **82-84 $^\circ$ /atm**, **84 $^\circ$ /atm**, **d** 22 **0.722 g/cm³**, **flash point -6 $^\circ$** . $\text{LiN}(\text{iso-Pr})_2$ is purified by refluxing over Na wire or NaH for 30 minutes and then distilled into a receiver under N_2 . Because of the low boiling point of the amide, a dispersion of NaH in mineral oil can also be used directly in this purification without prior removal of the oil. It is **HIGHLY FLAMMABLE**, and is decomposed by air and moisture. Solutions of LDA are also commercially available in hexanes (10 wt % suspension), in mineral oil (30 wt% suspension), and in THF/heptane/ethylbenzene (1.8–2M solution, **d** 25 0.836–0.812g/cm³), [Wittig & Hesse *Org Synth* **50** 66 1970, DOI: 10.15227/orgsyn.050.0066; *Beilstein* **4** H 154, **4** I 369, **4** II 630, **4** III 274, **4** IV 510; for applications see Fieser **1** 611, **2** 249, **3** 184, **4** 298, **5** 400, **6** 334, **7** 204, **8** 292, **9** 280, **10** 241, **11** 296, **12** 277, **13** 163, **15** 188, **16** 196, **17** 165.]

Lithium formate monohydrate [6108-23-2 (H_2O), 556-63-8 (anhydrous)] $\text{HCO}_2\text{Li} \cdot \text{H}_2\text{O}$, **M 70.0**, **m** **94 $^\circ$** , **d** 20

1.46. Crystallise HCO_2Li from hot water (0.5ml/g) by chilling. [*Beilstein* **2** III 22, **2** IV 13.]

Lithium methyle (lithium methoxide) [865-34-9] CH_3OLi , **M 38.0**. The most probable impurity is LiOH due to hydrolysis by moisture. It is important to keep the sample dry. It can be dried by keeping in a vacuum at $60\text{--}80^\circ$ under dry N_2 using an oil pump for a few hours. Store it under N_2 in the cold. It should not have bands above 3000cm^{-1} , and its IR (KBr) should have ν_{max} at 1078, 2790, 2840 and 2930cm^{-1} . 1M and 2.2M solutions in MeOH are also commercially available. [*J Org Chem* **21** 156 1956, DOI: 10.1021/jo01108a002; *Beilstein* **1** IV 1220, 1241.]

Lithium pentamethylcyclopentadienide (LiCp', 1,2,3,4,5-pentamethyl-2,4-pentadiene-1-yl lithium) [51905-34-1] $\text{C}_{10}\text{H}_{15}\text{Li}$, **M 142.2**, **m >230°**. In the necessary Schlenk equipment under argon or dry N_2 an equimolar amount of 4M *n*-BuLi in hexane is added dropwise to a stirred solution of equimolar pentamethylcyclopentadiene (Cp') in dry THF (30ml per g of Cp') at 0° which gives a pulpy pale yellow suspension. Magnetic stirring is continued for 2 hours while the temperature is allowed to rise to $\sim 25^\circ$; and this can be used directly for coordinating a Cp' ligand to metals. If LiCp' is to be isolated, then the solvent to be used is petroleum ether ($\sim 15\text{ml}$ per g of Cp'), addition of the *n*-BuLi solution is done within a few minutes without cooling as the reaction is slower than in THF, and stirring is continued for 24 hours. The solvent is removed *in vacuo*, or the colourless to pale grey solid (>90% yield) is filtered off, washed with petroleum ether under N_2 and dried *in vacuo*. [Kohl et al. in *Organometallic Synthesis* (R. Bruce King and J.J. Eisch) **vol 3** 381, Elsevier Amsterdam 1986]. LiCp' is air and moisture sensitive, and should be stored and weighed in an inert atmosphere. [King & Bisnette *J Organomet Chem* **8** 287 1967, DOI: 10.1016/S0022-328X(00)91042-8.] This reagent has been prepared also *in situ* from Cp' with MeLi/THF at 0° [White et al. *Synth Commun* **3** 425 1973, DOI: 10.1080/00397917308065936], with *n*-BuLi in dimethoxyethane [Manriquez & Bercaw *J Am Chem Soc* **96** 6229 1974, DOI: 10.1021/ja00826a071], or as a slurry of LiCp' in Et_2O at $\sim 25^\circ$ [Beachley et al. *Organometallics* **4** 1675 1985, DOI: 10.1021/om00128a033]. The metal-Cp' bond in LiCp' has a more covalent nature than that in KCp' and NaCp' which are more ionic [Jutzi et al. *Chem Ber* **118** 1959 1985, DOI: 10.1002/cber.19851180520].

Lithium picrate (lithium 2,4,6-trinitrophenolate) [18390-55-1] $\text{C}_6\text{H}_2\text{N}_3\text{O}_7\text{Li}$, **M 235.0**. Recrystallise the picrate three times from EtOH and dry it under vacuum at 45° for 48 hours [D'Aprano & Sesta *J Phys Chem* **91** 2415 1987, DOI: 10.1021/j100293a042]. [*Beilstein* **6** H 276, **6** II 263, **6** III 880, **6** IV 1390.] The necessary precautions should be taken in case of **EXPLOSION**.

Lithium salicylate [552-38-5] $\text{C}_7\text{H}_5\text{O}_3\text{Li}$, **M 144.1**, **m >350°**. Recrystallise the salicylate from EtOH (2ml/g) by partial evaporation. [*Beilstein* **10** H 59, **10** II 32, **10** III 93, **10** IV 126.]

Magnesium acetate [142-72-3 (anhydrous), 16674-78-5 ($4\text{H}_2\text{O}$)] $\text{C}_4\text{H}_6\text{O}_4\text{Mg}$, **M 142.4**, **m $80^\circ(4\text{H}_2\text{O})$, d $^{25} 1.45\text{g/cm}^3$** . Crystallise it from anhydrous acetic acid, then dry it under vacuum for 24 hours at 100° . [Nencollas *J Chem Soc* 744 1956, *Beilstein* **2** IV 113.]

Magnesium benzoate trihydrate [553-70-8] $\text{C}_{14}\text{H}_{10}\text{O}_4\text{Mg} \cdot 3\text{H}_2\text{O}$, **M 320.6($3\text{H}_2\text{O}$)**, **m $\sim 200^\circ$** . Crystallise it from water (6ml/g) between 100° and 0° . [*Beilstein* **9** III 376, **9** IV 280.]

Magnesium ethyle (magnesium ethoxide) [2414-98-4] $\text{C}_4\text{H}_{10}\text{O}_2\text{Mg}$, **M 114.4**. Dissolve *ca* 1g of solid in 12.8ml of absolute EtOH and 20ml of dry xylene, and reflux in a dry atmosphere (use CaCl_2 in a drying tube at the top of the condenser). Add 10ml of absolute EtOH and cool. Filter the solid under dry N_2 and dry it in a vacuum. Alternatively, dissolve it in absolute EtOH and pass it through molecular sieves (40 mesh) under N_2 , evaporate under N_2 , and store it in a tightly stoppered container. [Smith & Wiley *J Am Chem Soc* **68** 887 1946, DOI: 10.1021/ja01209a056; *Beilstein* **1** III 1283.]

Magnesium D-gluconate [3632-91-5, 59625-89-7 ($x\text{H}_2\text{O}$)] $\text{C}_{12}\text{H}_{22}\text{O}_{14}\text{Mg}$, **M 414.6(anhydr)**, **$[\alpha]_{\text{D}}^{20} +13.5$, $[\alpha]_{\text{D}}^{20} +11.3$ (c 1, H_2O)**. Crystallise it from dilute EtOH to give *ca trihydrate*, and then dry it at 98° in high vacuum. It is insoluble in EtOH, and the solubility in H_2O is 16% at 25° . [Prescott et al. *Ind Eng Chem* **45** 338

1953, DOI: 10.1021/ie50518a030; *Beilstein* 3 IV 1256.]

Magnesium ionophore I (ETH 1117), (*N,N'*-diheptyl-*N,N'*-dimethyl-1,4-butanediamide) [75513-72-3] $\text{C}_{20}\text{H}_{40}\text{N}_2\text{O}_2$, **M 340.6**. Purify it by flash chromatography (at 40 kPa) on silica and eluting with EtOH/hexane (4:1). Its IR (CHCl_3) has ν_{max} at 1630 cm^{-1} . [Erne et al. *Helv Chim Acta* **63** 2271 1980, DOI: 10.1002/hlca.19800630816.] It is a good Mg^{2+} *selectophore* compared with Na^+ , K^+ and Ca^{2+} [Lanter et al. *Anal Chem* **52** 2400 1980, DOI: 10.1021/ac50064a037].

Magnesium ionophore II (ETH 5214), [*N,N'*-octamethylene-bis(*N'*-heptyl-*N''*-methyl methylmalonamide)] [119110-37-1] $\text{C}_{32}\text{H}_{62}\text{N}_4\text{O}_4$, **M 566.9**. The reagent (*ca* 700mg) can be purified by flash chromatography on Silica Gel 60 (30g) and eluting with $\text{CH}_2\text{Cl}_2/\text{Me}_2\text{CO}$ (4:1). [Hu et al. *Anal Chem* **61** 574 1989, DOI: 10.1021/ac00181a015.]

Magnesium lactate [18917-37-1] $\text{C}_6\text{H}_{10}\text{O}_6\text{Mg}$, **M 202.5**. Recrystallise the salt from water (6ml/g) between 100° to 0° . [*Beilstein* 3 IV 636.] A food and beverages additive, and Mg supplement. **Magnesium succinate** [556-32-1] $\text{C}_4\text{H}_4\text{O}_4\text{Mg}$, **M 140.4** crystallises from water (0.5ml/g) between 100° and 0° . [*Beilstein* 2 IV 1912.] Used as a Mg supplement.

Magon [3-hydroxy-4-(hydroxyphenylazo)-2-naphthoyl-2,4-dimethylanilide, Xylidyl Blue II] [523-67-1] $\text{C}_{25}\text{H}_{21}\text{N}_3\text{O}_4$, **M 411.5, m 246-247°**. Suspend it in H_2O and add aqueous NaOH until it dissolves, filter and acidify with dilute HCl. Collect the red dye, dissolve it in hot EtOH (solubility is 100mg/L at *ca* 25°) concentrate to a small volume and allow to cool. The solubility of the Na salt in H_2O is 0.4mg/ml. The reagent is dried at 100° and is stable at $\sim 25^\circ$ indefinitely; and ethanolic solutions are stable for at least 2 months. [Mann & Yoe *Anal Chim Acta* **16** 155 1957, DOI: 10.1016/S0003-2670(00)89905-5; Mann & Yoe *Anal Chem* **28** 202 1956, DOI: 10.1021/ac60110a016.]

Manganese (II) acetylacetonate [14024-58-9] $\text{C}_{10}\text{H}_{14}\text{O}_4\text{Mn}$, **M 253.2, m 248-250°(dec), $\sim 250^\circ$, 261°(dec)**. Purify it by stirring 16g of reagent for a few minutes with 100ml absolute EtOH and filter by suction as rapidly as possible through coarse filter paper. Sufficient EtOH is added to the filtrate, to make up for the loss of EtOH and to redissolve any solid that separates. Water (15ml) is added to the filtrate, and the solution is evaporated with a stream of N_2 until reduced to half its volume. Cool for a few minutes and filter off the yellow crystals, dry them under a stream of N_2 , then in a vacuum at room temperature for 6-8 hours. These conditions are important for obtaining the *dihydrate*. A vacuum to several mm of Hg or much lower pressure for several days produces the *anhydrous complex*. The degree of hydration can be established by determining the loss in weight of 100g of sample after heating for 4 hours at 100° and $<20\text{mmHg}$. The theoretical loss in weight for $2\text{H}_2\text{O}$ is 12.5%. The material sublimates at $200^\circ/2\text{mm}$. It is soluble in heptane, MeOH, EtOH or $^*\text{C}_6\text{H}_6$ at 30° . [Charles et al. *Inorg Synth* **6** 164 1960, DOI: 10.1002/9780470132371.ch51; Fernelius & Bryant *Inorg Synth* **5** 105 1957, DOI: 10.1002/9780470132364.ch29; *Beilstein* 3 II 3122, 1 IV 3674.]

Manganese (0) decacarbonyl [10170-69-1] $\text{Mn}_2(\text{CO})_{10}$, **M 390.0, m 151-152°, 154-155°(sealed tube), $d^{25} 1.75\text{g/cm}^3$** . Golden yellow crystals which in the absence of CO begin to decompose at 110° , and on further heating yield a metallic mirror. In the presence of 3000psi of CO it does not decompose on heating to 250° . It is soluble in common organic solvents, insoluble in H_2O , not very stable in air, to heat or UV light. It dissolves in a lot of $^*\text{C}_6\text{H}_6$ and can be crystallised from it. It distils with steam at $92-100^\circ$. It can be purified by sublimation under reduced pressure ($<0.5\text{mm}$) at room temperature to give well-formed golden yellow crystals. If the sample is orange coloured, this sublimation leads to a mixture of golden-yellow and dark red crystals of the carbonyl and carbonyl iodide, respectively, which can be separated by hand picking under a microscope. Separation followed by resublimation provides the pure compounds. **POISONOUS**. [Brimm et al. *J Am Chem Soc* **76** 3831 1954, DOI: 10.1021/ja01643a071; Closson et al. *J Am Chem Soc* **80** 6167 1958, DOI: 10.1021/ja01556a005; Podall et al. *J Am Chem Soc* **82** 1325 1960, DOI: 10.1021/ja01491a013.]

Manganous acetate tetrahydrate [6156-78-1 ($4\text{H}_2\text{O}$), 638-38-0 (*anhydrous*)] $\text{C}_4\text{H}_6\text{O}_4\text{Mn} \cdot 4\text{H}_2\text{O}$, **M 245.1($4\text{H}_2\text{O}$), m $80^\circ(4\text{H}_2\text{O})$, $210^\circ(\text{anhydr})$, $d^{25} 1.74\text{g/cm}^3(\text{anhydr})$, $d^{25} 1.59\text{g/cm}^3(4\text{H}_2\text{O})$, $\text{pK}^{25} 10.59$ (for Mn^{2+} hydrolysis). Crystallise it from water acidified with acetic acid. It is also soluble in MeOH EtOH and AcOH. [*Beilstein* 2 IV 120.]**

Manganous lactate trihydrate [6505-50-6 ($3\text{H}_2\text{O}$), 51877-53-3 ($x\text{H}_2\text{O}$)] $\text{C}_6\text{H}_{10}\text{O}_6\text{Mn} \cdot 3\text{H}_2\text{O}$, **M 287.1**($3\text{H}_2\text{O}$). Recrystallise the lactate from water. [IR: Ranade & Biswas *J Indian Chem Soc* **44** 314 1967; for X-ray structure of the *trihydrate* see Kemmitt et al. *Aust J Chem* **54**(1) 37 2001, DOI: 10.1071/CH01012; *Beilstein* **3** IV 637.]

Mercuric(II) acetate $[\text{Hg}(\text{OAc})_2]$ [1600-27-7] **M 318.7, m 178-180°, 179-182°, pK_1^{25} 2.47, pK_2^{25} 3.49 (for Hg^{2+} hydrolysis).** It is prepared from HgO and acetic acid. Recrystallise it from glacial acetic acid. Its solubility in H_2O is 1g/2.5ml at $\sim 20^\circ$ and 1g/ml at 100° , and it is soluble in EtOH. The solid is light sensitive and is hydrolysed slowly in H_2O to yield a yellow oxide. Store it in a tightly stoppered bottle away from light. It is a reagent for *mercuration of alkenes* (and aromatic compounds, cf. mercury trifluoroacetate below) [Larock *Tetrahedron* **38** 1713 1982, DOI: 10.1016/0040-4020(82)80245-7]; *de-mercuration* being achieved with NaBH_4 . Mercuric acetate in 10% aqueous AcOH (reflux for ~ 20 hours) has been used to dehydrogenate cyclic secondary amines, e.g. 2-methyl, 2,5-dimethyl and 2,4,4-trimethyl (but not 2,2-dimethyl or 2,2,3-trimethyl) pyrrolidines to the respective Δ^1 pyrrolines; the reaction being followed by the precipitation of mercurous acetate [Bonnett et al. *J Chem Soc* 2087 1959, DOI: 10.1039/JR9590002087]. [Wagenknecht & Juza in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **II** p 1120-1121 1965, *Beilstein* **2** IV 114; for applications see Fieser **1** 644, **2** 264, **3** 194, **4** 319, **5** 424, **6** 358, **10** 252, **15** 198, **17** 176.] **POISONOUS.**

Mercury(II) bis(cyclopentadienyl) (HgCp_2 , mercurocene) [18263-08-6, 12083-67-9] $\text{C}_{10}\text{H}_{10}\text{Hg}$, **M 330.8 starts to dec at $\sim 60^\circ$ but melts at $83-85^\circ$.** HgCp_2 has been prepared by metathesis between TiCp [34822-90-7, see above] and HgCl_2 at -40° in THF [see Cradock & Duncan, below]. The pale yellow crystals are purified by low-temperature (-78°) recrystallisation from Et_2O . Store it at low temperature in the dark as it decomposes slowly at room temperature even in the dark. It reacts with FeCl_2 in tetrahydrofuran to give ferrocene quantitatively. It is soluble in Et_2O , $^*\text{C}_6\text{H}_6$, CHCl_3 , CCl_4 , and most organic solvents. [Campbell & Fyfe *J Am Chem Soc* **94** 8387 1972, DOI: 10.1021/ja00779a017; Wilkinson & Piper *J Inorg Nucl Chem* **2** 32 1956, DOI: 10.1016/0022-1902(56)80101-2; Piper & Wilkinson *J Inorg Nucl Chem* **3** 104 1956, DOI: 10.1016/0022-1902(56)80073-0; *Beilstein* **16** IV 1702.] [For photoelectron spectrum of Cp_2Hg see Cradock & Duncan *JCS Faraday Trans 2* 194 1978, DOI: 10.1039/F29787400194]. **POISONOUS.**

Mercury dibromofluorescein disodium salt {mercurochrome, merobromin, [2',7'-dibromo-4'-(hydroxy-mercurio)-fluorescein di-Na salt]} [129-16-8] $\text{C}_{20}\text{H}_8\text{Br}_2\text{Na}_2\text{O}_6\text{Hg}$, **M 750.6, 804.8**($3\text{H}_2\text{O}$), **m > 300°**. The green Na salt is dissolved in the minimum volume of H_2O , or the free acid suspended in H_2O and dilute NaOH is added to cause it to dissolve, filter and acidify it with dilute HCl. Collect the precipitate, wash it with H_2O by centrifugation and dry it in a vacuum. The *di Na salt* can be purified by dissolving it in the minimum volume of H_2O and is precipitated by adding EtOH, filter, wash it with EtOH or Me_2CO and dry it in a vacuum. Its solubility in 95% EtOH is 2% and in MeOH it is 16%. [White *J Am Chem Soc* **42** 2355 1920, DOI: 10.1021/ja01456a031.] Useful disinfectant.

Mercury Orange [1-(4-chloromercuriophenylazo)-2-naphthol] [3076-91-3] $\text{C}_{16}\text{H}_{11}\text{ClN}_2\text{OHg}$, **M 483.3, m 291.5-293°(corr) with bleaching.** Wash the dark red powder several times with boiling 50% EtOH and recrystallise it from 1-butanol (0.9g/l of boiling alcohol). The fine needles are insoluble in H_2O but slightly soluble in cold alcohols, CHCl_3 and soluble in aqueous alkalis. [Chaikin *J Am Chem Soc* **70** 3522 1948, DOI: 10.1021/ja01190a600.]

Mercury(II) trifluoroacetate [13257-51-7] $\text{C}_2\text{F}_3\text{O}_2\text{Ag}$, **M 426.6, m 171-173°**. It is prepared from HgO and trifluoroacetic acid and is recrystallised from trifluoroacetic anhydride/trifluoroacetic acid. It is a useful reagent for mercuration of aromatic compounds [Lau & Kochi *J Am Chem Soc* **108** 6720 1986, DOI: 10.1021/ja00281a043]. It is soluble in THF, DME, and dioxane but insoluble in hexane. Store it in a tightly stoppered bottle away from light. [cf. Aylett in *Comprehensive Inorganic Chemistry* (Bailar et al. eds) Vol **3** p187 1973, Kočovský & Luxenburger *e-EROS Encyclopedia of Reagents for Organic Synthesis* 14 March 2008, DOI: 10.1002/047084289X.rm046.pub2; *Beilstein* **2** IV 458; for applications see Fieser **1** 659, **3** 195, **4** 325, **5** 429, **6** 360, **9** 294, **13** 175, **16** 377.] It is very **TOXIC**, can be absorbed through the skin and is *hygroscopic*.

Methylmercuric chloride [115-09-3] CH_3ClHg , **M 251.1, m 167°, 170°**. Recrystallise it from absolute EtOH

(20ml/g). Its UV has λ_{\max} at 206nm (ϵ 1.37). [See EtHgCl above; Breiting & Kress. *J Organomet Chem* **256** 217 1983, DOI: 10.1016/S0022-328X(00)99198-8; Slotta et al. *J Prakt Chem* **120** 249 1929, Waugh et al. *J Phys Chem* **59** 395 1955, DOI: 10.1021/j150527a004; *Beilstein* **16** IV 1729.]

Molybdenum hexacarbonyl [13939-06-5] $\text{Mo}(\text{CO})_6$, **M 264.0**, **$\sim 150^\circ$ dec without melting**, **b $156^\circ/\text{atm}$** , **d²⁵ $1.96\text{g}/\text{cm}^3$** . Sublime it in a vacuum before use [Connor et al. *JCS Dalton Trans* 511 1986, DOI: 10.1039/DT9860000511]. Useful catalyst, alone or with other ligands, for alkyne metathesis [Marradi *Synlett* (spotlight 119) 1195 2005, DOI: 10.1055/s-2005-865206.] **TOXIC**.

Monoperoxyphthalic acid magnesium hexahydrate salt (MMPP) [84665-66-7, 78948-87-5] $\text{C}_{16}\text{H}_{10}\text{O}_{10}\text{Mg} \cdot 6\text{H}_2\text{O}$, **M 494.7**, **m $\sim 93^\circ(\text{dec})$** . MMPP is a *safer reagent* than *m*-chloroperbenzoic acid for oxidation reactions because it is not as explosive and has advantages of solubility. It is soluble in H_2O , low-molecular-weight alcohols, *i*-PrOH and DMF. The product of reaction, Mg phthalate, is soluble in H_2O . It has been used in aqueous phase to oxidise compounds in e.g. CHCl_3 and using a phase transfer catalyst, e.g. methyltriocetylammmonium chloride [Brougham et al. *Synthesis* 1015 1987, DOI: 10.1055/s-1987-28153]. The 749xidizing activity can be checked (as for perbenzoic acid in Silbert et al. *Org Synth Coll Vol* **5** 904 1973, DOI: 10.15227/orgsyn.044.0081], and if found to be low it would be best to prepare afresh from phthalic anhydride (1mol), H_2O_2 (1mol) and MgO at $20\text{--}25^\circ$ to give MMPP. [Hignett, European Pat Appl 27 693 1981, *Chem Abstr* **95** 168810 1981, *Beilstein* **9** IV 3260.]

Neopentoxo lithium [3710-27-8] $\text{C}_5\text{H}_{11}\text{Oli}$, **M 94.1**. Recrystallise the alkoxide from hexane [Kress & Osborn *J Am Chem Soc* **109** 3953 1987, DOI: 10.1021/ja00247a020]. [*Beilstein* **1** H 406, **1** I 201, **1** II 435. **1** III 1448, **1** IV 1690, for neopentyl alcohol and Al and Fe neopentoxides.]

New Methylene Blue N (2,8-dimethyl-3,7-bis(ethylamino)phenothazinium chloride 0.5ZnCl_2) [6586-05-6] $\text{C}_{18}\text{H}_{22}\text{ClN}_3\text{S} \cdot 0.5\text{ZnCl}_2$, **M 416.1**, **m $>200^\circ(\text{dec})$** , **λ_{\max} 591nm and 630nm**, **pK₁ 3.54, pK₂ 4.82**. Crystallise the dye from *benzene/MeOH (3:1). [*Beilstein* **27** IV 5163.]

Nickel (II) acetate ($4\text{H}_2\text{O}$) [6018-89-9] $\text{C}_4\text{H}_6\text{O}_4\text{Ni} \cdot 4\text{H}_2\text{O}$, **M 248.9**, **d²⁵ $1.798\text{g}/\text{ml}$** , **pK₁²⁵ 8.94 (from Ni^{2+} hydrolysis)**. The acetate crystallises from aqueous AcOH as the green *tetrahydrate salt*. It is soluble in 6 parts of H_2O . It forms *lower hydrates* and should be kept in a well-sealed container. [Hardt & Pohlmann *Z Anorg Allgem Chem* **343** 92 1966, DOI: 10.1002/zaac.19663430112; *Beilstein* **2** IV 120.] Polynuclear-nickel polyoxotungstate duster compounds prepared from nickel acetate are ideal for the design of nuclear magnets [Clemente-Juan et al. *Inorg Chem* **38** 55 1999, DOI: 10.1021/ic9807902].

Nickelocene [bis-(cyclopentadienyl)nickel II] [1271-28-9] $\text{C}_{10}\text{H}_{10}\text{Ni}$, **M 188.9**, **m $173\text{--}174^\circ(\text{under } \text{N}_2)$** . Dissolve it in Et_2O , filter and evaporate in a vacuum. Purify it rapidly by recrystallisation from petroleum ether using a solid $\text{CO}_2/\text{Me}_2\text{CO}$ bath. It has **m $171\text{--}173^\circ$** (in an evacuated tube). Also purify by vacuum sublimation. [Wilkinson et al. *J Am Chem Soc* **76** 1970 1954, DOI: 10.1021/ja01636a080; *Beilstein* **16** IV 1831.]

Nickel (II) phthalocyanine [14055-02-8] $\text{C}_{32}\text{H}_{16}\text{N}_8\text{Ni}$, **M 571.3**, **m $>300^\circ$** , **λ_{\max} 670nm**. Wash it well with H_2O and boiling EtOH and sublime it at high vacuum in a slow stream of CO_2 . A special apparatus is used (see reference), with the phthalocyanine being heated to red heat. The *sublimate* is in the form of needles with an extremely bright *red lustre*. The powder is *dull greenish blue* in colour. [Barrett et al. *J Chem Soc* 1719 1936, DOI: 10.1039/JR9360001719; *Beilstein* **26** III/IV 4255.]

Nickel 5,10,15,20-tetraphenylporphyrin (NiTPP) [14172-92-0] $\text{C}_{44}\text{H}_{28}\text{N}_8\text{Ni}$, **M 671.4**, **λ_{\max} 414(525)nm**. Purify it by chromatography on neutral (Grade I) alumina, followed by recrystallisation from $\text{CH}_2\text{Cl}_2/\text{MeOH}$ [Yamashita et al. *J Phys Chem* **91** 3055 1987, DOI: 10.1021/j100295a082]. [*Beilstein* **26** III/IV 1960.]

Phenylmercuric acetate (PMA, PhHgOAc) [62-38-4] $\text{C}_8\text{H}_8\text{O}_2\text{Hg}$, **M 336.7**, **m $148\text{--}151^\circ$, 149° , $151.8\text{--}152.8^\circ$** . This acetate forms small colourless lustrous prisms from EtOH. Its solubility in H_2O is 0.17%, but it is

more soluble in EtOH, Me₂CO and *C₆H₆. [Maynard *J Am Chem Soc* **46** 1510 1925, DOI: 10.1021/ja01671a024; Coleman et al. *J Am Chem Soc* **59** 2703 1937, DOI: 10.1021/ja01291a067; Grave et al. *J Am Pharm Assoc* **25** 752 1936, DOI: 10.1002/jps.3080250906; *Beilstein* **16** IV 1720; Fieser **6** 458.] Used as a preservative in paints, and as a disinfectant. See PhHgOH below.

Phenylmercuric hydroxide (PhHgOH) [100-57-2] C₆H₆O₂Hg, M **294.7**, m **195-203°**, pK²⁵ **10 (~4.5)**. Crystallise it from dilute aqueous NaOH. [Waugh et al. *J Phys Chem* **59** 395 1955 DOI: 10.1021/j150527a004; *Beilstein* **16** IV 1721.] **Phenylmercuric nitrate (PhHgNO₃)** [8003-05-2] C₆H₅NO₃Hg.C₆H₆O₂Hg, M **634.4**, m **175-185°**, **178-188°**. When recrystallised from water or aqueous EtOH, it has m **191-191.5°**, and from 95% EtOH it has m **186.5°** [Challenger & Rothstein *J Chem Soc* 1258 1934, DOI: 10.1039/JR9340001258]. When recrystallised from *C₆H₆ or CHCl₃, it has m **131-131.5°** [Morton et al. *J Am Chem Soc* **69** 908 1947, DOI: 10.1021/ja01196a051.] [*Beilstein* **16** III 1348, **16** IV 1721.]

Phthalocyanine [574-93-6] C₃₂H₁₈N₈, M **514.6**, m **>300°(dec)**. Purify phthalocyanine by sublimation (two to three times) in an argon flow at 300-400Pa; and similarly for the Cu(II), Ni(II), Pb(II), VO(II) and Zn(II) phthalocyanine complexes. It is suitable as a laser dye with λ_{max} 670nm. [*Beilstein* **26** III/IV 4255.]

Platinum (II) acetate (diacetatoplatinum) [3375-31-3, 38928-89-1] C₄H₆O₄Pt, M **313.2 (monomer)**, m **245°(dec)**. Purify the purple crystals by dissolving them in CHCl₃, filtering, mixing with half its volume of glacial acetic acid and set aside to evaporate at room temperature. The resulting large, **almost black**, crystals are collected, washed with H₂O and dried in air. It dissolves in CHCl₃, *C₆H₆ and toluene to give **purple solutions**. Its molecular weights (ebullioscopically) in CHCl₃, *C₆H₆ and chlorobenzene are 950 (±2), 937 and 888, respectively, indicate that it is **trimeric in solution**. [Stephenson et al. *J Chem Soc* 3632 1965, DOI: 10.1039/JR9650003632.]

Platinum (II) acetylacetonate [15170-57-7] C₁₀H₁₄O₄Pt, M **393.3**, m **249-252°**. It crystallises from *C₆H₆ as yellow crystals and is dried in air or in a vacuum desiccator. It is soluble in CHCl₃, and Me₂CO, but insoluble in H₂O. [Werner *Chem Ber* **34** 2584 1901, DOI: 10.1002/cber.190103402206; Fernelius & Bryant *Inorg Synth* **5** 105 1957, DOI: 10.1002/9780470132364.ch29; *Beilstein* **1** H 783, **1** IV 3678.]

Potassium antimonyltartrate trihydrate [28300-74-5 (3H₂O), 331753-56-1 (xH₂O)] C₈H₄O₁₂K₂Sb₂, M **613.8(anhydr)**, m **>300°(dec)**, d²⁵ **2.6g/ml**, [α]_D²⁰ **+141 (c 2, H₂O)**. Crystallise '**tartar emetic**' from water between 100°(3ml/g) and 0°(12ml/g). Insol in EtOH. Dry it at 100°. [Chinoporos & Papathanasopulos *J Phys Chem* **65** 1643 1961, DOI: 10.1021/j100905a512; *Beilstein* **3** IV 1227.] It is an expectorant and a parasitide.

Potassium benzoate [582-25-2] C₇H₅O₂K, M **160.2**, m **>300°(dec)**, d²⁵ **1.5g/cm³**. Crystallise it from water (1ml/g) between 100° and 0°. Its solubility in H₂O (g/100ml) is 69.87 (17.5°), 73.83 (25°), 79 (33.3°), 88.33 (20°), and it is soluble in EtOH, slightly soluble in MeOH and insoluble in Et₂O. [*Beilstein* **9** III 375, **9** IV 279; Fieser **1** 49.] **Potassium 4-aminobenzoate** [138-84-1] C₁₇H₆NO₂K, M **175.2**. Crystallise the aminobenzoate from EtOH. [*Beilstein* **14** II 246, **14** IV 1128.]

Potassium tert-butoxide [865-47-4] C₄H₉OK, M **112.2**. It sublimes at 220°/1mm. The last traces of *tert*-BuOH are removed by heating at 150-160°/2mm for 1 hour. It is best prepared afresh as likely impurities are *tert*-BuOH, KOH and K₂CO₃ depending on its exposure to air. Its solubility at 25-26° in hexane, toluene, Et₂O, and THF is 0.27%, 2.27%, 4.34% and 25.0%, respectively. Solutions in *tert*-BuOH (1M, d²⁵ 0.82g/ml) and in THF (1M, d²⁵ 0.902g/ml) are commercially available. [Feuer et al. *J Am Chem Soc* **78** 4364, DOI: 10.1021/ja01598a045; Doering & Urban *J Am Chem Soc* **78** 5938 1956, DOI: 10.1021/ja01603a060; *Beilstein* **1** IV 1612.]

Potassium citrate tribasic monohydrate (tripotassium citrate monohydrate) [6100-05-6] C₆H₅O₇K₃.H₂O, M **324.4**, m **275°(dec)**. Its solubility in H₂O is 154g/100ml and it loses H₂O at 180°. [*Beilstein* **3** III 1091.]

Potassium dihydrogen citrate [866-83-1] C₆H₇O₇K, M **230.2**. It crystallises from H₂O with a solubility of 11.5% at 100°. Dry it at 80°, or in a vacuum desiccator over Sicapent. [*Beilstein* **3** I 209, **3** III 336, **3** IV 402.]

Potassium hydrogen D-glucarate (D-tetrahydroxyadipic acid mono K salt, D-glucosaccharic acid mono K salt) [18404-47-2] $\text{C}_6\text{H}_9\text{O}_8\text{K}$, **M 286.3, 248.2, m 188°(dec)**, $[\alpha]_{\text{D}}^{20} +4.95$ (c 1.2, H_2O), $\text{pK}_{\text{a}}^{25} 5.0$ (free acid). Crystallise it from water. [Wolfrom & Neely *J Am Chem Soc* **75** 2778 1953, DOI: 10.1021/ja01107a519; *Beilstein* **3** H 579, **3** II 378, **3** III 1119, **3** IV 1291.]

Potassium hydrogen malate [4675-64-3] $\text{C}_4\text{H}_5\text{O}_5\text{K}$, **M 172.2**. A saturated aqueous solution at 60° is decolorised with activated charcoal, and filtered. The filtrate is cooled in a water-ice bath, and the salt is precipitated by adding EtOH. After five recrystallisations from ethanol-water mixtures, it is dried overnight at 130° in air. Assaying by titration found that after the first crystallisation it was 99.69% pure, and after the third recrystallisation it was **99.99% pure**; and was used for dissociation constants measurements after the fifth recrystallisation. [For the resolution of the dissociation constants of *dl*-malic acid from 0° to 50° see Eden & Bates *J Res Nat Bur Stand* **62** 161 1959, <http://dx.doi.org/10.6028/jres.062.028>]. [*Beilstein* **3** III 915, **3** IV 1123.]

Potassium hydrogen oxalate (H_2O) [127-95-7] $\text{C}_2\text{HO}_4\text{K}$, **M 137.1, m ~300°(dec)**, $\text{d}^{25} 2.0\text{g/cm}^3$. Crystallise the white acid oxalate from boiling H_2O [solubility in w/v% is 2.5 (~25°) and 16.7 (boiling)], filtering and cooling to 25°. The crystals, after washing three or four times with water, are allowed to dry in air. [*Beilstein* **2** III 1552.] It is an ink stain remover, a mordant in dyeing and used for scouring metal. [see K tetaoxalate below.]

Potassium hydrogen phthalate (KHP) [877-24-7] $\text{C}_8\text{H}_5\text{O}_4\text{K}$, **M 204.2, m 295°, d²⁵ 1.636g/cm³, pK²⁵ 5.4**. Crystallise it first from a dilute aqueous solution of K_2CO_3 , then H_2O (3ml/g) between 100° and 0°. Before being used as a standard in volumetric analysis, analytical grade potassium hydrogen phthalate should be dried at 120° for 2 hours, then allowed to cool in a desiccator. [*Beilstein* **9** IV 3169.] Buffering agent for pH electrode calibration at *ca* pH 4.

Potassium hydrogen *d*-tartrate [868-14-4] $\text{C}_4\text{H}_5\text{O}_6\text{K}$, **M 188.2, d²⁵ 1.954g/ml, n_D²⁵ 1.511, $[\alpha]_{\text{D}}^{20} +10$ (c 2, 10% HCl), $[\alpha]_{546}^{20} +37.5$ (c 10, M NaOH)**. It crystallises from water (17ml/g) between 100° and 0°. Dry it at 110°. [*Beilstein* **3** IV 1222.] It is a product of fermentation of grape juice, and can precipitate in bottles of wine. In cooking it is known as '*cream of tartar*'.

Potassium laurate (potassium dodecanoate) [10124-65-9] $\text{C}_{12}\text{H}_{23}\text{O}_2\text{K}$, **M 338.4**. Recrystallise potassium laurate three times from EtOH [Neto & Helene *J Phys Chem* **91** 1466 1987, DOI: 10.1021/j100290a037]. [*Beilstein* **2** H 360, **2** I 156, **2** II 318, **2** III 880, **2** IV 186.] It is a detergent. See purification of sodium dodecanoate below.

Potassium nonafluorobutane sulfonate [29420-49-3] $\text{C}_4\text{F}_9\text{O}_3\text{SK}$, **M 338.2, m >300°**. Wash it with H_2O and dry it *in vacuo*. When the K salt is distilled with 100% H_2SO_4 , it gives the *free acid* which can be distilled (**b** 105°/22mm, 210-212°/760mm) and then converted back into the pure K salt by careful neutralisation with aqueous KOH. [Gramstad & Haszeldine *J Chem Soc* 2640 1957, DOI: 10.1039/JR9570002640; *Beilstein* **2** IV 818.]

Potassium oleate [143-18-0] $\text{C}_{18}\text{H}_{33}\text{O}_2\text{K}$, **M 320.6**. Recrystallise potassium oleate from EtOH (1g/ml). It can appear as a white-yellow waxy solid or a clear amber liquid with a soapy odour, and dissolves in H_2O slowly. [*Beilstein* **2** H 465, **2** I 196, **2** I 202, **2** II 436, **2** III 1404, **2** IV 1646.]

Potassium oxalate monohydrate [6487-48-5] $\text{C}_2\text{O}_4\text{K}_2 \cdot \text{H}_2\text{O}$, **M 184.2, m 160°(hydrate, dec), 365°(anhydr)**, $\text{d}_4^{20} 2.13$. Recrystallise the oxalate from hot water. Its solubility in H_2O is 36.4g/100ml at 20°. It forms various K + oxalic acid salts [e.g. potassium hydrogen oxalate $\text{KH}(\text{C}_2\text{O}_4)$ —see above, and $\text{KH}_3(\text{C}_2\text{O}_4)_2$ which is, surprisingly, called potassium tetraoxalate—see below]. [*Beilstein* **2** I 224, **2** II, 485, **2** III 1552, **2** IV 1823.]

Potassium pentamethylcyclopentadienide (KCp', 1,2,3,4,5-pentamethyl-2,4-pentadiene-1-ylpotassium) [94348-92-2] $\text{C}_{10}\text{O}_{15}\text{K}$, **M 174.3, m >230°**. In an efficient fume cupboard, liquid NH_3 (1.3L) is condensed into a flask containing freshly cut K metal (11.7g, 300mg-atoms) under an argon or N_2 atmosphere, stirred at -78°, and a small crystal of $\text{Fe}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$ is added to the blue coloured solution. The mixture is made to warm to room temperature allowing refluxing until the blue solution turns into a light grey suspension. Cp' (40.9g,

300mmol) is added *via* a syringe, the mixture is stirred at reflux (MeOH condenser cooled at -78°) for 4 hours, and the NH_3 is allowed to evaporate using a stream of N_2 . The residue is dried *in vacuo* and stored in a N_2 atmosphere as it is air and moisture sensitive, and unlike other Cp^*M s it can be spontaneously flammable in air. It is more soluble in THF than Cp^*Li but less soluble than Cp^*Na . It is a Cp^* ligand source for complexing with a large number of metals and non-metal halides. [Kohl et al. in *Organometallic Synthesis* (R. Bruce King and J.J. Eisch eds) **vol 3** 381, Elsevier Amsterdam 1986, DOI: 10.1080/00945718708059453].

Potassium phthalimide (phthalimide K salt) [1074-82-4] $\text{C}_8\text{H}_4\text{NO}_2\text{K}$, **M 185.2, m $>300^{\circ}$** . The solid may contain phthalimide and K_2CO_3 from hydrolysis. If too much hydrolysis has occurred (this can be checked by extraction with cold Me_2CO in which the salt is insoluble, evaporate the Me_2CO and weigh the residue), it would be better to prepare it afresh. If little hydrolysis has occurred, then recrystallise it from a large volume of EtOH, and wash the solid with a little Me_2CO and dry it in a continuous vacuum to constant weight. [Salzerg & Supniewski *Org Synth Coll Vol 1* 119 1941, DOI: 10.15227/orgsyn.007.0008; Raman & IR: Hase *J Mol Struct* **48** 33 1978, DOI: 10.1016/0022-2860(78)87220-2; Dykman *Chem Ind (London)* 40 1972, IR, NMR: Assef et al. *Bull Soc Chim Fr II* 167 1979, *Beilstein* **21/10** V 270.] This salt is useful for transforming allyl- and alkyl halides into phthalimido protected primary amines which react with hydrazine to provide the respective primary amine and insoluble phthalazin-2,4-dione [Lu et al. *J Am Chem Soc* **129** 7768 2007, DOI: 10.1021/ja072844p].

Potassium picrate [573-83-1] $\text{C}_6\text{H}_2\text{N}_3\text{O}_7\text{K}$, **M 267.2, m 250° , (detonates at 331° , before boiling), $d^{25} 1.852\text{g/cm}^3$** . Recrystallise it from water or 95% EtOH, and dry it at room temperature in vacuum. It is soluble in 200 parts of cold water and 4 parts of boiling water. Store wet. [*Beilstein* **3** III 880, **3** IV 1390.] **THE DRY SOLID EXPLODES WHEN STRUCK OR HEATED.**

Potassium propionate [327-62-8] $\text{C}_3\text{H}_5\text{O}_2\text{K}$, **M 112.2, m $>300^{\circ}$, 410°** . Recrystallise $\text{C}_2\text{H}_5\text{CO}_2\text{K}$ from water (30ml/g) or 95% EtOH. [*Beilstein* **2** II 217, **2** III 516, **2** IV 701.]

Potassium sodium tartrate tetrahydrate [6381-59-5 ($4\text{H}_2\text{O}$), 304-59-6 (*R,R*)] $\text{C}_4\text{H}_4\text{O}_6\text{KNa} \cdot 4\text{H}_2\text{O}$, **M 282.3, m 75° (hydrate), b $220^{\circ}/\text{atm}$ (becoming anhydrous at $\sim 130^{\circ}$), $d^{25} 1.79\text{g/cm}^3$** . Recrystallise the tartrate from distilled water (1.5ml/g) by cooling to 0° . Solubility in H_2O is 26g/100ml (0°) and 66g/100ml (26°). [*Beilstein* **3** IV 1223.] **Potassium d-tartrate monohydrate** [921-53-9] $\text{C}_4\text{H}_4\text{O}_6\text{KNa} \cdot \text{H}_2\text{O}$, **M 235.3, m loses H_2O at 150° , $d_4^{20} 1.98$** . Recrystallise it from pure water (solubility: 0.4ml/g at 100° , 0.7ml/g at 14°). [*Beilstein* **3** IV 1223.]

Potassium tetraoxalate dihydrate [oxalic acid hemipotassium salt] [6100-20-5 ($2\text{H}_2\text{O}$), 127-96-8] $\text{C}_4\text{H}_{10}\text{O}_{10}\text{K}$, **M 254.2, b $365.1^{\circ}/760\text{mm}$, $d^{25} 1.84\text{g/cm}^3$** . Crystallise it from water below 50° (solubility is $\sim 0.1\text{M}$ at 20° , and 8.3g at 100°). Dry it $<60^{\circ}/760\text{mm}$. [See potassium acid oxalate above, *Beilstein* **2** IV 1823.]

Potassium trifluoroacetate [2923-16-2] $\text{C}_2\text{F}_3\text{O}_2\text{K}$, **M 152.1, m $140-142^{\circ}$, $d^{25} 1.49\text{g/ml}$, $\text{pK}^{25} 0.52$ (for $\text{CF}_3\text{CO}_2\text{H}$)**. To purify it, dissolve the salt in trifluoroacetic acid with *ca* 2% of trifluoroacetic anhydride (CARE), filter it and evaporate it carefully to dryness (avoid over heating), and finally dry it in a vacuum at 100° . It can be recrystallised from trifluoroacetic acid (solubility in the acid is *ca* 50.1%). [Simons & Lorentzen *J Am Chem Soc* **74** 4746 1952, DOI: 10.1021/ja01139a006; Hara & Cady *J Am Chem Soc* **76** 4285 1954, DOI: 10.1021/ja01646a009; Watt & Muga *J Inorg Nucl Chem* **9** 166 1959, DOI: 10.1016/0022-1902(59)80077-4; *Beilstein* **2** IV 458, *Fieser* **11** 557.]

Pyridinium chlorochromate (PCC) [26299-14-9] $\text{C}_5\text{H}_6\text{ClNO}_3\text{Cr}$, **M 215.6, m $205^{\circ}(\text{dec})$, $205-208^{\circ}(\text{dec})$** . Dry it in a vacuum for 1 hour. It is not hygroscopic and can be stored for extended periods at room temperature without change. If the purity is very suspect, it can be readily prepared. It is soluble in CH_2Cl_2 , Et_2O , Me_2CO , MeCN, THF and $^*\text{C}_6\text{H}_6$. [Corey & Scuggs *Tetrahedron Lett* 2647 1975, DOI: 10.1016/S0040-4039(00)75204-X; Piancatelli et al. *Synthesis* 245 1982, DOI: 10.1055/s-1982-29766.] [Applications see *Fieser* **6** 498, **7** 308, **8** 425, **9** 397, **10** 334, **11** 450, **12** 417, **13** 262, **14** 269, **15** 267.] It is a stable versatile oxidising agent [Piancatelli et al. *Synthesis* 245 1982, DOI: 10.1055/s-1982-29766], used for mild and selective oxidation of alcohols [Kasmai et al. *J Org Chem* **60** 2267 1995, DOI: 10.1021/jo00112a059], and the oxidation of an allylic methylene group in a synthesis of α,β -unsaturated lactones from D-glucose [Srikanth et al. *Tetrahedron* **62** 11165 2006, DOI: 10.1016/j.tet.2006.09.016]. (Possible **CARCINOGEN**.) § Available commercially on a polymer support.

Pyridinium dichromate [20039-37-6] $\text{C}_{10}\text{H}_{10}\text{N}_2^{2+} \cdot \text{Cr}_2\text{O}_7^{2-}$, **M 376.2, m 145-148°, 152-153°**. Dissolve it in the minimum volume of H_2O and add 5 volumes of cold Me_2CO and cool to -20° . After 3 hours the orange crystals are collected, washed with a little cold Me_2CO and dried in a vacuum. It is soluble in dimethylformamide (0.9g/ml at 25°), and in H_2O , and has a characteristic IR with ν_{max} at 930, 875, 765, 730 and 730 cm^{-1} . [Corey & Schmidt *Tetrahedron Lett* 399 1979, DOI: 10.1016/S0040-4039(01)93515-4; Coats & Corrigan *Chem Ind (London)* 1594 1969.] (Possible **CARCINOGEN**.)

§ Available commercially on silica gel and on a polymer support.

Pyronin B [di-(3,6-bis(diethylamino)xanthylium chloride) di FeCl_5 complex] [2150-48-3] $\text{C}_{42}\text{H}_{54}\text{Cl}_8\text{N}_4\text{O}_2\text{Fe}_2$, **M 1042.2, m 176-178° (diFe complex), CI 45010, λ_{max} 553nm, $\text{pK}^{25}_{\text{H}} 7.7$** . Recrystallise it from EtOH. It forms an Fe stain. It is a biological stain used in the *methyl green-pyronin method* for differential coloration of nucleic acids. [For 'Analysis and testing of biological stains' see Penney et al. *Biotechnic and Histochemistry* 1 1 2002, PMID 12564600; Beilstein 18/10 V 182.]

Quinolinium chlorochromate [108703-35-1] $\text{C}_9\text{H}_7\text{N} \cdot \text{HCrO}_3\text{Cl}$, **M 265.6, m 127-130°**. It is a yellow-brown solid which is stable in air for long periods. If it has deteriorated or been kept for too long, it is best to prepare it freshly. Add freshly distilled quinoline (13ml) to a mixture of chromic acid (CrO_3) (10g) and $\sim 5\text{M}$ HCl (11ml of conc HCl and 10ml of H_2O) at 0° . A yellow-brown solid separates, it is filtered off on a sintered glass funnel, dried for 1 hour in a vacuum, and can be stored for extended periods without serious loss in activity. It is a good oxidant for primary alcohols in CH_2Cl_2 . [Singh et al. *Chem Ind (London)* 751 1986, method of Corey & Suggs *Tetrahedron Lett* 2647 1975, DOI: 10.1016/S0040-4039(00)75204-X; for structure and solvent dependent kinetic study of the oxidation of substituted benzaldehydes see Jeyanthi & Elango *Int J Chem Kinet* 35 154 2003, DOI: 10.1002/kin.10112; Beilstein 20/7 V 276.]

Resorufin (7-hydroxy-3H-phenoxazine-3-one Na salt) [635-78-9] $\text{C}_{12}\text{H}_7\text{NO}_3$, **M 213.2, m $>300^\circ$, λ_{max} 573nm. $\text{pK}^{30}_1 6.93$, $\text{pK}^{30}_2 9.26$, $\text{pK}^{30}_3 10.0$** . Wash Resorufin with water and recrystallise it several times from EtOH. **Resazurin** {the N-10-oxide of resorufin, [550-82-3, 62758-13-8 (Na salt)] $\text{C}_{12}\text{H}_7\text{NO}_4$, **M 229.2**} is a blue dye (above pH 6.5) that is weakly fluorescent but can be reversibly reduced to the pink coloured (below pH 3.8) and *highly red fluorescent resorufin*. It has been used as an oxidation-reduction indicator in cell viability assays. [González-Pinzón et al. *J Geophys Res* 117 (G3) G00N06, Bibcode:2012JGRG.117.0N06G 2012, DOI: 10.1029/2012JG001965; Beilstein 27 III/IV 2263; see detail in 'Physiologically Active Compounds', Chapter 6]

Rhodium (I) carbonyl chloride (di- μ -chloro-tetracarbonyl dirhodium I) [14532-22-9] $[\text{Rh}(\text{CO})_2\text{Cl}]_2$, **M 388.8, $d^{25}_{\text{4}} 2.708\text{g/cm}^3$, m 121°, 124-125°**. This catalyst is soluble in most organic solvents, but not petroleum ethers, and forms orange-red crystals from hexane. It sublimes at $80^\circ/0.1\text{mm}$ to a red solid. It decomposes on exposure to air when in organic solvents but the solid is stable in dry air. It is moisture sensitive and should be stored in ampoules under N_2 or argon. It catalyses ring-opening silylformylation of olefins. [McCleverty & Wilkinson *Inorg Synth* 8 211 1966, DOI: 10.1002/9780470132463; Dahl et al. *J Am Chem Soc* 83 1761 1961, DOI: 10.1021/ja01468a049; Cramer et al. *Inorg Synth* 15 14 1974, DOI: 10.1002/9780470132463.ch4; Fukumoto et al. *J Org Chem* 58 4187 1993, DOI: 10.1021/jo00068a004; Colton et al. *Aust J Chem* 23 1351 1970, DOI: 10.1071/CH9701351.]

Rhodizonic acid sodium salt (5,6-dihydroxycyclohex-5-ene-1,2,3,4-tetraone di-Na salt) [523-21-7] $\text{C}_6\text{O}_6\text{Na}_2$, **M 214.0, m $>300^\circ$, $\text{pK}^{25}_1 4.378 (\pm 0.009)$, $\text{pK}^{25}_2 4.652 (\pm 0.014)$, $\text{pK}^{30}_1 4.1 (4.25)$, $\text{pK}^{30}_2 4.5 (4.72)$** . The *free acid* is obtained by acidifying and extracting with Et_2O , drying (MgSO_4), filtering, evaporating and distilling in a vacuum (**b 155-160°/14mm**). The *free acid* solidifies on cooling, and the colourless crystals can be recrystallised from tetrahydrofuran/petroleum ether or $^*\text{C}_6\text{H}_6$. It forms a *dihydrate* **m 130-140°, 131-132°, [118-76-3] $\text{C}_6\text{H}_2\text{O}_6 \cdot 2\text{H}_2\text{O}$** . The *pure di Na salt* is formed by dissolving the acid in 2 equivalents of NaOH and evaporating in a vacuum. It forms violet crystals which give an orange solution in H_2O that is unstable for extended periods even at 0° , and should be prepared freshly before use. Salts of rhodizonic acid **cannot** be purified by recrystallisation without great loss due to conversion to crotonate, so that the original material must be *prepared anew* if pure salt is required. It can be washed with NaOAc solution, then EtOH, to

remove excess NaOAc, dried under vacuum and stored in the dark. Rhodizonates have various shades of red, from yellow to purple with respective colours of their solutions depending on the pH, and transmitting reflected light with a greenish lustre. [UV and tautomerism: Schwarzenbach & Suter *Helv Chim Acta* **24** 617 1941, DOI: 10.1002/hlca.19410240183; Polarography: Preisler & Berger *J Am Chem Soc* **64** 67 1942, DOI: 10.1021/ja01253a016; Souchay & Taibouet *J Chim Phys* **49** C108 1952, *Beilstein* **8** H 535, **8** II 572, **8** III 4214, **8** IV 3609.]

Ruthenium (III) acetylacetonate [14284-93-6] ($\text{C}_5\text{H}_7\text{O}_2$)₃Ru, **M 398.4, m 240°(dec)**. Purify the complex by recrystallisation from *benzene. [Wilkinson *J Am Chem Soc* **74** 6146 1952, DOI: 10.1021/ja01143a538; *Beilstein* **1** IV 3677.] Ru(acac)₃ complex catalyses the reduction of dimethyl oxalate to ethylene glycol under mild hydrogenation conditions [Teunissen & Elsevier *JCS Chem Commun* 667 1997, DOI: 10.1039/A700862G].

Ruthenocene [bis-(π -cyclopentadienyl)ruthenium] [1287-13-4] $\text{C}_{10}\text{H}_{10}\text{Ru}$, **M 231.2, m 195.5°, 199-210°**. If it has to be prepared *ab initio*, then in a flask (500ml), under continuous flushing with N₂, and stirring, dry 1,2-dimethoxyethane (300ml) is added followed by Na (7.2g, 0.312 g.atom) cut in small pieces then cyclopentadiene (31.0ml, 0.376 mole; see [542-92-7], prepared from the dimer, cf. Wilkinson *Org Synth Coll Vol* **4** 473 1963, DOI: 10.15227/orgsyn.036.0031] is added dropwise. After H₂ evolution stops, the mixture is kept just below boiling for 1-2 hours until all the Na has dissolved. If some metal remains undissolved, cool, and add a few milliliters of cyclopentadiene, and heat again until complete dissolution of the metal occurs. To this solution of sodium pentadienide [see also 4989-82-1], under N₂ and stirring, is added a mixture of RuCl₃ (14.6g, 0.07 mole) and Ru metal (2.4g, 0.024 g.atom), and heated just below reflux for 80 hours. The solvent is removed under reduced pressure, the flask is filled with N₂, the solid residue is transferred (use a dry-box) to a sublimator, and sublimed at 0.1mm and 130° (bath temperature). It is advantageous to use a Dry-ice sublimation finger. The sublimate is dissolved in *C₆H₆, passed through a column of activated Al₂O₃ (2.5 x 30cm), and evaporated to give pure **non-pyrophoric ruthenocene** (12.2-15.1g, 56-69%), m 199-200°. The residue from sublimation is, however, **pyrophoric**, but this residue can be destroyed by adding H₂O under N₂. It can be further sublimed in high vacuum at 120°. It forms yellow crystals which can be recrystallised from CCl₄ as transparent plates. [Bublitz et al. *Org Synth Coll Vol* **5** 1001 1973, DOI: 10.15227/orgsyn.041.0096; Wilkinson *J Am Chem Soc* **74** 6146 1952, DOI: 10.1021/ja01143a538; *Beilstein* **16** IV 1833.] It is generally less reactive than ferrocene towards electrophiles.

Two other alternative syntheses have been adopted using Schlenk techniques in an atmosphere of N₂ or argon, with dry and de-oxygenated solvents. In the **first**, [Sn-*n*-Bu₃(C₅H₅)] (32.0g, 90mmol, prepared from Sn-*n*-Bu₃Cl, 1461-22-9, and LiC₅H₅ as per Davison & Rakita *Inorg Chem* **9** 289 1970, DOI: 10.1021/ic50084a022) is added to a suspension of [RuCl₂(COD)_n] (8.4g, 30mmol, see [50982-12-2]) in EtOH (150ml), and the mixture is stirred and refluxed for 48 hours whereby the Ru polymer dissolved. The hot solution is filtered in air, and on cooling the **pale yellow ruthenocene** that crystallises out is filtered off, washed thoroughly with cold EtOH (2 x 20ml) then Et₂O (2 x 20ml), and recrystallised to analytical purity (5.2g, 75%) from hot EtOH or Me₂CO. Its ¹H NMR (500MHz, CDCl₃) has δ at 4.55 (s, C₅H₅), and its ¹³C NMR (125MHz, CDCl₃) has δ at 20.1 (s, C₅H₅) (any *n*-butyl signals from impurities would be at their usual shifts). In the **second**, [RuCl₂(COD)_n] (1.4g, 50mmol) and cyclopentadienyl thallium (2.7g, 10mmol, see [34822-90-7]) in dimethoxyethane (~50ml) are refluxed for 1 hour, filtered hot (in air), the solvent is removed *in vacuo*, the residue is extracted with Et₂O (4 x 20ml), filtered, evaporated, and the residue is recrystallised and/or sublimed, as above, to give pure **ruthenocene** (0.9g, 78%). [Albers et al. *Organometallics* **5** 2321 1986, DOI: 10.1021/om00142a025.] The Raman spectrum of the melt has ν_{max} at 160w, 325s(p), 396m, 590vw, 815m(p), 830w, 1060m, 1098s(p), 1195vw 1360w, 1410m cm⁻¹ (p is with polarized light) [Aleksanyan et al. *J Organomet Chem* **124** 293 1977, DOI: 10.1016/S0022-328X(00)92593-2].

Silver acetate [563-63-3] C₄H₃O₂Ag, **M 166.9, pK²⁵ >11.1 (for aquo Ag⁺ hydrolysis)**. Shake AgOAc with acetic acid for three days, and the process is repeated with fresh acid. The solid is then dried in a vacuum oven at 40° for 48 hours. It has also been recrystallised from water containing a trace of acetic acid, and dried in air. Store it in the dark. [*Beilstein* **2** IV 112; Fieser **1** 1002, **2** 362, **6** 511, **8** 440.] Effective cycloaddition reactions of isocyanoacetates with a variety of olefins are catalysed by silver acetate [Grigg et al. *Tetrahedron* **55** 2025 1999, DOI: 10.1016/S0040-4020(98)01216-2], which is also used in a novel preparation of highly reflect-

ive and conductive silvered polymer films [Southward et al. *Chem Mater* **11** 501 1999, DOI: 10.1021/cm981014v].

Silver lactate [128-00-7] $\text{C}_3\text{H}_5\text{O}_3\text{Ag}$, **M 196.9**, **m $\sim 100^\circ$, 120-122 $^\circ$** . Recrystallise it from H_2O by adding EtOH. The solid is collected, washed with EtOH, then Et_2O , and dried at 80° to give the *dihydrate*. It is a white powder soluble in 15 parts of H_2O but only slightly soluble in EtOH. Store it in the dark. [Engelhardt & Maddrell *Justus Liebigs Ann Chem* **63** 83 1847, DOI: 10.1002/jlac.18470630104; Karrer et al. *Helv Chim Acta* **2** 242 1919, DOI: 10.1002/hlca.19190020126; *Beilstein* **3** III 464.] It is used as a binder in compresses and is an antiseptic.

Silver tetrafluoroborate [14104-20-2] AgBF_4 , **M 194.7**, **m 70-73 $^\circ$, dec $>200^\circ$ (to form a silver mirror)**. The anhydrous salt has been freshly prepared under dry N_2 , by suspending AgF (34g, 268mmol) in dry MeNO_2 (30.5ml) while BF_3 is passed through with stirring. The temperature rises to 60° , and is maintained by occasional dipping into an ice bath. After about 30 minutes most of the AgF dissolves and ~ 280 mmoles of BF_3 will have dissolved. Excess BF_3 is removed by bubbling dry N_2 for 1 hour through the murky solution which is then filtered (through a sintered glass frit, under N_2), and the MeNO_2 is distilled off at $70^\circ/5\text{mm}$ into a liquid N_2 trap. The off-white solid is ground in a mortar under dry *n*-pentane in a glove box flushed with dry N_2 , and the solvent is evaporated under N_2 at $\sim 25^\circ$. The residual salt (237mmole, 88.5%) is pumped dry at 70° and stored under *n*-pentane. [Olah & Quin *J Inorg Nucl Chem* **14** 295 1960, DOI: 10.1016/0022-1902(60)80279-5.] The synthesis has been carried out in $^*\text{C}_6\text{H}_6$ with which it forms a **1:2 complex** [Heyns & Paulsen *Angew Chem* **72** 349 1960, DOI: 10.1002/ange.19600721009], although a **1:1 complex** [Meerwein et al. *Arch Pharm* **291** 541 1958, DOI: 10.1002/ardp.19582911103] and **2:3 complexes** [Sharp & Sharpe *J Chem Soc* 1855 1956, DOI: 10.1039/JR9560001855] were reported. AgBF_4 is very soluble in H_2O , Et_2O , toluene, MeNO_3 , moderately soluble in $^*\text{C}_6\text{H}_6$, cyclohexene but insoluble in pentane and cyclohexane. The solubility in unsaturated hydrocarbons is attributed to π -bond complexing. A solution obtained by dissolving Ag_2O (1.0g) in 45% fluoroboric acid (7.2g) has been used for the synthesis of *Nezukone* (4-isopropyltropone) [Birch & Keeton *J Chem Soc (C)* 109 1968, DOI: 10.1039/J39680000109]. The salt is used for making tetrafluoroborate salts of complexes and other salts (see examples in this chapter) by displacing chloride from precious metal complexes, e.g. $\text{Ru}(\text{Me}_2\text{SO}_4)\text{Cl}_2$ with excess AgBF_4 under N_2 in refluxing EtOH (*ca* 1 hour) and repeatedly recrystallising from $\text{H}_2\text{O}/\text{EtOH}$ to give a 60% yield of $[\text{Ru}(\text{Me}_2\text{SO}_4)(\text{H}_2\text{O})_2] \text{BF}_4$, which is used for the oxidation of alcohols by persulfate [Bressan et al. *J Mol Catal* **79** 85 1993, DOI: 10.1016/0304-5102(93)85093-9]. It provides a silver-ion template for improved macrolactamisation of linear dipeptides, e.g. to form cyclohexadepsipeptides [Yanjie et al. *Synlett* 1901 2007, DOI: 10.1055/s-2007-984510]. [For applications see Fieser **1** 1015, **2** 365, **3** 250, **4** 428, **5** 587, **6** 519, **8** 443, **9** 414, **11** 471, **12** 434, **13** 273.]

Silver trifluoroacetate [2966-50-9] $\text{C}_2\text{F}_3\text{O}_2\text{Ag}$, **M 220.9**, **m 251-255 $^\circ$, 251-260 $^\circ$ (dec)**. Extract the salt (Soxhlet) with Et_2O . The extract is filtered and evaporated to dryness, then the powdered residue is completely dried in a vacuum desiccator over silica gel. Its solubility in Et_2O is 33.5g in 750ml. It can be recrystallised from $^*\text{C}_6\text{H}_6$ (solubility is: 1.9g in 30ml of $^*\text{C}_6\text{H}_6$, and 33.5g will dissolve in 750ml of anhydrous Et_2O). [Traynham & Dehn *J Org Chem* **23** 1545 1958, DOI: 10.1021/jo01104a039; Haszeldine *J Chem Soc* 584 1951, DOI: 10.1039/JR9510000584.] Store it in the dark. It is also soluble in trifluoroacetic acid (15.2% at 30°), toluene, *o*-xylene and dioxane [Hara & Cady *J Am Chem Soc* **76** 4285 1954, DOI: 10.1021/ja01646a009]. [*Beilstein* **2** IV 461; For applications see Fieser **1** 1018, **7** 323, **8** 444, **10** 355, **11** 557, **12** 529, **16** 301, **17** 313.]

Sodium acetate (anhydrous) [127-09-3, 6131-90-4 ($3\text{H}_2\text{O}$)] $\text{C}_4\text{H}_3\text{O}_2\text{Na}$, **M 82.0**, **m 58 $^\circ$ ($3\text{H}_2\text{O}$), $>300^\circ$ (dec), 324 $^\circ$ (anhydr), **b 881.4 $^\circ$ /atm (anhydr), 122 $^\circ$ /atm ($3\text{H}_2\text{O}$, dec), $d^{20} 1.528\text{g}/\text{cm}^3$ (anhydr), $d^{20} 1.45\text{g}/\text{cm}^3$ ($3\text{H}_2\text{O}$), $n_D^{25} 1.464$. Crystallise it from acetic acid and keep it under vacuum for 10 hours at 120° . Alternatively, it crystallises from aqueous EtOH as the *trihydrate*. This material can be converted to *anhydrous* salt by heating slowly in a porcelain, nickel or iron dish, so that the salt liquefies. Steam is evolved and the mass again solidifies. Heating is now increased so that the salt melts again. (*Note*: if it is heated too strongly, the salt can char; avoid this.) After several minutes, the salt is allowed to solidify and is cooled to a convenient temperature (in a desiccator) before being powdered and bottled. The water content should now be less than 0.02%. It is a white *deliquescent* powder. The solubility (g/100ml) in H_2O is 119 (0°), 123.3 (20°), 137.2 (60°) and 162.9 (100°) for the *anhydrous salt*, and 32.9 (-10°), 36.2 (0°), 46.4 (20°) and 82 (50°) for the *trihydrate*; in****

MeOH it is 16 (15°) and 16.55 (67.7°) for *anhydrous salt*; in EtOH it is 5.3 (25°) for the *trihydrate*; and in Me₂CO it is 0.05 (15°). [*Beilstein* 2 II 113, 2 III 184, 2 IV 109; Fieser 1 1024, 5 591.]

Sodium acetylacetonate [Na(acac), 2,4-pentanedione ion (1-) sodium (1:1)] [15435-71-9] C₅H₇O₂Na, M 122.1, m 217-219° and is quite stable to heating below 191°, d₄²⁰ 1.213. It is prepared by two different procedures. In the *first*, 2,4-pentanedione (100g, 1.0 mole) in H₂O or EtOH is slowly added to NaOH (40g, 1.0 mole) in the minimum amount of H₂O while keeping the temperature below 70°. On cooling Na(acac) (98g, 80%) crystallises out. Note that addition of solid NaOH resulted in a product containing NaOH. Recrystallisation from aqueous EtOH, Me₂CO or EtOAc gives pearly plates of the *dihydrate* (most probably a coordinated compound) that is soluble EtOAc but not in toluene. It gives off H₂O on heating, and the *anhydrous* form (salt form ?) is no longer soluble in EtOAc. In the *second* procedure, 2,4-pentanedione (50g, 0.5 mole) is added to Na metal (11.5g, 0.5 mole) in toluene (100ml) while the temperature is maintained at 100° to keep the metal in the molten state (**CARE**, as the experiment may be dangerous; N₂ should be flushing through the apparatus until the metal has dissolved). Na(acac) (101g, 83%) separates during the reaction. **Note** that using lower temperatures provide Na(acac) contaminated with Na. The product can be recrystallised from EtOH (see above) or by dissolving in EtOH and adding *C₆H₆. Its solubility in *n*-hexane, cyclohexane, or *C₆H₆ is less than 0.01% at 70°. However, although quite insoluble in *C₆H₆ or toluene, a suspension of Na(acac) will dissolve in these solvents on warming if excess of 2,4-pentanedione, or other related dicarbonyl compounds, e.g. benzoylacetone or ethyl acetoacetate, are added. **Note** that on heating the anhydrous or hydrated material decomposition and charring occurs if the sample in a soft glass tube is used. However, if the tube is placed in a bath at ~212° and the temperature is raised slowly, a sharp and *reproducible melting point* (217-219°) can be obtained with very little decomposition. [Hatch & Sutherland *J Org Chem* 13 249 1948, DOI: 10.1021/jo01160a012; Sidgwick & Brewer *J Chem Soc* 127 2379 1925, DOI: 10.1039/CT9252702379.] The ratios of mole of decomposition product/mole of 'salt' after 50 hours when heated under N₂ at 27° and 191° are both 0.07, i.e. it is reasonably stable at high temperatures. [Charles & Pawlikowski *J Phys Chem* 62 440 1958, DOI: 10.1021/j150562a017.]

Its UV (EtOH) has λ_{max} at 277 (ε 800 x 10³) nm [Hatch & Sutherland *J Org Chem* 13 249 1948, DOI: 10.1021/jo01160a012]; and its electrical molecular conductivities (dilution) are: 5.794 (8), 8.254 (16), 11.187 (32), 14.426 (64), 18.116 (128), 21.82 (256), 25.636 (512), 29.927 (1024), and do not approach complete dissociation when nearing maximum dilution as hydrolysis may well be occurring [White *J Chem Soc* 1413 1928, DOI: 10.1039/JR9280001413]. Like K(acac) and Cs(acac), Na(acac) is unstable in aqueous solution or in moist conditions and is decomposed by hot H₂O into Me₂CO and the alkali acetate [Morgan & Moss *J Chem Soc* 105 189 1914, DOI: 10.1039/CT9140500189]. Store it in a dry atmosphere. [Marchi *Inorg Synth* 2 10 1946, DOI: 10.1002/9780470132333.ch4; Fernelius & Bryant *Inorg Synth* 5 105 1957, DOI: 10.1002/9780470132364.ch29.]

Sodium acetylide [1066-26-8] C₂HNa, M 48.0. It disproportionates at ca 180° to sodium carbide. It sometimes contains diluents, e.g. xylene, butyl ether or dioxane that can be removed by filtration followed by a vacuum at 65-60°/5mm. *Alternatively*, the acetylide is purged with HC≡CH at 100-125° to remove diluent. NaC₂H adsorbs 2.2x, 2.0x and 1.6x its wt of xylene, butyl ether and dioxane, respectively. Powdered NaC₂H is yellow or yellow-grey in colour and is relatively stable. It can be heated to ca 300° in the absence of air. Although no explosion or evolution of gas occurs, it turns brown due to disproportionation. At 170-190° in air it ignites slowly and burns smoothly. At 215-235° in air it 'flash-ignites' and burns quickly. It can be dropped into a *slight excess* of H₂O without flashing or burning, but vigorous evolution of HC≡CH (**HIGHLY FLAMMABLE IN AIR**) occurs. A sample had been stored in the absence of air for one year without deterioration. Due to the high flammability of HC≡CH, the salt should be stored dry and should be treated with care. After long storage, NaC≡CH can be redissolved in liquid NH₃ and used for the same purposes as the fresh material. However, it may be slightly turbid due to the presence of moisture. [Rutledge *J Org Chem* 22 649 1957, DOI: 10.1021/jo01357a016; Greene *J Am Chem Soc* 77 5013 1955, DOI: 10.1021/ja01624a019; Greenlee et al. *Inorg Synth* 2 75 1946, DOI: 10.1002/9780470132333.ch21; Campbell & Campbell *Org Synth* 30 15 1950, DOI: 10.15227/orgsyn.0300015; *Beilstein* 1 H 238; Fieser 1 1027.] It is available commercially under N₂ in Sure/Seal bottles as an 18 wt% solution in xylene/mineral oil. See 'Aliphatic Compounds', Chapter 3, for its preparation.

Sodium alginate (Algin) [9005-38-3] ($\text{C}_6\text{H}_8\text{O}_6$)_n Na_m, **M 48,000-186000**. This is the Na salt of *alginic acid* {see [9005-32-7] in Carbohydrates Chapter 7}. Free Algin from heavy metal impurities by treatment with ion-exchange resins (Na^+ -form), or with a dilute solution of the sodium salt of EDTA. *Alternatively*, dissolve it in 0.1M NaCl, centrifuge and fractionally precipitate it by gradual addition of EtOH or 4M NaCl. The resulting gels are centrifuged off, washed with aqueous EtOH or acetone, and dried under vacuum. [Büchner et al. *J Chem Soc* 3974 1961, DOI: 10.1039/JR9610003974.]

Sodium 4-aminobenzoate [555-06-6] $\text{C}_7\text{H}_6\text{NO}_2\text{Na}$, **M 159.1**. Recrystallise it from water. [Hermann *Helv Chim Acta* **9** 785 1926, DOI: 10.1002/hlca.192600901103; *Beilstein* **14** II 247.]

Sodium 4-aminosalicylate dihydrate [6018-19-5] $\text{C}_7\text{H}_6\text{NO}_3\text{Na} \cdot 2\text{H}_2\text{O}$, **M 175.1, m 250°**. Recrystallise it from water at room temperature (2ml/g) by adding acetone and cooling. [*Beilstein* **14** III 1436, **14** IV 1969.]

Sodium antimonyl tartrate (Stibnal, sodium tartar emetic) [34521-09-0] $\text{C}_8\text{H}_4\text{O}_{12}\text{Na}_2\text{Sb}_2$, **M 581.6, alternatively, $\text{C}_4\text{H}_4\text{O}_7\text{NaSb}$, M 308.8**. It crystallises from water. [*Beilstein* **3** III 1014, **3** IV 1227.]

Sodium barbitone (sodium 5,5-diethylbarbiturate) [144-02-5] $\text{C}_8\text{H}_{11}\text{N}_2\text{O}_3\text{Na}$, **M 206.2, pK₁²⁵ 3.99, pK₂²⁵ 12.5 (barbituric acid)**. Crystallise it from water (3ml/g) by adding an equal volume of EtOH and cooling to 5°. Dry it under vacuum over P_2O_5 . It is a useful buffer. [*Beilstein* **24** III/IV 1904.]

Sodium benzoate [532-32-1] $\text{C}_7\text{H}_5\text{O}_2\text{Na}$, **M 144.1, m >300°, 410°, d²⁵ 1.497g/cm³**. Crystallise the white powder from EtOH (12ml/g). Its solubility (g/100ml) in H_2O is 62.69 (0°), 62.87 (30°) and 71.11 (100°); in MeOH it is 8.22 (15°) and 7.55 (67.2°); in EtOH it is 2.3 (25°) and 8.3 (78°); and in 1,4-dioxane it is 0.082 (25°). It is soluble in liquid NH_3 and pyridine. [*Beilstein* **9** IV 27; Fieser **1** 1044, **2** 377.] It is used as a food additive and preservative.

Sodium tert-butoxide [865-48-5] $(\text{CH}_3)_3\text{CONa}$, **M 96.1**. It sublimes at 180°/1mm. Its solubility in *tert*-BuOH is 0.208M at 30.2° and 0.382M at 60°, and it is quite soluble in tetrahydrofuran (32g/100g). It should not be used if it has a brown colour. [Feuer et al. *J Am Chem Soc* **78** 4364 1956, DOI: 10.1021/ja01598a045; Hurd et al. *Inorg Synth* **1** 87 1939, DOI: 10.1002/9780470132326.ch30; IR: Seubold *J Org Chem* **21** 156 1956, DOI: 10.1021/jo01108a002; *Beilstein* **1** IV 1609.] It is a strong non-nucleophilic base.

Sodium butyrate [156-54-7] $\text{C}_4\text{H}_7\text{O}_2\text{Na}$, **M 110.1, m 250-253°**. Prepare it by neutralising the acid with Na_2CO_3 , and recrystallising it from EtOH. [*Beilstein* **2** IV 779.] It is physiologically active: decreases Ca^{2+} release from intracellular stores, inhibits histone deacetylase and induces apoptosis in several cell lines.

Sodium carboxymethylcellulose [9004-32-4]. Dialyse it for 48 hours against distilled water and freeze-dry if a solid is required. It is commercially available in Av. M_w from ~90,000 (m 274° dec) to ~700,000 (m 270° dec).

Sodium decanoate (sodium caproate) [1002-62-6] $\text{C}_{10}\text{H}_{19}\text{O}_2\text{Na}$, **M 194.2**. Neutralise sodium hydroxide by adding a slight excess of free decanoic acid and recovering the excess acid by Et_2O extraction. The salt is recrystallised from the aqueous solution by adding pure acetone and repeating the steps several times, then drying the salt in an oven at ca 110° [Chaudhury & Awuwallia *Trans Faraday Soc* **77** 3119 1981, DOI: 10.1039/F19817703119]. [*Beilstein* **2** IV 1041.]

Sodium deoxycholate monohydrate [302-95-4, 145224-92-6] $\text{C}_{24}\text{H}_{39}\text{O}_4\text{Na} \cdot \text{H}_2\text{O}$, **M 432.6, [α]_D²⁰ +48 (c 1, EtOH), +44 (c 1, H_2O)**. Recrystallise it from EtOH and dry it in an oven at 100°. The solution is freed from soluble components by repeated extraction with acid-washed charcoal. [*Beilstein* **10** IV 1608.] It is a useful protein solubilising agent

Sodium diethyloxaloacetate [63277-17-8, 40876-98-0] $\text{C}_8\text{H}_{11}\text{O}_5\text{Na}$, **M 210.2, m 188-190°**. Extract it several times with boiling Et_2O (until the solvent remains colourless), and then the residue is dried in air. [*Beilstein* **3** H 789.]

Sodium diformylamide [18197-26-7] $(\text{CHO})_2\text{NNa}$, **M 95.0**. Grind the amide under dry tetrahydrofuran (fumehood), filter and wash it with this solvent, then dry it *in vacuo*. It is soluble in EtOH and H_2O but insoluble in Et_2O and petroleum ether. [IR and preparation: Rakshit *J Chem Soc* **103** 1557 1913, DOI: 10.1039/CT9130301557; Yinglin & Hongwen *Synthesis* 122 1990, DOI: 10.1055/s-1990-26805; Allenstein &

Beyl *Chem Ber* **100** 3551 1967, DOI: 10.1002/cber.19671001110; Allenstein et al. *Chem Ber* **102** 4089 1969, DOI: 10.1002/cber.19691021216; *Beilstein* **2** II 22.]

Sodium *p*-dimethylaminoazobenzene-*o*'-carboxylate (Methyl Red Na salt) [845-10-3] $C_{15}H_{14}N_3O_2Na$, M **291.2.** It can be precipitated from aqueous solution as the *free acid* which is recrystallised from 95% EtOH, then reconverted to the sodium salt. In H_2O it is pink at pH 4.2 and yellow at pH 6.2 and has UV with λ_{max} at 437nm. **Methyl Red Hydrochloride** [63451-28-5] has M **305.8**, m **175°(dec)** and UV with λ_{max} at 493nm. [*Beilstein* **16** H 329, **16** II 164.]

Sodium *p*-dimethylaminoazobenzene-*p*'-carboxylate [845-46-5] $C_{15}H_{14}N_3O_2Na$, M **291.3.** It has been precipitated from aqueous solution as the free acid which was recrystallised from 95% EtOH, then reconverted to the sodium salt. [*Beilstein* **16** H 329.]

Sodium dodecanoate (sodium laurate) [629-25-4] $C_{12}H_{23}O_2Na$, M **222.3 m **244-246°**. pK^{20} **5.3** (COOH).** Neutralise it by adding a slight excess of dodecanoic acid and removing it by ether extraction. The salt is recrystallised from the aqueous solution by adding pure Me_2CO and repeating the process (see sodium decanoate). It has also been recrystallised from MeOH. It is a detergent. [*Beilstein* **2** IV 1085.]

Sodium ethoxide [141-52-6] C_2H_5ONa , M **68.1.** It is a *hygroscopic* powder which should be stored under N_2 in a cool place. A likely impurity is EtOH which can be removed by warming at 60-80° under high vacuum. It is hydrolysed by H_2O to yield NaOH and EtOH. Other impurities, if kept in air for long periods are NaOH and Na_2CO_3 . In this case the powder cannot be used if these impurities affect the reactivity, and a fresh sample would be acquired. It is prepared by adding Na to absolute EtOH under N_2 and evaporated *in vacuo*, or used *in situ*. [IR: Seubold *J Org Chem* **21** 156 1956, DOI: 10.1021/jo01108a002]. [*Beilstein* **1** IV 1289; Fieser **1** 1065, **3** 265, **4** 451, **6** 540.] Commercially available as a 21wt% solution in EtOH (d^{25} **0.868g/ml³**, n_D^{20} **1.385**).

Sodium ethylmercurithiosalicylate (Thimerosal) [54-64-8] $(C_2H_5HgS)C_6H_4CO_2Na$, M **404.8, m **~230°**.** Recrystallise this antibacterial from EtOH/Et₂O. It is also used as an antifungal preservative. **HIGHLY TOXIC**. [Trikojus *Nature* **158** 472 1946, DOI: 10.1038/158472a0; *Beilstein* **10** III 213, **10** IV 272.]

Sodium fluoroacetate (FCH_2CO_2Na) [62-74-8] $C_2H_2FO_2Na$, M **100.0, m **200-205°(dec)**.** It is a free-flowing white **HIGHLY POISONOUS** powder which is purified by dissolving it in *ca* 4 parts of H_2O , and the pH is checked. If it is alkaline, add a few drops of FCH_2CO_2H to make the solution just acidic. Evaporate (fumehood) on a steam bath until crystals start to separate, cool and filter the solid off. More solid can be obtained by adding EtOH to the filtrate. Dry it at 100° *in vacuo*. The *p*-nitrobenzyl ester crystallises from EtOH with m **76°**. The *free acid* interferes with the citric acid cycle. [Saunders & Stacey *J Chem Soc* 1773 1948, DOI: 10.1039/JR9480001773; *Beilstein* **2** IV 446.] **VERY TOXIC**, lethal oral human dose is 2–10mg/Kg.

Sodium formate (anhydrous) [141-53-7] CHO_2Na , M **68.0, m **253°**, **259-262°**, d^{25} **1.92g/cm³**.** A saturated aqueous solution at 90° (0.8ml water/g) is filtered and allowed to cool slowly. (The final temperature should be above 30° to prevent formation of the hydrate.) After two such crystallisations, the crystals are dried in an oven at 130°, then under high vacuum. [Westrum et al. *J Phys Chem* **64** 1553 1960, DOI: 10.1021/j100839a052; Roecker & Meyer *J Am Chem Soc* **108** 4066 1986, DOI: 10.1021/ja00274a035.] The salt has also been recrystallised twice from 1mM DTPA (diethylenetriaminepentaacetic acid, which was recrystallised 4x from MilliQ water and dried in a vacuum), then twice from water [Bielski & Thomas *J Am Chem Soc* **109** 7761 1987, DOI: 10.1021/ja00259a026]. [*Beilstein* **2** IV 3, Fieser **8** 458]

Sodium D-gluconate [527-07-1] $C_6H_{11}O_7Na$, M **218.1, m **200-205°dec**, $[\alpha]_{546}^{20}$ **+14**, $[\alpha]_D^{20}$ **+12** (c 20, H_2O).** Crystallise it from a small volume of H_2O (solubility is 59g/100ml at 25°), or dissolve it in H_2O and add EtOH since it is sparingly soluble in EtOH. It is insoluble in Et₂O. It forms a Cu complex in alkaline solution and a complex with Fe in neutral solution. [Sawyer & Bagger *J Am Chem Soc* **81** 5302 1959, DOI: 10.1021/ja01529a014; *Beilstein* **3** I 188, **3** IV 255.]

Sodium glycochenodeoxycholate [640-79-9, 16564-43-5] $C_{26}H_{42}NO_5Na$, M **471.6, m **245-250°**, $[\alpha]_D^{23}$ **+41.2****

(**c 1, H₂O**). Dissolve it in EtOH, filter it and concentrate it to crystallisation, and recrystallise from a little EtOH. It also recrystallises from EtOH/Et₂O. [*Beilstein* **10** IV 1611.] [For behaviour with sodium glycocholate in binary mixed micelles of bile salt and non-ionic surfactant see Takamura et al. *Colloids and Surfaces B: Biointerfaces* **7**(5) 239 1996, DOI: 10.1016/0927-7765(96)01304-5.]

Sodium glycocholate (N-cholylglycine sodium salt hydrate) [863-57-0] **C₂₆H₄₂NO₆Na.xH₂O, M 487.6(anhydr), m 210-215°, 230-240°, micellar avg wt 1000, CMC 7mM (20-25°), [α]_D²³ +29.2 (c 1, H₂O).** Dissolve this bile salt in EtOH, filter it and concentrated to crystallisation, and recrystallise from a little EtOH. It also crystallises from 95% EtOH/Et₂O. [*Beilstein* **3** IV 573, **10** IV 2077.]

Sodium glycolate dihydrate (sodium hydroxyacetate) [2836-32-0] **C₂H₃O₃Na. 2H₂O, M 98.0(anhydr), m 210-218°, pK_a²⁵ 3.83 (for acid).** Precipitate it from aqueous solution by adding EtOH and dry it in air. Also recrystallise it from H₂O, where its solubility is 38% at 0° and 61% at 100°. [*Beilstein* **3** III 370, **3** IV 573.]

Sodium hydrogen diglycollate (2,2'-oxydiacetic acid monosodium salt) [50795-24-9] **C₄H₅O₅Na, M 156.1, pK₁²⁰ 2.70, pK₂²⁰ 4.22 (for acid).** Crystallise it from hot water (7.5ml/g) by cooling to 0° with constant stirring. The crystals are filtered off on to a sintered-glass funnel and dried at 110° overnight. Its solubility in water is 2.6% (at 0°) and 20% (at 90°). [*Beilstein* **3** III 377, **3** IV 577.] [For use as a reference buffer see Keyworth & Hahn *Talanta* **1** 41 1958, DOI: 10.1016/0039-9140(58)80006-5.]

Sodium hydrogen oxalate monohydrate [1186-49-8] **C₂H₂O₄Na. H₂O, M 130.0, m 100°(loses H₂O), b 200°(dec).** Crystallise it from hot water (5ml/g) by cooling. Its solubility in H₂O is 1.7% at 15.5° and 21.3% at boiling. [Souchay & Lenfsen *Justus Liebigs Ann Chem* **99** 31 1856, DOI: 10.1002/jlac.18560990104; for twin structure by X-ray crystallography see Komunjer & Lefaucheux *Crystal Res Tech* **30**(4) 463 1995, DOI: 10.1002/crat.2170300407; *Beilstein* **2** H 513, **2** I 223, IV 1817.]

Sodium hydrogen succinate [2922-54-5] **C₄H₅O₄Na, M 140.0.** Crystallise it from hot water and dry it at 110°. Its solubility in H₂O is 17% at 0°, 40% at 25° and 86% at 75°. [Marshall & Bain *J Chem Soc* **97** [I] 1074 1910, DOI: 10.1039/CT9109701074; *Beilstein* **2** H 606, **2** I 262, **2** III 16557, **2** IV 1911.]

Sodium hydrogen d-tartrate [526-94-3] **C₄H₅O₆Na, M 172.1, m 100°(loses H₂O), m 253°dec, [α]_D²⁰ +26 (c 1, H₂O) and [α]_D²⁰ +24 (c 1, H₂O).** It crystallises from warm water (10ml/g) by cooling to 0°. [*Beilstein* **3** IV 1219.]

Sodium 2-hydroxy-4-methoxybenzophenone-5-sulfonate [6628-37-1] **C₁₄H₁₁O₆SNa, M 330.3.** Crystallise it from MeOH and dry it under vacuum.

Sodium p-hydroxyphenylazobenzene-p'-sulfonate [2623-36-1] **C₁₂H₁₃N₂O₆SNa. 2H₂O, M 336.3.** Recrystallise it from 95% EtOH.

Sodium ionophore I (ETH 227) (N,N',N''-triheptyl-N,N',N''-trimethyl-4,4',4''-propylidene-tris(3-oxabutylamide)) [61183-76-4] **C₃₆H₇₁N₃O₆, M 642.0.** It is purified (ca 200mg) by TLC on Kieselgel F₂₅₄ with CHCl₃/Me₂CO (1:1) as solvent, followed by HPLC (50mg) with an octadecyltrimethylsilane modified column (Mercksorb SI 100, 10μm) [IR, NMR, MS: Güggi et al. *Helv Chim Acta* **59** 2417 1976, DOI: 10.1002/hlca.19760590716]. [See Simon & Carafoli *Methods Enzymol* **56** 439 1977, DOI: 10.1016/0076-6879(79)56043-1.]

Sodium ionophore V (ETH 4120) [2,2'(4-octadecanoyloxymethyl-1,2-phenylenedioxy) N,N',N',N'-tetracyclohexyldiacetamide] [129880-73-5] **C₅₃H₈₈N₂O₆, M 849.3.** Purify it (ca 1.5g) by flash chromatography (120g of silica gel 60) with hexane/EtOAc (7:3) and recrystallisation from hexane/EtOAc [Prepn and properties: Géhrig et al. *Anal Chim Acta* **233** 295 1990, DOI: 10.1016/S0003-2670(00)83491-1].

Sodium ionophore VI {bis[(12-crown-4)methyl]dodecyl methyl malonate} [80403-59-4] **C₃₄H₆₂O₁₂, M 662.9.** Purify it by gel permeation or column chromatography. [Preparation and NMR data: Shono et al. *J Electroanal Chem* **132** 99 1982, DOI: 10.1016/0022-0728(82)85009-2; Cram & Cram *Science* **183** 803 1974, DOI: 10.1126/science.183.4127.803.]

Sodium RS-mandelate [114-21-6] **C₈H₇O₃Na, M 174.1, b 321.8°/760mm, pK_a²⁵ 3.41 (for the acid).** It crystallises from 95% EtOH and is dried in a vacuum. [Ross & Morrison *J Chem Soc* 1016 1933, DOI: 10.1039/

JR9330001016; Banks & Davies *J Chem Soc* 73 1938, DOI: 10.1039/JR9380000073; *Beilstein* 10 III 450.]

Sodium 2-mercaptoethanesulfonate (MESNA) [19767-45-4] $\text{C}_2\text{H}_5\text{NaO}_3\text{S}_2$, M 164.2, $\text{pK}_1^{20} < 0$, pK_2^{20} 9.53 (SH). Recrystallise it from H_2O and has $m > 250^\circ$. Alternatively, purify it by passing a 2M solution through Amberlite IRA-120 (H^+ form), evaporate the eluate *in vacuo*. The viscous oily residue is then carefully neutralised with aqueous NaOH and evaporated *in vacuo*. Recrystallise the salt residue from H_2O . [Schramm et al. *J Am Chem Soc* 77 6231 1955, DOI: 10.1021/ja01628a052; *Beilstein* 4 IV 85.] It is a detoxifying agent.

Sodium methoxide [124-41-4] CH_3ONa , M 54.0. It behaves like sodium ethoxide. It is *hygroscopic* and hydrolyses in moist air to NaOH and MeOH. Should be kept and used under N_2 . If erratic results are obtained, it should be freshly prepared thus: Clean Na (37g) cut in 1-3g pieces is added in small portions to stirred MeOH (800ml) in a 2L three-necked flask under a condenser with a drying tube. After all the Na has dissolved, the MeOH is distilled off *in vacuo*, and the residual NaOMe is dried at 150° *in vacuo* and kept under dry N_2 until required [Burness *Org Synth* 39 49 1959, DOI: 10.15227/orgsyn.039.0049]. [*Beilstein* 1 IV 1220; for applications see Fieser 1 1091, 2 385, 3 259, 4 457, 5 617, 6 545, 8 463, 9 430, 11 489.]

Sodium monensin [22373-78-0] $\text{C}_{36}\text{H}_{61}\text{O}_{11}\text{Na}$, M 692.8. Crystallise this polyether antibiotic from EtOH/ H_2O [Cox et al. *J Am Chem Soc* 107 4297 1985, DOI: 10.1021/ja00300a037]. It is a Na^+ ionophore, blocks glycoprotein secretion, and could induce catecholamine secretion from chromaffin cells. It is used in potentiometric and spectroscopic studies of alkali metal ion complexes.

Sodium oleate (sodium *cis*-9-octadecenoate) [143-19-1] $\text{C}_{18}\text{H}_{33}\text{O}_2\text{Na}$, M 304.3, has m 205° , 233 - 235° , and is recrystallised from EtOH, and dried at 100° . [*Beilstein* 2 IV 1645.] It is a soap and is an anti-foaming agent. It also activates protein kinase C in hepatocytes. **Sodium oxalate** [62-76-0] $\text{C}_2\text{O}_4\text{Na}$, M 134.0, has m 250 - $270^\circ(\text{dec})$, d_4^{20} 2.34, and crystallises from hot water (16ml/g). For volumetric analysis, analytical sodium oxalate should be dried at $120^\circ/2$ hours then cool in a desiccator. [*Beilstein* 2 IV 1819.] **Sodium palmitate** [408-35-5] $\text{C}_{16}\text{H}_{31}\text{O}_2\text{Na}$, M 278.4, has m 283 - 290° , 285 - 201° and crystallises from EtOH. [*Beilstein* 2 IV 1157.]

Sodium pentamethylcyclopentadienide (NaCp', 1,2,3,4,5-pentamethyl-2,4-pentadiene-1-ylsodium) [40585-51-1] $\text{C}_{10}\text{H}_{15}\text{Na}$, M 158.2, $m > 230^\circ$, d^{25} 0.905g/cm³. It is prepared, handled and stored as for KCp' but is not as flammable. It is a Cp' ligand source for complexing with a large number of metals and non-metal halides. [Kohl et al. in *Organometallic Synthesis* (R. Bruce King and J.J. Eisch eds) Vol 3 381, Elsevier Amsterdam 1986]. A 0.5M solution in THF is commercially available.

Sodium phenoxide [139-02-6, 156150-40-2 ($3\text{H}_2\text{O}$)] $\text{C}_6\text{H}_5\text{ONa} \cdot 3\text{H}_2\text{O}$, M 170.1, m 61 - 64° . The ground powder is washed with Et_2O , then heated at $60^\circ/1\text{mm}$ for 12 to 24 hours to remove any free phenol and solvent. [Kornblum & Lurie *J Am Chem Soc* 81 2705 1959, DOI: 10.1021/ja01520a030; *Beilstein* 6 I 718.]

Sodium phenylacetate [114-70-5] $\text{C}_8\text{H}_7\text{O}_2\text{Na}$, M 158.1. Its aqueous solution is evaporated to crystallisation on a steam bath; the crystals are washed with absolute EtOH and dried under vacuum at 80° . [*Beilstein* 9 IV 1614.]

Sodium *o*-phenylphenolate tetrahydrate [132-27-4] $\text{C}_{12}\text{H}_9\text{ONa} \cdot 4\text{H}_2\text{O}$, M 264.3. Crystallise the salt from acetone and dry it under vacuum at room temperature. This salt is a preservative, mould inhibitor and a fungicide [*Beilstein* 16 IV 4600.]

Sodium phenylpyruvate [114-76-1] $\text{C}_9\text{H}_7\text{O}_3\text{Na}$, M 186.1, $m > 300^\circ$. The salt should have no OH broad bands in the IR at $\sim 3000\text{cm}^{-1}$. If so then dry the solid thoroughly in a vacuum over P_2O_5 . Otherwise wash the salt well with Et_2O till free of acid and dry it *in vacuo*. Alternatively, add a slight excess of the *free acid* (see [156-06-9]) in EtOH to ethanolic NaOH, evaporate to dryness and extract excess acid from the salt with dry Et_2O . [*Beilstein* 10 I 325.]

Sodium phytate monohydrate [*myo*-inositolhexakis(H_2PO_4) Na salt] [14306-25-3] $\text{C}_6\text{H}_{18}\text{O}_{24}\text{P}_6\text{Na}_x \cdot x\text{H}_2\text{O}$, M 857.9. Crystallise sodium phytate from hot water. [*Beilstein* 6 IV 7927.]

Sodium polyacrylate (NaPAA) [9003-04-7] $[\text{C}_3\text{H}_3\text{O}_2\text{Na}]_n$. Commercial polyacrylamide is first neutralized with an aqueous solution of NaOH, and the polymer is precipitated with acetone. The precipitate is redissolved in a small amount of water and freeze-dried. The polymer is then repeatedly washed with EtOH and water to

remove traces of low-molecular-weight material, and finally dried in vacuum at 60° [Vink *JCS Faraday Trans 1* **75** 1207 1979, DOI: 10.1039/F19797501207]. It has also been dialysed overnight against distilled water, then freeze-dried. It is commercially available as a 35 wt% solution in H₂O of polymer with $M_w \sim 8,000$ (d_{25}^{25} 1.32g/cm³, n_D^{20} 1.428), and a 45 wt% solution in H₂O of polymer with $M_w \sim 15,000$ (d_{25}^{25} 1.25g/cm³, n_D^{20} 1.405).

Sodium poly(α-L-glutamate) (PLGNa) [C₅H₆NO₃Na]_n, M 1.1x10⁵. After saponification of poly(γ-methyl-α-L-glutamate), PLGNa is obtained by washing it with acetone, drying it in a vacuum, dissolving it in water and precipitating it with isopropanol at 5°. Impurities and low-molecular-weight fractions are removed by electrodialysis of the aqueous solution for 50 hours, followed by ultrafiltration through a Diafilter G 10 T impermeable to polymers of molecular weights greater than 10⁴. The polymer is recovered by freeze-drying and stored in a desiccator until used. The molecular weight was determined by viscosity measurements. [Mori et al. *JCS Faraday Trans 1* 2583 1978, DOI: 10.1039/F19787402583.]

Sodium propionate [137-40-6] C₃H₅O₂Na, M 96.1, m 285-286°, 287-289°. Recrystallise it from H₂O (solubility 10%) and dry by heating at 100° for 4 hours. The solubility of the anhydrous salt in MeOH is 13% at 15° and 13.77% at 68°. It is insoluble in *C₆H₆ and Me₂CO. [Henstock *J Chem Soc* 1340 1934, DOI: 10.1039/JR9340001340; *Beilstein 2* IV 701.]

Sodium stearate [822-16-2] C₁₈H₃₅O₂Na, M 306.6, m 245-255°, d_{25}^{25} 1.02g/cm³. It is better to prepare it by adding a slight excess of octadecanoic acid to ethanolic NaOH, evaporating and extracting the residue with dry Et₂O to remove free acid. It is a very common soap. [*Beilstein 2* III 1003.]

Sodium R-(+)-tartrate dihydrate [6106-24-7] C₄H₄O₆Na₂·2H₂O, M 230.1, m 120°(loses H₂O), d_4^{20} 1.82, [α]_D²⁰ +26 (c 1, H₂O). It crystallises from warm dilute aqueous NaOH on cooling. [*Beilstein 3* H 524.]

Sodium trifluoroacetate [2923-18-4] C₂F₃O₂Na, M 136.0, m 205-207°(dec), 206-210°(dec), pK²⁵ 0.52 (for CF₃CO₂⁻). A possible contaminant is NaCl. The solid is treated with CF₃CO₂H and evaporated twice. Its solubility in CF₃CO₂H is 13.1% at 29.8°. The residue is crystallised from dilute EtOH, and the solid is dried in vacuum at 100°. [Hara & Cady *J Am Chem Soc* **76** 4285 1954, DOI: 10.1021/ja01646a009.] It can be precipitated from EtOH by adding dioxane, then recrystallising several times from hot absolute EtOH. Dry it at 120-130°/1mm. [*Beilstein 2* IV 461; *Fieser 11* 557.]

Stannous bis-cyclopentadienyl (stannocene, SnCp₂) [1294-75-3, 26078-96-6] C₁₀H₁₀Sn, M 248.9. SnCp₂ was prepared from sodium cyclopentadienide [4894-81-2, see Chapter 5, Catalysis Part 1] and anhydrous SnCl₂ in THF [general procedure: Wilkinson *Org Synth* **36** 31 1956, DOI: 10.15227/orgsyn.036.0031; Wilkinson et al. *J Inorg Nuclear Chem* **2** 95 1956, DOI: 10.1016/0022-1902(56)80004-3]. Purify it by vacuum sublimation. **It is sensitive to moisture in solvents.** Handle and store it under dry N₂. The related thallium and indium compounds are similarly purified. [For preparation and properties, IR, UV & NMR see Dave, Evans and Wilinon *J Chem Soc* 3684 1959, DOI: 10.1039/JR9590003684; for photoelectron spectra see Cradock & Duncan *JCS Faraday Trans 2* 194 1978, DOI: 10.1039/F29787400194].

Strontium acetate [543-94-2] (C₂H₃O₂)₂Sr, M 205.7, d_{25}^{25} 2.1g/ml, pK²⁵ 13.0 (for aquo Sr²⁺ hydrolysis). Crystallise it from AcOH, then dry it under vacuum for 24 hours at 100°. [*Beilstein 2* II 91.]

Strontium lactate trihydrate [29870-99-3] (C₃H₅O₃)₂Sr·3H₂O, M 319.8(3H₂O), m 120°(loses 3H₂O). It crystallises from aqueous EtOH. Its solubility in H₂O is 33g/100ml at ~25° and 200g/100ml at 100°, and is slightly soluble in EtOH. [*Beilstein 3* IV 633.]

Strontium oxalate monohydrate [814-95-9] C₂O₄Sr·H₂O, M 193.6(H₂O), m 150°. It crystallises from hot water on cooling. The solubility at ~25° in H₂O is 1g/20L, and in 3.5% AcOH and 25% AcOH it is 1g/1.9L and 1g/1.1L respectively. **IRRITANT.** [*Beilstein 2* H 515, *2* IV 1826.]

Strontium salicylate [526-26-1] (C₇H₆O₃)₂Sr, M 361.8, b 336.3°/atm. It crystallises from hot water (4ml/g) or EtOH. [*Beilstein 10* IV 125.]

Strontium tartrate [868-19-9] C₄H₆O₆Sr, M 237.7. It crystallises from hot water. Crystals of a **pentahydrate** can be grown from silica gel impregnated with L-tartaric acid and using Sr(NO₃)₂. The SEM, XRD and FTIR of the crystals have been measured. The **pentahydrate** is thermally stable up to 105° but decomposes on further heating liberating H₂O at various stages and finally reduced to strontium oxide [Firdous et al. *Cryst Res Technol* **43**(10) 1015 2008, DOI: 10.1002/crat.200800115]. [*Beilstein 3* IV 1219.]

Tantalum pentaethoxide [6074-84-6] ($\text{C}_2\text{H}_5\text{O}$)₅Ta, M 406.3, m 21°, b 147°/0.2mm, 155°/0.01mm, 202°/10mm, d²⁵ 1.566g/ml, n_D²⁰ 1.487, pK₁²⁵ 9.6, pK₂²⁵ 13.0 (for tantalic acid). Purify it by distillation under reduced pressure. It aggregates in *C₆H₆, EtOH, MeCN, pyridine and diisopropyl ether. Store it in a dry atmosphere. [Bradley et al. *J Chem Soc* 726 1955, DOI: 10.1039/JR9550000726; Bradley et al. *J Chem Soc* 5 1956, DOI: 10.1039/JR9560000005; *Beilstein* 1 IV 1312.]

Tetraallyltin (tetraallylstannane) [7393-43-3] (C₃H₅)₄Sn, M 283.0, b 52°/0.2mm, 69-70°/15mm, d²⁵ 1.179/ml, n_D²⁰ 1.539. Possible contaminants are allyl chloride and allyltin chloride. Check the ¹H NMR and IR [Fishwick & Wallbridge *J Organomet Chem* 25 69 1970, DOI: 10.1016/S0022-328X(00)86206-3], and if impure, dissolve it in Et₂O and shake it with a 5% aqueous solution of NaF which precipitates allyltin fluoride. Separate the Et₂O layer, dry (MgSO₄), and distil it at ~0.2mm. It decomposes slightly on repeated distillation. [O'Brien et al. *Inorg Synth* 13 73 1972, DOI: 10.1002/9780470132449.ch14; Fishwick & Wallbridge *J Chem Soc (A)* 57 1971, DOI: 10.1039/J19710000057; *Beilstein* 4 III 1922.]

Tetrabutylammonium borohydride [33725-74-5] (C₄H₉)₄.NBH₄, M 257.3, m 124-128°, 128-129°. Purify it by recrystallisation from EtOAc followed by careful drying under vacuum at 50-60°. Samples purified in this way showed no signs of loss of *active H* after storage at room temperature for more than 1 year. Nevertheless samples should be stored at ca 6° in tightly stoppered bottles if kept for long periods. It is soluble in CH₂Cl₂. [Raber & Guida *J Org Chem* 41 690 1976, DOI: 10.1021/jo00866a022; Brändström et al. *Tetrahedron Lett* 3173 1972, DOI: 10.1016/S0040-4039(01)93995-4.]

Tetrabutyl orthotitanate monomer (titanium IV tetra-*n*-butoxide) [5593-70-4] (C₄H₉O)₄Ti, M 340.4, b 142°/0.1mm, 134-136°/0.5mm, 160°/0.8mm, 174°/6mm, 189°/13mm, d²⁰ 1.00/ml, d₄³⁵ 0.993, n_D²⁰ 1.491. Dissolve it in *C₆H₆, filter if solid is present, evaporate and vacuum fractionate through a Widmer 24inch column. The ester hydrolyses when exposed to air to give hydrated ortho-titanic acid. The titanium content can be determined thus: weigh a sample (ca 0.25g) into a weighed crucible and cover it with 10ml of H₂O and a few drops of concentrated HNO₃. Heat (hot plate) carefully till most of the H₂O has evaporated. Cool and add more H₂O (10ml) and concentrated HNO₃ (2ml), and evaporate carefully (no spillage) to dryness and ignite the residue at 600-650°/1 hour. Weigh the residual TiO₂. [Bradley et al. *J Chem Soc* 2773 1952, DOI: 10.1039/JR9520002773; Speer *J Org Chem* 14 655 1949, DOI: 10.1021/jo01156a019; *Beilstein* 1 II 398, 1 III 1515, 1 IV 1415, Fieser 15 316.] It was used to prepare nanosised titania in the anatase form *via* controlled hydrolysis of titanium alkoxide [Chen et al. *J Mater Sci* 31 3497 1996, DOI: 10.1007/BF00360754], and for preparing nanocrystalline TiO₂ powders at room temperature [Wang et al. *Mater Lett* 43 87 2000, DOI: 10.1016/S0167-577X(99)00236-0].

Tetrabutyl tin (tin tetrabutyl) [1461-25-2] (C₄H₉)₄Sn, M 347.2, m -97°, b 94.5-96°/0.28mm, 145°/11mm, 245-247°/atm, d²⁵ 1.0559/ml, n_D²⁰ 1.473. Dissolve it in Et₂O, dry it over MgSO₄, filter, evaporate and distil it under reduced pressure. Although it does not crystallise easily, once the melt has crystallised, then it will recrystallise more easily. It is soluble in Et₂O, Me₂CO, EtOAc and EtOH, but insoluble in MeOH and H₂O, and shows no apparent reaction with H₂O. [Johnson & Fritz *J Org Chem* 19 74 1954, DOI: 10.1021/jo01366a014; Staveley et al. *J Chem Soc* 1992 1954, DOI: 10.1039/JR9540001992; Van der Kerk & Luijten *Org Synth Coll Vol* 4 881 1963, DOI: 10.15227/orgsyn.036.0086; *Beilstein* 4 III 1920, 4 IV 4312.]

Tetraethyl lead [78-00-2] (C₂H₅)₄Pb, M 323.5, m -136°, b 84-85°/15mm, 200°/atm 227.7°/atm, dec), d²⁵ 1.653/ml, n_D²⁰ 1.519. Its more volatile contaminants can be removed by exposure to a low pressure (by continuous pumping) for 1 hour at 0°. Purify it by stirring with an equal volume of H₂SO₄ (d 1.40), keeping the temperature below 30°, repeating this process until the acid layer is colourless. It is then washed with dilute Na₂CO₃ and distilled water, dried with CaCl₂ and fractionally distilled at low pressure under H₂ or N₂ [Calingaert *Chem Rev* 2 43 1926, DOI: 10.1021/cr60005a002]. It prevents 'knocking' in petrol combustion engines. [Milde & Beatty *Adv Chem Res* 23 306-318 1959, DOI: 10.1021/ba-1959-0023.ch029 *Beilstein* 4 H 639.] A 50 wt% solution in xylene (b 147°/atm, d²⁵ 1.108/ml, n_D²⁰ 1.502) is available commercially. **VERY POISONOUS.**

Tetraisopropyl orthotitanate (titanium IV tetraisopropoxide) [546-68-9] (*i*-C₃H₇O)₄Ti, M 284.3, m 18.5°,

18-20°, b 80°/2mm, 78°/12mm, 228-229°/755mm, d²⁰ 0.96/ml, n_D²⁰ 1.464. Dissolve it in dry *C₆H₆, filter if a solid separates, evaporate and fractionate. It is hydrolysed by H₂O to give solid Ti₂O(*iso*-OPr)₂ **m ca 48°**. [Bradley et al. *J Chem Soc* 2027, 1952, DOI: 10.1039/JR9520002027; Bradley et al. *J Chem Soc* 469 1957, DOI: 10.1039/JR9570000469; *Beilstein* 1 II 328, 1 IV 1469S; for applications see Fieser 6 11, 10 404, 11 374, 12 19, 12 504, 13 13, 13 311, 14 247, 14 311, 15 322, 16 399, 17 347.] This propoxide was used for *trans-esterifications* with various alcohols under neutral conditions [Imwinkelried et al. *Org Synth* 65 230 1987, DOI: 10.15227/orgsyn.065.0230], in the *Sharpless asymmetric epoxidation* of allylic alcohols [Katsuki & Martin *Org React* 48 1 1996, DOI: 10.1002/0471264180.or048.01], was applied in the formation of *a heterosupramolecule* consisting of a TiO₂ nanocrystalline-viologen electron acceptor complex whose light-induced electron transfer was demonstrated [Cusak et al. *Chem Mater* 9 1765 1997, DOI: 10.1021/cm9605173], and was used for *making porous titanosilicates*, potential ion-exchange materials, for the selective removal and safe storage of radioactive ¹³⁷Cs and ⁹⁰Sr nuclear wastes [Behrens et al. *Chem Mater* 8 1236 1996, DOI: 10.1021/cm950534c].

Tetrakis(diethylamino) titanium [(titanium IV tetrakis(diethylamide)] [4419-47-0] [(C₂H₅)₂N]₄Ti, M 336.4, b 85-90°/0.1mm, 112°/0.1mm, d²⁵ 0.931/ml, n_D²⁰ 1.536. Dissolve it in *C₆H₆, filter if a solid separates, evaporate under reduced pressure and distil it. It is an orange liquid which reacts violently with alcohols. [Bradley et al. *J Chem Soc* 3857 1960, DOI: 10.1039/JR9600003857; *Beilstein* 4 IV 313.]

Tetramethyloxorhenium [(CH₃)₄ReO] [53022-70-1] (CH₃)₄ORE, M 262.4, m ~45° (no dec <150°). It is best prepared by adding a 1M solution of MeLi in Et₂O (80ml, 80mmol) slowly to a stirred suspension of ReOCl₃(PPh₃)₂ (8.32g, 10mmol, cf. [17442-18-1]) in Et₂O (120ml) at -78°, and the mixture is allowed to warm slowly to ~25° to form a dark brown solution which is stirred further for 0.5 hours. The excess of MeLi is destroyed by dropwise addition, at -78°, of H₂O (~10ml), and allowed to warm to ~25°, and then cooled again, to -30°; and H₂O₂ (2.5g, 30% diluted in 20ml of H₂O) is added slowly dropwise with vigorous stirring. The colour of the solution changes to red rapidly and stirring is continued for 0.5 hours after the temperature has risen to ~25°. This mixture is then cooled to -78°, the Et₂O layer is filtered free from ice, dried first with CaCl₂ at ~25° then with molecular sieves, cooled again to -78°, filtered, and the solvent is carefully evaporated at -50°. The red-purple (carmine) crystalline residue is sublimed onto a probe at -78°, in a 1mm vacuum, to give Me₄ReO (1.26g, ~48%; but can be as high as 70% by very careful removal of solvent). Lower yields are obtained if due care is not taken during the removal of solvent *in vacuo*. The low melting solid is extremely volatile subliming readily even at -35°/10⁻³mm; it co-distils at low temperatures with Et₂O or petroleum ether, and always occurs during removal of solvents (see earlier), hence the temperature has to be kept as low as possible, and a cold probe should be included in the system to avoid losses. In an *alternative method* ReOCl₄ (3mmol) is reacted with MeLi (12mmol) in Et₂O at -78° to give a 20% yield of Me₄ReO, although this low yield may well be due to evaporative losses.

Me₄ReO is thermally stable and can be stored indefinitely *in vacuo*. It is very soluble in petroleum ethers, Et₂O, CS₂, CCl₄, but these solutions slowly decompose with time even in the absence of air. However, solutions in THF and aromatic hydrocarbons are more stable. It is surprisingly unreactive at ~25° in Et₂O, H₂O, mineral acids, alcohols, H₂, CO, CO₂, SO₂, HCl gas, butadiene, tertiary phosphines, amines, tetramethylthiuram disulfide and 1,1-diphenyl-2-picrylhydrazyl. It is however, highly reactive with air and oxygen producing instantly white fumes and crystals of MeReO₃ (methyltrioxorhenium, **MTO**, the carrier catalyst, see [70197-13-6]). In *C₆H₆ solution with air, it slowly oxidises to form yellow-orange products (cf. yellow cis-Me₃ReO₃ [56090-01-8]). In the mass spectrum, ions of Re isotopes are identified (¹⁸⁵Re 37%, ¹⁸⁷Re 63%) with the natural isotopic abundance. The solid state IR (at ~ -250° co-condensed with argon) has ν_{max} at 2974s(CH str), 1231s(CH def), 1018s(Re=O str), 1004sh,br, 746m(Me rock), 552m and 529s(ReC str) cm⁻¹, and IR (de-oxygenated CS₂) has ν_{max} at 2980s, 1370m, 1002s, 749w, 520m cm⁻¹. [Mertis et al. *JCS Chem Commun* 93a 1974, DOI: 10.1039/C3974000093A; Mertis, Williamson and Wilkinson *JCS Dalton Trans* 607 1975, DOI: 10.1039/DT9750000607; Mertis & Wilkinson *JCS Dalton Trans* 1488 1976, DOI: 10.1039/DT9760001488; Beattie & Jones *Inorg Chem* 18 2318 1979, DOI: 10.1021/ic50198a056.]

Tetramethyltin (tetramethylstannane, tin tetramethyl) [594-27-4] (CH₃)₄Sn, M 178.8, m -55°, b 76.6°/748mm, 74-75°/atm, 78°/atm, d²⁵ 1.291g/ml, n_D²⁰ 1.441. Organotin compounds are **HIGHLY TOXIC**, and all due precautions should be taken when handling them. Me₄Sn is an ethereal smelling liquid that is highly

flammable with a flash point of less than 20°, and should be stored at low temperatures. An improved preparation of Me₄Sn has been achieved by carrying out the reaction in the higher boiling di-*n*-butyl ether, and preparing the Grignard reagent *in situ*. Under strictly anhydrous conditions, to Mg turnings (50g, 2.06g.atoms) in dry *n*-Bu₂O (600ml, containing a few crystals of I₂) is added dropwise a solution of MeI (225g, 1.59moles, freshly distilled) in *n*-Bu₂O (600ml) with gentle stirring. After the initial reaction has started, the MeI solution is added at such a rate as to keep the solution refluxing gently, requiring *ca* 3 hours. Cool the solution to ~25° and add to it, dropwise, anhydrous SnCl₄ (75g, 0.29 moles) while sustaining gentle reflux (*ca* 2-2.5 hours). The mixture is then refluxed steadily (85-95°) for 1 hour, and set aside at ~25° for several hours. The reflux condenser is replaced by a Claisen head and the mixture is distilled; whereby Me₄Sn and *n*-Bu₂O distil together at ~85-95°/atm. It dissolves stopcock silicone greases. The Me₄Sn is purified by fractionation through a Todd column (35-40 plates) to give *pure stannane* with **b 76.6°/748mm** in 91% yield. [Edgell & Ward *J Am Chem Soc* **76** 1169 1954, DOI: 10.1021/ja01633a073.] Its IR (liquid film) has ν_{\max} at 2920 (vs) and 3000 (vs) C-H str, 1443 (s) and 1198 (s) CH₃ deformation, 770 (vs) CH₃ rocking, **528 (s) C-Sn str cm⁻¹**, and from Raman spectrum 160 (vs) and 145 (vs) cm⁻¹ for **C-Sn-C** bending [Taimsalu & Wood *Trans Farad Soc* **59** 1754 1963, DOI: 10.1039/TF9635901754; Edgell & Ward *J Am Chem Soc* **77** 6486 1955, DOI: 10.1021/ja01629a014]. The ¹¹⁹Sn NMR (15.5 MHz, at 28°C, 12mm tube not spinning, neat SnCl₄ as external standard) has δ at -147.8 ($J_{\text{H}^{119}\text{Sn}} = 54.0\text{Hz}$ from 60MHz proton spectrum) [Lassigne & Wells *Can J Chem* **55** 927 1977, DOI: 10.1139/v77-130; for ¹H and ¹³C NMR spectra see Singh *J Organomet Chem* **99** 251 1975, DOI: 10.1016/S0022-328X(00)88454-5.] [Beilstein **4** H 631, **4** I 583, **4** II 1010, **4** III 1917, **4** IV 4307.]

Tetraphenyltin [595-90-4] (C₆H₅)₄Sn, **M 427.1, m 224-225°, 224-227°, 226°, b >420°**. SnPh₄ forms yellow crystals from CHCl₃, petroleum ether (b 77-120°), xylene or *benzene/cyclohexane, and is dried at 75°/20mm. [Gilman & Rosenberg *J Am Chem Soc* **74** 531 1952, DOI: 10.1021/ja01122a074; Beilstein **16** IV 1592.]

Tetra-*n*-propylammonium perruthenate (TPAP, tetrapropyl tetraoxoruthenate) [114615-82-6] (C₃H₇)₄NRuO₄, **M 351.4, m 160°(dec)**. It is a dark green solid that is stable at ~25° for long periods without much deterioration, and is best kept in a refrigerator. It is a *mild oxidant*, and should *not* be heated directly as it may explode. It can be used stoichiometrically or catalytically with a *co-oxidant* [Ley et al. *Aldrichimica Acta* **23** 13 1990, *Synthesis* 639 1994, DOI: 10.1055/s-1994-25538]. Small amounts decompose in a flame at ~155° in air. It can be washed with aqueous *n*-propanol, then H₂O and dried over KOH in a vacuum. It is soluble in CH₂Cl₂ and MeCN. Generally, at ambient temperatures in these solvents it is most useful, when in conjunction with *N-methylmorpholine N-oxide* (NMO), for the oxidation of primary and secondary alcohols, including allylic and benzylic alcohols, lactol oxidation, heteroatom oxidation, and some cleavage reactions. [Dengel et al. *Transition Met Chem* **10** 98 1985, DOI: 10.1007/BF00618458; Griffith et al. *JCS Chem Commun* 1625 1987, DOI: 10.1039/C39870001625; for a review see Ley et al. *Synthesis* 639 1994, DOI: 10.1055/s-1994-25538; and for its catalytic activity see chapter 5 in 'Catalysts'.] The **TPAP/NMO** system has been used to oxidise *N, N'*-dihydroxyimidazolines to the respective nitronyl-nitroxides free radicals in CH₂Cl₂ (~25°, 1-12hrs) in 44-90% yields [Gorini et al. *Synlett* 948 2006, DOI: 10.1055/s-002-4768]. In conjunction with NMO, Swern oxidation or Dess-Martin periodinane conditions, it promotes the oxidation of hydroxyl-substituted tri-*n*-butylammonium trifluoroarylborates without cleavage of the C—B bond [Molander & Petrillo *J Am Chem Soc* **128** 9634 2006, DOI: 10.1021/ja062974i]. [For applications see Fieser **14** 302, **16** 325.]

§ Polymer supported reagent is available commercially.

Thallium (I) acetate (TlOAc) [563-68-8] CH₃CO₂Tl, **M 263.4, m 131°**. Thallous (I) acetate crystallises from EtOH (needles) or Me₂CO. The salt forms white deliquescent crystals (specific gravity 3.68) which are soluble in H₂O and EtOH. Store it in sealed containers. It forms TlOAc·AcOH crystals which melt at 64°. [Beilstein **2** H 115, **2** II 119, **2** III 195.] **POISONOUS — HANDLE CAREFULLY WITH GLOVES, and work in a well-ventilated fume hood.**

Thallium (III) triacetate [Tl(OAc)₃] [2570-63-0] (C₂H₃O₂)₃Tl, **M 381.5, m 182°(dec)**. The salt is prepared by stirring thallic oxide (50g) in AcOH (300ml) at 65° until all the brown-black solid has dissolved (~24 hours), then filter and cool to obtain the salt as white crystals which are collected by decantation and dried over P₂O₅ *in vacuo* as they turn brown in air. A second crop can be obtained by concentrating the filtrate (total yield 73g, 99%). On a large scale (457g) of the brown trioxide was treated with AcOH (2L) containing Ac₂O (110ml) by

stirring at 80-90°, the walls of the flask were washed down with a further volume of AcOH (500ml). Heating (90-100°) and stirring was continued for 2 hours (all the brown oxide had dissolved), the hot solution was filtered using a sintered glass funnel, cooled to ~20°, and collected onto a glass frit. The solid was dissolved in the minimum volume of AcOH (~800ml) at 80-90°, filtered through a glass frit and allowed to cool first to ~20° then to ~15°, and collected by filtration. The $\text{Ti}(\text{OAc})_3$ was dried by spreading it in a large dish in a desiccator over KOH pellets overnight to give ~97% pure salt (467g, 61% yield). It promotes the quantitative photochemical decarboxylation of a variety of aliphatic acids to yield a high proportion of the corresponding alkanes with some alkenes, dialkyls and other minor products at wavelength 3500Å; whereas at wavelength 2537Å the alkyl dimers are formed in high yields. [Kochi & Bethea *J Org Chem* **33** 75 1968, DOI: 10.1021/jo01265a015.] [For use in the oxidation of olefins see Moriarty & Gopal *Tetrahedron Lett* 347 1972, DOI: 10.1016/S0040-4039(01)84321-5]. [*Beilstein* **2** H 115, **2** III 195.] **POISONOUS — HANDLE CAREFULLY WITH GLOVES, and work in a well-ventilated fume hood.**

Thallium (III) trifluoroacetate [$\text{Ti}(\text{tfl})_3$, $\text{Ti}(\text{OCOCF}_3)_3$] [23586-53-0] ($\text{C}_2\text{F}_3\text{O}_2$)₃Tl, **M 543.4, m dec slowly >100°, 213° (dec.)**. This *thallation reagent* is prepared in 90-100% yield by heating a suspension of thallium(III) oxide in TFA containing 10-20% of H₂O until clear. This colourless solution can be used directly in thallation reactions or a granular solid salt can be isolated by evaporation *in vacuo*, and then used in inert solvents such as MeCN. The salt does not have a sharp melting point and decomposes slowly >100°. It is *water sensitive* but can be stored at ~25° in a stoppered container away from light without appreciable decomposition for long periods. A solution of the salt in TFA, however, can tolerate up to ~20% of H₂O to give a 0.8M solution before serious hydrolysis of the salt occurs. It thallated aromatic rings (e.g. PhCl, PhF, toluene, xylene, *o*-benzoic acid) in TFA to produce the corresponding $\text{ArylTi}(\text{tfl})_2$ at low temperatures which can be isolated in 70-100% yields, or reacted further (e.g. with KI) to displace the $\text{Ti}(\text{tfl})_2$ group and form the respective iodides. [McKillop et al. *J Am Chem Soc* **93** 4841 1971, DOI: 10.1021/ja00748a029; Taylor et al. *Org Synth Coll Vol* **6** 709 1988, DOI: 10.15227/orgsyn.055.0070; *Beilstein* **2** II 186; Fieser **3** 286, **4** 498, **5** 658, **6** 579, **7** 365, **8** 478, **13** 295.] Useful oxidant for the synthesis of S-substituted cysteine peptides, and proteins containing disulfide bonds [Yajima et al. *Tetrahedron* **44** 805 1988, DOI: 10.1016/S0040-4020(01)86118-4]. **POISONOUS — HANDLE CAREFULLY WITH GLOVES, and work in a well-ventilated fume hood.**

Thallous (I) ethoxide (TIOEt) [20398-06-5] $\text{C}_2\text{H}_5\text{OTl}$, **M 249.4, d²⁵ 3.522g/ml, n_D²⁰ 1.676**. The alkoxide is prepared by refluxing dry EtOH in a modified Soxhlet extractor whereby the hot alcohol leaches thallium shot or turnings (see below) placed in the vapours of refluxing alcohol while oxygen flows through the apparatus which is protected from moisture with a soda lime drying tube at the top of the condenser. The apparatus devised by Fieser and Fieser can be used. Thallium shot (m 303°) is made by holding a clean piece of metal with tongs and heated with a blow torch allowing the molten metal to drip in ~2L of cold H₂O. *Alternatively*, thallium turnings made with a pencil sharpener from thallium rods can be used. A Soxhlet flask (500ml) containing dry EtOH (300ml) is placed under Tl shot (100ml, dried by pressing with filter paper) in the crucible above, and the EtOH is refluxed while a rapid stream of dry O₂ is made to flow just beneath the metal. The volume of boiling EtOH in the flask is maintained at ~300ml by further addition of dry EtOH. When all the metal has been converted to TIOEt (12-16 hours), and all the oily alkoxide has run into the flask which will form a separate heavy oily liquid layer (note that TIOEt had d = 3.522), boiling is stopped and the volume of the alkoxide solution is adjusted to 300ml which would give a saturated solution containing 9g of TIOEt /100ml. Store away from moisture. *Alternatively*, the oily layer can be sucked out with a vacuum into a container, but great care should be taken to avoid contact with moisture. Thallium in the ethoxide solution can be determined as TIOH by cooling it in an ice bath without separating the excess of EtOH (to avoid partial conversion of TIOH to black Tl_2OH), and adding an equal volume of boiled and re-cooled H₂O (i.e. free from O₂) and evaporating *in vacuo* to give yellow crystalline TIOH . [See Fieser **2** 407, **4** 501, **5** 656, **6** 577, **7** 362, **10** 395; Dönges in *Handbook of Preparative Inorganic Chemistry* (Ed. Brauer) Academic Press Vol **I** p 877-878 1963, Freudenberg & Uthemann *Chem Ber* **52** 1508 1919, DOI: 10.1002/cber.19190520813.] **POISONOUS — use gloves, work in a good fume hood.**

Tin (II) acetate (stannous acetate) [638-39-1] (CH_3CO_2)₂Sn, **M 236.8, m 180-182°, 182.55-183°, b 239-241°/atm(some dec under N₂), d²⁵ 2.31g/ml**. It is prepared by dissolving blue-black SnO₂ (25g) in refluxing 50%v/v AcOH (200ml), evaporating *in vacuo* (over KOH) and subliming the white residue *in vacuo* at 150-155° (~96% yield of white orthorhombic crystals). *Alternatively*, finely divided Sn is refluxed in glacial AcOH for

80-90 hours [not less as $\text{Sn}(\text{OAc})_2 \cdot 2\text{AcOH}$ is formed] and isolated as before. It hydrolyses slowly in H_2O to give blue Black SnO_2 , Me_2CO , CO_2 and H_2 ; and can be stored for long periods under N_2 , and is soluble in EtOH (2.9%), Me_2CO (1.4%) and 2N AcOH (33%). It is a useful *reducing (and acetylating) agent*. [Donaldson et al. *J Chem Soc* 5942 1964, DOI: 10.1039/JR9640005942.]

Titanium (IV) methoxide [992-92-7] $(\text{CH}_3\text{O})_4\text{Ti}$, **M 172.0, m 200-210°, b 243°/52mm**. It is extremely sensitive to moisture. Dissolve it in H_2O -free $^*\text{C}_6\text{H}_6$, filter, evaporate and distil it *in vacuo* under N_2 . It is **FLAMMABLE** and **TOXIC**. [for $\text{Ti}(\text{OEt})_4$ see: Bradley et al. *J Chem Soc* 721 1955, DOI: 10.1039/JR9550000721; Bradley et al. *Metal Alkoxides* Academic Press 1978, ISBN 0121242501.] [*Beilstein* 1 II 274.] This ethoxide was used for preparing polyoxotitanates [Clegg et al. *JCS Dalton Trans* 681 1996, DOI: 10.1039/DT9960000681].

Titanocene dichloride [1271-19-8] $\text{C}_{10}\text{H}_{10}\text{Cl}_2\text{Ti}$, **M 248.9, m 260-280°(dec), 289.2°, 298-291°, d²⁵ 1.6g/ml**. It forms bright red crystals from toluene or xylene/ CHCl_3 (1:1) and sublimes at 190°/2mm. It is moderately soluble in EtOH and insoluble in Et_2O , $^*\text{C}_6\text{H}_6$, CS_2 , CCl_4 , petroleum ether and H_2O . The crystalline *dipicrate* explodes on melting at 139-140°. [Wilkinson et al. *J Am Chem Soc* 75 1011 1953, DOI: 10.1021/ja01100a527; IR: Wilkinson & Birmingham *J Am Chem Soc* 76 4281 1954, DOI: 10.1021/ja01646a008; NMR and X-ray: Glivicky & McCowan *Can J Chem* 51 2609 1973, DOI: 10.1139/v73-394; *Beilstein* 16 IV 1769; for applications see Fieser 6 48, 10 130, 12 168, 13 102, 14 120, 16 116.] This titanocene catalyses a one-pot synthesis of α -methylene- γ -butyrolactones from benzaldehydes and bromomethylacrylates [Paira et al. *Tetrahedron Lett* 48 3205 2007, DOI: 10.1016/j.tetlet.2007.03.036].

Tri-*n*-butyl tin chloride (TBTC) [1461-22-9] $(\text{C}_4\text{H}_9)_3\text{SnCl}$, **M 325.5, b 98-100°/0.4mm, 140-152°/10mm, 172°/25mm, d²⁵ 1.21g/ml, n_D²⁰ 1.492**. Fractionate it in an inert atmosphere and seal it in small aliquots in glass ampoules. It is sensitive to moisture. [Jones et al. *J Chem Soc* 1446 1947, DOI: 10.1039/JR9470001446; *Beilstein* 4 III 1926, 4 IV 4330; for applications see Fieser 6 604, 7 378, 13 315.]

Tributyl tin hydride [688-73-3] $(\text{C}_4\text{H}_9)_3\text{SnH}$, **M 291.1, b 76°/0.7mm, 81°/0.9mm, d²⁵ 1.082g/ml, n_D²⁰ 1.473**. Dissolve it in Et_2O , add quinol (500mg for 300ml, to stabilise it), dry over Na_2SO_4 , filter, evaporate and distil it under dry N_2 . It is a clear liquid if dry and decomposes very slowly. In the presence of H_2O , traces of tributyl tin hydroxide are formed in a few days. Store it in sealed glass ampoules in small aliquots. It is estimated by reaction with aqueous NaOH when H_2 is liberated. **CARE:** stored samples may be under pressure due to liberated H_2 . [Ono et al. *Tetrahedron* 41 4013 1985, DOI: 10.1016/S0040-4020(01)97180-7; Neuman *Synthesis* 665 1987, DOI: 10.1055/s-1987-28044; Curran *Synthesis* 417 1988, DOI: 10.1055/s-1988-27600; *Beilstein* 4 IV 4312; for applications see Fieser 1 1192, 2 424, 3 294, 4 518, 5 685, 6 604, 7 379, 8 497, 9 476, 10 411, 11 545, 12 516, 13 316, 14 312, 15 325, 16 343, 17 351.] This hydride is a *radical reagent* for reductive cleavage [Neumann *Synthesis* 665 1987, DOI: 10.1055/s-1987-28044], radical dehalogenation and intramolecular radical cyclisation [Curran *Synthesis* 417 1988, DOI: 10.1055/s-1988-27600], radical prompted intramolecular cyclisation [Singh & Batra *Tetrahedron Lett* 47 7043 2006, DOI: 10.1016/j.tetlet.2006.07.106], and is a tin fragment used in a synthesis of (+)-*panepophenanthrin* [Comméiras et al. *Tetrahedron* 62 9892 2006, DOI: 10.1016/j.tet.2006.08.010].

Triethyltin hydroxide [994-32-1] $(\text{C}_2\text{H}_5)_3\text{SnOH}$, **M 222.9, m 49-50°, b 153-155°/20mm**. Treat it with HCl , followed by KOH , and filter it to remove diethyltin oxide [Prince *J Chem Soc* 1783 1959, DOI: 10.1039/JR9590001783]. [*Beilstein* 4 H 633, 4 I 585, 4 II 1012, 4 III 1924, 4 IV 4325.]

Triiron dodecacarbonyl [17685-52-8] $\text{Fe}_3(\text{CO})_{12}$, **M 503.7, m 165°(dec)**. The dark black-green solid usually contains 10% by weight of MeOH as stabiliser. This can be removed by keeping it in a vacuum at 0.5mm for at least 5 hours. It can be sublimed slowly at high vacuum and is soluble in organic solvents. [Landesberg et al. *J Org Chem* 37 930 1972, DOI: 10.1021/jo00972a002; Case & Whiting *J Chem Soc* 4632 1960, DOI: 10.1039/JR9600004632; King et al. *Inorg Synth* 7 193 1963, DOI: 10.1002/9780470132388.ch52; McFarlane et al. *Inorg Synth* 8 181 1966, DOI: 10.1002/9780470132395.ch47] **TOXIC as it is a source of carbon monoxide.**

Trimethyloxonium tetrafluoroborate [420-37-1] $(\text{CH}_3)_3\text{O}^+\text{BF}_4^-$, **M 147.9, m 141-143°(sinters, open capil-**

lary), 179.6-180.0°(dec), 210-220°(dec). The salt must be a white crystalline solid **m** ~ **179.6-180.0°** (dec, sealed tube). Under a N₂ atmosphere (e.g. Dry Box), wash it twice with CH₂Cl₂, then twice with Na-dried Et₂O, and dry by passing dry N₂ over the salt until free from Et₂O [Curphey *Org Synth Coll Vol* **6** 1019 1988, DOI: 10.15227/orgsyn.051.0142]. The oxonium salt, purified in this way, can be handled in air for short periods. The sample kept in a desiccator (Drierite) for 1 month at -20° had an unaltered melting point, and samples stored in this way for >1 year are satisfactory for alkylations. ¹H NMR in liquid SO₂ in a sealed tube had a single peak at δ 4.54 (impurities have δ at 3.39). [Meerwein *Org Synth Coll Vol* **5** 1096 1973, DOI: 10.15227/orgsyn.046.0120.] If the sample looks good, dry it in a vacuum desiccator for 2 hours (25°/1mm) and store it under N₂ at -20°. The melting point depends on heating rate. [Beilstein **1** IV 1248; for applications see Fieser **1** 1232, **2** 438, **3** 314, **4** 541, **13** 327.]

Cis-Trimethylrhenium dioxide [*cis*-trimethyldioxorhenium(VII), *cis*-(CH₃)₃ReO₂] [56090-01-8] (CH₃)₃ReO₂, **M 263.3, m 10-11°.** This yellow dioxide is best prepared, in ~60% yield, from Me₄ReO (preceding oxide [53022-7-1], e.g. 0.5g) in Et₂O (e.g. 40ml) at -78° by exposing the solution to nitric oxide when the red colour disappears while allowing to warm to ~25° with stirring. After 1 hour (without isolating the nitroso intermediate), the Et₂O is removed at -40° *in vacuo* and the yellow crystalline residue is **sublimed** at 10⁻³mm onto a probe at -78° to give *cis*-(CH₃)₃ReO₂, which can be recrystallised from petroleum ether solution on cooling to -78°, or distilled *in vacuo*. It is reasonably stable at room temperature, and like the starting material it fumes in air forming needles of **MTO**, the carrier catalyst (see above). Store it under N₂ or argon in a sealed ampoule. Its IR spectrum is consistent with the trigonal bipyramidal structure, where bands due to methyl vibrations are present together with two strong bands at ν_{max} 992 and 951 cm⁻¹ attributable to *cis*-ReO₂. This structure is also consistent with the ¹H NMR in CS₂ or deuteriotoluene which has two sharp resonances at δ 2.07 and 2.50 in the ratio 1:2 in which the higher field band is assumed to be from the axial methyl group. The lines do not broaden on cooling to -78° or heating to 85° in toluene solution which is consistent with a rigid molecule. The MS is as predicted from the two isotopes of Re (see above). [Mertis & Wilkinson *JCS Dalton Trans* 1488 1976, DOI: 10.1039/DT9760001488; Beattie & Jones *Inorg Chem* **18** 2318 1979, DOI: 10.1021/ic50198a056.]

Trimethyltin chloride (chlorotrimethylstannane) [1066-45-1] (CH₃)₃SnCl, **M 199.3, m 37.5-39.5°, 42°, b 45-47°/10mm, 152°/760mm, 154-156°/atm.** Me₃SnCl forms colourless needles that have **HIGHLY TOXIC** vapours and is best purified by distillation at atmospheric pressure or in a vacuum. It has been prepared by dropwise addition of a solution of Et₂O saturated with dry HCl (20ml) to freshly distilled (*dimethylamino*)-*trimethylstannane* (2.21g, **b 126°/atm**, d²⁵ 1.274, n_D²⁰ 1.463 [993-50-0]) in dry Et₂O (10ml). Me₂NH.HCl precipitated out, was filtered, the solvent was evaporated at <20°/15mm, and the residue was distilled at 760mm providing the colourless crystalline solid **Me₃SnCl** (1.80g, 89.7%). It sublimes slowly at ~25°/20mm. [Jones & Lappert *J Organomet Chem* **3** 295 1965, DOI: 10.1016/S0022-328X(00)84649-5; for ¹H and ¹³C NMR see Singh *J Organomet Chem* **99** 251 1975, DOI: 10.1016/S0022-328X(00)88454-5.] It is also available commercially as 1.0M solutions in hexanes or THF. Like the reaction of Me₂N-SnMe₃, the **Me₃SnCl** reacts with H₂O on heating to give **Me₃SnOH** (almost quantitatively, [56-24-6]) as white crystals **m 118°**, which sublime at their melting points. It has IR (Nujol mull) with ν_{max} at 3620 (m), 2985 (m), 2915 (m), 1405 (w), 1195 (m), 920 (s), 765 (s), **540 (s) Sn-C str**, 370 (s) and 316 (w) cm⁻¹; it exists as a **dimer [Me₃SnOH]₂** in the solid state (IR mull) and in *C₆H₆, CCl₄ and in CHCl₃ solutions (ebullioscopy) [Okawara & Yasuda *J Organomet Chem* **1** 356 1964, DOI: 10.1016/S0022-328X(00)80063-7].

Triphenylantimony III (triphenylstibine) [603-36-1] (C₆H₅)₃Sb, **M 353.1, m 52-54°, b 377°/atm, d²⁵ 1.53g/cm³.** Recrystallise Ph₃Sb from acetonitrile [Hayes et al. *J Am Chem Soc* **107** 1346 1985, DOI: 10.1021/ja00291a039; Hiers *Org Synth Coll Vol* **1** 550 1941, DOI: 10.15227/orgsyn.007.0080]. [Beilstein **16** H 891, **16** IV 1198.]

Triphenylbismuth (bismuth triphenyl) [603-33-8] (C₆H₅)₃Bi, **M 440.3, m 75-76°, 77-78°, 78.5°, b 124°/7mm, d²⁵ 1.6427(melt).** Dissolve Ph₃Bi in EtOH, precipitate it with H₂O, extract with Et₂O, dry and evaporate till the residue crystallises. It has been recrystallised from EtOH and Et₂O/EtOH and is a stable compound. [Forward et al. *J Chem Soc suppl* p121 1949, DOI: 10.1039/JR949000S121; Pfeiffer et al. *Chem Ber* **37** 4620 1904, DOI: 10.1002/cber.19040370468; Gilman & Yablunsky *J Am Chem Soc* **62** 665 1940, DOI: 10.1021/ja01860a068; UV: Jaffé *J Chem Phys* **22** 1430 1954, DOI: org/10.1063/1.1740411; Beilstein **4** III

1841.] Controlled hydrolysis of Ph_3Bi in supercritical CO_2 using a high-pressure cold wall reactor provided conformational bismuth oxide films [O'Neil & Watkins *Chem Mater* **19** 5460 2007, DOI: 10.1021/cm070288s].

Triphenyltin chloride (Fentin, chlorotriphenylstannane, triphenylchlorostannane) [639-58-7] (C_6H_5)₃ SnCl , M 385.5, m 103-106°(dec), 108°(dec), b 240°/13.5mm. Purify it by distillation, followed by recrystallisation from MeOH by adding petroleum ether (b 30-60°), m 105-106° [Kozeschow et al. *Chem Ber* **67** 1348 1934, DOI: 10.1002/cber.19340670810], or by crystallisation from Et_2O , or 5 parts of EtOH and a small volume of petroleum ether. [Krause *Chem Ber* **51** 912 1918, DOI: 10.1002/cber.191805101112.] It **sublimes** in a vacuum. [Beilstein **16** H 914, **16** I 540, **16** II 625, **16** III 1240, **16** IV 1606; Fieser **10** 451.] **VERY TOXIC.**

Triphenyltin hydroxide [76-87-9] (C_6H_5)₃ SnOH , M 367.0, m 122-123.5°, 124-126°. West, Baney and Powell [*J Am Chem Soc* **82** 6269 1960, DOI: 10.1021/ja01509a018] purified a sample which was grossly contaminated with tetraphenyltin and diphenyltin oxide by dissolving it in EtOH, most of the impurities remaining behind as an insoluble residue. Evaporation of the EtOH extract gave the crude hydroxide which was converted to **triphenyltin chloride** (above) by grinding in a mortar under 12M HCl, then evaporating the acidic solution. The residual **chloride**, after crystallisation from EtOH, had m 104-105°. It was dissolved in Et_2O and converted to the hydroxide by stirring with excess aqueous ammonia. The ether layer was separated, dried, and evaporated to give triphenyltin hydroxide which, after crystallisation from EtOH (or MeCN) and drying under vacuum, was in the form of white crystals (m 119-120°), which retained some cloudiness in the melt above 120°. The hydroxide retains water (0.1-0.5 moles of water per mole) tenaciously. [Glidewell & Liles *Acta Cryst* (B) **34** 129 1978, DOI: 10.1107/S0567740878002435; Beilstein **16** I 540, **16** II 625, **16** III 1240, **16** IV 1606.]

Tris(2,2'-bipyridine)ruthenium(II) dichloride ($6\text{H}_2\text{O}$) [14323-06-9 (anhydrous), 50525-27-4 ($6\text{H}_2\text{O}$)] $\text{C}_{30}\text{H}_{24}\text{Cl}_2\text{N}_6\text{Ru} \cdot 6\text{H}_2\text{O}$, M 748.6, m >300°. Recrystallise the red solid from water, then from MeOH [Ikezawa et al. *J Am Chem Soc* **108** 1589 1986, DOI: 10.1021/ja00267a032]. Soluble in Me_2CO . [For synthesis see Broomhead & Young *Inorg Synth* **28** 338 1990, DOI: 10.1002/9780470132593.ch86.]

Tris(η^5 -2,4-cyclopentadien-1-yl)samarium III [SmCp_3] [1298-55-1] $\text{C}_{15}\text{H}_{15}\text{Sm}$, M 345.6, m 365°. SmCp_3 was prepared by stirring SmCl_3 with cyclopentadienyl sodium [see 4989-82-1] in THF, the solvent is removed and the residue is heated at 200-250° in a high vacuum. Crystalline SmCp_3 is stable up to 400° and sublimes at 220°/10⁻⁴mm (also 145°/10⁻³mm) to give orange or yellow-orange crystals that are soluble in THF and are hydrolysed in H_2O . The IR is similar to that of SmCp_2 . [Wilkinson & Birmingham *J Am Chem Soc* **76** 6210 1954, DOI: 10.1021/ja01652a114; Birmingham & Wilkinson *J Am Chem Soc* **78** 42 1956, DOI: 10.1021/ja01582a009; Watt & Gillow *J Am Chem Soc* **91** 775 1969, DOI: 10.1021/ja01031a061.] It is used in the selective reduction of specific double bonds, i.e. has regioselectivity [Qian et al. *J Organomet Chem* **344** 175 1988, DOI: 10.1016/0022-328X(88)80477-7]. The **acetonitrile SmCp_3 complex** [100439-84-7] M 386.7 forms bright yellow crystals soluble in MeCN, the **cyclohexylisocyanide SmCp_3 complex** [37299-04-0] is soluble in C_6H_6 and has m 148°, and the **pyridine SmCp_3 complex** [114269-11-3] M 424.7 is obtained as a yellow crystalline solid by dissolving it in pyridine and evaporating to dryness and is soluble in pyridine and MeCN. [Deacon et al. *Aust J Chem* **40** 895 1987, DOI: 10.1071/CH9870895.]

Tris(d,d-dicampholylmethanato) europium (III) [$\text{Eu}(\text{dcm})_3$] [52351-64-1] $\text{C}_{63}\text{H}_{105}\text{EuO}_6$, M 1110.5, m 220-227.5°, 229-232°, [α]_D²⁵ +28.6 (c 5.4, CCl_4 , and varies markedly with concentration). Dissolve it in pentane, filter it from any insoluble material, evaporate to dryness and dry the residue (white powder) at 100°/0.1mm for 36 hours. Its IR has ν_{max} at 1540cm⁻¹. [McCreary et al. *J Am Chem Soc* **96** 1038 1974, DOI: 10.1021/ja00811a016.] Useful **chiral shift reagent** for determining enantiomeric ratios by NMR spectroscopy, and gives the highest chemical shift difference compared to other shift reagents.

Tris-[(3-trifluoromethylhydroxymethylene)-d-camphorato] europium (III) [$\text{Eu}(\text{tfc})_3$] [34830-11-0] $\text{C}_{36}\text{H}_{42}\text{EuF}_9\text{O}_6$, M 893.6, m 195-299° (dec), ~220°, [α]_D²⁴ +152 (c 2, CCl_4 , and varies markedly with concentration). Purify it by extraction with pentane, filter and the filtrate is evaporated and the residual bright yellow amorphous powder is dried at 100°/0.1mm for 36 hours. A sample purified by fractional molecular distillation at 180-200°/0.004mm gave a liquid which solidified and softened at ~130°, melted at ~180° and was analytically pure. Its IR (CCl_4) has ν_{max} at ν_{max} 1630-1680 cm⁻¹ and the ¹H NMR (CCl_4) has δ at -1.3 to 0.5 (br), -0.08 (s), 0.41 (s), 1.6-2.3 and 3.39 (s). [McCreary et al. *J Am Chem Soc* **96** 1038 1974, DOI: 10.1021/

ja00811a016; Goering et al. *J Am Chem Soc* **93** 5913 1971, DOI: 10.1021/ja00751a065.] Optically active NMR shift reagent. Used with Ag(fod) for enantiomeric resolution of *sec*-butylisothiuronium chloride [Wenzel & Zala *Anal Chem* **59** 562 1987, DOI: 10.1021/ac00131a006].

Trisodium citrate dihydrate [68-04-2 (Anhydrous), 6132-04-3 (2H₂O), 6858 (5H₂O)] C₆H₅O₇Na₃·2H₂O, M **294.1(2H₂O)**, m **150°(loses H₂O) and dec >300°**. Crystallise the salt from warm water by cooling to 0°. Solubility in H₂O is 92g/100ml at 25°. [Beilstein **3** III 1100, **3** IV 1274.] Useful antacid, and effective in removing scale from boilers and radiators.

Tungsten hexacarbonyl [14040-11-0] W(CO)₆, M **351.9**, m **150°**, d²⁵ **2.65g/cm³**. Sublime it *in vacuo* before use [Connor et al. *JCS Dalton Trans* 511 1986, DOI: 10.1039/DT9860000511]. Catalyst in the photochemical cyclisation of silyloxytrienes *via* either a 6-endo pathway or a **Cope rearrangement** [Miura et al. *J Organomet Chem* **692** 562 2007, DOI: 10.1016/j.jorganchem.2006.08.037], a precursor to a **Fischer carbene complex** which reduces Pd(II) to nanoparticulate Pd(0) that is catalytically active in Hiyama cross-coupling [Srimani et al. *Org Lett* **9** 3639 2007, DOI: 10.1021/ol7015143]. **TOXIC due to possible release of carbon monoxide.**

Vanadium (III) acetylacetonate (vanadyl tris[2,4-pentadienato-O,O'], V[AcAc]₃) [13476-99-8] (C₅H₇O₂)₃V, M **348.3**, m **181-184°, 185-190°, pK₁²⁵ 2.92, pK₂²⁵ 3.5(for aquo V³⁺ hydrolysis)**. It crystallises from acetylacetone in brown plates. It can be distilled in small quantities without decomposition. It is soluble in CHCl₃ and *C₆H₆, and evaporation of a CHCl₃ solution yields brown crystals which are washed with cold EtOH and dried in vacuum or at 100° in a CO₂ atmosphere. Under moist conditions it readily oxidises [V(AcAc)₃ to V(AcAc)₂O]. [Fernelius & Bryant *Inorg Synth* **5** 105 1957, DOI: 10.1002/9780470132364.ch29; McKaveney & Freiser *Anal Chem* **30** 526 1958, DOI: 10.1021/ac60136a023; Beilstein **1** IV 3672.]

Vanadyl (IV) acetylacetonate (vanadyloxa bis[2,4-pentadienato-O,O'], VO[AcAc]₂) [3153-26-2] (C₅H₇O₂)₂VO, M **265.2**, m **235°(dec), 256-259°, 258°, d²⁵ 1.50g/cm³**. It provides blue-green crystals from acetone. It is soluble in CHCl₃, CH₂Cl₂, MeOH, EtOH and *C₆H₆. [cf. Fernelius & Bryant *Inorg Synth* **5** 105 1957, DOI: 10.1002/9780470132364.ch29; Rowe & Jones *Inorg Synth* **5** 113 1957, DOI: 10.1002/9780470132364.ch30; Beilstein **1** IV 3672; for applications see Fieser **2** 256, **3** 331, **11** 47.]

Vinylferrocene (ferrocenylethene) [1271-51-8] C₁₂H₁₂Fe, M **212.1**, m **51-52.5°, 51-53°, b 80-85°/0.2mm**. Dissolve vinylferrocene in Et₂O, wash it with H₂O and brine, dry (Na₂SO₄), and evaporate to a small volume. Purify it through an Al₂O₃ (Spence grade H) column by eluting the yellow band with petroleum ether (b 40-60°). The low melting orange crystals can be sublimed. The **tetracyanoethylene adduct** [49716-63-4] crystallises from *C₆H₆/pentane and has m **137-139°(dec)**. [Horspool & Sutherland *Can J Chem* **46** 3453 1968, DOI: 10.1139/v68-574; Berger et al. *J Org Chem* **39** 377 1974, DOI: 10.1021/jo00918a012; Rausch & Siegel *J Organomet Chem* **11** 317 1968, DOI: 10.1016/0022-328X(68)80054-3; Beilstein **16** III 1787.] Useful for the preparation of 2-aminoethylferrocenes and lithium amides [Joly et al. *Tetrahedron* **63** 761 2007, DOI: 10.1016/j.tet.2006.10.078].

Vinyltributylstannane (vinyltributyltin) [7486-35-3] (C₅H₇O₂)₂VO, M **317.1**, b **104-106°/3.5mm**, d²⁵ **1.085g/ml**, n_D²⁰ **1.478**. Fractionate the stannane under reduced pressure and taking the middle fraction to remove impurities such as (*n*-Bu)₃SnCl. [Seyferth & Stone *J Am Chem Soc* **79** 515 1957, DOI: 10.1021/ja01560a003; Beilstein **16** III 1279, **4** IV 4315] Useful vinyl nucleophile for bromoacetylenes [Yoshida et al. *Org Lett* **6** 1979 2004, DOI: 10.1021/ol049438k] and bromoaromatics [Occhiato et al. *J Med Chem* **47** 3546 2004, DOI: 10.1021/jm031131o].

Zinc acetate dihydrate [5970-45-6] (CH₃CO₂)₂Zn·2H₂O, M **219.5**, m **100°(loses 2H₂O), 237°, d₄²⁰ 1.74, pK²⁵ 8.96 (for hydrolysis of Zn²⁺ to ZnOH⁺)**. It crystallises (in poor yield) from hot water or, better, from EtOH. [Beilstein **2** III 193, **2** IV 114.]

Zinc acetonilacetate [Zn(acac)₂] [14024-63-6, 108503-47-5 (xH₂O)] (C₅H₇O₂)₂Zn·xH₂O, M **263.6(anhydr), m **135-138°, 138°, 138°**. The zinc complex crystallises from hot 95% EtOH. [Fernelius &**

Bryant *Inorg Synth* **5** 105 1957, DOI: 10.1002/9780470132364.ch29; Wakita et al. *J Organometal Chem* **301** C17 1986, DOI: 10.1016/0022-328X(86)80017-1; *Beilstein* **2** IV 144.]

Zinc caprylate (zinc n-octanoate) [557-09-5] $C_{16}H_{30}O_4Zn$, **M 351.8, m 135-136°, 136°, 142°**. It crystallises from boiling EtOH in plates, but is sparingly soluble in boiling H_2O . Originally precipitated by addition of an aqueous solution of zinc sulfate to an ammoniacal solution of caprylic acid. Store dry as it as it hydrolyses in moist air liberating caprylic acid. It has fungicidal activity. [Van Renesse *Justus Liebigs Ann Chem* **171** 380 1874, DOI: 10.1002/jlac.18741710224; *Beilstein* **2** H 284.]

Zinc formate dihydrate [557-41-5, 5970-62-7] $(CHO_2)_2Zn \cdot 2H_2O$, **M 191.4(2H₂O), m 140°(loses H₂O), d²⁰ 2.21g/cm³**. It crystallises from hot water (solubility is 5.2g/100ml at 20°). [*Beilstein* **2** H 16, **2** I 14, **2** II 21, **2** III 25, **2** IV 17.]

Zinc RS-lactate trihydrate [554-05-2, 16039-53-5 (2H₂O), 51120-75-3 (DL- 3H₂O), 312619-27-5 (L- H₂O)] $C_6H_{12}O_7Zn$, **M 297.5, m 272-274° (L- H₂O)**. It crystallises from water (6ml/g). [*Beilstein* **3** I 107, **3** II 204, **3** III 464, **3** IV 637.]

Zincon (o-[1-(2-hydroxy-5-sulfo)-3-phenyl-5-formazono]-benzoic acid) [135-52-4] $C_{20}H_{16}N_4O_6S$, **M 440.4**. Main impurities are inorganic salts which can be removed by treatment with dilute acetic acid. Organic contaminants are removed by refluxing with ether. It can be recrystallised from dilute H_2SO_4 and it complexes with Zn ions (see below). [Fichter & Schiess *Chem Ber* **33** 747 1900, DOI: 10.1002/cber.190003301131; *Beilstein* **16** IV 421.]

Zincon disodium salt (o-[1-(2-hydroxy-5-sulfo)-3-phenyl-5-formazono]-benzoic acid di-Na salt) [135-52-4, 56484-13-0] $C_{20}H_{15}N_4O_6Na$, **M 462.4, m ~250-260° (dec), λ_{max} 523nm, 490nm**. Zincon solution is prepared by dissolving 0.13g of the powder in aqueous N NaOH (2ml diluted to 100ml with H_2O). This gives a deep red colour which is stable for one week. It is a good reagent for zinc ions but also forms stable complexes with Cu ions as well in colorimetry. Used in chelate titration for Ca, Cd, Hg, Pb as well as Zn ions. [UV-VIS: Rush & Yoe *Anal Chem* **26** 1345 1954, DOI: 10.1021/ac60092a024; Hunter & Roberts *J Chem Soc* 820 1941, DOI: 10.1039/JR9410000820; Platte & Marcy *Anal Chem* **31** 1226 1959, DOI: 10.1021/ac60151a048;] The **free acid** (see above) has been recrystallised from dilute H_2SO_4 . [Fichter & Scheiss *Chem Ber* **33** 747 1900, DOI: 10.1002/cber.190003301131; *Beilstein* **16** IV 421.]

Zinc phthalocyanine [14320-04-8] $C_{32}H_{16}N_8Zn$, **M 577.9, λ_{max} 701nm**. Sublime it in oxygen-free N_2 . [For solvent effects on photochemical and fluorescence properties of zinc phthalocyanines see Ogunsipe et al. *J Mol Struct* **650(1)** 131 2003, DOI: 10.1016/S0022-2860(03)00155-8]. [*Beilstein* **26** III/IV 4257.]

Zinc 5,10,15,20-tetraphenylporphyrin [14074-80-7] $C_{44}H_{28}N_4Zn$, **M 678.1, λ_{max} 418nm and 556nm**. Purify the porphyrin by chromatography on neutral (Grade I) alumina, followed by recrystallisation from CH_2Cl_2 /MeOH [Yamashita et al. *J Phys Chem* **91** 3055 1987, DOI: 10.1021/j100295a082]. [*Beilstein* **26** IV 1959.]

Zirconium (IV) acetylacetonate [17501-44-9] $(C_5H_7O_2)_4Zr$, **M 487.7, m 190-193° (sealed capillary) 194.5-195°, 192-195°**. The zirconium complex crystallises from hot 95% EtOH (**m** 190-191°) or petroleum ether and sublimes *in vacuo* at 140° with slight decomposition. The **decahydrate** effloresces in air and dehydrates in a 0.1mm vacuum. Its solubility in 1L at 20° is 30g in CS_2 , 47g in CCl_4 , 44g in ethylene dibromide, 56g in acetylacetone and 200g in $*C_6H_6$. [Young, Arch et al. *Inorg Synth* **2** 121 1964, DOI: 10.1002/9780470132333.ch35; *Beilstein* **1** H 782, **1** II 835, **1** III 3121, **1** IV 3672; see Fieser **13** 351]

Zirconium (IV) propoxide [23519-77-9] $(C_3H_7O)_4Zr$, **M 327.6, b 198°/0.03mm, 208°/0.1mm, d²⁵ 1.044g/ml, n_D²⁰ 1.451**. Although it was reported that it could not be crystallised or sublimed even at 150°/10⁻⁴mm [Bradley & Wardlaw *J Chem Soc* 280 1951, DOI: 10.1039/JR9510000280], the propoxide, when properly prepared, has been purified by distillation in a high vacuum [Bradley et al. *J Chem Soc* 2025 1953, DOI: 10.1039/JR9530000205]. [*Beilstein* **1** IV 1420; for applications see Fieser **11** 605, **13** 352.] This propoxide is the common precursor for the preparation of Zirconia (ZrO_2) [Sui et al. *Langmuir* **22(9)** 4390 2006, DOI: 10.1021/la053513y; Heine et al. *Combustion and Flame* **144(4)** 809820 2006, DOI: 10.1016/j.combustflame.2005.09.012].

CHAPTER 5

CATALYSTS

INTRODUCTION

Catalysts, and the process of catalysis, are almost as old as the chemical and physical sciences. Over the centuries a large number of metals and their derivatives, and a large variety of organic compounds, have been found to catalyse reactions successfully. The importance of catalysis in the sciences can be judged by the number of Nobel Prizes that have been awarded over the years for discoveries in this field of research. During the past two or so decades much research emphasis has concentrated on combining metals with organic ligands, by complexation, in order to achieve greater metal solubility in a variety of solvents, not only for more efficient catalysis but also, by using chiral ligands, to achieve high, if not almost complete, regioselectivity and/or stereoselectivity. This is evidenced by the award of the 2001, 2005 and 2010 Nobel Prizes for Chemistry respectively for *Catalytic Asymmetric Synthesis* (to W. S. Knowles, R. Noyori and K. B. Sharpless), for *Catalytic Olefin Metathesis* (to Y. Chauvin, R. H. Grubbs and R. R. Schrock) and for *Catalytic Cross-Coupling Reactions* (to R. Heck, E.-I. Negishi and A. Suzuki). In addition to the many recently published reviews and monographs on various aspects of catalysis, several new journals have appeared, e.g. *ChemCatChem* (from 2009) and *Advanced Synthesis & Catalysis* (from 2001), and others that can be found on the internet. Original research papers, and reviews, on ‘cutting edge’ discoveries in chemical and enzyme catalysis can be published particularly in these periodicals, allowing proximity and cross communication between these fields of work.

Commercial suppliers of chemicals have availed themselves of this surge in new catalysts, and are now providing a large variety of recently discovered catalysts to the scientific community. The present chapter addresses a selection of these commercially available catalysts. Generally they can be used as supplied. However, some samples may have partially deteriorated, or have short shelf lives; so knowledge about their purification, or assessment of purity, is essential. Purification details are provided for these, and in many cases published details of their preparations have also been included. Knowledge of the preparations provides information about probable impurities, and may dictate the best purification procedures. Since several catalysts (and some ligands) have high and/or *not* well defined melting points, some spectroscopic data, and optical properties if they are chiral, have been included.

This chapter is divided into two sections. The first section includes heterogeneous and homogeneous catalysts. The second section contains ligands and reagents used for the preparation of these catalysts and of these ligands, as well as a section on phase transfer catalysts, and a section containing a small number of chiral auxiliaries. There are some substances mentioned in this chapter, but not included here, whose purification procedures are described in Chapters 3, 4 and 6. Their purification procedures are **not** repeated here. However, they can be located *via* their Chemical Abstracts Registry Numbers (CASRN), also used for cross referencing, and their place in the book can be found from the CASRN Index which provides their respective page entries. **Abbreviations** of titles of periodicals are defined as in the Chemical Abstracts Service Source Index (CASSI), but without punctuation. Fieser & Fieser’s *Reagents for Organic Synthesis* references will be shortened to Fieser throughout.

It should be pointed out that great care must be taken to ascertain that the catalyst is indeed the metal or reagent used, and not traces of unsuspected contaminating metal. This point has been clearly made in the article by J.M. Crow entitled ‘When is a catalyst not a catalyst’ (Chemistry World 8 46 2011). He draws attention to the papers of Buchwald & Bolm Angew Chem Int Ed 48 5586 2009, DOI: 10.1002/anie.200902237; Larsson et al. Angew Chem Int Ed 48 5691 2009, DOI: 10.1002/anie.200902236; Leadbeater & Marco Angew Chem Int Ed 42 1407 2003, DOI: 10.1002/anie.200390362; Arvela et al. J Org Chem 70 161 2005, DOI: 10.1021/jo048531j; and the measurement of trace impurities at the ppb (parts per billion) and/or ppt (parts per trillion) levels in Gonda, Tolnai and Novák Chem Eur J 16 11822 2010, DOI: 10.1002/chem.201001880; and the possibility of co-catalysis.

Purification of Laboratory Chemicals.

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CATALYSTS—PART 1

HETEROGENEOUS METAL CATALYSTS

Copper Chromite (Lazier catalyst, copper chromium oxide) [12053-18-8] $2\text{CuO} \cdot \text{Cr}_2\text{O}_3$, **M 311.1**. It is also known as **Adkins catalyst** (with very slight difference in preparation). The catalyst is prepared by adding $\text{Cu}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$ (109g) to a warm solution of $\text{Ba}(\text{NO}_3)_2$ (13g) in H_2O (140ml) at 70° and stirring until clear. Ammonium chromate solution, made by dissolving ammonium dichromate (63g) in H_2O (300ml) and adding concentrated aqueous ammonia (75ml, sp.gr. 0.90), is stirred in a thin stream into the solution of the above nitrates, stirred for 5 minutes further and the red-brown precipitate of copper barium ammonium chromate is filtered off, washed with H_2O , drained well and dried at 110° . The salt is placed in a suitable container and heated in a muffle furnace at $350\text{--}450^\circ$ for 1 hour. **Note** that a spontaneous exothermic reaction occurs on heating with evolution of gas. The ignited blue-black residue ($\sim 80\text{g}$), composed of copper and barium chromates and CuO , is pulverised thoroughly, suspended in aqueous 10% AcOH (600ml), stirred for 10 minutes, allowed to settle during 15 minutes, the supernatant is decanted, and the process is repeated. Filter off the solid, dry it at 110° and grind it in a mortar to give the catalyst ($\sim 70\text{g}$) as a fine black powder. It is not affected by air or moisture and no special storing precautions are necessary. Other than the use of $\text{Na}_2\text{Cr}_2\text{O}_7$ (Adkin) instead of $(\text{NH}_4)_2\text{Cr}_2\text{O}_7$ (Lazier), there seems to be very little difference in the preparation procedure or the activity of the catalyst [see Fieser **2** 82]. Note that the inclusion of barium is for protection against sulfate which is a catalyst poison, and stabilises it against reduction. If a **barium free** catalyst is required, then it is omitted in the preparation and the amount of $\text{Cu}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$ should be increased to 121g without altering the preparation procedure. [Lazier & Arnold *Org Synth Coll Vol* **2** 142 1943, DOI: 10.15227/orgsyn.019.0031; Adkins et al. *J Am Chem Soc* **72** 2626 1950, DOI: 10.1021/ja01162a079]. It is an effective catalyst for the reduction of aldehydes, ketones and esters to the corresponding alcohols, aliphatic but not aromatic double bonds, the reduction of amides to amines, and for the decarboxylation of carboxylic acids [see Fieser **1** 186]. These reactions however may have to be carried out in stainless steel bombs or autoclaves as several reactions require high temperatures ($100\text{--}260^\circ$) and pressures (up to 5000psi).

Nickel [7440-02-0] **Ni**, **M 58.7**, **m 1455°**, **b 2730°**, **d²⁵ 8.90**. The catalytic use of the metal can be divided into (a) the Raney nickel type and (b) the non-Raney nickel type.

Raney Nickel type: This catalyst was introduced by the Raney Catalyst Division of Grace & Co and is made from a Ni-Al alloy ($\sim 1:1$, e.g. by fusion at $1200\text{--}1500^\circ$). The catalyst is prepared by treating the alloy with alkali, which dissolves the Al, and the alkali is washed off as thoroughly as possible. The method was developed by Adkins and coworkers at the University of Wisconsin (USA) and the catalyst was designated as **W-1** to **W-6** (chronological, not activity, order) [Billica & Atkins *Org Synth Coll Vol* **3** 176 1955, DOI: 10.15227/orgsyn.029.0024; and for comparison of Ws see Adkins & Krsek *J Am Chem Soc* **70** 412 1948, DOI: 10.1021/ja01181a501]. The catalyst has been used very extensively for reduction reactions, as well as for replacing sulfur by hydrogen in compounds; and there is a large volume of literature on this subject [see Fieser **1** 440, 723, **2** 289, 293, **4** 267, **5** 263, 381, 570, 736, **6** 502, **7** 321, **8** 433, **9** 405, **10** 339, **11** 457, **12** 175, 218, 338, 422, **13** 265, **14** 270, **16** 278, **17** 296]. The reactions normally require stirring or shaking at ambient temperatures, and pressures of H_2 not greater than 100psi. A variety of equipment for this purpose is available commercially. The catalyst is essentially prepared by adding the Ni-Al alloy (150g) in small portions to aqueous sodium hydroxide (190g in 750ml of H_2O) with stirring in an ice bath at 10° at such a rate that the temperature is kept below 25° , ca 2 hours. *n*-Octanol may be added to prevent excessive foaming. This reaction should be carried out in an efficient fume cupboard as H_2 is liberated during the reaction (with the possibility of igniting), and it is preferable to use magnetic stirring to avoid sparking from an electric motor. When addition of the alloy is complete, allow the temperature of the mixture to rise to $\sim 25^\circ$, and then heat it gradually (avoiding frothing over) on a boiling water bath until evolution of H_2 is complete (8-12 hours). Add distilled H_2O to restore the original volume, stir, allow the Ni to settle down, then decant the supernatant liquid. Transfer the Ni to a graduated cylinder with H_2O , allow it to settle, decant the supernatant liquid, add aqueous NaOH (25g, in 250ml H_2O), shake the mixture thoroughly, allow the Ni to settle again and decant off the supernatant alkaline liquid. Wash the Ni with H_2O ($\sim 750\text{ml}$ each time) by shaking, allow it to settle and decant the supernatant H_2O . Repeat

the process until the washings are neutral to litmus and then a further 10 times. This may require 30 to 40 washings. Repeat the washing process three times with 95% EtOH, then again three times with absolute EtOH. Store the Ni catalyst in bottles which are filled with absolute EtOH to the brim and stopper them. Keep them in a cold store room. Yield of Raney Ni is usually ~75g. The metal should be kept moist, or under liquid, at all times because the dry solid is highly **pyrophoric**. The more active the catalyst, the more readily it ignites. For this reason it is best to spoon the catalyst out from the settled material. When prepared as above it contains ~0.6g of Ni per ml of settled solid, and a level tablespoonful contains ~3g of Ni. [Mozingo *Org Synth Coll Vol 3* 181 1955, DOI: 10.15227/orgsyn.021.0015]. The activity of Raney Ni deteriorates slowly with time, but if prepared properly and stored as above it should be quite active for about 6 months. Its activity can be enhanced immensely by addition of a very small amount of triethylamine and platinum chloride [Levering & Lieber *J Am Chem Soc* **71** 1515 1949, DOI: 10.1021/ja01172a528]. It has been used both as a **catalytic oxidation catalyst** (with an appropriate donor), as well as a **reduction catalyst** [Kleiderer & Kornfeld *J Org Chem* **13** 455 1948, DOI: 10.1021/jo01161a022].

Non-Raney Nickel type: This includes a catalyst of the composition Ni/Al₂O₃ formed by treating Ni-Al alloy with H₂O at 70° [Tyman *Chem Ind (London)* 404 1964], and the commercially available Ni on silica (with 60 wt% loading on Kieselguhr) and Ni on silica/alumina (with ~65 wt% loading). All **Ni coordination compounds** which have catalytic activity can come under this heading.

Palladium [7440-05-3] Pd, M 106.4, m 1555°, b 3167°/atm, d₄²⁰ 0.951, electrical resistivity at 0° is μΩ-cm. Pd is a silver-white metal which also occurs as a black powder or in a compressible spongy form. It is reactive towards HNO₃, H₂SO₄, HCl and HClO₃, particularly in the presence of air or oxygen; and forms dihalides with halogen at high temperatures. It has the capacity of adsorbing and absorbing large volumes of H₂. It is extremely useful in chemical reactions as displayed in a limited way in this chapter. Apart from its extensive value in metal coordination chemistry it has out-classed many related metals in the field of catalysis. As well as its use in catalysis as the free metal in precipitated form, pellet or granular form, it has found extensive use in catalysis when diluted with C [7440-44-0], BaSO₄, CaSO₄, SrSO₄, Kieselguhr, and Al₂O₃ (see below). The following catalyst preparations have been used for many decades:

Palladium on charcoal (5% Pd)— A clear solution of PdCl₂ (8.2g, 46mmol) in concentrated HCl (20ml, 240mmol) and H₂O (50ml), obtained by heating on a steam bath for 2 hours or until complete dissolution, is added with stirring to a suspension of charcoal (93g, HNO₃—washed)[†] in H₂O (1.2L) at 80° followed by 37% formaldehyde solution (80ml, 100mmol). The solution is made slightly alkaline to litmus (pH ~8.0) by careful addition of aqueous 30% sodium hydroxide with stirring which is continued further for 5 minutes longer. The catalyst is filtered off and washed with H₂O (10 x 250ml), drained well then dried at 25°, first in air then over KOH or CaCl₂ in a desiccator. The dry catalyst (93-98g), which is in a reduced state is stored in a tightly closed vessel. [Do not dry at high temperatures as ignition may occur.]

[†] Darco G-60, Norit or other carbons can be used and should be heated on a steam bath with 10% aqueous HNO₃ for 2-3 hours, washed free from acid (check pH of washings), and dried at 100-110° before use (see chapter on Aliphatic Compounds).

A comparison of the rates of hydrogenation with 5% Pd/C, 5% Pt/C, 5% Rh/C and 5% Ru/C catalysts towards reduction of C=C, C=O, C-OH and benzene rings in various solvents at 25° and initial H₂ pressure of 1 atmosphere were made by Breitner, Roginski & Rylander [*J Org Chem* **24** 1855 1959, DOI: 10.1021/jo01094a004.]

Note that palladised charcoal has also been used for **dehydrogenation**, e.g. tetralin is converted to naphthalene by boiling for 4 hours in the presence of Pd/C in a slow stream of CO₂, and α-tetralone is dehydrogenated to naphthalene by boiling (1 hour, with internal temperature of 235°) in the presence of Pd/C and a slow stream of CO₂ in 75% yield.

Palladium on charcoal (10% Pd)— A clear solution of PdCl₂ (8.3g) in concentrated HCl (5.5ml) and H₂O (40ml), prepared as above, is poured into a solution of NaOAc.3H₂O (135g) in H₂O (500ml) in a hydrogenation bottle, and charcoal (45g, HNO₃—washed, see above[†]) is added and the mixture is hydrogenated until absorption ceases (1-2 hours). The catalyst is collected on a filter funnel, washed with H₂O (5 x 400ml), drained well, dried in air and then in a desiccator over KOH or CaCl₂; store the reduced catalyst (48-50g) as above.

Palladium chloride on charcoal (5% Pd)— A clear solution of PdCl₂ (8.2g, 46mmol) in concentrated HCl (20ml, 240mmol) and H₂O (50ml), obtained by heating on a steam bath for 2 hours or until complete dissolution, is diluted with H₂O (140ml) and poured over charcoal (92g, HNO₃—washed, see above[†]) in a 20cm

evaporating dish, thoroughly mixed and dried, firstly on a steam bath and then in an oven at 100° with occasional stirring until the catalyst (98-100g) is completely dry and is then stored in a stoppered bottle until required. When required, the desired aliquot of catalyst is transferred to a hydrogenation bottle (or flask) and reduced in the solvent used for hydrogenation by shaking with H₂ in the hydrogenation apparatus. When absorption of H₂ is complete, the catalyst is filtered off, washed with fresh solvent to remove HCl, returned to the hydrogenation bottle (which should have been rinsed with the same fresh solvent), the material for reduction is added and hydrogenation is carried out in the usual way.

Palladium on barium sulfate (5% Pd)— To a rapidly stirred hot (80°) solution of reagent grade Ba(OH)₂·8H₂O (126.2g, 400mmol) in H₂O (1.2L), 6N H₂SO₄ (120ml, 360mmol) is added all at once then more 6N H₂SO₄ is added to the BaSO₄ suspension that is formed in order to be just acidic to litmus (pH ~5). A clear solution of PdCl₂ (8.2g, 46mmol) in concentrated HCl (20ml, 240mmol) and H₂O (50ml), obtained by heating on a steam bath for 2 hours or until complete dissolution, is subsequently added to the hot (80°) stirred BaSO₄ suspension[#] followed by 37% formaldehyde solution (8ml, 100mmol). The suspension is made alkaline to litmus (pH ~8.0) by careful addition of aqueous 30% sodium hydroxide with stirring, which is continued further for 5 minutes longer, and allowed to settle. The clear colourless supernatant liquid is decanted off, replaced by H₂O, resuspended, and washed ten times by decantation. The catalyst is collected on a 90mm medium porosity sintered glass funnel, drained until the filter cake begins to crack, washed again with H₂O (95 x 250ml), drained as completely as possible and placed in an oven at 80° until a dry catalyst (93-98g) is obtained. Store it as above. [This catalyst is particularly useful for *Rosenmund catalytic reductions* of acid chlorides (RCOCl) to aldehydes (RCHO), cf. Hershberg & Cason *Org Synth Coll Vol 3* 626 1955, DOI: 10.15227/orgsyn.023.0063.]

[#]An equivalent weight of precipitated BaCO₃ (93g) may be used instead for BaSO₄; but the volume of concentrated HCl in preparing the PdCl₂ solution should be reduced to 8.2ml (to avoid decomposition of the carbonate) to give **5% Pd on calcium carbonate** catalyst. [Mozingo *Org Synth Coll Vol 3* 685 1955, DOI: 10.15227/orgsyn.026.0077.]

Note that at the completion of hydrogenations with these catalysts, the used catalysts that are filtered off should be kept moist with solvent or H₂O as they may be *pyrophoric* when dry. Also, with the catalysts that have been prepared using the formaldehyde (HCHO) procedure, the spent catalysts from reduction reactions in which MeOH is used as solvent are particularly flammable, possible because MeOH/HCHO form an *oxidation-reduction couple* since there is no guarantee that the catalyst is completely free from HCHO.

Other commonly used supports for Pd include **CaCO₃** [Busch & Stöve *Chem Ber* **49** 1063 1916, DOI: 10.1002/cber.191604901114], **SrCO₃** [Martin & Robinson *J Chem Soc* 491 1943, DOI: 10.1039/JR9430000491], **Al₂O₃** [Alan C. Johnson US pat 2,366,409, *Chem Abstr* **39** 2001 1945], **Kieselguhr** [Rosenmund & Langer *Chem Ber* **56** 2262 1923, DOI: 10.1002/cber.19230561012; Sabalitschka & Moses *Chem Ber* **60** 786 1927, DOI: 10.1002/cber.19270600333], **silica-gel** [Fester & Brude *Chem Ber* **56** 2245 1923, DOI: 10.1002/cber.19230561007], **asbestos** [Zelinsky & Borisoff *Chem Ber* **57** 150 1924, DOI: 10.1002/cber.19240570126; Zelinsky & Turowa-Pollak *Chem Ber* **58** 1292, 1298 1925, DOI: 10.1002/cber.19250580717, DOI: 10.1002/cber.19250580718] and **polyvinyl alcohol** [Kavanagh & Nord *J Am Chem Soc* **66** 2126 1944, DOI: 10.1021/ja01240a506].

Platinum [7440-06-4] **Pt**, **M 195.1**, **m 1772°**, **b 3827°**, **d²⁰ 21.45**. Platinum, when used directly or on a carbon, alumina or silica support, or *via* its oxide, is generally an excellent catalyst for the reduction of compounds. Hydrogenations using Pt catalysts are usually carried out at ambient temperatures, and atmospheric pressure of H₂, and the reaction can be followed by using a manometer and noting the absorption of gas. They may require, as in the reduction of aromatic rings to alicyclic rings, somewhat elevated pressures (a few to several atmospheres) and temperatures. Specialised equipment is used in such cases (see Ni catalysts above).

Platinum Black: It is obtained by reducing a solution of chloroplatinic acid [16941-12-1] with alkaline formaldehyde and shaking vigorously to coagulate the colloidal metal formed [Feulgen *Chem Ber* **54** 360 1921, DOI: 10.1002/cber.19210540225]. In addition to its use as a catalyst, it destroys H₂O₂ in liquids or solvents by shaking them with the metal until liberation of O₂ ceases [Cope & Ciganek *Org Syn Coll Vol 4* 612 1963, DOI: 10.15227/orgsyn.039.0040].

Platinum Sponge: Ammonium hexachloroplatinate is precipitated when recovered Pt residues are dissolved in *aqua regia* (see below), evaporated to dryness then dissolved in the minimum volume of H₂O and treated with saturated ammonium chloride. The (NH₄)₂PtCl₆ is filtered off and dried at 100°. Ignition of this salt gives Pt sponge. [Wickers *J Am Chem Soc* **43** 1268 1921, DOI: 10.1021/ja01439a008.]

Brown & Brown Catalyst: This is usually prepared *in situ* by adding a solution of NaBH₄ in EtOH stabilised

with NaOH to a suspension of charcoal (see palladium catalysts above) in a solution of chloroplatinic acid in EtOH (or aqueous EtOH) to form the supported catalyst. Excess HCl is added, followed by the substrate for reduction, then NaBH₄ in EtOH stabilised with NaOH is added dropwise at such a rate as to maintain atmospheric pressure of H₂. Here the supported catalyst and H₂ are generated *in situ* resulting in more effective reductions. [Brown et al. *J Org Chem* **28** 214 1963, DOI: 10.1021/jo01036a501.]

Platinum on support: The catalyst is prepared from a cooled solution of the metal (5.0g, or PtCl₄ or H₂PtCl₆) in H₂O (50ml) and concentrated HCl (5ml) and support [11g, acid washed charcoal (see Pd above), neutral alumina, silica or asbestos (Gooch asbestos, termolyte not chrysolite, boil with concentrated HNO₃, filter, wash free from acid with H₂O and dry at 100°)] in a freezing mixture and treat with formalin (50ml, 40% formaldehyde). The mixture is then stirred while aqueous KOH (50g in 50ml of H₂O) solution is added slowly and keeping the temperature below 5°. The temperature is then allowed to rise to 60° in 15 minutes. The catalyst is washed thoroughly with H₂O by decantation, finally with dilute AcOH, collected onto a suction filter and washed with H₂O until free from chloride ions and alkali, and drained well. Dry the solid at 100° and store it in a desiccator. This method gives a catalyst containing *ca* 30-40% of metal, but the amount can be adjusted by varying the ratio of metal to support. [Brown & Brown *J Am Chem Soc* **84** 2827, 2829 1962, DOI: 10.1021/ja00873a037, DOI: 10.1021/ja00873a039.]

A comparison of the rates of hydrogenation with 5% Pd/C, 5% Pt/C, 5% Rh/C and 5% Ru/C catalysts towards reduction of C=C, C=O, C-OH and benzene rings in various solvents at 25° and initial H₂ pressure of 1 atmosphere were made by Breitner, Roginski & Rylander [*J Org Chem* **24** 1855 1959, DOI: 10.1021/jo01094a004].

Platinum (IV) oxide (Adams catalyst) [*1H₂O* 12137-21-2; *xH₂O* 52785-06-5] **PtO₂, M 227.1, M 245.1 (1 H₂O), m >450°.** Adams catalyst can be prepared in three ways.

Firstly, from ammonium hexachloroplatinate (IV) [*16919-58-7*] [3.0g, (NH₄)₂PtCl₆] and KNO₃ (30g, Analytical Grade; NaNO₃ yields slightly less active catalyst) which are thoroughly mixed in a porcelain dish (**fume cupboard, use a head shield for protection**) and heated gently at first until the rapid evolution of gas subsides, then strongly until the temperature reaches 300° (15 minutes, no splattering). The liquid mass is heated further at 500-530° for 30 minutes and allowed to cool. Add H₂O (50ml) to the residue, stir and allow the brown platinum oxide (PtO₂, H₂O) to settle. Wash it once or twice with H₂O by decantation, collect it on a filter, wash it free from nitrates (stop washing if the solid starts to become *colloidal*;; traces of NaNO₃ do not affect catalytic activity) and dry the oxide in a desiccator. Store it in small aliquots. # If *colloidal Pt* filters through, it can be checked by testing a small volume of the filtrate for Pt by acidifying it with HCl, adding a few drops of stannous chloride when a yellow or brown colour results depending on the amount of Pt present. The yellow colour is soluble in Et₂O rendering the **test more sensitive**. If Pt is present in the filtrate then heat the filtrate with excess formalin and aqueous NaOH, cool, and the platinum black which separates is filtered off and worked up with other Pt residues. The weight of PtO₂ obtained is about equal to half the weight of (NH₄)₂PtCl₆.

Secondly, purest possible chloroplatinic acid [3.5g, H₂PtCl₆] in H₂O (10ml) is mixed with KNO₃ (35g, Analytical Grade; NaNO₃ yields slightly less active catalyst) in a porcelain dish, evaporate gently to dryness and heat at 350-370° for 10 minutes (use a strong heat source, **fume cupboard, use a head shield for protection**) when brown oxides of nitrogen evolve and brown PtO₂ separates. Stir more vigorously if frothing occurs and direct an additional burner at the top of the mixture. [Take care not to allow the top of the mixture to solidify, e.g. by removing the burner that is underneath, otherwise frothing over occurs]. The temperature is raised to 400° in the following 15 minutes, after which liberation of fumes decreases considerably, and the temperature is raised further to 500-550° (*ca* 2 minutes); at this stage evolution of brown gases practically ceases but there is still a gentle evolution of gas. Maintain at this temperature (with burner at full heat) for 30 minutes by which time fusion is complete. Allow the container to cool slowly to ~20-25°, add H₂O (50ml) and isolate the catalyst as above.

Thirdly, Pt metal and or Pt metal residues are dissolved in *aqua regia* [**prepared by adding slowly concentrated HNO₃ (1 volume) to concentrated hydrochloric acid (3 volumes) in a glass container in an efficient fume cupboard and head protection**] using gentle heat if necessary, evaporate several times almost to dryness with concentrated HCl, dissolve the final residue in a small volume of H₂O and precipitate the metal as (NH₄)₂PtCl₆ with saturated aqueous NH₄Cl. Filter off the precipitate, dry it at 100° and continue as in the first procedure (see above). [Adams et al. *Org Synth Coll Vol* **1** 463, 466 1941, DOI: 10.15227/orgsyn.008.0092; Bruce *J Am Chem Soc* **58** 687 1936, DOI: 10.1021/ja01295a501.]

Reactions catalysed by Adams catalyst can be carried out in alcoholic solution which is sometimes enhanced by

the presence of HCl; a variety of other organic solvents have been used which include EtOAc, EtOAc+15% AcOH, EtOAc+8%EtOH, and glacial AcOH; and neat $\text{CF}_3\text{CO}_2\text{H}$ is a particularly good solvent for the reduction of $\text{C}=\text{N}$ in heterocyclic compounds.

PtO_2 on silicic acid has been prepared in much the same way as above (*fume cupboard and head protection*) from chloroplatinic acid (7g) in H_2O (25ml) and silicic acid (20g, 200 mesh) in a borosilicate beaker (100ml) which are stirred into a smooth paste and heated on a burner until desiccated. Heat to $\sim 360^\circ$ and add powdered NaNO_3 (70g) in portions while stirring and mixing vigorously with a glass rod (evolution of brown fumes occurs at each addition with frothing) and heat until gas evolution ceases. Add further molten NaNO_3 (20g) to the beaker with heating while scratching its sides. Cool, and transfer to a 2L beaker containing H_2O (1.5L), stir for 2 hours, allow to settle, decant the supernatant and collect on a medium porosity sintered glass funnel. Wash the solid well with H_2O , then with 95% EtOH (2 x) and Et_2O , dry it in air for 1 hour then in a vacuum desiccator (P_2O_5). The dry powder is then pulverised in a mortar and sieved (200 mesh) until all the solid has passed through (use a Camel's-hair brush to help the solid through) to give the catalyst as a light brown powder (21.0g) containing $\sim 0.14\text{g}$ of Pt per gram of solid. Store it in screw capped vials. It can be exposed to light and air without loss of activity. It gives very reproducible hydrogenation results and has been used analytically for the estimation of reducible groups [Vandenheuvel *Anal Chem* **28** 362 1956, DOI: 10.1021/ac60111a019]; and for the hydrogenation of methyl esters in EtOH without causing trans-esterification [Ackman & Burgher *J Lipid Res* **5** 130 1964, PMID: 14173321].

Other applications: Prior to the development of catalysts in which Pt is coordinated to a plethora of ligands in order to achieve particular specificities (e.g. Pt with phosphine ligands below), some specificities were obtained by manipulating the metal. Examples include a **platinum-tin chloride** catalyst made from H_2PtCl_6 and SnCl_2 for specific reduction of olefins [Cramer et al. *J Am Chem Soc* **85** 1691 1963, DOI: 10.1021/ja00894a035], **$\text{PtO}_2/\text{Pt black}+\text{FeCl}_2/\text{Zn}(\text{OAc})_2$** for reduction of unsaturated aldehydes to unsaturated alcohols, e.g. cinnamaldehyde to cinnamyl alcohol [Tulley & Adams *J Am Chem Soc* **47** 3061 1925, DOI: 10.1021/ja01689a036], and **PtS_2/C** which is less active than the oxide but is insensitive to poisons (e.g. sulfur compounds) and will reduce a nitro group without affecting halogen substituents, e.g. chloronitrobenzenes to chloroanilines [Dovell & Greenfield *J Am Chem Soc* **87** 2767 1965, DOI: 10.1021/ja01090a050].

Rhenium metal used in preparing the following catalysts has [7440-15-5] Re, M 186.2, m 3180°, b 5596°, d²⁰ 21.02.

Rhenium(II) oxide ($\text{ReO} \cdot 2\text{H}_2\text{O}$) [12143-03-2] $\text{ReO} \cdot 2\text{H}_2\text{O}$, M 238.2. This catalyst is prepared in a fume cupboard by the reduction of ammonium perrhenate with either Na/liqNH_3 , Li/liqNH_3 , or Li/EtNH_2 which give consistently $\text{ReO} \cdot 2\text{H}_2\text{O}$. In dried apparatus used typically for Na/liqNH_3 reduction, liquid NH_3 or EtNH_2 (50-75ml) and NH_4ReO_4 (1-2g, weighed) are stirred in a dry atmosphere until the salt dissolves; then Na or Li (15-30g) are added in chunks as rapidly as possible. The colourless solution becomes typically blue-black and is stirred until the colour disappears (15 minutes to 2 hours). The NH_3 or EtNH_2 are allowed to evaporate while the total volume is kept constant by addition of EtOH. The residue is transferred to a fritted glass thimble in a Soxhlet apparatus and extracted with dilute HCl ($\sim 2\text{N}$) for 24 hours (the pot acquiring a brown colour), then with EtOH for an equal time. When the brown acid extract, that decolorises aqueous KMnO_4 , is basified with aqueous NH_4OH black $\text{ReO} \cdot 2\text{H}_2\text{O}$ precipitates, and the supernatant now does not decolorise aqueous KMnO_4 . The solid is collected, washed with cold EtOH, and dried at 100° *in vacuo* over P_2O_5 (conditions under which it is stable and does not dehydrate) to provide the hydrated oxide (75-85%) as a very finely divided black powder. **Note** that any solid left in the thimble has the same composition as this oxide. It is stored as a suspension in EtOH which tends to settle rather readily, or as a suspension in distilled H_2O which requires more than 24 hours to settle out completely. Aliquots of suspensions (with or without evaporating to dryness) are used as the heterogeneous catalyst. The amount of rhenium can be determined as described by Broadbent and Selin [*J Org Chem* **28** 2343 1963, DOI: 10.1021/jo01044a045].

$\text{ReO} \cdot 2\text{H}_2\text{O}$ catalyses the hydrogenation of a wide variety of organic substrates in high yields with unusual selectivity, being particularly efficient for carboxylic acids. Hydrogenations are carried out in a rocking Parr reactor under pressures of ~ 200 atmospheres and temperatures varying from $\sim 20^\circ$ to $\sim 200^\circ$, and reduction times from 0.5 to ~ 10 hours. The catalyst prepared by the Li/liqNH_3 method is only marginally better than by the other methods. [Broadbent & Seegmiller *J Org Chem* **28** 2347 1963, DOI: 10.1021/jo01044a047.]

Rhenium(III) oxide (Re_2O_3) [12060-05-8] Re_2O_3 , M 420.5. Re_2O_3 is prepared from a mixture of ammonium

perrhenate (2.04g, 160mmol, NH_4ReO_4), a solution of 8*N* NH_4Ac (20ml, 160mmol) and glacial AcOH (2ml, 35mmol) in H_2O (100ml) in an ice-water bath to which is added dropwise, with vigorous stirring, an ice-cold solution of NaBH_4 (3.4g, 90mmol) in H_2O (100ml) over a 1 hour period. After half of the NaBH_4 solution is added, a further amount of AcOH (2ml, 35mmol) is added in one lot. Stirring is continued for 15 minutes and the solution is centrifuged at full speed in an ultracentrifuge, the supernatant is decanted off, the oxide is washed twice by suspension in H_2O and finally re-suspended in H_2O , and stored as such before use. The fine solid remains in suspension for 12-24 hours. Analysis indicated that reduction was ~85% complete giving a yield of 70-90% (~1.35g). The activity is checked by its ability to effect hydrogenation of styrene at room temperature in 6-10 hours. It catalyses a range of substrates including aliphatic olefins, maleic acid, benzene, nitrobenzene, bromonitrobenzene, naphthalene, pyridine, ketones and nitriles mostly in yields approaching 100% with (e.g. H_2O or EtOH), and without solvents. Typically the substrate (0.2mol, with or without solvent) and catalyst (0.2g of Re) in a 420ml rocking Parr bomb under hydrogen pressures up to 3000psi (204 atmospheres) are heated (40-50° increments) until the pressure starts to drop. Reductions occur in the temperature range of 25-250°. [Broadbent & Johnson *J Org Chem* **27** 4400 1962, DOI: 10.1021/jo01059a065.]

Rhenium(IV) oxide [12036-09-8] $\text{ReO}_2 \cdot 2.5\text{H}_2\text{O}$, **M 218.2**. A vigorously stirred mixture of a solution of ammonium perrhenate (1.2g) in H_2O (150ml) and an excess of granular zinc (5-10g, previously treated with a little dilute acid, to remove any surface oxide, and washed thoroughly to clean the surface) is treated dropwise over a period of several hours with 3*N* H_2SO_4 (64ml) at such a rate that black finely divided rhenium oxide is suspended into the solution (which is periodically decanted from the zinc); then the oxide is centrifuged off, the supernatant is returned to the reaction flask and addition of acid is resumed. The acid should not be added rapidly as the finely divided rhenium oxide coats the zinc and slows (or prevents) further reaction, and does not produce a Zn-free product. At the end of the reaction all the rhenium oxide residues from centrifugation are combined and washed with H_2O by decantation until the washings are free from sulfate. Conversion from perrhenate to rhenium (IV) oxide is >60%, and nearer 100% if an excess of Zn is maintained in the reaction by further addition of the metal. $\text{ReO}_2 \cdot 2.5\text{H}_2\text{O}$ analyses as a hydrate, is very stable, resists dehydration at 95° *in vacuo* over P_2O_5 , and loses H_2O only very slowly at 250°; however when the hydrated oxide is heated at 250° in $^*\text{C}_6\text{H}_6$ under hydrogenation conditions anhydrous ReO_2 can be isolated from the experiment. It exists in two crystallographic forms: *the α -form* is obtained when synthesised below 300° ($\text{Re} + 2\text{ReO}_3 = 3\text{ReO}_2$), and is irreversibly transformed into the *orthorhombic β -form* which is obtained when synthesised above 300° [Rogers et al. *Inorg Synth* **13** 135, 142-145 1972, DOI: 10.1002/9780470132449.ch27; Rogers et al. *Inorg Synth* **30** 96 1995, DOI: 10.1002/9780470132616.ch20].

$\text{ReO}_2 \cdot 2.5\text{H}_2\text{O}$ has catalytic activity for the hydrogenation of a variety of organic substrates, and although generally less active than Nickel or palladium catalysts (i.e. requires higher temperatures and/or pressures), it is however much more effective than these for the reduction of the carboxylic acid function. It has a potentially valuable selectivity, and very few other elements exhibit this characteristic to such a degree. [Broadbent & Selin *J Org Chem* **28** 2343 1963, DOI: 10.1021/jo01044a045.]

Rhenium(VI) oxide (ReO_3) [1314-28-9] ReO_3 , **M 234.2**, **b 750°**, **d²⁰ 6.9-7.4**. ReO_3 is prepared from the heptoxide (1.0g, see below) by adding it directly to *p*-dioxane (10-25ml) or tetrahydrofuran and warming gently to produce a colourless to dark green solution, and the excess solvent is removed *in vacuo* to give a sticky, tarry, black complex. The complex decomposes when carefully heated to *ca* 145° to give gaseous products and a pure, red, crystalline residue of ReO_3 (84-95%yield). This is finely ground and stored dry in a screw-cap vial. It forms non-hygroscopic, red/maroon-coloured crystals (with a green lustre) which are air stable below 100°, and disproportionates in a vacuum to rhenium(IV) and rhenium(VII) oxides at 400°. It is inert to H_2O , dilute alkali, and is thus more convenient to handle and store than the hygroscopic heptoxide which also has high catalytic activity. Analysis showed it to have the formula of the anhydrous oxide ReO_3 . It is oxidised by HNO_3 to HReO_4 . It has catalytic activity when used as such or when prepared *in situ* by using Re_2O_7 and allowing it to be reduced to ReO_3 during the hydrogenation process. This oxide, when used directly or prepared *in situ*, is the most effective hydrogenation catalyst of all the Re catalysts. It is very efficient in reducing carboxylic and carboxamide groups without reducing the aromatic ring, with the latter producing amines. Benzoic acid, ethyl benzoate, benzaldehyde, and *m*-nitrobenzaldehyde are reduced to the respective carbinols without hydrogenolysis to the toluenes as in most catalytic hydrogenations. Like the preceding rhenium oxide catalysts hydrogenations are carried out in shaking Parr-Bomb reactors under pressures of *ca* 200 atmospheres and temperatures ranging from ~100° to ~200° depending on when the substrates begin to absorb H_2 at a reasonable rate. [Broadbent & Bartley *J Org Chem* **28** 2345 1963, DOI: 10.1021/jo01044a046; Nechamkin et al. *Inorg*

Synth **3** 186 1950, DOI: 10.1002/9780470132340.ch49.] It is also obtained in quantitative yield from Re_2O_7 (purified by sublimation) in a CO atmosphere by heating at 175° until the oxide is blue, the temperature is raised slowly to 225° , then increased to 280° after the colour of the oxide has turned to red [Melaven et al. *Inorg Synth* **3** 188 1950, DOI: 10.1002/9780470132340.ch50].

Rhenium(VII) oxide (rhenium heptoxide) [1314-68-7] Re_2O_7 , **M 484.4**, **m 300.3°**, **b 360.3°**, **d²⁰ 6.103**. Dirhenium heptoxide is prepared by heating the metal or its lower oxides in a stream of oxygen or air at 400° to 425° . It forms canary-yellow crystals which are as deliquescent as P_2O_5 , and sublimes at $250^\circ/760\text{mm}$ and melts at 297° in a sealed tube [Ogawa *Bull Chem Soc Jpn* **7** 265 1932, DOI: 10.1246/bcsj.7.265]. Great care should be taken to avoid moisture during handling and storage. It is very soluble in H_2O to form perrhenic acid HReO_4 . It is also soluble in most organic solvents such as ethers, esters, alcohols, dioxane and pyridine. [Melaven et al. *Inorg Synth* **3** 188 1950, DOI: 10.1002/9780470132340.ch50.]

Reduction of Re_2O_7 during hydrogenation reactions generates **rhenium ‘blacks’** which mainly consist of ReO_3 and is among the best catalyst for reducing a variety of carboxylic acids to alcohols (at $\sim 160^\circ/\text{ca } 200$ atmospheres/few hours) in very good yields. The alcohols are occasionally accompanied with esters; unreduced acids rarely survive, amides are reduced to the respective amines and the usual functional groups that are reduced by Ni and Pt catalysts are also reduced with this rhenium catalyst. [Broadbent et al. *J Org Chem* **24** 1847 1959, DOI: 10.1021/jo01094a003.]

It is a precursor for organorhenium oxides which catalyse olefin oxidation, metathesis and other transformations. It is a **Lewis-acid metal oxide** which forms adducts of the formula $\text{Re}_2\text{O}_7 \cdot n\text{L}$ with a variety of O, N and S ligands such as 1,2-dimethoxyethane, 4,4'-di-*tert*-butyl-2,2'-bipyridine, and 1,4,7-trithiacyclononane respectively [Herrmann et al. *Inorg Chem* **34** 4701 1995, DOI: 10.1021/ic00123a001].

Rhenium(VII) sulfide (rhenium heptasulfide) [12038-67-4] Re_2S_7 , **M 596.9**. The heptasulfide is obtained by bubbling H_2S through a 3% solution of potassium perrhenate in 5-6 N HCl maintained at the boiling point until precipitation is complete. The suspension is allowed to stand overnight, then it is filtered through a sintered glass funnel, the solid is washed with 1:4 aqueous HCl, H_2O , dried for several days in a desiccator over CaCl_2 , powdered in a mortar and stored in a common screw capped vial. The catalyst is not adversely affected by occasional exposure to air, is prepared with high catalytic reproducibility, is insoluble in concentrated non-oxidising acids, is stable under all the conditions of hydrogenation used (1000-3600psi/ 25° to $335^\circ/0.5$ to 3 hours), and is not reduced to form H_2S . It is remarkably **resistant to ‘poisoning’**. It catalyses the reduction of a variety of organic substrates in high yields including sulfides, with desulfurisation [e.g. diphenylsulfide to benzene (94%), and thiophenol to a mixture of benzene (60%) and cyclohexane (40%)]; and without desulfurisation [e.g. allyl phenyl sulfide to propyl phenyl sulfide (100%), and of thiophene to thiophane (70%)]. [Broadbent et al. *J Am Chem Soc* **76** 1519 1954, DOI: 10.1021/ja01635a016.] When the heptasulfide is heated for 16 hours at $350\text{--}400^\circ$ in a tube furnace under a stream of CO_2 to blow out sulfur vapours, and cooled in a desiccator, it gives **rhenium disulfide** [12038-63-0]. The catalytic activity of ReS_2 parallels that of Re_2S_7 but to a lesser extent, and is more active than related molybdenum or cobalt catalysts. [Broadbent et al. *J Am Chem Soc* **76** 1519 1954, DOI: 10.1021/ja01635a016.]

Rhodium metal used in preparing the following catalysts has [[7440-16-6] Rh, **M 102.9**, **m 1966°**, **b 3727°**, **d²⁰ 12.41**].

Rhodium on alumina (5% Rh loading). $\text{Rh}-\text{Al}_2\text{O}_3$ has been used as a general catalyst for the complete reduction of the aromatic ring in phenols, e.g. gallic acid to *cis*-3,4,5-trihydroxycyclohexane carboxylic acid [Burgstahler & Bithos *Org Synth* **42** 62 1962, DOI: 10.15227/orgsyn.042.0062] requiring high pressures (2200psi) and temperatures ($\sim 100^\circ$) for 8-12 hours [see also Smith & Stump *J Am Chem Soc* **83** 2739 1961, DOI: 10.1021/ja01473a032; Kaye & Matthews *J Org Chem* **28** 325 1963, DOI: 10.1021/jo01037a012], the hydrogenation of 1-naphthol to *cis*-1-hydroxydecalin [6psi, 25° , 12 hours; Meyers et al. *J Org Chem* **29** 3427 1964, DOI: 10.1021/jo01034a520], resorcinol gives cyclohexane-1,3-dione [50psi, 25° , 16-18 hours; Sircar & Meyers *J Org Chem* **30** 3206 1965, DOI: 10.1021/jo01020a506], D-mandelic acid gives pure D-hexahydromandelic acid but L-mandelic acid gives (\pm)-hexahydromandelic acid [3-4 atmospheres, 25° , 1.5 hours, AcOH is mandatory; Stocker *J Org Chem* **27** 2288 1962, DOI: 10.1021/jo01053a077], reduction of vinylic and allylic double bonds [500psi, 100° , 15 minutes; Ham & Coker *J Org Chem* **29** 194 1964, DOI: 10.1021/jo01024a045], reduction of oximes or aliphatic nitriles to primary amines [1-2 atmospheres, $\sim 25^\circ$, 1-2 hours; Freifelder et al. *J Org Chem* **27** 2209 1962, DOI: 10.1021/jo01053a504; Freifelder *J Am Chem Soc*

82 2386 1960, DOI: 10.1021/ja01494a067], hydrogenolysis of ketals to the corresponding dialkyl ethers [Howard & Brown *J Org Chem* **26** 1026 1961, DOI: 10.1021/jo01063a010], hydrogenation of heterocyclic compounds e.g. 1,2,3,4-tetrahydroisoquinoline-3-carboxylic acid to *cis*-decahydroisoquinoline-3-carboxylic acid [40psi, ~25°, 3 hours; Rapala et al. *J Am Chem Soc* **79** 3770 1957, DOI: 10.1021/ja01571a042], nicotinic acid to hexahydronicotinic (nipecotinic) acid without decarboxylation [2 atmospheres, ~25°, 4 hours; Freifelder *J Org Chem* **27** 4046 1962, DOI: 10.1021/jo01058a501; Freifelder *J Org Chem* **28** 602, 1135 1963], complete hydrogenation of quinoxaline [Broadbent et al. *J Am Chem Soc* **82** 189 1960, DOI: 10.1021/ja01486a043], reduction of the 4,5-double bond of pyrimidine nucleosides and nucleotides [Cohn & Doherty *J Am Chem Soc* **78** 2863 1956, DOI: 10.1021/ja01593a063], dehydrogenation e.g. hydrogen transfer from hexahydrohexahelicene to benzene in order to provide hexahelicene and cyclohexane in 73% yield [Newman & Lednicer *J Am Chem Soc* **78** 4765 1956, DOI: 10.1021/ja01599a060; cf. also Anderson & Anderson *J Org Chem* **22** 1197 1957, DOI: 10.1021/jo01361a018], and hydrogenation of anilines to cyclohexylamine with little hydrogenolysis [Freifelder et al. *J Org Chem* **30** 2485 1965, DOI: 10.1021/jo01018a524].

Rhodium on carbon (5% and 10% loading), Rh/C. A 10%-Rh, 0.1%Pd on carbon catalyst has been prepared from a mixture of $\text{RhCl}_3 \cdot 3\text{H}_2\text{O}$ (5.26g), PdCl_2 (0.34g) and carbon (18g, e.g. Darco G-60, or any acid washed charcoal can be used) in H_2O (200ml) is rapidly stirred and heated to 80°. Lithium hydroxide hydrate (2.7g) dissolved in H_2O (10ml) is added in one lot and heating is stopped but the mixture is stirred overnight, filtered, and washed with 0.5% v/v aqueous AcOH (100ml) then dried *in vacuo* at 65° to give 20.6-21g of catalyst. One gram of this catalyst will absorb 0.0022-0.0028 mole of H_2 in aqueous suspension. It has been used, for example, in the reduction of *p*-aminobenzoic acid to a mixture of *cis*- and *trans*- 4-aminocyclohexanecarboxylic acid [at 50psi/~25°/~24 hours] which was converted to 2-azabicyclo[2.2.2]octan-3-one (3-isoquinuclidone) in 81-84% yield. *The point was made that when fresh catalyst is added to the reaction vessel, it should be moistened with the solvent; and that hydrogen should be removed by evacuation at the end of the reduction in order to avoid having a hydrogen-air atmosphere that could be explosive.* [Pearlman *Org Synth Coll Vol* **5** 670 1973, DOI: 10.15227/orgsyn.049.0075; Pearlman *Tetrahedron Lett* **8** 1663 1967, DOI: 10.1016/S0040-4039(00)70335-2.] It is a useful catalyst (even without the trace of Pd) for the reduction of nitriles to primary amines, and the hydrogenation of aromatic and pyridine rings, aldehydes and sugars. It is superior to Rh- Al_2O_3 for reduction of the pyridine ring and larger amounts of catalyst have to be added to overcome the poisoning effect of the piperidine products [Freifelder et al. *J Org Chem* **27** 284 1962, DOI: 10.1021/jo01048a505]. It is useful for the stepwise reduction of 3-*H*-pyrrolizine to 3,4-dihydropyrrolizine, and then to the fully reduced pyrrolizidine [Schweizer & Light *J Am Chem Soc* **86** 2963 1964, DOI: 10.1021/ja01068a059]; and in boiling norbornadiene complete conversion to a mixture of dimers and trimers occurs [Mrowca & Katz *J Am Chem Soc* **88** 4012 1966, DOI: 10.1021/ja00969a021]. A comparison of the rates of hydrogenation with 5% Pd/C, 5% Pt/C, 5% Rh/C and 5% Ru/C catalysts towards reduction of C=C, C=O, C-OH and benzene rings in various solvents at 25° and initial H_2 pressure of 1 atmosphere were made by Breitner, Roginski & Rylander [*J Org Chem* **24** 1855 1959, DOI: 10.1021/jo01094a004].

Ruthenium metal used in preparing the following catalysts has [7440-18-8] Ru, M 101.1, m 2310°, b 3900°, d²⁰ 12.5].

Ruthenium-aluminium oxide/hydroxide catalyst (~2.5wt% Ru loading). The preparation of this catalyst is described in Chapter 7 because it consists of Ru(0) nanoparticles encapsulated in an aluminium oxy-hydroxide matrix and is superior as a catalyst to the commercially available material. It is a recyclable catalyst for the efficient oxidant-free alcohol dehydrogenation (compare with $\text{RuCl}_3 \cdot 3\text{H}_2\text{O}$ or xH_2O above) of a wide range of alcohols including benzylic alcohols, secondary long chain fatty alcohols, secondary allylic alcohols, 3-pyridylmethanol, 2-thiophenylmethanol and 4-methylthiobenzyl alcohol in dry toluene to the respective carbonyl compounds under argon at dehydrogenation temperatures of 80° to 110°, using 3-6 mol% of Ru, in yields of >90%. It is 5 times superior to the commercial Ru/ Al_2O_3 catalyst and is re-usable, after washing with Me_2CO and drying in air, at least 10 times without loss of activity. [Kim et al. *Org Lett* **8** 2543 2006, DOI: 10.1021/ol060750z.]

Ruthenium on alumina (5% Ru/ Al_2O_3 reduced loading). Ru/ Al_2O_3 can be freshly prepared by treating alumina powder with a ruthenium compound, e.g. $\text{RuCl}_3 \cdot \text{xH}_2\text{O}$ in H_2O , $\text{Ru}(\text{acac})_3$ or $\text{Ru}_3(\text{CO})_{12}$ in *benzene (to give a solid that is ~5wt% in Ru), followed by evaporating the solvent and reducing the solid in a stream of H_2 at 500°. This catalyst will oxidise dibenzothiophene or dibenzothiophene oxide (Ru molar ratio 11:1) in * C_6H_6

to dibenzothiophene sulfone at 100° in air (70 atmospheres) for 12 hours in 97% yield. **Slow poisoning** of Ru occurs during the oxidation. The activity of the catalyst depends on the preparation, but is much more active than the following catalysts in the sequence $\text{Ru}_3(\text{CO})_{12} > \text{Ru}(\text{acac})_3 > \text{RuCl}_3 \cdot x\text{H}_2\text{O}$. Ru/C or Ru/SiO₂, and the commercially available Ru/Al₂O₃ are less active than the above preparation. Pt/Al₂O₃ or Ir/Al₂O₃ gives only a trace of the sulfone, and Pd/Al₂O₃, Rh/Al₂O₃, Os/Al₂O₃ or Cu/Al₂O₃ are totally inactive. It should be noted that with this catalyst oxidation of *n*-Bu₂S (0.2M in *C₆H₆; in the ratio of Ru/sulfide 21:1) provides 44 and 9% molar yields of *n*-Bu₂SO and *n*-Bu₂SO₂ respectively, with full recovery of unreacted *n*-Bu₂S. [Ledlie & Howell *Tetrahedron Lett* **17** 785 1976, DOI:10.1016/S0040-4039(00)77951-2.]

Ruthenium on carbon (5% Ru/C loading). A comparison of the rates of hydrogenation with 5% Pd/C, 5% Pt/C, 5% Rh/C and 5% Ru/C catalysts towards reduction of C=C, C=O, C-OH and benzene rings in various solvents at 25° and initial H₂ pressure of 1 atmosphere were made by Breitner, Roginski & Rylander [*J Org Chem* **24** 1855 1959, DOI: 10.1021/jo01094a004.] Ru/C is a superior catalyst than Pd/C and Pd/C for the hydrogenation of ketones in basic and neutral media and in some cases (e.g. tetramethylcyclobutan-1,3-dione at 1000-1500psi of H₂/125°/1 hour) >90% yields of diols were obtained [Hasek et al. *J Org Chem* **27** 700 1961, DOI: 10.1021/jo01062a013], C=C (with some selectivity), and acetylenes were reduced to alkanes [Berkowitz & Rylander *J Org Chem* **24** 708 1959, DOI: 10.1021/jo01087a610], nitrobenzene was reduced to hydrazobenzene in alcoholic KOH in 80% yield without over-reduction to aniline [Peitra & Res *Ann Chim (Rome)* **48** 299 1958], and 10% Ru/C was better than other catalysts for reducing phenolic rings in polycyclic compounds [Walton et al. *J Am Chem Soc* **78** 4760 1956, DOI: 10.1021/ja01599a059].

Ruthenium(IV) oxide [12036-10-1] RuO₂, M 133.1, d²⁵ 6.97. The dioxide is prepared by heating Ru metal in a silica boat in a silica combustion tube placed horizontally in a furnace, and a slow stream of dry oxygen is allowed to flow over it while it is being heated at 1000° for 24 hours. During heating a little of the oxide volatilises and small crystals may condense on the walls at the downstream end of the tube. A 90-95% yield of polycrystalline RuO₂ remains in the boat. It forms tetragonal dark blue crystals which are insoluble in organic solvents, H₂O and acids but soluble in fused alkali. [Schäfer & Heitland *Z Anorg Allgem Chem* **304** 249 1960, DOI: 10.1002/zaac.19603040502; Schäfer et al. *Z Anorg Allgem Chem* **319** 327 1963, DOI: 10.1002/zaac.19633190514; Rogers et al. *Inorg Synth* **13** 135 1972, DOI: 10.1002/9780470132449.ch27; **30** 96 1995, DOI: 10.1002/9780470132616.ch20.] It is a good catalyst, no doubt by being converted into the metal *in situ* which promotes catalysis. It has been used for the reduction of aromatic rings to cyclohexane rings [e.g. at 1350psi/147°/16 hours, or 70psi/90°/1 hour; Freifelder & Stone *J Am Chem Soc* **80** 5270 1958, DOI: 10.1021/ja01552a063; Freifelder & Stone *J Org Chem* **26** 3805 1961, DOI: 10.1021/jo01068a043; Freifelder & Stone *J Org Chem* **27** 3568 1962, DOI: 10.1021/jo01057a041; Johnson et al. *J Am Chem Soc* **84** 2181 1962, DOI: 10.1021/ja00870a033], and phenolic compounds to the respective cyclohexanols [Rapala & Farkas *J Org Chem* **23** 1404 1958, DOI: 10.1021/jo01103a628; Counsell *Tetrahedron* **15** 202 1961, DOI: 10.1016/0040-4020(61)80027-6; Ireland & Schiess *J Org Chem* **28** 6 1963, DOI: 10.1021/jo01036a002]; but with some enones only the double bond was reduced without affecting the carbonyl function [Rapala & Farkas *J Am Chem Soc* **80** 1008 1958, DOI: 10.1021/ja01537a074].

HOMOGENEOUS METAL CATALYSTS

Allylpalladium(II) chloride dimer [di-μ-chloro-η³-allylpalladium(II)] [12012-95-2] [Pd(η³-C₃H₅)Cl]₂, M 365.9, m 155-156°(dec.), ~160°(dec.). It is prepared from allyl chloride, PdCl₂, and NaCl in the presence of carbon monoxide and is extracted into CHCl₃, washed with H₂O, dried (CaCl₂), filtered and evaporated to dryness to give the complex as yellow crystals (use a very efficient fume cupboard). These can be used as such, but by recrystallisation from a mixture of CH₂Cl₂ and hexane analytically pure catalyst can be obtained (**m 155-156° dec.**). It also crystallises from benzene and is soluble in MeOH, Et₂O, Me₂CO and CHCl₃. The ¹H NMR (CDCl₃) has two doublets at δ_H 3.03 (anti CH₂, *J* = 12.0Hz) and 4.10 (syn CH₂, *J* = 7.1Hz), and a triplet at δ_H 5.48 (CH) 2:2:1. [Tatsuno et al. *Inorg Synth* **28** 342 1990, DOI: 10.1002/9780470132616.ch20; Shaw *Proc Chem Soc* 247 1960, **notes start on p 233**, DOI: 10.1039/PS9600000233; Hüttel et al. *Chem Ber* **94** 766 1961, DOI: 10.1002/cber.19610940329; Dent et al. *J Chem Soc* 1585 1964, DOI: 10.1039/JR9640001585.]

It is a labile Pd complex, but is very useful for preparing a variety of stable catalytic Pd(0) complexes with a wide selection of ligands to perform stereoselective reactions. A particularly useful ligand is *cis,cis,cis*-1,2,3,4-tetrakis(diphenylphosphinomethyl)cyclopentane [Tedicyp] [Laurenti et al. *J Org Chem* **66** 1633 2001, DOI: 10.1021/jo001146j]. The catalyst with this ligand is typically prepared under argon, in a Schlenk type system,

by dissolving $[\text{Pd}(\eta^3\text{-C}_3\text{H}_5)\text{Cl}]_2$ (4.2mg, 11.6 μmol) and Tedicyp (20mg, 23.2 μmol , see [333380-86-2] in Part 2) in dry tetrahydrofuran (10ml) and stirring for 10 minutes before use. It gives a solution of 2.32 $\mu\text{mol}/\text{ml}$ of catalyst, which has ^{31}P NMR (162MHz, CDCl_3) with δ at 25 (vbrs, *width* 80Hz), 19.4 (vbrs, *width* 110Hz). It is very efficient in catalysing allylic substitutions with high turnover numbers (up to 9,800,000) and frequencies (up to 190,000 h^{-1}).

$[\text{Pd}(\eta^3\text{-C}_3\text{H}_5)\text{Cl}]_2$ is used for cross-coupling reactions [Wallow & Novak *J Org Chem* **59** 5034 1994, DOI: 10.1021/jo00096a056], the Heck reactions with olefins [Fall et al. *Synthesis* 1683 2007, DOI: 10.1055/s-2007-966063] and enol ethers [Battace et al. *Eur J Org Chem* 3122 2007, DOI: 10.1002/ejoc.200700152], and the formation of ethyl cyclopropane carboxylates with ethyl diazoacetate and olefins [Armstrong *J Org Chem* **31** 618 1966, DOI: 10.1021/jo01340a522] to name a few.

Bis(acetonitrile)dichloropalladium(II) [14592-56-4] $\text{Pd(II)Cl}_2(\text{MeCN})_2$, **M 259.4, dec > 200°**. It is prepared by mixing anhydrous PdCl_2 (1mol) and MeCN (2mols) until clear and precipitating the complex with petroleum ether. Alternatively, MeCN (3ml) is added to a solution of K_2PdCl_4 (2g) in H_2O (40ml) and stirred at $\sim 25^\circ$ until separation of yellow prisms is complete (could take up to 8 days). Filter these off, wash them with H_2O , EtOH, Et_2O , and dry them *in vacuo*. Further purification, if necessary, can be achieved by dissolving it in anhydrous solvents (which are not likely to displace the ligands) such as MeOH, toluene, THF, precipitating it with light petroleum, and drying the solid *in vacuo*. It is also soluble in $^*\text{C}_6\text{H}_6$ to the extent of 100mg/4.0ml. In addition to its use as a source of metal for palladium complexes and catalysts such as $\text{PdCl}_2(\text{R})\text{-BINAP}$ [115826-95-4] (see Ozawa et al. *Organometallics* **12** 4188 1993, DOI: 10.1021/om00034a064), it also catalyses Overman rearrangements (aza-Claisen rearrangement, Overman & Garpenter *Org React* **66** 1 2005, DOI: 10.1002/0471264180.or066.01; see below). A particularly interesting example of the rearrangement being the high diastereoselectivity which is directed by a δ -methoxymethylether group in δ -substituted allylic trichloroacetamides, where the ether oxygen atoms coordinate with Pd to direct the orientation of the reaction [Jamieson & Sutherland *Tetrahedron* **63** 2123 2007, DOI: 10.1016/j.tet.2006.12.067]. The IR spectrum shows, from the RCN and M-X vibrations, that the MeCN groups have a *trans*-planar configuration, unlike related Pt complexes which have the *cis*-configuration. For IR see reference (Nujol) has ν_{max} at 1417m, 1358m, $\sim 1160\text{wbr}$, 1030m, 970wbr cm^{-1} [Walton *Spectrochim Acta* **21** 1795 1965, DOI: 10.1016/0371-1951(65)80091-1].

Bis(benzonitrile)dichloropalladium(II) [14220-64-5] $\text{Pd(II)Cl}_2(\text{PhCN})_2$, **M 383.6, m 131°**. It is prepared and purified in the same manner as the $\text{Pd(II)Cl}_2(\text{MeCN})_2$ complex above. Alternatively, anhydrous PdCl_2 (2g) suspended in benzonitrile (50ml) is stirred at 100° . Most of the PdCl_2 dissolves after 20 minutes to give a red solution which is filtered while hot, and the hot filtrate is poured slowly into petroleum ether (b 40-60°). The light yellow complex that crystallises out is collected, washed with petroleum ether and dried *in vacuo* (4.0g, 93%). It can be purified further by dissolving it in the minimum volume of $^*\text{C}_6\text{H}_6$, and the hot red solution is filtered and diluted with petroleum ether to allow the complex to crystallise out. [Doyle et al. *Inorg Synth* **6** 216 1960, DOI: 10.1002/9780470132371.ch69; Kharasch et al. *J Am Chem Soc* **60** 882 1938, DOI: 10.1021/ja01271a035.] $\text{Pd(II)Cl}_2(\text{PhCN})_2$ is an effective catalyst for the regiospecific cycloaddition cross-coupling reactions of aziridines with carbodiimides to form imidazolidin-2-imines in 40-94% yields [Baeg & Alper *J Org Chem* **57** 157 1992, DOI: 10.1021/jo00027a030]. In the presence of a **Buchwald ligand** (e.g. 2-di-*t*-butyl-2'4'6'-tri-*i*-propylbiphenyl) in CH_2Cl_2 , it catalyses the stereoselective formation of α -O-glycosides from protected glycalimides with a variety of phenols or protected glycosides in high yields at room temperature [Schuff, Mercer and Nguyen *Org Lett* **9** 3173 2007, DOI: 10.1021/ol071268z]. Also, in the presence of Xantphos {4,5-bis(diphenylphospheno)-9,9-dimethylxanthene, see [161265-03-8]}, it promotes Stille-type cross coupling reactions between alkynyl- SnBu_3 and α -halo esters or amides under neutral conditions in THF to provide 3-alkynoates or alkynoamides in high yields without homocoupling of the alkynyltin reagents [Shi et al. *Chem Commun* 2342 2007, DOI: 10.1039/B703221H]. The IR spectrum shows, from the RCN and M-X vibrations, that the PhCN groups have a *trans*-planar configuration, unlike related Pt complexes which have the *cis*-configuration. The IR (Nujol) has ν_{max} at 3096w, 3057w, 3033w, 3002vwm 2287s, 1480m-w, 1440s, 760s, 678s cm^{-1} [Walton *Spectrochim Acta* **21** 1795 1965, DOI: 10.1016/0371-1951(65)80091-1].

Bis(cyclopentadienyl)cobalt(II) (cobaltocene, Cp_2Co) [1277-43-6] $\text{C}_5\text{H}_5\text{CoC}_5\text{H}_5$, **M 189.1, m 173-174° (under N_2), 176-180°(dec)**. Cp_2Co is prepared by adding anhydrous CoCl_2 (0.5 mol, prepared by dehydrating the salt at 200° in a vacuum) to a cold solution of 1 mol of sodium cyclopentadienide [4984-82-1] in THF (see

below), stirring for 2 hours, removing the solvent and subliming the product directly from the mixture at 150-200°/10⁻⁴ mm in 75-80% yields; or by using CoBr₂ and refluxing for 3 hours. *Alternatively*, (method of Wilkinson et al. *J Inorg Nucl Chem* **2** 95 1956, DOI: 10.1016/0022-1902(56)80004-3), cyclopentadiene (42ml, 0.5mol) and Et₂NH (~100ml, 1mol) are added slowly to CoBr₂ (0.25mol) with cooling, and then stirring at ~25° for 6-8 hours; the solvent is removed *in vacuo* and the residue is extracted with petroleum ether, filtered, and the solvent is evaporated to give dark violet monoclinic crystals m 171-173°.

When anhydrous **cobalticenium chloride** (5g, prepared from sodiocyclopentadienide and anhydrous CoCl₂ in THF as above but stirring at the boiling point for 2 hours, evaporating the solvent and adding 1:1 hydrochloric acid) in dry 1,2-dimethoxyethane (150ml) is treated with excess of NaBH₄ (5g, or LAH) in small portions while stirring (H₂ is evolved), then warm to 60°, or add H₂O (2ml), the mixture becomes deep wine-red in colour. After 1 hour the solution is filtered, the solid is washed with Et₂O and the filtrate is evaporated to a few mls *in vacuo*, diluted with petroleum ether (b ~40-60°), concentrated again and applied on to an alumina column (20 x 2cm). The first red band that elutes with petroleum ether is evaporated, and the residue is sublimed *in vacuo* onto a cold finger at -70° to give wine-red crystals of **π -cyclopentadienyl(cyclopentadiene)cobalt [π -C₅H₅CoC₅H₆]** (yield 80%) with **m 98-99°**, has **M 193** (ebulioscopic in *C₆H₆), and a UV with λ_{\max} nm(ϵ) at 263.8 (1.78 x 10⁵), 327 (9.27 x 10³), and 395 (5.27 x 10³). It has a **camphorous odour** and should be kept under argon as it is stable in air for only a few minutes, but for a few hours in degassed solutions of organic solvents. [Green et al. *J Chem Soc* 3753 1959, DOI: 10.1039/JR9590003753.]

Sodium cyclopentadienide [4984-82-1] **C₅H₅, C₅H₅Na, M 88.1**, is prepared under N₂ by slowly adding cyclopentadiene (37ml) to a cooled suspension of sodium (10g) in THF (150ml) when H₂ evolves rapidly and a solution of sodium cyclopentadienide is formed. The colour of the solution depends on the purity of the solvent and the amount of air in the system. Clear pale orange or red solutions are obtained with good conditions, but dark red to purple colours are obtained in the presence of traces of air without, however, seriously decreasing the yields of the complexes formed. [Wilkinson et al. *J Inorg Nucl Chem* **2** 95 1956, DOI: 10.1016/0022-1902(56)80004-3; Reynolds & Wilkinson *J Inorg Nuclear Chem* **9** 86 1959, DOI: 10.1016/0022-1902(59)80015-4; Beilstein **16** IV 1698.] It is available commercially as a 2.0M solution in THF as sodium cyclopentadienylide. The dry solid is **pyrophoric**.

Cp₂Co catalyses the condensation of acetylene and mono-substituted acetylenes (but not di-substituted acetylenes) with nitriles at 150° to give 30-70% yields of substituted pyridines [Wakatsuki & Yamazaki *Synthesis* 26 1976, DOI: 10.1055/s-1976-23943]. This synthesis has been applied successfully in a 2+2+2 cycloaddition reaction between di(trimethylsilyl)propargyl ether and acetonitrile to prepare a pyridine intermediate which was used to obtain pyridoxine (vitamin B₆) [Geiger et al. *Helv Chim Acta* **67** 1274 1984, DOI: 10.1002/hlca.19840670513].

The above are general procedures and they have been used for preparing **Bis(cyclopentadienyl)chromium (II) [Cp₂Cr]** [1271-24-5] **C₅H₅CrC₅H₅, M 182.2, m 168-170°** [Wilkinson et al. *J Inorg Nucl Chem* **2** 95 1956, DOI: 10.1016/0022-1902(56)80004-3; Beilstein **16** IV 1774], **Bis(cyclopentadienyl)magnesium(II) [Cp₂Mg]** [1284-72-6] **C₅H₅MgC₅H₅, M 154.5, m 176-178°, 180°(dec)** [Wilkinson et al. *J Inorg Nucl Chem* **2** 95 1956, DOI: 10.1016/0022-1902(56)80004-3; Barber *J Inorg Nucl Chem* **4** 373 1957, DOI: 10.1016/0022-1902(57)80026-8; Beilstein **16** IV 1695], and **Bis(cyclopentadienyl)vanadium(II) [Cp₂V]** [1277-47-0] **C₅H₅VC₅H₅, M 181.1, m 165-167°** [Wilkinson et al. *J Inorg Nucl Chem* **2** 95 1956 DOI: 10.1016/0022-1902(56)80004-3; Beilstein **16** IV 1771].

Bis(dibenzylideneacetone)palladium(0) [Pd(dba)₂] [32005-36-0] **C₃₄H₂₈O₂Pd, M 575.3**. The precise nature of this complex is not clear, but a product with reproducible stoichiometry [C₃₄H₂₈O₂Pd] is prepared first by stirring PdCl₂ (8.87g, 50mmol) and NaCl (2.92g, 50mmol) in MeOH (250ml) at 25° for 16 hours, filtering through a cotton wool plug, diluting the filtrate to ~1.5L with MeOH, and heating to 60° to form Na₂[PdCl₆]. Dibenzylideneacetone (36.5g, 150mmol, see [538-58-9], cf. 'Organic Compounds' in Chapter 3) is then added to this warm solution stirred solution of Na₂[PdCl₆], and stirring for a further 15 minutes and anhydrous NaOAc (75g) is added. The reaction begins immediately, heating is stopped but stirring is continued until the reaction cools to room temperature. The dark brown precipitate that is formed is washed with MeOH (2 x 25ml), H₂O (5 x 50ml), Me₂CO (5 x 15ml) and dried in air to constant weight (923g, 80% yield) [Rettig et al. *Inorg Synth* **17** 134 1977, DOI: 10.1002/9780470132487.ch37]. See the related complexes of **Pd(dm-dba)₂** below for their structures in CDCl₃ solution. Catalysis with these complexes occurs at room temperature in the

presence the ligand DIPHOS [dpe, 1,2-bis-[diphenylphospheno]ethane (see [1663-45-2] in ‘Metal-Organic Compounds’ in Chapter 4) which is more efficient than with the less stable $\text{Pd}(\text{PPh}_3)_4$. It is used for the synthesis of allylic substituted cyclopentadienes [Fiaud & Malleron *Tetrahedron Lett* **21** 4437 1980, DOI: 10.1016/S0040-4039(00)92193-2] and is a homogeneous catalyst [Fiaud & Malleron *Tetrahedron Lett* **22** 1399 1981, DOI: 10.1016/S0040-4039(01)90332-6; Black *Aldrichimica Acta* **15** 13 1982] which catalyses the alkylation of a variety of nucleophiles under mild conditions [Ferroud et al. *Tetrahedron Lett* **25** 4379 1984, DOI: 10.1016/S0040-4039(01)81443-X.] Together with cyclic thiourea ligands such as *1N,3N*-bis(1,4-di-*tert*-butylphen-2-yl)imidazoline-2-thione, it catalyses efficient aerobic oxidations of alcohols to aldehydes and ketones in toluene in yields >95% [Yang et al. *Synlett* 3057 2006, DOI: 10.1055/s-2006-951501.]

Bis(3,5,3',5'-dimethoxybenzylideneacetone)palladium(0) {bis[1,3-bis(3',5'-methoxybenzylidene)acetone palladium(0), $[\text{Pd}(\text{dm-dba})_2]$ } [811862-77-8] $\text{C}_{42}\text{H}_{44}\text{O}_{10}\text{Pd}$, **M 815.2, m 170-178°**. This Pd catalyst with the dimethoxy-dba ligand is prepared and purified as for $\text{Pd}(\text{dba})_2$ above but using 1,5-bis(3',5'-dimethoxyphenyl)penta-1*E,4E*-diene-3-one (dm-dba, [39777-58-7]),* instead of dba. Alternatively, Fairland's method can be used, whereby NaCl (3.3g, 56mmol) is added to PdCl_2 (5g, 28mmol) in MeOH (140ml) and stirred under argon at 25° for 24 hours, filtered and evaporated *in vacuo* to half its volume. To this solution at 60° is added dm-dba (20.6g, 88mmol), stirred at 60° for 15 minutes, anhydrous NaOAc (42.3g) is added and allowed to cool to 25°. The mixture is stirred at 25° for 2 hours as the dark red precipitate of the complex separates. The solid is collected, washed with MeOH (2 x 100ml), H_2O (2 x 100ml) then Me_2CO (2 x 20ml), partially dried by suction, placed in a Schlenk flask and stirred under a flow of N_2 overnight to give the complex as a maroon/purple microcrystalline solid (11g 68%). It was found to have higher activity than those of the complex with the unsubstituted ligand [i.e. $\text{Pd}(\text{dba})_2$] in Suzuki-Miyaura cross-coupling reactions of organic halides with arylboronic acids. **Note** that a solution of the complex, as in the case of $\text{Pd}(\text{dba})_2$ above, in CDCl_3 , was found to contain one dm-dba ligand coordinated to Pd and one free in solution. Addition of two equivalents of Ph_3P to the complex in THF and $(\text{CD}_3)_2\text{CO}$ resulted in the formation of the η^2 -(dm-dba) $\text{Pd}^{(0)}(\text{Ph}_3\text{P})_2$ complex as shown by the presence of two phosphorous doublets in the ^{31}P NMR spectrum. [Fairlamb et al. *Org Lett* **6** 4435 2004, DOI: 10.1021/ol048413i; cf. *Handbook of Organopalladium for Organic Synthesis* Negishi ed. Wiley, Hoboken NJ 2002, ISBN 0-471-31506-0.] For $\text{Pd}_2(p,p'$ -dimethoxy-dba) $_3$ see $\text{Pd}_2(\text{dba})_3(\text{CDCl}_3)$ below.

Bis(di-*tert*-butyl(4-dimethylaminophenyl)phosphine)dichloropalladium(II) [(A-*ta*Phos) $_2\text{PdCl}_2$] [887919-35-9] $\text{C}_{32}\text{H}_{56}\text{Cl}_2\text{N}_2\text{P}_2\text{Pd}$, **M 708.1**. The catalyst is prepared by mixing one equivalent of di-*tert*-butylphosphine, 1 equivalent of 4-dimethylaminophenyl bromide and 1 mol% of Pd_2dba_3 at 90° for 12 hours. The mixture is cooled to 25°, filtered through a short silica gel plug, the plug is rinsed with toluene and the combined filtrate is evaporated under vacuum. The residue is dissolved in tetrahydrofuran and 0.35 equivalents of solid $\text{PdCl}_2(\text{COD})$ are added followed by stirring at 25° for 12 hours. The product is filtered off, and the yellow solid collected is washed with pentane and dried *in vacuo* to give ~85% yield of (A-*ta*Phos) $_2\text{PdCl}_2$ which provides the correct microanalysis for C and H. It has ^1H NMR (CD_2Cl_2) with δ at 7.5 (s br, 4 H, ArH), 6.42 (d, $J = 8.3\text{Hz}$, 4H, ArH), 2.75 (s, 12H, NMe $_2$), 1.38 (t, $J_{\text{PH}} = 6.6\text{Hz}$, 36H, *tert*-Bu); and ^{31}P NMR (CDCl_3) with δ at 52.1. Similarly **bis[di-*tert*-butyl(*p*-trifluoromethylphenyl)phosphine)dichloropalladium(II)** can be prepared which has ^1H NMR (CDCl_3) with δ at 8.0 (s br, 4 H, ArH), 7.59 (d, $J = 7.9\text{Hz}$, 4H, ArH), 1.62 (t, $J_{\text{PH}} = 7.0\text{Hz}$, 36H, *tert*-Bu); ^{31}P NMR (CDCl_3) with δ at 55.7; and ^{19}F NMR (CDCl_3) with δ at -63.4. Also **bis[di-*tert*-butyl[phenyl]phosphine)dichloro-palladium (II)** was prepared in the same way and has ^1H NMR (CDCl_3) with δ at 7.9-7.8 (m, 4 H, ArH), 7.36-7.30 (m, 6H, ArH), 1.56 (t, $J_{\text{PH}} = 6.9\text{Hz}$, 36H, *tert*-Bu); ^{31}P NMR (CDCl_3) with δ at 54.4. These are **air-stable active catalysts** for Suzuki-Miyaura cross-coupling with aryl halides including 5- and 6-membered heteroaryl halides in the presence of bases such as K_2CO_3 , Na_2CO_3 , KOAc, K_3PO_4 with high turn over numbers (TON) of 100 to 100,000 [Guram et al. *Org Lett* **8** 1787 2006, DOI: 10.1021/ol060268g].

2-[Bis(2,4-di-*tert*-butylphenoxy)phosphinoxy]-3,5-di(*tert*-butyl)phenyl-palladium(II) chloride dimer {Bedford's Catalyst, 2[bis(2,4-di-*tert*-butylphenoxy)phosphino- κP -oxy)-3,5-di-*tert*-butylphenyl- κC]di- μ -chloro-dipalladium} [217189-40-7] $\text{C}_{84}\text{H}_{124}\text{Cl}_2\text{O}_6\text{P}_2\text{Pd}_2$, **M 1575.6**. The thoroughly characterised Bedford catalyst, an *ortho*-metallated dimer, is prepared in 96% yield from the bulky tris(2,4-di-*tert*-butylphenyl)phosphite and PdCl_2 . The catalyst is remarkably stable to moisture and air, showing no signs of deterioration after several weeks, can be kept in air for at least 6 months and is not decomposed on heating at 130° in toluene for 24 hours. It is very efficient for coupling aryl halides with either arylboronic acids (**Suzuki**

reaction) or aryltin compounds (*Stille reaction*) in solvents such as toluene or tetrahydrofuran to form biaryls with very high turn over numbers (TONs up to 1,000,000 moles of product/mol of Pd) at even very low Pd catalyst concentrations, and in moderate to high yields [Albisson et al. *JCS Chem Commun* 2095 1998, DOI: 10.1039/A806041J]. *Bedford's catalyst* also proved to be extremely active in the arylation (with e.g. aryl bromides) of alkenes (e.g. acrylate, styrene) with turn over numbers of up to 5,750,000 (mole product/mol Pd), and turn over frequencies of 300,000 (mol product/mol Pd/hour) [Albisson et al. *Tetrahedron Lett* **39** 9793 1998, DOI: 10.1016/S0040-4039(98)02175-3].

***R,R*-(+) and *S,S*-(-) *N,N'*-Bis(3,5-di-*tert*-butylsalicylidene)-1,2-*trans*-cyclohexanediamine Manganese(III) Chloride** [Jacobsen's catalyst,) [*R,R*-138124-32-0, *S,S*-135620-04-1] $C_{36}H_{52}ClN_2O_2Mn$, **M 635.2, m 330-332°, 324-326°, $[\alpha]_D^{20} \pm 580$ (c 0.01, EtOH)**. This complex is commonly referred to as the Mn(salen) catalyst. If this enantioselective catalyst for the epoxidation of olefins requires purification, the brown powder (*ca* 6g) is dissolved in CH_2Cl_2 (100ml), extracted with H_2O (2 x 100ml), the organic phase is dried (Na_2SO_4), filtered, evaporated to dryness and the solid residue is dried in a vacuum desiccator over $CaSO_4$. [Jacobsen et al. *J Am Chem Soc* **113** 7063 1991, DOI: 10.1021/ja00018a068]. For large scale preparation see Larrow et al. *J Org Chem* **59** 1939 1994, DOI: 10.1021/jo00086a062; and Deng & Jacobsen *J Org Chem* **57** 4320 1992, DOI: 10.1021/jo00041a054. For epoxidation of indenenes see Hughes et al. *J Org Chem* **62** 2222 1997, DOI: 10.1021/jo961735i; and for the epoxidation of isoflavones see Adam et al. *Tetrahedron: Asymmetry* **9** 1121 1998, DOI: 10.1016/S0957-4166(98)00102-5; Jacobsen et al. *Tetrahedron* **50** 4323 1994, DOI: 10.1016/S0040-4020(01)89369-8; Jacobsen's Asymmetric Catalytic Epoxidation of Unfunctionalized Olefins in *Catalytic Asymmetric Synthesis Vol I*, p. 159, Ojima Ed. VCH, NY 1993 (see also Bibliography in Part 2), and for enantioselective hydrolysis of epoxides see Furrow et al. *J Org Chem* **63** 6776 1998, DOI: 10.1021/jo981332d.

On a large scale, the brown *R,R*-(+)- catalyst (300-400g) is washed with hot H_2O (2 x 800ml, at 50°), and dried on the filter for 2 hours then under vacuum (50-100mm) at 60-70° for 12 hours, **m 324-326°** (product of acceptable purity has *m* ~320°), $[\alpha]_D^{23} +580$ (c 0.01, EtOH), and IR with ν_{max} (KBr) at 1535, 1612, 2950-2958 cm^{-1} . [Larrow et al. *J Org Chem* **59** 1939 1994, DOI: 10.1021/jo00086a062.]

The related chiral complexes of **Al(III)-Cl** [*R,R* 250611-13-3; *S,S* 307926-51-8], of **Cr(III)-Cl** [*R,R* 164931-83-3; *S,S* 219143-92-7] and of the **Co(II)-Cl** [*R,R*-(+) 176762-62-5; *S,S*-(-) 188264-84-8] (Tokunaga et al. *Science* **277** 936 1977, DOI: 10.1126/science.277.5328.936; Jacobsen et al. *Tetrahedron Lett* **38** 773 1997, DOI: 10.1016/S0040-4039(96)02414-8) are similarly prepared and are available commercially. [See also Schaus et al. *J Am Chem Soc* **124** 1307 2002, DOI: 10.1021/ja016737i; Brandes & Jacobsen *Tetrahedron Asymm* **23** 3927 1977, Gurar et al. *Heterocycles* **48** 1471 1998.]

NOTE: It was shown that addition of pyridine *N*-oxides, and particularly 4-(3-phenylpropyl)pyridine-1-oxide (P_3NO , see [34122-28-6] in Part 2), stabilise and improve the catalytic activity of Mn(salen) catalysts [Larrow & Jacobsen *J Am Chem Soc* **116** 12129 1994, DOI: 10.1021/ja00105a094; Srinivasan et al. *J Am Chem Soc* **108** 2309 1986, DOI: 10.1021/ja00269a029; Senanayake *Aldrichimica Acta* **31** 3 1998.] Thus in a typical epoxidation $MnLCl$ (0.56mmol, salen Mn catalyst) and P_3NO (2.32mmol) in chlorobenzene (10ml) are added to a 2M solution of NaOCl (26.5mmol) cooled to 0° under N_2 , and stirred at 0° for 15 minutes, followed by simultaneous addition of indene (10ml) and further NaOCl (79.5mmol) (syringe pumps) during 30 minutes. The reaction is complete within 1 hour at 0°, and can be performed on a multi-kilogram scale to provide indene epoxide in 89% yield and with an optical purity of 88% e.e. [Senanayake et al. *Tetrahedron Lett* **37** 3271 1996, DOI: 10.1016/0040-4039(96)00565-5.]

A dimeric Jacobsen ligand, formed by linking two units with a methylene bridge has been prepared and crystallised from CH_2Cl_2 /pentane. Its **bis-Mn catalyst** exhibited improved retention in a poly-dimethylsiloxane membrane for the asymmetric epoxidation of olefines [Janssen et al. *Tetrahedron: Asymmetry* **8** 3481 1997, DOI: 10.1016/S0957-4166(97)00465-5].

[(*R*)-(+)-2,2'-Bis(diphenylphosphino)-1,1'-binaphthyl]palladium(II) dichloride (**[*R*-BINAP] $_2PdCl_2$**) [115826-95-4] $C_{44}H_{32}Cl_2P_2Pd$, **M 800.0, m 255-260° (dec)**, $[\alpha]_D^{24} +682$ (c 0.5, $CHCl_3$). This catalyst is prepared by dissolving a mixture of *R*-BINAP (500mg, 0.803mmol, see [76189-55-4]), η -allyl- η -cyclopentadienyl-Pd(II) (84.2mg, 0.396mmol, see [1271-03-0], cf Tatsuno et al. *Inorg Synth* **19** 220 1979, DOI: 10.1002/9780470132500.ch51) in C_6H_6 (5ml), in a Schlenk tube under N_2 (dried over P_2O_5), and heated at 80° for 3 hours to give a deep red solution which is then evaporated to dryness. The red solid is then dissolved

in CH_2Cl_2 (10ml), filtered and the filtrate is evaporated *in vacuo* down to ca 4ml, diluted with Et_2O (40ml) and cooled to 4° whereby red-purple crystals of $[\text{R-BINAP}]_2\text{Pd}(0)$ separate, which on washing with cold Et_2O and drying *in vacuo* give analytically pure $[\text{R-BINAP}]_2\text{Pd}(0)\text{-Et}_2\text{O}$ (480mg, 90%). The 1:1 ratio of Et_2O to Pd is confirmed by ^1H NMR spectroscopy. It has ^1H NMR (90MHz, $^*\text{C}_6\text{D}_6$, TMS) with δ_{H} at 5.88 (t, $J = 7.6\text{Hz}$, 8H), 6.19 (t, $J = 7.3\text{Hz}$, 4H), 6.8-7.6 (m, 40H), 7.83 (d, $J = 8.2\text{Hz}$, 4H), 8.36 (brs, 8H); and the $^{31}\text{P}\{^1\text{H}\}$ NMR (36MHz, $^*\text{C}_6\text{D}_6$, referenced to external 85% H_3PO_4 at δ 0.0) has δ_{P} at 27.3. This Pd(0) complex catalyses asymmetric arylation in the Heck reaction. $[\text{R-BINAP}]_2\text{PdCl}_2$ is obtained by stirring *R*-BINAP (249mg, 0.40mmol) in $^*\text{C}_6\text{H}_6$ (4.0ml) with $\text{PdCl}_2(\text{MeCN})_2$ (104mg, 0.40mmol) in $^*\text{C}_6\text{H}_6$ (4.0ml) overnight. The yellow precipitate is collected by filtration, washed with $^*\text{C}_6\text{H}_6$ and dried *in vacuo*. It is crystallised by dissolving in Me_2CO , layering with hexane and setting aside at $\sim 25^\circ$ to give red crystals of $[\text{R-BINAP}]_2\text{PdCl}_2$ (238mg, 75%, **m 255-260**) of analytical purity, and suitable for X-ray crystallographic structure determination which confirmed its structure. [Ozawa et al. *Organometallics* **12** 4188 1993, DOI: 10.1021/om00034a064; cf. review by Shimizu, Nagasaki & Saito *Tetrahedron* **61** 5405 2005, DOI: 10.1016/j.tet.2005.03.022].

Bis(trifluoromethanesulfonyl)amine metal salts $[\text{NTf}_2]$, bis-(trifluoromethanesulfonyl)amide metal salts, $(\text{CF}_3\text{SO}_2)_2\text{NMetal}$ see [82113-65-3]. NTf_2 readily forms salts, e.g. with Li, Ba, Ca, Al, Zn, Al and lanthanides, which act as **Lewis Acids** in effectively catalysing Diels-Alder reactions [Kobayashi et al. *Chem Lett* 307 1995, DOI: 10.1246/cl.2003.654; Handy et al. *Synlett* 565 1995, DOI: 10.1055/s-1995-5289] and acylation reactions [Mikami et al. *Synlett* 171 1996, DOI: 10.1055/s-1996-5363]. It forms an ***N*-trimethylsilyl derivative** (TMSNTf_2 prepared from allylTMS and NTf_2 , see above) which catalyses Diels-Alder reactions between methyl acrylate and various dienes [Mathieu & Ghosez *Tetrahedron Lett* **38** 5497 1997, DOI: 10.1016/S0040-4039(97)01208-2], as well as Friedel-Crafts acylation reactions of anisole, and allylation and bis-allylation of carbonyl derivatives [Ishii et al. *Synlett* 1145 1997, DOI: 10.1055/s-1997-995]. NTf_2 is a weakly coordinating counter ion which confers stability to catalytic metal complexes such as the gold catalyst $[\text{Ph}_3\text{PAu}(\text{I})]^+ [\text{NTf}_2]^-$, see [866395-16-6].

***R*(-)- and *S*(+)- 4,12-Bis(diphenylphosphino)-[2.2]-paracyclophane {*R*- and *S*- PHANEPhos, (*R*)- and (*S*)- 5,11-bis(phenylphosphino)-tricyclo[8.2.2.2^{4,7}]hexadeca-hexaene}** [(*R*) 364732-88-7; (*S*) 192463-40-4] **M C₄₀H₃₄P₂, 576.6, m 222-226 $^\circ$, 224-276 $^\circ$, $[\alpha]_{\text{D}}^{25}$ *R* -34, *S* +34 (c 1, CHCl_3)**. These are chiral ligands for the rhodium precatalysts $(\text{COD})\text{Rh}\{(\text{R})\text{-[2.2]PHANEPhos}\}^+ \text{OTf}^-$ and $(\text{COD})\text{Rh}\{(\text{S})\text{-[2.2]PHANEPhos}\}^+ \text{OTf}^-$ (see preparation and purification below) which are highly enantioselective in the hydrogenation of dehydroamino acid methyl esters generally, and also of methyl 1-Ac-4-Boc-1,4,5,6-tetrahydropyrazine-1-carboxylate which is difficult to reduce. When they are pre-reduced in MeOH, they form $(\text{MeOH})_2\text{Rh}\{(\text{R})\text{-[2.2]PHANEPhos}\}^+ \text{OTf}^-$ whose activity is more pronounced and reduction is complete in less than 60 minutes by passing H_2 gas through the MeOH solution at temperatures between -45° to $+50^\circ$, with high substrate to catalyst ratios. The preparations of these ligands (described briefly here for convenience) involve di-bromination of *p*-cyclophane, separation of the *meso*- from the *racemic*- isomers (latter are more soluble in triglyme), reaction of the *rac*-isomer (1 equivalent) with *tert*-BuLi (4.2 equivalents), and $\text{MgBr}_2\cdot\text{Et}_2\text{O}$ (2.3 equivalents) in THF then diphenylphosphinyl chloride (Ph_2POCl , 2.2 equivalents.) to give *rac*-4,12-bis(diphenylphosphinyl)-[2.2]-paracyclophane (76% yield). Optical resolution of this racemate (in CHCl_3) with (+)-dibenzoyl-D-tartaric acid (in EtOAc) at 60° , then cooling to $\sim 25^\circ$ over 18 hours, gives a white precipitate from which the ***R*(-)-enantiomer** (>99.5% 'ee') is isolated. The absolute configuration was obtained from the X-ray crystallographic structure determination of the *R*-dibenzoyl-D-tartrate salt (for definition of the **chiral descriptor notation** see R.S. Chan, C.K. Ingold and V. Prelog *Angew Chem Int Ed* **5** 385 1966, DOI: 10.1002/anie.196603851). From the mother liquors the ***S*(+)-enantiomer** (>99.5% 'ee') was isolated by washing the diastereomeric tartrate salt (in CHCl_3) with 0.5M aqueous NaOH. They are white solids that are separated by SFC (supercritical fluid chromatography) using a 25cm Hewlett Packard Chiracel AD column with 15% MeOH/supercritical CO_2 at 1.0ml/minute) with retention times of 13.0 minutes for the *R*-isomer and 17.8 minutes for the *S*-isomer, and they have $[\alpha]_{\text{D}}^{25}$ -104 and $[\alpha]_{\text{D}}^{25}$ +104 (c 0.1, EtOH) respectively. When *R*-4,12-bis(diphenylphosphinyl)-[2.2]-paracyclophane (4.0g, 6.58mmol) is treated with Cl_3SiH (10ml, 99mmol, 10025-78-2) in *p*-xylene (80ml), and slowly heated to 140° , kept at this temperature for 18 hours, a further aliquot of Cl_3SiH (5ml, 49.5mmol) is added, and heating is continued for 24 hours, the reduction appears complete. The mixture is then cooled to -10° , quenched with 30% aqueous NaOH (60ml, care violent reaction), extracted with EtOAc (3 x 80ml), the extract is dried (MgSO_4), filtered and evaporated to dryness. Degassed MeOH (30ml, by sonication) is added to the residue which

is filtered and dried to give the pure diphosphino ***R*-[2.2]PHANEPhos** as a white solid. It has ^1H NMR (400MHz, CDCl_3 , with residual CHCl_3 as reference at δ 7.27) with δ_{H} at 7.54, (m, 2H), 7.45 (m, 3H), 7.41 (m, 2H), 7.24 (m, 3H), 6.62 (*ortho* and *meta*, 2H), 6.58 (dd, $J = 7.6$ and 5.0Hz, 1H), 3.13 (m, 1H), 3.03 (*ortho* and *meta*, 2H), 2.65 (m, 1H); for ^{13}C NMR see references, and ^{31}P NMR (160MHz, CDCl_3 , referenced to external 85% H_3PO_4 at δ 0.0) with δ_{P} at -0.53. The enantiomeric ***S*-[2.2]PHANEPhos** is similarly obtained.

$\text{Rh}(\text{COD})\{(\text{R})\text{-[2.2]PHANEPhosS}\}^+ \text{OTf}^-$ is prepared using Schlenk equipment under N_2 from the red solution of ***R*-[2.2]PHANEPhos** (630mg, 1.09mmol) and $[(\text{Rh COD})]_2^+ \text{OTf}^-$ (512mg, 1.09mmol, see 12092-47-6 for chloride, cf. Catalysts-Part 2), in CH_2Cl_2 (10ml) which is stirred for 30 minutes, evaporated *in vacuo*, $\text{MeO-}t\text{-Bu}$ (10ml) is added to the residue, sonicated for 30 minutes, stirred vigorously for 30 minutes, filtered and the solid is dried under N_2 to give the orange *R*- precatalyst (860mg, 85%). It has ^1H NMR (400MHz, CDCl_3 , with residual CHCl_3 as reference at δ 7.27) with δ_{H} at 8.58 (m, 2H), 7.85 (m, 1H), 7.31 (m, 2H), 7.59 (m, 1H), 7.43 (m, 1H), 7.36 (m, 2H), 7.19 (m, 2H), 6.55 (br d, $J = 8.0\text{Hz}$, 1H), 6.43 (dd, $J = 8.0$ and 4.0Hz, 1H), 4.50 (br s, 2H), 2.77 (m, 1H), 2.67 (m, 1H), 2.54 (m, 1H), 2.48 (m, 1H), 2.20 (om, 3H), 2.03 (m, 1H); for ^{13}C NMR see references; and ^{31}P NMR (160MHz, CDCl_3 , referenced to external 85% H_3PO_4 at δ 0.0) has δ_{P} at 32.7 (d, $J_{\text{P-Rh}} = 146.1\text{Hz}$). The enantiomeric **$\text{Rh}(\text{COD})\{(\text{S})\text{-[2.2]PHANEPhosS}\}^+ \text{OTf}^-$** is similarly prepared. [Pye, Rossen et al. *J Am Chem Soc* **119** 6207 1997, DOI: 10.1021/ja970654g.] For the catalytic hydrogenation of β -keto esters to β -hydroxy esters using **$\text{Ru}(\text{II})\{[\text{2.2]PHANEPhos}\}(\text{trifluoroacetate})_2$** with up to 96% ‘ee’, see Pye, Rossen et al. *Tetrahedron Lett* **39** 4441 1998, DOI: 10.1016/S0040-4039(98)00842-9.

***R*(-)- and *S*(+)- 4,12-Bis[di(3,5-xylyl)phosphino]-[2.2]-paracyclophane {*R*- and *S*- xylyl-PHANEPhos, (*R*)- and (*S*)-5,11-bis[di(3,5-xylyl)phenylphosphino]-tricyclo-[8.2.2.2^{4,7}]hexadeca-hexaene} [(*R*) 325168-89-6; (*S*) 325168-88-5] $\text{C}_{48}\text{H}_{50}\text{P}_2$, **M 688.9, m 234-238°**, $[\alpha]_{\text{D}}^{22}$ ***R* -61.0, *S* +61.0 (c 0.1, EtOH)**. These chiral ligands are prepared and purified in much the same way as for [2.2]PHANEPhos above except that 3,5-dimethylphenylphosphinyl chloride is used to introduce the phosphorous group into the *p*-cyclophane. When these enantiomers are complexed with ruthenium and chiral ethylenediamine-type bases, they form powerful precatalysts that in the presence of H_2 , will catalyse the reduction of a range of simple aromatic, heteroaromatic as well as α,β -unsaturated ketones with high ‘ee’ values. Only one example, **$\text{R}\{[\text{xylyl-PHANEPhos}]\text{-Ru}_2\text{Cl}(\text{S,S})\text{-DPEN}\}$** is described here. It is prepared from the red solution of $[(\text{C}_6\text{H}_6)\text{RuCl}_2]_2$ (218mg, 0.436mmol, see [37366-09-9]), *R*-xylyl-PHANEPhos (0.60mg, 0.871mmol) in dry DMF (6ml) by heating at 100° for 4 hours, to which is then added ***S,S*-1,2-diphenylethylene-1,2-diamine** (185mg, 0.871mmol, *S,S*-DPEN, [29841-69-8]), and heating, and stirring is continued at 100° for 1.5 hours. The mixture is filtered, the filtrate is evaporated *in vacuo*, treated with Et_2O (10ml) and MeOH (100), the yellow catalyst is filtered off, washed with MeOH (12ml) and dried (420mg, 45% yield). It has ^1H NMR (400MHz, CDCl_3) with δ_{H} at 1.78 (ddd, $J = 15.0, 10.5, 4.5\text{Hz}$, 1H), 2.07 (s, 6H), 2.27 (s, 6H), 2.59 (s, 1H), 2.78 (m, 1H), 4.01 (m, 1H), 4.40 (m, 1H), 4.49 (m, 1H), 6.36 (d, $J = 8.0\text{Hz}$), 6.42 (d, $J = 8.0\text{Hz}$), 6.60 (m, 2H), 6.79 (m, 1H), 6.96 (m, 2H), 7.06 (s, 1H), 7.13 (m, 3H), 8.04 (m, 2H), 8.21 (m, 1H); and ^{31}P NMR (162MHz, CDCl_3 , referenced to external 85% H_3PO_4 at δ 0.0) with δ_{P} at +46.06 (s). The crude catalysts performed equally as well as the **purified precatalysts**. [Burk et al. *Org Lett* **2** 4173 2000, DOI: 10.1021/ol000309n.]**

{1,2-Bis[(4*S*)-4-isopropyl-2-oxazolin-2-yl]benzene}zinc(II) Chloride [131380-93-3] $\text{C}_{18}\text{H}_{24}\text{N}_2\text{O}_2\text{ZnCl}_2$, **M 436.7, m 202°**. The zinc complex is obtained by adding a solution of the free ligand (300mg, 1mmol, see [131380-80-8]) in dry THF (5ml) to a solution of fused anhydrous ZnCl_2 (136mg, 1mmol, dried by melting in high vacuum and cooling under argon) in dry THF (10ml) and collecting the crystals, but if crystals are not readily formed, evaporate to dryness and recrystallise the residue from EtOH to give the colourless complex (420mg, 98%). It has IR (KBr) with ν_{max} at 1655 ($\text{C}=\text{N}$) cm^{-1} ; the ^1H NMR (300MHz, CD_3CN) has δ at 0.84 (d, $J = 6.9\text{Hz}$, 6H, $\text{CH}(\text{CH}_3)_2$), 0.95 (d, $J = 7.1\text{Hz}$, 6H, $\text{CH}(\text{CH}_3)_2$), 2.52-2.62 (m, 2H, $\text{CH}(\text{CH}_3)_2$), 4.50-4.62 (m, 6H, OCH_2CHN), 7.77-7.81 (m, 2H, 4-H and 5-H), 7.85-7.89 (m, 2H, 3-H and 6-H) from TMS; for the ^{13}C NMR see reference. [Bolm et al. *Chem Ber* **124** 1173 1991, DOI: 10.1002/cber.19911240532.] Like many chiral bis(oxazolin-2-yl) complexes it can be involved in a variety of metal catalysed asymmetric synthesis such as allylation, aziridination, cyclopropanation, Diels-Alder and retro Diels Alder, Mukaiyama aldol condensation and hydrosilylation [see reviews by Ghosh et al. *Tetrahedron: Asymmetry* **9** 1 1998, DOI: 10.1016/S0957-4166(97)00593-4; Pflatz *Acta Chem Scand* **50** 189 1996, DOI: 10.3891/acta.chem.scand.50-0189; Johnson & Evans *Acc Chem Res* **33** 325 2000, DOI: 10.1021/ar960062n; Jørgensen et al. *Acc Chem Res* **32** 605 1999, DOI: 10.1021/ar970347f.]

1,3-Bis(2,4,6-trimethylphenyl)-imidazol-2-ylidenegold(I) chloride (IMesAuCl) [852445-81-9] $\text{C}_{21}\text{H}_{25}\text{N}_2\text{AuCl}$, **M 537.8**, **m ~280-286°**. The gold complex is prepared by adding Ag_2O (0.5mmol) to IMes^+Cl^- (1mmol) in CH_2Cl_2 and the suspension is stirred for 3 hours at $\sim 25^\circ$ whereby it became clear; then a solution of $\text{Au}(\text{Me}_2\text{S})\text{Cl}$ (1mmol, see [29892-37-3]) in CH_2Cl_2 is added dropwise and stirred for a further 4 hours. After filtration through Celite, the filtrate is evaporated to a small volume and hexane is added to allow the complex to separate. It is collected and dried (49% yield). Its ^1H NMR (400MHz, CDCl_3 , TMS) has δ_{H} at 7.08 (s, 2H), 6.97 (bs, 4H), 2.33 (s, 6H), 2.09 (s, 12H), for ^{13}C NMR and HRMS-FAB see reference. It catalyses the intramolecular [4+2] cycloadditions of 1-en-6-yne, or aralkynes with alkenes, in the presence of AgSbF_6 ; also the related **1-methyl-3-(2,4,6-trimethylphenyl)-imidazol-2-ylgold(I) chloride** is almost equally effective [Nieto-Oberhuber *J Am Chem Soc* **127** 6178 2005, DOI: 10.1021/ja042257t]. [For X-ray structure of dimeric PF_6 salt see Hu et al. *Organometallics* **23** 755 2004, DOI: 10.1021/om0341855.]

The saturated 4,5-dihydro- analogue **1,3-bis(2,4,6-trimethylphenyl)-imidazolin-2-ylidenegold(I) chloride (ISMeAuCl)** [852445-82-0] $\text{C}_{21}\text{H}_{227}\text{N}_2\text{AuCl}$, **M 539.9**, behaves in much the same manner as IMesAuCl, chemically and catalytically. It has recently been shown that both IMesAuCl and SIMesAuCl can be made by metal exchange from the cheaper respective IMesCuCl and SIMesCuCl (see below) in 71-90% yields. Thus a mixture of $[\text{AuCl}(\text{SMe}_2)]$ (0.4-0.47mmol, [29892-37-3]) and $[\text{CuCl}(\text{NHC})]$ (1 equivalent) in CH_2Cl_2 (5ml) is stirred at 40° for 1-2 hours, then filtered through Celite and the solvent is removed *in vacuo*. CH_2Cl_2 (3ml) and petroleum ether (10ml) are added to the residue, the **almost pure Au complex** is filtered off, washed with petroleum ether (3 x 5ml) until colourless, and dried *in vacuo*. Their catalytic activities from these different preparations are the same. [Furst & Cazin *Chem Commun* **46** 6924 2010, DOI: 10.1039/C0CC02308F.]

Chloro[(1,1'-biphenyl-2-yl)di-tert-butylphosphine]gold(I) (JohnPhos-gold(I) chloride, JohnPhos-AuCl) [854045-93-5] $\text{C}_{20}\text{H}_{27}\text{PAuCl}$, **M 530.8**, **m 237-240°**. The gold complex is made from sodium tetrachloroaurate(III) dihydrate [13874-02-7] or hydrogen tetrachloroaurate(III) [see 16961-25-4] (1 mmol) in H_2O and the yellow solution is cooled in ice, to which is added **bis(2-hydroxyethyl)sulfide** [see 111-48-8] (3mmol) dropwise over ~ 45 minutes with stirring. Then a solution of JohnPhos [224311-51-7] (1mmol) in EtOH is added dropwise, whereby the gold complex separates as a white solid which is filtered off (43% yield), washed with MeOH and dried *in vacuo*. The ^1H NMR (400MHz, CDCl_3) has δ_{H} at 7.87 (td, $J = 7.7, 1.7\text{Hz}$, 1H), 7.51 (m, 5H), 7.31 (m, 1H), 7.13 (dd, $J = 8.0, 1.0\text{Hz}$, 2H), 1.41 (d, $J = 15.6\text{Hz}$, 18H); the ^{31}P NMR (160MHz, CDCl_3) has δ_{P} at 63.11; and for ^{13}C NMR and HRMS-ESI see references. [Nieto-Oberhuber et al. *J Am Chem Soc* **127** 6168 2005, Al-Sa'Ady et al. *Inorg Synth* **23** 191 1985, DOI: 10.1002/9780470132548.ch39.]

The acetonitrile complex salt derivative **(acetonitrile)(2-biphenyl)-di-tert-butylphosphinogold(I) hexafluoroantimonate** [866641-66-9] **M 772.1**, has **m 198-203°(dec)**. It catalyses a highly efficient intramolecular addition of β -ketoamide to unactivated olefins leading to highly substituted lactams in the presence of AgOTf (e.g. *N*-allyl *N*-benzyl benzoylacetamide to *cis*-3-benzoyl-1-benzyl-4-methylpyrrolidin-2-one in 99% yield [Zhou & Che *J Am Chem Soc* **129** 5828 2007, DOI: 10.1021/ja070027j]). JohnPhos-AuCl and related complexes, in the presence of AgSbF_6 (see SbF_6^- salts), catalyse intramolecular [4+2] cycloadditions of 1,3-enynes or arylalkynes and alkenes very efficiently [Nieto-Oberhuber et al. *J Am Chem Soc* **127** 6178 2005, DOI: 10.1021/ja042257t].

Chloro[2-dicyclohexyl(2',4',6'-tri-isopropylbiphenyl)phosphine]gold(I) [X-Phos-gold(I) chloride, X-Phos-AuCl] [854045-94-6] $\text{C}_{33}\text{H}_{49}\text{PAuCl}$, **M 709.1**, **m 243-250°**. This complex is prepared in much the same way as JohnPhos-AuCl but using a solution containing 1mmol of X-Phos [564483-18-7] in EtOH instead and giving a 61% yield of the X-Phos-AuCl as a white solid. The ^1H NMR (400MHz, CDCl_3) has δ_{H} at 7.57-7.55 (m, 2H), 7.23 (m, 1H), 7.05 (s, 2H), 2.95 (hept, $J = 7.0\text{Hz}$, 1H), 2.21 (hept, $J = 6.7\text{Hz}$, 2H), 2.05-2.01 (m, 4H), 1.83-1.71 (m, 6H), 1.67-1.61 (m, 2H), 1.53-1.42 (m, 3H), 1.35 (s, 3H), 1.34 (s, 3H), 1.28 (s, 3H), 1.26 (s, 3H), 1.23-1.13 (m, 7H), 0.92 (s, 3H), 0.91 (s, 3H); the ^{31}P NMR (160MHz, CDCl_3) has δ_{P} at 38.48; and for ^{13}C NMR and HRMS-ESI see references. [Nieto-Oberhuber et al. *J Am Chem Soc* **127** 6178 2005, DOI: 10.1021/ja042257t; Al-Sa'Ady et al. *Inorg Synth* **23** 191 1985, DOI: 10.1002/9780470132548.ch39.] Its catalytic activities are similar to those of JohnPhos-AuCl above.

Chloro(triphenylphosphine)gold(I) [14243-64-2] $[(\text{C}_6\text{H}_5)_3\text{P}]\text{AuCl}$, **M 495.7**, **m 240-241°, 242-243°**. Ph_3PAuCl is prepared under argon by adding a solution of hydrogen tetrachloroaurate (2.0g, 5.87mmol, $\text{H}[\text{AuCl}_4] \cdot n\text{H}_2\text{O}$, $n \sim 3$, 'brown gold chloride' see [16903-35-8]) in EtOH (10ml), which has been filtered to remove any insoluble Au-containing material, to a stirred solution of Ph_3P (3.08g, 11.7mmol) in absolute EtOH

(50ml of 100%, warm if necessary to dissolve). The solution immediately becomes colourless and after stirring for 15 minutes the white microcrystalline solid is collected (fritted glass filter, porosity 4), washed with EtOH (2 x 5ml), then Et₃O (3 x 15ml), and dried *in vacuo* (0.1mm) to give the analytically pure complex (1.9-2.3g 67-80%, or up to 92% if equipment is thoroughly dried, solvents degassed, and working in a strict inert atmosphere). It (~1.g) crystallises from CH₂Cl₂ (10ml) by slow addition of *n*-pentane (60ml) and cooling to -25° to give small slender white needles within a few minutes, and it also crystallises from CHCl₃ solution by addition of heptane. [Braunstein et al. *Inorg Synth* **27** 218 1990, DOI: 10.1002/9780470132586.ch42; Bruce et al. *Inorg Synth* **26** 324 1986, DOI: 10.1002/9780470132579.ch59; cf. JohnPhos-AuCl above.]. Ph₃PAuCl has also been prepared by adding Ph₃P (1.5g) in Me₂CO (5ml) to a solution of HAuCl₄ (1g, 0.5mol equivalents, 'chlorogold(III) acid' see [16903-35-8]) in Me₂CO at 20°. The volume of the solution is reduced to *ca* 5ml and the complex that separates (1.2g, 83%) is collected and dried or crystallised as before [Burgess et al. *JCS Dalton Trans* 1661 1983, DOI: 10.1039/DT9830001661]. **Alternatively**, to the orange solution of sodium tetrachloroaurate(III).2H₂O (1mmol, [13874-02-7]) dissolved in 2H₂O, cooled in ice, is slowly added 2,2'-thiodiethanol {3mmol, i.e. bis(2-hydroxyethyl)sulfide [see 111-48-8]} at 0° during ~45 minutes, followed dropwise by a solution of the Ph₃P ligand (1mmol, or other PX₃ ligand) in EtOH which causes a white solid to separate. This is filtered off, washed with MeOH, and dried *in vacuo* (0.1mm) to give Ph₃PAuCl (or other PX₃AuCl) as a white solid in ~88% yield. It is recrystallised as described above. [Li et al. *J Org Chem* **73** 4323 2008, DOI: 10.1021/jo8003875]. The ¹H NMR (400MHz, CDCl₃) has δ_H at 2.03-1.93 (m, 9H), 1.89-1.85 (m, 6H), 1.75-1.72 (m, 3H), 1.52-1.41 (m, 6H), 1.35-1.21 (m, 9H); the ³¹P NMR (160MHz, CDCl₃) has δ_P at 57.12 (34.224ppm reported also at 300MHz, CDCl₃, from external H₃PO₄); and for ¹³C NMR and HRMS-ESI see references. [Nieto-Oberhuber et al. *J Am Chem Soc* **127** 6178 2005, DOI: 10.1021/ja042257t; Al-Sa'Ady et al. *Inorg Synth* **23** 191 1985, DOI: 10.1002/9780470132548.ch39; Li et al. *J Org Chem* **73** 4323 2008, DOI: 10.1021/jo8003875.] Its catalytic activities are similar to those of JohnPhos-AuCl above. It has also been used in the cyclisation of *O*-propargyl carbamates to alkylideneoxazolidiones at room temperature, *via* a 5-exo-digonal pathway [Ritter et al. *Synlett* 3309 2006, DOI: 10.1055/s-2006-951555]. It is soluble in Me₂CO, CH₂Cl₂, CHCl₃, *C₆H₆ and THF, but **not** in EtOH or hexane, and is used for the preparation of a variety of Ph₃PAu salts and cluster complexes. It is stable indefinitely in air, and note that ³¹P NMR studies of Ph₃PAuCl showed no change in the chemical shift (at 33.8ppm in THF) after bubbling O₂ through the solution of 24 hours, indicating how stable it is compared with cationic Ph₃PAu⁺ species in the presence of O₂ [Liu et al. *J Am Chem Soc* **128** 11332 2006, DOI: 10.1021/ja062610q]. X-Ray crystallography showed that the molecule of Ph₃PAuCl is linear with a P-Au-Cl angle of 179.68° and Au-P distance of 2.235Å [Baenziger et al. *Acta Cryst* **B32** 962 1976, DOI: 10.1107/S0567740876004330].

Chlorotris(triphenylphosphine)rhodium(I) (Wilkinson's catalyst) [14694-95-2] [(C₆H₅)₃P]₃RhCl, **M 925.2, m 138°(dec), 140°(dec), 157-158°(dec)**. It is prepared by adding PPh₃ (12g, recrystallised from EtOH to remove any PPh₃O) in hot EtOH (350ml) to RhCl₃.3H₂O (2g) in 95% EtOH (70ml) in a flask purged with N₂, refluxed in the presence of N₂ for 2 hours, and the red solid that is formed is collected on a sintered glass funnel and washed with small volumes of dry Et₂O (50ml) to give the catalyst (6.25g, 88%). It forms dark burgundy crystals from hot EtOH after refluxing for 30 minutes. When the solution is heated for only 5 minutes, orange crystals are formed. Heating the orange crystals in EtOH yields red crystals. Crystallisation from Me₂CO gives the orange crystals. The dimorphic forms have similar IR spectra, but the X-ray diffraction patterns are slightly different. Excess of PPh₃ can be recovered by adding H₂O to the EtOH filtrates from recrystallisation and, collecting it on standing for 2-3 days. [Osborne et al. *J Chem Soc (A)* 1711 1966, DOI: 10.1039/J19660001711; Osborn et al. *Inorg Synth* **10** 67 1967, DOI: 10.1002/9780470132418.ch12; Bennett & Donaldson *Inorg Chem* **16** 655 1977, DOI: 10.1021/ic50169a033.] The solubilities are: in CH₂Cl₂ ~2% (25°), in toluene 0.2% (25°), but less so in Me₂CO, MeOH, BuOH and AcOH, and insoluble in petroleum ethers and cyclohexane. It reacts with donor solvents e.g. pyridine, DMSO and MeCN. Solutions of RhCl[(PPh₃)₃] are **air sensitive** absorbing O₂, and as well as the solid, should be stored under O₂-free N₂ or argon. On heating solutions in *C₆H₆, toluene or methyl ethyl ketone the chlorine-bridged **dimer** [(PPh₃)₃RhCl₂Rh(PPh₃)₃] is formed as salmon-red crystals which absorb oxygen even in the solid state, but can be converted to the **monomer** in refluxing EtOH containing PPh₃. Red solution of RhCl[(PPh₃)₃] absorb H₂ reversibly at 760mm/25°, become yellow in colour, and are effective in catalysing the reduction of C-C double bonds and triple bonds, often in a selective manner. [Osborn et al. *Inorg Synth* **28** 77 1990, DOI: 10.1002/9780470132593.ch17.] The stoichiometric and catalytic reactions of RhCl[(PPh₃)₃] were reviewed by Jardine [*Prog Inorg Chem* **28** 63 1981, DOI: 10.1002/9780470166291.ch2].

In the presence of 1.5 mol of Et_2Zn , 1 mol of 4-phenylbut-3-ene-2-one (and related α -ene-ones) reacts with 1.7 mol of CF_3Cl in THF at 5° to give 4-phenyl-3-(trifluoromethyl)butan-2-one (and related ketones) in moderate yields in the presence of catalytic amounts of $\text{RhCl}[(\text{PPh}_3)_3]$ (~ 0.02 mol) [Sato, Omote, Ado and Kumadaki *Org Synth* **83** 177 2006, DOI: 10.15227/orgsyn.083.0177]. § This catalyst is also available commercially on a styrene-divinylbenzene polymer support. It catalyses efficient cross-coupling of activated alkenyl tosylates with aromatic boronic acids [Wu et al. *Eur J Org Chem* 5260 2006, DOI: 10.1002/ejoc.200600469], and the cleavage of allyl phenolic ethers to phenols in the presence of DABCO [Aristegui et al. *Org Lett* **8** 5927 2006, DOI: 10.1021/ol062297x]. [See Fieser **1** 140, **2** 448, **3** 325, **4** 559, **5** 736, **6** 652, **8** 109, **9** 113, **15** 90, **16** 86.]

Chromium (III) acetylacetonate [$\text{Cr}(\text{acac})_3$] [21679-31-2] ($\text{C}_5\text{H}_7\text{O}_2$)₃Cr, M 349.3, m 212-216°, 216°, b 340°, pK²⁵ 4.0 (see chromic chloride). Purify it by dissolving 6g in hot C_6H_6 (20ml) and adding petroleum ether (75ml) slowly. Cool to room temperature then chill on ice, filter off and dry in air to give 2.9g. It also crystallises from EtOH. It is soluble in heptane, C_6H_6 , toluene and pentane-2,4-dione at 20-40°. It forms a 1:2 complex with CHCl_3 . [Fernelius et al. *Inorg Synth* **5** 130 1957, DOI: 10.1002/9780470132364.ch35; Steinbach & Burns *J Am Chem Soc* **80** 1839 1958, DOI: 10.1021/ja01541a018; *Beilstein* **1** H 782, **1** II 836, **1** IV 3673.] $\text{Cr}(\text{acac})_3$ catalyses the oxidation of methacrylic acid esters in the presence of H_2O_2 which provides a new route to pyruvic esters [Inoue et al. *Chem Lett (Jpn)* **18** 99 1989, DOI: 10.1246/cl.1989.99]. **Potential carcinogen.**

Cobalt Oxazoline Palladacycles (COPs) are organocobalt-palladium complexes which catalyse the asymmetric rearrangements of non-chiral allylic trichloroacetamides with very high enantiomeric selectivity (>90%) to provide chiral allylic amines [it is an aza-Claisen rearrangement, ‘*The Overman Rearrangement*’ Overman & Carpenter *Org React* **66** 2005, DOI: 10.1002/0471264180.or066.01; Kirsch, Overman and Watson *J Org Chem* **69** 8101 2004, DOI: 10.1021/jo0487092]; and in the presence of phenols stereospecific cross-coupling also occurs to provide chiral phenoxyallyl ethers with very high (>90%) enantiomeric selectivity [Kirsch, Overman and White *Org Lett* **9** 911 2007, DOI: 10.1021/ol070110b; Overman & Carpenter *Org React* **66** 2005, DOI: 10.1002/0471264180.or066.01].

(S)-(+)-COP-OAc dimer catalyst { $S(\text{COP-OAc})_2$, di- μ -acetobis[η^5 -(S)-(pR)-2-(2'-(4'-methylethyl)-oxazoliny)cyclopentadienyl, 1-C-3'-N)(η^4 -tetraphenylcyclobutadiene)cobalt[dipalladium]} [222400-03-5] $\text{C}_{82}\text{H}_{72}\text{N}_2\text{O}_6\text{Co}_2\text{Pd}_2$, M 1512.2, m 189-194° (250-257° dec.), [α]_D²⁴ +942 (c 0.215, CHCl_3). This COP is prepared in a Schlenk flask flushed with argon containing (η^5 -(S)-2-(4'-methylethyl)-oxazoliny)cyclopentadienyl-(η^4 -tetraphenylcyclobutadiene)cobalt (9.6g, 16.2mmol, see [22240-02-4]) in glacial acetic acid (96ml) to which is added palladium(II) acetate (3.6g, 16.2mmol, must be recrystallised from C_6H_6), and the red solution is heated at 95° for 30 minutes when an orange precipitate separates. After cooling to $\sim 25^\circ$, the solid is collected, washed with glacial acetic acid (50ml) and dried *in vacuo* to give **almost analytically pure** S -(COP-OAc)₂ (8.9g, 73%) as mustard coloured crystals. It has IR (thin film) with ν_{max} at 3061, 2961, 1583 (C=N), 1501, 1417 (acetate bridge), 1366, 1181, 1069 cm^{-1} ; ^1H NMR (500MHz, CDCl_3) with δ_{H} at -0.01 (d, J = 6.6Hz, 3H, CH_3), 0.46 (d, J = 7.1Hz, 3H, CH_3), 1.72-1.81 (m, 1H, CH), 1.96 (s, 3H, CH_3), 2.98 (td, J = 9.0, 3.1Hz, 1H, CH), 3.36 (t, J = 9.0Hz, 1H, CH_2), 4.08 (dd, J = 8.6, 3.9Hz, 1H, CH_2), 4.23 (t, J = 2.4Hz, 1H, CH), 4.62 (d, J = 1.4Hz, 1H, CH), 4.68 (d, J = 2.0Hz, 1H, CH), 7.20-7.29 (m, 12H, ArH), 7.64 (m, 8H, ArH); see ^{13}C NMR in refs.

The **(R)-(-)-COP-OAc dimer enantiomer** [849592-74-1] $\text{C}_{82}\text{H}_{72}\text{N}_2\text{O}_6\text{Co}_2\text{Pd}_2$, M 1512.2, m 241-251° dec. is prepared in the same way but using the enantiomeric starting material. [Stevens & Richards *Organometallics* **18** 1346 1999, DOI: 10.1021/om980812s; Anderson et al. *Org Synth* **84** 148 2007, DOI: 10.15227/orgsyn.084.0148.] The reactions of trichloroacetimidate derivatives of Z -2-alken-1-ols with phenolic nucleophiles in the presence of chiral COP-OAc dimer catalysts provide 3-aryloxy-1-alkenes in high yields (63-90%) and high enantiomeric purity (90-97% ‘ee’); and are compatible with the presence of base-labile substituents in either reactant [Kirsch, Overman and White *Org Lett* **9** 911 2007, DOI: 10.1021/ol070110b].

(S)-(+)-COP-Cl dimer catalyst { $S(\text{COP-Cl})_2$, di- μ -chlorobis[η^5 -(S)-(pR)-2-(2'-(4'-methylethyl)-oxazoliny)cyclopentadienyl, 1-C-3'-N)(η^4 -tetraphenylcyclobutadiene)cobalt[dipalladium]} [581093-92-7] $\text{C}_{78}\text{H}_{66}\text{Cl}_2\text{N}_2\text{O}_2\text{Co}_2\text{Pd}_2$, M 1464.98, m 204-205°, [α]_D²⁰ +1201 (c 0.1, CH_2Cl_2). It is obtained from the above S -(COP-OAc)₂ (8.9g, 5.9mmol) in acetone (59ml) by stirring vigorously at $\sim 25^\circ$ with aqueous NaCl (2M, 30ml) to give a homogeneous mixture until the reaction is complete (~ 4 hours, checked periodically by sampling a filtered aliquot and examining the ^1H NMR spectra). The yellow complex which separates is filtered off, washed

with H₂O (125ml) then Me₂CO (20ml) and dried *in vacuo* to give almost pure (COP-Cl)₂ (8.2g, 95%) as a mustard coloured solid. It can be purified further by filtering a solution in CH₂Cl₂ through a short plug of Celite and eluting with CH₂Cl₂ to give an *analytically pure* orange solid (97% recovery) which exists as a **1.0:0.7 mixture of dimers**. It has IR (thin film) with ν_{\max} at 3061, 2961, 1602, 1502, 1370, 1185 cm⁻¹; the ¹H NMR (500MHz, CDCl₃) has δ_{H} at 0.70-0.74 (m, 6H, CH₃), 0.76 (d, *J* = 6.8Hz, 3H, CH₃), 0.80 (d, *J* = 7.0Hz, 3H, CH₃), 2.20-2.33 (m, 2H, CH), 3.03-3.15 (m, 2H, CH), 3.34 (t, *J* = 9.0Hz, 1H, CH₂), 3.43 (t, *J* = 8.9Hz, 1H, CH₂), 4.16-4.22 (m, 2H, CH₂), 4.28 (t, *J* = 2.5Hz, 1H, CH), 4.40 (t, *J* = 2.5Hz, 1H, CH), 4.70 (d, *J* = 2.0Hz, 1H, CH), 4.73 (d, *J* = 2.0Hz, 1H, CH), 4.98 (d, *J* = 1.5Hz, 1H, CH), 4.99 (d, *J* = 1.5Hz, 1H, CH), 7.16-7.22 (m, 12H, ArH), 7.23-7.30 (m, 12H, ArH), 7.58-7.62 (m, 8H, ArH), 7.66-7.71 (m, 8H, ArH); and for ¹³C NMR see ref. The enantiomeric purity was determined by conversion into the acetylacetonate derivative **COP-acac** (see below) and the enantiopurity was shown to be >99% by HPLC [Diacel Chiralpak AD-H (0.46cm x 25cm) column, with 1.0ml/minute flow rate and eluted with 95:5 hexanes:*iso*-PrOH]. The retention time for *S*-COP-acac is 5.1 minutes whereas that of *R*-COP-acac would have been ~3.1 minutes [Anderson, Kirsch and Overman *Org Synth* **84** 148 2007, DOI: 10.15227/orgsyn.084.0148.]

(R)-(-)-COP-Cl dimer catalyst {R(COP-Cl)₂, di- μ -chlorobis[η^5 -(*R*)-(p*R*)-2-(2'-(4'-methylethyl)-oxazolinyl)cyclopentadienyl, 1-C-3'-N](η^4 -tetraphenylcyclobutadiene)cobalt[dipalladium]} [612065-00-6] C₇₈H₆₆Cl₂N₂O₂Co₂Pd₂, M 1464.98, m 205-208°, [α]_D²⁰ +1240 (c 0.1, CH₂Cl₂) is obtained and purified as for its *S*-enantiomer above but using the *R*-enantiomeric intermediates. These planar chiral COP-Cl dimers catalyse the [3+3]-sigmatropic rearrangements of a wide range of non-chiral *E*-allylic trichloroacetimidates into the corresponding transposed chiral allylic trichloroacetamides (from which the free base can be obtained) in high yields and very high enantioselectivities [Anderson & Overman *J Am Chem Soc* **125 12412 2003, DOI: 10.1021/ja037086r; Anderson, Overman and Watson *Org Synth* **82** 134 2005, DOI: 10.15227/orgsyn.082.0134]. The reactions of trichloroacetimidate derivatives of *Z*-2-alken-1-ols with phenolic nucleophiles in the presence of chiral COP-Cl dimer catalysts yield 3-aryloxy-1-alkenes in high yield (63-90%) and high enantiomeric purity (90-97% 'ee'); and are compatible with the presence of base-labile substituents in either reactant [Kirsch, Overman and White *Org Lett* **9** 911 2007, DOI: 10.1021/ol070110b].**

(S)-(+)-COP-acac monomer catalyst {acetylacetonato[η^5 -(*S*)-(p*R*)-2-(2'-(4'-methylethyl)-oxazolinyl)cyclopentadien-yl, 1-C-3'-N](η^4 -tetraphenylcyclobutadiene)cobalt[palladium]} [805315-09-7] C₄₄H₄₀NO₃CoPd, M 796.1, m 100-104°(dec.), [α]_D²⁸ +246.1, [α]₅₇₇²⁸ +250.2, [α]₅₄₆²⁸ +175.5, [α]₄₃₅²⁸ -59.8, [α]₄₀₅²⁸ -91.4 (c 1.00, CHCl₃). (*S*)-(+)-COP-acac is obtained as a monomer from the above dimer *S*-(COP-OAc)₂ (1.0g, 0.66mmol), sodium acetylacetonate (954mg, 6.8mmol, see [15435-71-9]), Me₂CO (6.6ml) and H₂O (3.3ml) by stirring vigorously for 24 hours at ~25°. The mixture is then extracted with CH₂Cl₂ (10ml), the extract is separated, dried (MgSO₄), filtered and concentrated to give almost pure (*S*)-(+)-COP-acac (1.0g, quantitative) as an orange solid. It can be further purified by filtering through a silica-gel column and eluting with *iso*-PrOH:hexanes (5:95) and evaporating the orange band to give an analytically pure complex. See above *S*-(COP-Cl)₂ for HPLC data; and it has R_F 0.59 (silica gel, hexanes-EtOAc, 80:20). It has IR (thin film) with ν_{\max} at 3058, 2960, 1597, 1579, 1508, 1399, 1265, 1183, 1067 cm⁻¹; the ¹H NMR (400MHz, CDCl₃) has δ_{H} at 7.64 (d, *J* = 7.6Hz, 8H), 7.20-7.30 (m, 12H), 5.26 (s, 1H), 5.17 (s, 1H), 4.89 (d, *J* = 1.6Hz, 1H), 4.47 (s, 1H), 4.27 (dd, *J* = 8.4, 5.6Hz, 1H), 3.67 (t, *J* = 9.0Hz, 1H), 3.29-3.31 (m, 1H), 2.26-2.27 (m, 1H), 2.00 (s, 3H), 1.94 (s, 3H), 0.84 (d, *J* = 7.6Hz, 3H), 0.82 (d, *J* = 8.0Hz, 3H); and for ¹³C NMR see references. [Kirsch et al. *J Org Chem* **69 8101 2004, DOI: 10.1021/jo0487092; Anderson et al. *Org Synth* **84** 148 2007, DOI: 10.15227/orgsyn.084.0148.]**

The isomer, **(R)-(-)-COP-acac monomer catalyst {acetylacetonato[η^5 -(*R*)-(p*R*)-2-(2'-(4'-methylethyl)-oxazolinyl)cyclopentadien-yl, 1-C-3'-N](η^4 -tetraphenylcyclobutadiene)cobalt[palladium]} [CASRN is the same as preceding *S*-(+)-enantiomer] C₄₄H₄₀NO₃CoPd, M 796.1, m 100-104°(dec), [α]_D²⁸ -246.1, [α]₅₇₇²⁸ -250.2, [α]₅₄₆²⁸ -175.5, [α]₄₃₅²⁸ +59.8, [α]₄₀₅²⁸ +91.4 (c 1.00, CHCl₃) can be obtained and purified as for its *S*-enantiomer above but using the *R*-enantiomeric intermediates. These monomeric COPs catalyse the asymmetric rearrangements of non-chiral allylic trichloro-acetimidates into chiral transposed allylic trichloroacetamides in good yield with high asymmetric induction (>90% 'ee'). These are more soluble catalysts than the COP dimers, and because they are soluble in a much wider variety of solvents, the reactions can be carried out at high substrate concentrations (e.g. at ~2.6M) [Kirsch et al. *J Org Chem* **69** 8101 2004, DOI: 10.1021/jo0487092.]. The reactions of trichloroacetimidate derivatives of *Z*-2-alken-1-ols with phenolic nucleophiles in the presence of**

chiral COP-acac monomer catalysts yield 3-aryloxy-1-alkenes in high yield (63-90%) and high enantiomeric purity (90-97% 'ee'); and are compatible with the presence of base-labile substituents in either reactant [Kirsch, Overman and White *Org Lett* **9** 911 2007, DOI: 10.1021/ol070110b].

(S)-(+)-COP-hfacac {hexafluoroacetylacetonato[η^5 -(*S*)-(p*R*)-2-(2'-(4'-methylethyl)oxazolinyl)-cyclopentadienyl,1-*C*-3'-*N*)(η^4 -tetraphenylcyclobutadiene)cobalt[palladium]} [805315-08-6]

C₄₄H₃₄F₆NO₃ CoPd, M 904.0, m 108-110°(dec.) [α]_D²⁸ +271.2, [α]₅₇₇²⁸ +277.6, [α]₅₄₆²⁸ +214.8, [α]₄₃₅²⁸ -57.7, [α]₄₀₅²⁸ -86.8 (c 1.02, CHCl₃). (S)-(+)-COP-hfacac is obtained as a monomer from the above dimer S-(COP-OAc)₂ (1.0g, 0.66mmol), sodium hexafluoroacetylacetonate (1.6g, 6.8mmol, see [22466-49-5]), Me₂CO (6.6ml) and H₂O (3.3ml) by stirring vigorously at ~25° for 9-11 hours (check periodically by sampling a filtered aliquot and examining the ¹H NMR spectra until completion). The solid that separates is filtered off, washed with H₂O (10-20ml) and dried *in vacuo* over P₂O₅ to give (S)-(+)-COP-hfacac monomer (1.1g, 91%). A pure sample can be obtained by filtration through a short plug of silica gel with CH₂Cl₂ as eluent (in an 89% yield). It has R_F 0.63 (silica gel, hexanes-EtOAc, 80:20). It has IR (thin film) with ν_{\max} at 3061, 2964, 1629, 1598, 1509, 1475, 1258, 1208, 1150 cm⁻¹; the ¹H NMR (500MHz, CDCl₃) has δ_H at 7.54-7.56 (m, 8H, ArH), 7.19-7.29 (m, 12H, ArH), 5.95 (s, 1H, CH), 4.68 (d, *J* = 2.3Hz, 1H, CH), 4.90 (d, *J* = 2.3, 1H, CH), 4.53 (t, *J* = 2.3Hz, 1H, CH), 4.33 (*J* = 8.6, 5.3Hz, 1H, CH₂), 3.72 (t, *J* = 9.4Hz, 1H, CH₂), 3.45 (td, *J* = 9.4, 5.1Hz, 1H, CH), 2.05-2.08 (m, 1H, CH), 0.83 (d, *J* = 7.0Hz, 3H, CH₃), 0.79 (d, *J* = 6.9Hz, 3H, CH₃); and the ¹³C NMR (125MHz, CDCl₃) has δ_C at 14.9, 18.4, 29.3, 65.3, 72.3, 76.7, 78.9, 84.1, 84.8, 87.7, 90.1, 97.6, 116.6, 118.8, 126.5, 127.9, 128.9, 135.5, 173.3, 173.6, 174.1. [Kirsch et al. *J Org Chem* **69** 8101 2004, DOI: 10.1021/jo0487092; Anderson et al. *Org Synth* **84** 148 2007, DOI: 10.1522/orgsyn.084.0148.]

Enantiomer **(R)-(-)-COP-hfacac** {hexafluoroacetylacetonato[η^5 -(*R*)-(p*R*)-2-(2'-(4'-methylethyl)oxazolinyl)-cyclopentadienyl,1-*C*-3'-*N*)(η^4 -tetraphenylcyclobutadiene)cobalt[palladium]} [CASRN is given the same as preceding S-(+)-enantiomer] **C₄₄H₃₄F₆NO₃ CoPd, M 904.0, m 108-110°(dec.)**, [α]_D²⁸ -271.2, [α]₅₇₇²⁸ -277.6, [α]₅₄₆²⁸ -214.8, [α]₄₃₅²⁸ +57.7, [α]₄₀₅²⁸ +86.8 (c 1.02, CHCl₃) can be obtained and purified as for its S-enantiomer above but using the R-enantiomeric intermediates. These COP-hfacac complexes, like the related COP-acac complexes, are monomeric and more soluble than the dimeric COPs above. They can be used in a variety of solvents, e.g. THF, MeCN, which allow catalytic asymmetric allylacetimidate rearrangements to be performed at high concentrations and using as little as 1 mole% of catalyst. [Anderson et al. *Org Synth* **84** 148 2007, DOI: 10.1522/orgsyn.084.0148; Kirsch, Overman and Watson *J Org Chem* **69** 8101 2004, DOI: 10.1021/jo0487092]. The reactions of trichloroacetimidate derivatives of Z-2-alken-1-ols with phenolic nucleophiles in the presence of chiral COP-acac catalysts yield 3-aryloxy-1-alkenes in high yield (63-90%) and high enantiomeric purity (90-97% 'ee'); and are compatible with the presence of base-labile substituents in either reactant [Kirsch, Overman and White *Org Lett* **9** 911 2007, DOI: 10.1021/ol070110b].

Copper(I) [1,3-bis(2,6-di-iso-propylphenyl)imidazol-2-ylidene] chloride [1,3-bis(2,6-di-iso-propylphenyl)imidazol-2-ylidene copper(I) chloride, (IPr)=CuCl] [578743-87-0] **C₂₇H₃₆ClN₂Cu, M 487.59, m >300°**. This *NHC* (*N-heterocyclic carbene*) precatalyst is readily prepared in analytical purity in a Schlenk flask and under an argon atmosphere. Dry THF (100ml) is added to a mixture of freshly prepared Cu(I)Cl (1.0g, 10.1mmol), 1,3-bis(2,6-di-iso-propylphenyl)imidazolium chloride (4.29g, 10.1mmol, IPr.HCl see [250285-32-6]) and Na *tert*-butoxide (0.97g, 10.1mmol), and stirred mechanically under argon at ~25° for 20 hours. The mixture is then filtered through a plug of Celite, and the filtrate is evaporated *in vacuo* to give the complex as a white powder (4.59g, 9.40mmol, 94%), m >300°. Suitable crystals were grown from CH₂Cl₂/hexane solutions and subjected to single-crystal diffraction studies. The molecular structure was as expected with the imidazole C2, Cu and Cl atoms in the same plane, with the C2-Cu bond length of 1.953Å, the Cu-Cl bond length 2.089Å, and the C2—Cu—Cl angle of 180.00°, i.e. is linear. [Kaur et al. *Organometallics* **23** 1157 2004, DOI: 10.1021/om034285a]. It has IR (neat on a DiComp probe) with ν_{\max} at 2968, 2927, 2871, 1680, 1647, 1470, 1457, 809, 764, and 743 cm⁻¹; the ¹H NMR (500MHz, Me₂CO-*d*₆, downfield from TMS) has δ_H at 7.26 (s, 2H), 7.57-7.54 (dd, *J* = 15.5 and 8.0Hz, 2H), 7.43-7.41 (d, *J* = 8.0Hz, 4H), 2.70-2.65 (m, 4H), 1.32-1.31 (d, *J* = 7.0Hz, 12H), 1.27-1.25 (d, *J* = 6.5 Hz, 12H); and for ¹³CNMR and HRMS (EI) see ref. has: found 389.2962, and calculated for C₁₄H₁₆O: 389.2951 (M -CuCl) [Jurkauskas et al. *Org Lett* **5** 2417 2003, DOI: 10.1021/ol034560p.](IPr)=CuCl catalyses the conjugate reduction of α,β -unsaturated carbonyl compounds with catalytic amounts of the Cu complex, *tert*-BuONa and poly(methylhydrosiloxane) as the stoichiometric reduct-

ant in >81% yields [Jurkauskas et al. *Org Lett* **5** 2417 2003, DOI: 10.1021/ol034560p], it catalyses the methenylation of a variety of aliphatic and aromatic aldehydes and ketones in the presence of trimethylsilyldiazomethane (TMSCHN₂), Ph₃P and 2-propanol efficiently, and is a cheaper alternative to the corresponding rhodium catalyst [Lebel et al. *J Org Chem* **72** 144 2007, DOI: 10.1021/jo061781a], it catalyses the efficient hydrosilylation of ketones with Et₃SiH to yield the corresponding saturated triethylsilylethers [Kaur et al. *Organometallics* **23** 1157 2004, DOI: 10.1021/om034285a]. It catalyses the transfer of the ‘CHCO₂Et’ group (from ethyl diazoacetate) to saturated and unsaturated substrates, e.g. olefins, amines or alcohols, in very high yields [Frutos et al. *J Am Chem Soc* **126** 10846 2004, DOI: 10.1021/ja047284y], and has been used for the total synthesis of (-)-Angelastin A by catalysing the aziridinylation (using TsN=IPh) [55962-05-5] of a difficult rather electron-deficient cyclopentene intermediate [Trost & Dong *J Am Chem Soc* **128** 6054 2006, DOI: 10.1021/ja061105q].

NOTE: More recently it has been shown that the above IPr=CuCl (IPrCuCl) and its 4,5-dihydro-derivative SIPrCuCl, are readily prepared by refluxing ~1.9-2.3mmol of 1,3-bis-(2,6-diisopropylphenyl)imidazolium chloride (IPrCl, see [250285-32-6]) or 1,3-bis-(2,6-diisopropylphenyl)imidazolinium chloride (SIPrCl, see [25278-25-0]) with Cu₂O (~1.4-1.5mmol, ~95% purity, ca 0.65 equivalents) for 24 hours, then excess of Cu₂O is removed to provide the following respective yields in the solvents indicated: IPrCuCl (78% from toluene, 88% conversion; 94% from H₂O with 96% conversion; 74% from dioxane) or SIPrCuCl (88% from toluene, 99% conversion; 72% from H₂O, 78% conversion). [Cazin and coworkers: Citadelle et al. *JCS Dalton Trans* **39** 4489 2010, DOI: 10.1039/C0DT00128G; Son and coworkers: Chun et al. *Organometallics* **29** 1518 2010, DOI: 10.1021/om900768w].

Similarly the analogous [1,3-bis(2,4,6-trimethylphenyl)imidazol-2-ylidene] copper(I) chloride, [(IMes)=CuCl] and the 4,5-dihydro derivative [1,3-bis(2,4,6-trimethylphenyl)imidazolin-2-ylidene] copper(I) chloride [(SIMes)=CuCl, S is for saturated] can be prepared from the respective *IMesCl* and *SIMesCl* with 0.65 equivalents of Cu₂O by refluxing in toluene for 24 hours (in 86% and 71% yields respectively, with 100% conversion), and by refluxing in H₂O for 24 hours (in 98% and 99% yields respectively, with 100% conversion). [Cazin and coworkers: Citadella et al. *JCS Dalton Trans* **39** 4489 2010, DOI: 10.1039/C0DT00128G]. These are slightly less catalytically specific than the IPr analogues because the latter have slightly greater steric hindrance.

Copper(I) bis-[1,3-bis(2,6-di-iso-propylphenyl)imidazol-2-ylidene] tetrafluoroborate [(IPr)₂Cu⁺ BF₄⁻] [886061-48-9] C₅₄H₇₂N₄CuBF₄, M 927.5, m >300°. This copper catalyst is prepared in an oven dried septum screw capped vial (in a glove box) by mixing tetrakis(acetonitrile)copper (I) tetrafluoroborate (186mg, 0.5mmol), IPr.Cl (157mg, 0.5mmol, see [250285-32-6]) and *ter*-BuONa (96mg, 1mmol), sealing, then injecting THF (10ml) and stirring outside the box at ~25° for 6 hours. The mixture is filtered through a plug of Celite (THF), and the Cu complex salt is precipitated by adding excess hexane, filtered off, and dried *in vacuo* to give the pure white tetrafluoroborate (452mg, 92%). It has ¹H NMR (400MHz, Me₂CO-*d*₆) with δ at 7.54 (t containing a singlet at 7.55, *J* = 7.5 Hz, 8H, *p*-CH + NCH), 7.25 (d, *J* = 7.5 Hz, 8H, *m*-CH), 2.40 (septet, *J* = 6.8Hz, 8H, CH(CH₃)₂), 1.04 (d, *J* = 6.8Hz, 24H, CH₃), 0.94 (d, *J* = 6.8Hz, 24H, CH₃); and for ¹³CNMR see references. The hexafluorophosphate salt is similarly prepared in 96% yield starting from tetrakis(acetonitrile)copper (I) hexafluorophosphate [64443-05-6]. The structure of these salts was shown by X-ray crystal analysis (crystals grown from CH₂Cl₂-methyl *tert*-butyl ether solution) to consist of two *N*-heterocyclic carbenes (NHCs) attached to one Cu atom. These salts are air- and moisture- stable and are highly active catalysts for the hydroxysilylation of ketones with varying steric constraints, aldehydes and enolisable aldehydes, and esters in THF at ~25°, in the presence of Et₃SiH and *ter*-BuONa in >90% yields [Díez-González et al. *Organometallics* **25 2355 2006, DOI: 10.1021/om0600487].**

Diacetato[(*R*)-(+)- and (*S*)-(-)- 2,2'-Bis(diphenylphospheno)-1,1'-binaphthyl]ruthenium(II) [*R*- and *S*-(BINAP)₂ Ru(II) (OAc)₂] [*(R)* 325146-81-4 diacetato, (*S*) 261948-85-0 diacetato; (*R*) 132071-87-5 dichloro, (*S*) 134524-84-8; dichloro] C₄₈H₃₈O₄P₂Ru, M 841.8, m 185-187°(dec.), 186-188°(dec.), 188-190°(dec.), *R*-λ_{max} 335nm (Δε/mol⁻¹.dcm³.cm⁻¹ +4.5) and λ_{max} 450nm (Δε/mol⁻¹. dcm³.cm⁻¹ +2.5); *S*- λ_{max} 335nm (Δε/mol⁻¹.dcm³.cm⁻¹ -4.5) and λ_{max} 450nm (Δε/mol⁻¹ dcm³ cm⁻¹ -2.5); (c 6.5 x 10⁻⁵ M, CH₂Cl₂). Typically, the *S*-enantiomer is prepared in a Schlenk tube under dry argon, by dissolving (*S*)-BINAP (1.37g, 2.20mmol, see [76189-56-5]) and [RuCl₂(COD)]_n (0.56g, 2.0mmol, calc based on monomer weight, see [50982-12-2]) in dry toluene (55ml) containing Et₂NH (1.2ml, 8.6mmol), and the brown suspension is refluxed with stirring for 12 hours. The clear reddish brown solution is cooled to ~25°, the solvent is removed *in vacuo* and the

brown solid residue is dissolved in CH_2Cl_2 (40ml), filtered through a Celite pad and the filtrate is evaporated *in vacuo*. A solution of anhydrous NaOAc (0.88g, 11mmol) in *t*-BuOH (107ml) is added to the residue, stirred under reflux for 12 hours in an argon atmosphere, the solvent is removed *in vacuo* and the residue is extracted with absolute Et_2O (3 x 20ml) and evaporated again *in vacuo*. The brown residue is extracted several times with absolute EtOH (70ml total) and evaporated *in vacuo* to give a yellow-brown solid (1.58g) which is recrystallised from toluene (12ml)/hexane (30ml) mixture to afford purer complex (1.23g, 68% yield). This sample is pure enough for providing good catalytic activity for asymmetric hydrogenations. Analytical purity (to give 0.9g, 50%, 188-190° dec.) can be achieved by further recrystallisation from toluene (14ml)/hexane (13ml), and another amount (0.3g, 17%) can be recovered from the mother liquors by evaporation and recrystallisation of the residue (from 8ml of toluene and 10ml of hexane). Crystals of analytical material were subjected to X-ray structure determination that gave the required **absolute configuration** of the complex, and the circular dichroism spectrum of it and its enantiomer were consistent with the X-ray structure. It has IR (CH_2Cl_2) with ν_{max} at 1452, 1518 cm^{-1} ; and ^1H NMR (400MHz, CDCl_3 , TMS) with δ_{H} at 1.80 (s, 2 OCOCH_3), 6.47-7.84 (m, aromatic H); and for $^{13}\text{C}\{^1\text{H}\}$ NMR see references. The **R-enantiomer** can be prepared and purified in the same way.

The enantiomers of this complex catalyse the asymmetric hydrogenation of pro-chiral ketones and olefins in high yields and ‘ee’ values. [Ohta, Takaya and Noyori *Inorg Chem* **27** 566 1988, DOI: 10.1021/ic00276a025; Kumobyashi et al. *Synlett* 1055 2001, DOI: 10.1055/s-2001-14625] RhBINAP complexes with ligands other than acetate, e.g. Cl_2 (see *dichloro* in title of entry), $^*\text{C}_6\text{H}_6$, $(\text{OCO-}t\text{-Bu})_2$, $(\text{OCOCMe=CHMe})_2$ also catalyse asymmetric hydrogenations, isomerisations and asymmetric Heck reactions [Shimizu, Nagasaki and Saito *Tetrahedron* **61** 5405 2005, DOI: 10.1016/j.tet.2005.03.022.] An excellent asymmetric hydrogenation catalyst for α -(acylimino)acrylic acids to give the corresponding **chiral α -aminoacylpropionic acids** with high ‘ee’-ratios is (S)-[(BINAP)Rh⁺(OMe)₂] ClO₄⁻. It is obtained (together with norbornane) when a solution of (S)-[(BINAP)Rh⁺(norbornadiene)] ClO₄⁻ in MeOH is exposed to H₂ at room temperature. Exactly 2mols are absorbed, and it is isolated as deep red prisms whose ^1H NMR (CD_2Cl_2 , TMS) has δ_{H} at 7.50 (m) and 6.82 (BINAP), 3.42 (s, CH_3O); and ^{31}P NMR (CD_3OD with 5% H_3PO_4 in CD_3OD as external standard) has δ at 53.1 (d, $J_{\text{Rh-P}} = 206\text{Hz}$). It loses MeOH *in vacuo* to give an **equally active catalyst**. [Myashita et al. *J Am Chem Soc* **102** 7932 1980, DOI: 10.1021/ja00547a020.]

Dichloro(1,5-cyclooctadiene)palladium(II) [$\text{Pd}(\text{COD})\text{Cl}_2$] [12107-56-1] $\text{C}_8\text{H}_{12}\text{PdCl}_2$, **M 285.5, m 210° (dec)**. This air-stable catalyst can be prepared by adding 1,5-cyclooctadiene (3ml, 2.2mol, [111-78-4]) to a cooled and filtered solution of PdCl_2 (2.0g, 1 mol) and concentrated HCl (150ml, warm) in EtOH (150ml), whereby the yellow complex separates immediately. After keeping for 10 minutes the solid is filtered off, washed with Et_2O (3 x 30ml) and dried *in vacuo* (3.1g, 96%). The yellow powder is recrystallised by boiling in CH_2Cl_2 (200ml), and evaporating until the onset of crystals. **Dibromo(1,5-cyclooctadiene)palladium(II)** [$\text{Pd}(\text{COD})\text{Br}_2$] [12145-47-0] $\text{C}_8\text{H}_{12}\text{PdBr}_2$, **M 374.4, m 213° (dec)**, is obtained from the dichloride and NaBr in Me_2CO , or as above from PdCl_2/HCl in the presence of NaBr before adding the diene. [Drew et al. *Inorg Synthesis*, Coll Vol **28** 346 1990, DOI: 10.1002/9780470132593.ch89; **13** 47 1972 DOI: 10.1002/9780470132449.ch11.] **Alternatively**, a solution of sodium chloropalladite tetrahydrate (2g) in MeOH (75ml) and the diene (2ml) can be used, and the pale orange-yellow complex is collected after 1 hour and is recrystallised from glacial AcOH.

[$\text{Pd}(\text{COD})\text{Br}_2$] can also be prepared from the dichloride (0.45g) and LiBr (0.1g) in Me_2CO (20ml), boiled under reflux for 2 hours, filtered and the filtrate is evaporated to dryness in a vacuum (15mm), the residue is washed with H_2O , dried (0.22g) and is recrystallised from AcOH (orange-red needles). [Chatt et al. *J Chem Soc* 3413 1957, DOI: 10.1039/JR9570003413.] The **dichloro-complex** is sparingly soluble in cold EtOH and $^*\text{C}_6\text{H}_6$, but is more soluble in hot $^*\text{C}_6\text{H}_6$, CHCl_3 , Me_2CO , MeEtCO , $(\text{EtO})_2\text{CO}$, tetrahydrothiophene 1,1-dioxide (sulfolane) and nitrobenzene. It reacts with Me_2SO to form $\text{Pd}(\text{II})\text{Cl}_2(\text{Me}_2\text{SO})_2$. The IR (Nujol) has ν_{max} at 1489, 1419, 1337, 1088, 999, 867, 825, 794, 768 325 and 295 cm^{-1} ; and the ^1H NMR (CDCl_3) has δ at 2.69 (CH_2 protons) and 6.32 (CH=CH protons). The **dibromo-complex** is similarly **air-stable** and has similar solubilities in solvents, and for IR (Nujol) see references. The ^1H NMR (CDCl_3) has δ at 2.60 (CH_2 protons) and 6.32 ($\text{C}_2\text{H}_2=\text{CH}$ protons).

Dichloro(1,5-cyclooctadiene)platinum(II) [$\text{Pt}(\text{COD})\text{Cl}_2$]* [12080-32-9] $\text{C}_8\text{H}_{12}\text{PtCl}_2$, **M 374.2, m 220-278° (dec), 285° (dec)**. The colourless air-stable dichloro-complex is obtained by adding 1,5-cyclooctadiene* (6ml) to a warm solution (75°) of hydrated chloroplatinic acid (5g) in AcOH (15ml), stirring, and cooling to ~25° then diluting with H_2O (50ml), and the black suspension is kept at ~25° for 1 hour. The black solid is collected,

washed with H₂O (50ml), Et₂O (100ml), then suspended in CH₂Cl₂ (400ml), heated to boiling and kept at this temperature for 5 minutes. The solution is cooled, mixed with chromatographic grade silica gel (5g) and allowed to settle. The supernatant should be colourless, otherwise portions (~1g) of silica gel should be added until the supernatant is colourless. The solid is filtered off, washed with CH₂Cl₂ (2x 50ml) and the combined CH₂Cl₂ solutions (~500ml) are evaporated until crystallisation (75ml) occurs. The hot CH₂Cl₂ solution is then poured into petroleum ether (b 60-70°) producing a fine white powder which is collected, washed with petroleum ether (50ml) and dried (2.55g, 80%). Recrystallisation in the same way (dissolving in boiling CH₂Cl₂ and evaporating till crystallisation sets in) yields white macroscopic crystals. It also crystallises from AcOH (charcoal) as white needles. Its solubilities in solvents are similar to those of Pd(COD)Cl₂ above, i.e. being insoluble in most organic solvents except boiling CH₂Cl₂, CHCl₃ and AcOH, and decomposes slowly in Me₂SO (cf. related Pd compound above). Its IR (Nujol) has ν_{\max} at 1334, 1179, 1333, 1009, 1083, 871, 834, and 782 cm⁻¹; and the ¹H NMR (CDCl₃) has δ at 2.71 (CH₂ protons) and 5.62 ($J_{\text{Pt-H}} = 65\text{Hz}$, CH=CH protons). [Drew et al. *Inorg Synthesis*, Coll Vol **28** 346 1990, DOI: 10.1002/9780470132593.ch89; **13** 47 1972, DOI: 10.1002/9780470132449.ch11.]

Dibromo(1,5-cyclooctadiene)platinum(II) [Pt(COD)Br₂] [12145-48-1] C₈H₁₂PtBr₂, M 463.1, m 220-278° (dec) can be obtained from sodium chloroplatinate hydrate (4g), diene (4ml) and LiBr (2g) in EtOH (80-100ml), kept at ~20° for 2 days, and the precipitate is crystallised from AcOH (charcoal) in pale yellow needles. It is slightly more soluble than the chloro-analogue in organic solvents. [Chatt et al. *J Chem Soc* 2496 1957, DOI: 10.1039/JR9570002496; *Beilstein* 5 IV 404.]

* **Dichloro(1,3,5,7-cyclooctatetraene)Platinum(II), dichloro(dicyclopentadiene)Platinum(II) and dichloro(2,5-norbornadiene)Platinum(II)** have been prepared successfully by this procedure using the respective alicycles.

Dichloro[(1,2,3,6,7,10,11,12- η)-2,6,10-dodecatriene-1,12-diyl]ruthenium(IV), [(dichloro-2,6,10-dodecatriene-1,12-diyl)ruthenium(IV)] [12170-97-7] C₁₂H₁₈RuCl₂, M 334.2, decomposes >200° with part melting at 220°. This complex is much more stable than the corresponding Ni complex, and is prepared by bubbling butadiene for 7 hours through a solution of RuCl₃·3H₂O (0.43g, [13815-94-6]) in 2-methoxyethanol (25ml) at 90° in a hot water bath. The deep red solution deposits yellow-brown prisms when cooled. These are filtered off and recrystallised from CH₂Cl₂/petroleum ether (b 40-60°) to give the *analytically pure complex* (0.30g, 54.5%) as orange crystals. The IR shows a medium intensity band at $\nu = 1522\text{ cm}^{-1}$ assigned to the *trans* C=C co-ordinated to the Ru atom; the ¹H NMR (60MHz, CDCl₃/TMS) has δ at ~5.4 (m, w = 65Hz, 6H, vinylic and non terminal allylic H), 4.9 (d, $J = 7.5\text{Hz}$, 2H, terminal allylic H), 3.76 (d, $J = 11.2\text{Hz}$, 2H, terminal allylic H), 3.1 (m, w = 20Hz, 4H, methylene H) and 2.37 (m, w = 35Hz, 4H, methylene H); and the measured molecular weight (osmometry in 1.45% *C₆H₆ solution) is 340. [Nicholson & Shaw *J Chem Soc (A)* 807 1966, DOI: 10.1039/J19660000807.] The ¹H NMR is consistent with a Ru atom enveloped by dodeca-2,6,10-triene and co-ordinated to the two terminal allylic double bonds, i.e. butadiene trimerised about the metal atom, and this structure is confirmed by an X-Ray determination. This showed that Ru is bipyramidal, with the Cl atoms at the apices of the pyramids with three co-ordinate bonds (to the allylic and the C6=C7 double bonds) in the central plane. [Lydon et al. *Proc Chem Soc* p 421 (first page 385) 1964, DOI: 10.1039/PS9640000385].

The **dichloro-dodecatriene-ruthenium complex** is a highly efficient catalyst for the *one-pot* internal redox process that converts allylic alcohols (RC=C-C(OH)) into carbonyl compounds (RC-C-C=O, ketones or aldehydes) in the presence of CsCO₃ under N₂ with high yields and turnover frequencies (TOF) in THF or H₂O [Cadierno et al. *Chem Commun* 232 2004, DOI: 10.1039/B313069J; see also van der Drift et al. *J Organomet Chem* **650** 1 2002, DOI: 10.1016/S0022-328X(02)01150-6]. It is also an efficient catalyst for the de-protection of *N*-allylic and *N*-diallylic substrates (with ~3mol% of Ru) in aqueous solution at 90° under N₂ in very high yields (typically >95%) to give the free amine and propionaldehyde (allyl product); and can be carried out on a preparative scale [Cadierno et al. *Chem Commun* 4086 2005, DOI: 10.1039/B506788J].

Dichloro(η^5 -pentamethylcyclopentadienyl)iridium(III) dimer {di- μ -chloro-dichlorobis(η^5 -pentamethylcyclopentadienyl)diiridium (III), [Rh(η^5 -C₅Me₅)Cl₂]₂} [12354-84-6] C₂₀H₃₀Ir₂Cl₄, M 796.7, m >230°. The complex is prepared by stirring IrCl₃·xH₂O (10g, 26mmol), pentamethylcyclopentadiene (5g, 36mmol, Cp' see [4045-44-7]) and MeOH (300ml), purging the apparatus with N₂ for 5 minutes then refluxing the solution also under N₂ for 48 hours. After cooling to room temperature the orange crystals are filtered off in air through a glass sinter, the filtrate is concentrated *in vacuo* to ~50ml and cooled to give a second crop of crystals which are combined with the first crop, washed with Et₂O (3 x 50ml) and air dried. The **Ir-dimer (10.7g, 85%) can be obt-**

ained as an orange microcrystalline solid in analytical purity by recrystallisation using the minimum volume of CHCl_3 to dissolve it, filtering off insoluble material if present, and adding slowly twice that volume of hexane. The complex is stable in air at room temperature without obvious decomposition for several years. It is soluble in chlorinated solvents, much less so in Me_2CO , EtOH or hydrocarbon solvents, but is somewhat soluble in H_2O . The halogen atoms undergo metathesis and can be replaced by PF_6 and MeCN to form complexes such as $[\text{Ir}(\eta^5\text{-C}_5\text{Me}_5)(\text{NCMe})_3][\text{PF}_6]_2$. Its reactions can be studied by following the ^1H NMR (CDCl_3) signal of the single C_5Me_5 resonance at δ 1.73s. [White et al. *Inorg Synth* **29** 228 1992, DOI: 10.1002/9780470132609.ch53.] The ***Ir-dimer*** directly catalyses the C3 alkylation of a variety of indoles using aliphatic and benzylic alcohols in >80% yields [Grigg et al. *Tetrahedron* **65** 4375 2009, DOI: 10.1016/j.tet.2009.03.065]. It is used as a precursor of *N*-3-substituted-*N*-1-pyrimidyl-imidazolinium-Ir-pentamethyl-cyclopentadiene which catalyses transfer hydrogenation from isoPrOH to ketones, e.g. cyclohexanone, and imines, e.g. benzylidenedianiline, to form the respective alcohols and amines [Gnanamgari et al. *Organometallics* **28** 321 2009, DOI: 10.1021/om800821q]. It has also been used as a precursor to efficient phosphine-free ‘Ir-Cp’ chiral catalysts, e.g. with *N*-sulfonyl *S,S*-1,2-diphenylethylenediamine, for the hydrogenation of 2-substituted quinolines to give chiral 2-substituted 1,2,3,4-tetrahydroquinolines in >90% yields and >99% ‘ee’ [Li et al. *Org Lett* **10** 5265 2008, DOI: 10.1021/ol802016w].

Dichloro(η^5 -pentamethylcyclopentadienyl)rhodium(III) dimer {di- μ -chloro-dichlorobis(η^5 -pentamethylcyclopentadienyl)dirhodium (III), $[\text{Rh}(\eta^5\text{-C}_5\text{Me}_5)\text{Cl}_2]_2$ [12354-85-7] $\text{C}_{20}\text{H}_{30}\text{Rh}_2\text{Cl}_4$, M 618.1, m >230°. The complex is readily prepared by stirring $\text{RhCl}_3 \cdot 3\text{H}_2\text{O}$ (10g, 42mmol), pentamethylcyclopentadiene (6g, 44mmol, Cp’ see [4045-44-7]) and MeOH (300ml), purging the apparatus with N_2 for 5 minutes then refluxing the solution also under N_2 for 48 hours. After cooling to room temperature the dark red crystals are filtered off in air through a glass sinter, the filtrate is concentrated *in vacuo* to ~50ml and cooled to give a second crop of crystals which are combined with the first crop, washed with Et_2O (3 x 50ml) and air dried to give the ***Rh-dimer*** (11.25g, 95%). It can be obtained in ***analytical purity*** by recrystallisation using the minimum volume of CHCl_3 to dissolve it, filtering off insoluble material if present, and adding slowly twice that volume of hexane. The complex is ***stable in air*** at room temperature without obvious decomposition for several years. It is soluble in chlorinated solvents, much less so in Me_2CO , EtOH or hydrocarbon solvents, and is somewhat soluble in H_2O . The halogen atoms undergo metathesis and can be replaced by PF_6 and MeCN to form complexes such as $[\text{Rh}(\eta^5\text{-C}_5\text{Me}_5)(\text{NCMe})_3][\text{PF}_6]_2$. Its reactions can be studied by following the ^1H NMR (CDCl_3) signal of the single C_5Me_5 resonance at δ 1.60s. [White et al. *Inorg Synth* **29** 228 1992, DOI: 10.1002/9780470132609.ch53.] In the presence of base the rhodium dimer is a good catalyst for the hydrogenation of olefins [Gill et al. *JCS Dalton Trans* 617 1978, DOI: 10.1039/DT9780000617], for hydrosilylation of olefins [Millan et al. *J Mol Catal* **26** 89 1984, DOI: 10.1016/0304-5102(84)85022-1; Millan et al. *JCS Chem Commun* 673 1981, DOI: 10.1039/C39810000673], and for the disproportionation of aldehydes [Cook et al. *JCS Dalton Trans* 2342 1981, DOI: 10.1039/DT9810002342].

Dicobalt octacarbonyl (cobalt carbonyl) [10210-68-1] $(\text{CO})_8\text{Co}_2$, M 341.9, m 51°, 51-52° (dec), d 1.87. $\text{Co}_2(\text{CO})_8$ has been identified as the cobalt catalyst in hydroformylation, hydrogenation and homologation reactions involving H_2 and CO (synthesis gas). The earlier preparation [Gilmont et al. *Inorg Synth* **2** 238 1946, DOI: 10.1002/9780470132333.ch76] has been improved due to the availability of steel pressure vessels and CO under pressure. Thus ***Raney Co*** (4 to 6g) was placed with Et_2O (145ml) under CO at 3200psi in a steel pressure bomb and heated with shaking for 5 or 6 hours at 150°. The pressure during the heating process dropped from 4900 to 4300psi and then to 2200psi when the bomb was cooled to ~25°. The insoluble material was removed by centrifugation, and the reddish-brown clear solution (160ml, which includes washings and transfers) contained 8.9g of $\text{Co}_2(\text{CO})_8$. It is kept in a closed pressure-adjusted Pyrex bottle and aliquots siphoned as required because it decomposes slowly to liberate CO in an open vessel and should be stored in the cold. The solubility of the catalyst in Et_2O at ambient temperature is 7.5g per 100ml of solution. Cobalt on Kieselguhr ‘Co-100 powder’ that contains 12 to 15% of Co can be substituted for ***Raney Co***. [Adkins & Kresk *J Am Chem Soc* **70** 383 1948, DOI: 10.1021/ja01181a119.] The above preparation can also be carried out in $^*\text{C}_6\text{H}_6$ which has the advantage of being less volatile than Et_2O , and in which reactions could be carried out at lower temperatures. It has been kept at a concentration of 1.0×10^{-2} M in 50ml of $^*\text{C}_6\text{H}_6$. [Adkins & Kresk *J Am Chem Soc* **71** 3051 1949, DOI: 10.1021/ja01177a032.] It forms orange-brown ***air-sensitive*** crystals on recrystallisation from *n*-hexane under a carbon monoxide atmosphere. It has been sublimed *in vacuo* (orange platelets), and is available commercially as a solid moistened with 5-10% of hexane. It is insoluble in H_2O , but soluble in organic solvents

such as alcohols, Et₂O, *C₆H₆, CS, slowly decomposed by HCl, and H₂SO₄, but rapidly by HNO₃. [Wender et al. *J Am Chem Soc* **71** 4160 1949, DOI: 10.1021/ja01180a520; Ojima et al. *J Am Chem Soc* **109** 7714 1987, DOI: 10.1021/ja00259a020; see Hileman in *Preparative Inorganic Reactions*, Ed. Jolly, Vol **1** p101 1987].

Carbon monoxide is **VERY POISONOUS** as it complexes with haemoglobin. Great care should be exercised when using the catalyst as well as CO, and all operations should be carried out in an efficient fume cupboard. The **TOXIC LEVEL** of CO is 50ppm (~55mg/m³). **The ANTIDOTE should be at hand and always available in laboratories using CO; and staff should be trained to administer it.** On completion of reactions the autoclave should be filled with H₂ at 1000-1300psi and heated at 110-135° for 45 minutes where cobalt compounds are transformed to the metal. If the product of reaction does not contain oxidisable compounds, the mixture is heated in air on a steam bath when copious evolution of CO occurs and the metal is deposited as a cobalt mirror. A third procedure to decompose Co₂(CO)₈ and related CO compounds is to shake or stir the mixture with dilute H₂SO₄ until evolution of gas ceases. [Wender et al. *J Am Chem Soc* **72** 4375 1950, DOI: 10.1021/ja01166a012.]

2,6-Di-iso-propylphenylimino-neophylidene[(S)-(-)-BIPHEN]molybdenum(VI) {(S)-Schrock-Hoveyda Catalyst; Mo(*N*-2,6-di-iso-Pr₂C₆H₃)(CHCMe₂Ph)(*syn*-(S)-(-)-*tert*-Bu₂Me₄(BIPHEN); molybdenum, [(*S*- or *R*-) 3,3'-bis(1,1'-dimethylethyl)-5,5',6,6'-tetramethyl[1,1'-biphenyl]-2,2'-diolato(2-)-κO,κO'] [2,6-bis(1-methylethyl)benzenaminato(2-)](2-methyl-2-phenylpropylidene)- } [(*S*) 205815-80-1; (*R*) 329735-77-5; (*RS*) 300344-02-9] C₄₆H₆₁NO₂Mo, **M 755.9**. The key intermediate is **2,6-di-iso-propylphenylimino-neophylidene-bistriflate-dimethoxyethane-molybdenum (VI) complex** {molybdenum, [2,6-bis(1-methylethyl)benzeneaminato(2-)] [1,2-dimethoxy-κO]ethane} [2-methyl-2-phenylpropylidene]bis(1,1,1-trifluoromethanesulfonato-κO)-; [Mo(=N-2,6-isoPr₂C₆H₃)(=CHMePh)-(OTf)₂-DME] [126949-63-1] which is prepared in three steps. **Firstly** (preferred method): A solution of MoO₂Cl₂(THF)₂ (10.2g, 29.0mmol, see [13637-68-8, 556907-19-8; 12081-12-8]) in DME (200ml) at -30° is stirred vigorously while the following are added sequentially (i) a solution of 2,6-lutidine (12.4g, 116mmol) in DME (10ml at -30°) during 3-5 minutes, (ii) a solution of Me₃SiCl (31.5g, 290mmol, see [75-77-4]) in DME (40ml, at -30°) over a period of 3-5 minutes, and (iii) a solution of 2,6-di-iso-propylaniline (10.3g, 58mmol, see [24544-04-5]) in DME (15ml at -30°) during 15 minutes. The colour of the solution alters from pale yellow to bright red-orange to deep red-orange as a solid separated. The mixture is stirred as it warmed to room temperature during 6 hours. The mixture is then heated to 50° for 5 hours, filtered through Celite, to remove 2,6-lutidine hydrochloride, which is washed with Et₂O until the washings are clear; all filtrates are combined and evaporated *in vacuo* to give **bis(2,6-di-iso-propylphenylimino)dichloro-dimethoxyethane molybdenum** [Mo(=N-2,6-isoPr₂C₆H₃)₂Cl₂DME] (16.7g, 27.5 mmol, 95%) as a brick red solid which is almost of analytical purity (elemental C, H, N and Cl for MoC₂₈H₄₄Cl₂N₂O₂), and used in the next step. Its ¹H NMR (400MHz, C₆D₆, TMS) has δ at 7.01 (d, 4, arom H_m), 6.89 (t, 2, arom H_p), 4.29 (sept, 4, CHMe₂), 3.44 (s, 6, MeOCH₂CH₂OMe), 3.18 (s, 4, MeOCH₂CH₂OMe), 1.25 (d, 24, CHMe₂) and for ¹³C NMR see references. **Secondly**: Neophyl magnesium chloride (100ml, 0.5M, 50mmol, see [35293-35-7]) in Et₂O is added dropwise to a stirred solution of the preceding complex [Mo(=N-2,6-isoPr₂C₆H₃)₂Cl₂DME] (15.1g, 25mmol) in Et₂O (250ml) at -30° (all the complex need not have dissolved). The colour of the solution alters from red to orange as MgCl₂ separates, the mixture is allowed to thaw to 25° and is stirred at this temperature for 3 hours. The mixture is filtered through Celite, the filtrate is concentrated *in vacuo*, kept at -40° to provide orange crystals (usually three crops) of **almost analytically pure** (elemental C, H and N for MoC₄₄H₆₀N₂) **bis(2,6-di-iso-propylphenylimino)-bis(neophyl) molybdenum** [Mo(=N-2,6-isoPr₂C₆H₃)₂(-CH₂C Me₂Ph)₂] (14.3g, 80%). Its ¹H NMR (400MHz, C₆D₆, TMS) has δ at 7.45 (d, 4H, aromatic), 7.24 (t, 4H, aromatic), 7.10 (t, 2H, aromatic), 7.00-6.92 (m, 6H, aromatic), 3.65 (sept, 4H, CHMe₂), 1.72 (s, 4H, CH₂CMe₂Ph), 1.49 (s, 12H, CH₂CMe₂Ph), 1.11 (d, 24H, CHMe₂) and for ¹³C NMR see references. **Thirdly**: A solution of triflic acid (4.42g, 29.45mmol) in DME (15ml) is added slowly to an orange solution of the preceding Mo(=N-2,6-isoPr₂C₆H₃)₂(-CH₂C Me₂Ph)₂ (14.2g, 20mmol) in DME (150ml) at -30°, then allowed to thaw to ~25°, stirred overnight, and the deep yellow solution is evaporated to dryness *in vacuo*. The yellow residue is extracted with chilled (~0°) toluene (100ml), filtered, evaporated *in vacuo*, and the dark yellow residue is recrystallised from Et₂O to provide the desired **almost analytically pure** (elemental C, H, and N for MoC₂₈H₃₉F₆NO₈S₂) key intermediate **2,6-di-iso-propylphenylimino-neophylidene-bis(triflate)-dimethoxy-ethane molybdenum(IV) complex** [Mo(=N-2,6-isoPr₂C₆H₃)(=CHMePh)(OTf)₂DME] (5.94g, 76%, in three crops). Its ¹H NMR (400MHz, C₆D₆, TMS) has δ at 14.45 (s, 1H, CHCMe₂Ph), 7.57 (d, 2H, aromatic), 7.18 (t, 2H, aromatic), 6.97-6.89 (m, 4H, aromatic), 3.84 (sept, 2H, CHMe₂), 3.73 (s, 3H, OCH₃), 3.18 (M, 2H, OCH₂), 2.84-2.78 (m, 5H,

OCH₃, OCH₂), 1.91 (s, 6H, CHCMe₂Ph), 1.37 (d, 6H, CHMe₂), 1.21 (d, 6H, CHMe₂) and the ¹³C NMR (100MHz, C₆D₆, TMS) has δ at 328.4, 152.1, 151.8, 148.7, 130.6, 128.7, 126.6, 124.3, 72.8, 70.0, 65.7, 61.9, 58.8, 31.1, 28.3, 25.6, 22.8. [Schrock et al. *J Am Chem Soc* **112** 3875 1990, DOI: 10.1021/ja00166a023.]

The **Schrock-Hoveyda catalyst** is prepared in a glove box or in Schlenk equipment under an argon or N₂ atmosphere by adding solid KH (360mg, 9mmol, 3 equivalents) in portions to a stirred solution of H₂[S-BIPHEN] (1.06g, 3mmol, see [*R*- 329735-68-4, *S*- 205927-03-3, *RS*- 101203-31-0] in Part 2) in THF (100ml), whereby H₂ gas evolves while the K₂[BIPHEN] salt is being formed. After stirring for 18 hours at ~25°, the suspension is filtered through Celite to remove excess of KH, and the key intermediate above [Mo(=N-2,6-isoPr₂C₆H₃)(=CHMePh)(OTf)₂DME] (2.24g, 2.83mmol, 0.94 equivalents) is added as a solid during 2 minutes. The mixture is stirred for 3 hours at ~25°, the volatiles are removed *in vacuo*, and the residual red powder is treated with *C₆H₆ (40ml), and the slurry is filtered through Celite to remove insoluble CF₃SO₃K and washed with *C₆H₆ until the washings are no longer red in colour. The red *C₆H₆ solutions are combined, evaporated *in vacuo*, and the spongy red solid is recrystallised from Et₂O (~4ml) to give the desired catalyst, mainly in the *syn* configuration at the alkylidene group, as dark orange red crystals (1.35g, 64%). Its ¹H NMR (400MHz, *C₆D₆, TMS) has δ at 10.98 (s, 1H, *J*_{CH} = 123Hz, *syn* alkylidene C=CHCMe₂Ph), 7.42 (m, 3H, biphenyl and Ph CHs), 7.16 (m, 3H, biphenyl and Ph CHs), 7.05 (br t, *J* = 7.6Hz, 1H, aromatic), 6.92 (s, 3H, aromatic), 3.70 (heptet, *J* = 7.0Hz, 2H, *iso*-CHMe₂), 2.13 (s, 3H), 2.15 (s, 3H), 1.85 (s, 3H), 1.74 (s, 3H), 1.66 (s, 3H) {these signals 2.13–1.66 ppm are from 4 biphenyl-Mes and one methyl from CMeMePh}, 1.59 (s, 9H, *tert*-CMe₃), 1.54 (s, 9H, *tert*-CMe₃), 1.14 (d, *J* = 7.0Hz, 6H, *iso*-CHMe₂), 1.13 (s, 3H, CMeMePh), 0.90 (d, *J* = 7.0Hz, 6H, *iso*-CHMe₂), and for ¹³C{¹H} NMR see references. [Alexander et al. *J Am Chem Soc* **120** 4041 1998, DOI: 10.1021/ja974353i.] See also a *slightly modified preparation* in which the red spongy powder is dissolved in Et₂O (18ml), transferred to a 20ml vial, kept uncapped in a well purged glovebox (with N₂) until the volume decreased to 5ml, the red solution is decanted from the red crystal blocks which were washed with cold Et₂O and dried *in vacuo* to provide a 72% yield of *analytically pure* (elemental C, H, and N for MoC₄₆H₆₁NO₂) *syn*-(*S*)-catalyst. The X-ray crystal structure proved the absolute configuration of the biphenolate ligand. Related complexes in which the 2,6-di-*iso*-propylphenylimino ligand is replaced by various other ligands have been similarly prepared. [Alexander et al. *Organometallics* **19** 3700 2000, DOI: 10.1021/om000336h.] Schrock-Hoveyda catalysts with *R*-BIPHEN and *racemic*-BIPHEN also form red crystals, and have been similarly prepared. All are *air and moisture sensitive*, and have to be stored cold and in the dark. NMR data revealed that the *syn*-alkylidene conformer predominates in solution with a *syn-anti* exchange rate of ~1 s⁻¹. However, evidence from ROM (ring-opening metathesis) polymerisation suggests that reactions with the *anti* conformer are catalytically more active than with the *syn* conformer by some orders of magnitude [Oskam & Schrock *J Am Chem Soc* **115** 11831 1993, DOI: 10.1021/ja00078a023].

The related catalyst where the *tert*-butyl groups in the biphenol ligand are replaced by 1-adamantyl groups Mo(*N*-2,6-di-*iso*-Pr₂C₆H₃)(CHCMe₂Ph)(*syn*-(*S*)-(-)-*tert*-Bu₂Me₄(BIAD)) has been prepared in 54% yield as orange crystals from the above key intermediate [Mo(=N-2,6-isoPr₂C₆H₃)(=CHMePh)(OTf)₂DME] and (*S*)-H₂(BIAD) (see in Part 2) except that benzylpotassium was used instead of KH to prepare the phenolate salt. X-ray crystallography of the catalyst also confirmed the *S*- absolute configuration of (-)-H₂(BIAD). [Alexander et al. *Organometallics* **19** 3700 2000, DOI: 10.1021/om000336h; and for the X-ray crystallographic structures of related Mo-2,2'-dihydroxy-1,1'-biaryls see Totland et al. *Macromolecules* **29** 6114 1996, DOI: 10.1021/ma960351r.]

These and related *chiral molybdenum complexes* are remarkable catalysts that effect efficient asymmetric enantioselective ring-closing olefin metathesis (RCM) [Alexander et al. *J Am Chem Soc* **120** 4041 1998, DOI: 10.1021/ja974353i; Weatherhead et al. *Tetrahedron Lett* **41** 9553 2000, DOI: 10.1016/S0040-4039(00)01690-7], ring-opening metathesis/cross metathesis (ROM/CM) [La et al. *J Am Chem Soc* **123** 7767 2001, DOI: 10.1021/ja010684q], as well as ring-opening olefin metathesis (ROM) polymerisation [Totland et al. *Macromolecules* **29** 6114 1996, DOI: 10.1021/ma960351r] reactions. These complexes catalyse olefin metathesis reactions in which the starting olefin is a racemic mixture, and in the reaction process *kinetic resolution* occurs, whereby the products, and any unreacted starting material, are of one chirality, i.e. the asymmetric centre in the racemic mixture is converted into one enantiomer at the expense of the other enantiomer. The above Mo-(*S*)-[BIPHEN] complex also catalyses efficient and enantioselective *desymmetrisation* reactions, e.g. (±)-3-allyloxy-2,4-dimethylpenta-1,4-diene and its 1,5-dimethyl derivative (4-allyloxy-3,5-dimethyl-hepta-2,5-diene) are cyclised to 2-(prop-2'-en-2'-yl)-3(*R*)-methyl-2,5-dihydrofuran and 2-(but-2-en-2-yl)-3(*R*)-methyl-2,5-dihydro-furan in the absence of solvent, 1-2 mol% catalyst, at 22° in 5 minutes

in 85% and 93% yields, and 93% ‘ee’ and 99% ‘ee’ respectively [La et al. *J Am Chem Soc* **120** 9720 1998, DOI: 10.1021/ja9821089]. [For olefin metathesis review see Grubbs *Tetrahedron* **60** 7117 2004, DOI:10.1016/j.tet.2004.05.124; and R.H. Grubbs (Ed.), *Handbook of Metathesis*, Vols 1-3, Wiley-VCH, 2003. ISBN 3527306161.]

Ferric (III) acetylacetonate [Fe(acac)₃, iron(III) tris(2,4-pentadionate)] [14024-18-1] (C₅H₇O₂)₃Fe, **M 353.2, m 180-182°, 181.3-182.3°, d²⁵ 5.24g/ml**. When recrystallised twice from *benzene/petroleum ether, it has **m 181.3-182.3°** corr [Finn et al. *J Chem Soc* 1254 1938, DOI: 10.1039/JR9380001254]. However, when recrystallised from EtOH or Et₂O it has **m 179°** [Hantzsch & Desch *Justus Liebigs Ann Chem* **323** 1 1902, DOI: 10.1002/jlac.19023230102]. Recrystallisation from absolute EtOH also gives material with **m 159.5°** [Emmert & Jacob *Chem Ber* **67** 286 1934, DOI: 10.1002/cber.19340670225]. Dry it for 1 hour at 120°. [Beilstein **1** I 404, **1** II 836, **1** IV 3675; for applications see Fieser **4** 268, **6** 304.] Fe(acac)₃ catalyses a large variety of chemical reactions such as aromatic substitution, cross-coupling, Friedel-Crafts etc [see B. Pleitker (ed), *Iron Catalysis in Organic Chemistry* Wiley-VCH, 2008, ISBN 978-3-527-31927].

Ferrous (II) acetylacetonate [Fe(acac)₂, iron(II) bis(2,4-pentadionate)] [14024-17-0] (C₅H₇O₂)₂Fe, **M 254.1, m 175°**. For its catalytic properties see B. Pleitker (ed), *Iron Catalysis in Organic Chemistry* Wiley-VCH, 2008, ISBN 978-3-527-31927, and for the preparation see Chapter 4, ‘Metal-Organic Compounds’. [Beilstein **1** III 3122, **1** IV 3676]

Iridium(I) bis(1,5-cyclooctadiene) tetrafluoroborate complex [bis(1,5-cyclooctadiene)iridium (I) tetrafluoroborate, Ir(COD)₂⁺. BF₄⁻] [35138-23-9] (C₈H₁₂)₂Ir⁺. BF₄⁻, **M 495.4, m ~190° (dec)**. This dark red iridium complex is prepared and purified exactly as for the corresponding Rh(COD)₂⁺.BF₄⁻ [35138-22-8, below] from the dimer {[Ir(COD)Cl]₂} [12112-67-3] below, Herde et al. *Inorg Synth* **15** 18 1974, DOI: 10.1002/9780470132463.ch5}] with COD and AgBF₄ in 93% yield. [Schenck et al. *Inorg Chem* **24** 2334 1985, DOI: 10.1021/ic00209a003.]

Iridium(I) chloride 1,5-cyclooctadiene complex dimer {chloro(1,5-cyclooctadiene)iridium dimer, di-μ-chlorobis[(1,2,5,6-η)-1,5-cyclooctadiene]diiridium (I), [Ir(COD)Cl]₂} [12112-67-3] (C₁₆H₂₄)Ir₂Cl₂, **M 671.7, m 205°(dec)**. The complex is an orange-red, air-stable solid that is soluble in *C₆H₆ and CHCl₃, less so in Me₂CO and insoluble in Et₂O. It can be prepared from IrCl₃·3H₂O (3g) in 95% EtOH (34ml), H₂O (17ml) and cycloocta-1,5-diene (6ml), through which is bubbled (with magnetic stirring) a slow stream of N₂ while boiling under reflux for 24 hours, during which time a brick-red product precipitates. The mixture is cooled and Ir(COD)Cl₂ is filtered off, washed with ice-cold MeOH to remove unreacted COD and dried *in vacuo* at 25° for 8 hour (yield 1.5g, 72%, decomp >200°). [Herde et al. *Inorg Synth* **15** 18 1974, DOI: 10.1002/9780470132463.ch5.] Alternatively, to a mixture of H₂O (100ml), isoPrOH (35ml) and 1,5-cyclooctadiene (18ml) is added (NH₄)₂IrCl₆ (20g), and the mixture is refluxed under N₂ for 18 hours when the colour changes to orange-red, and a red or orange solid separated on cooling. This is filtered off (frit), and washed with EtOH (2 x 5ml at 0°) to give the **complex in high purity** (14.0g, 92%). Recrystallisation (if required) is best carried out by slowly adding an equal volume of EtOH to a saturated stirred solution of the solid in CH₂Cl₂, followed by gentle removal of half of the mixed solvent under reduced pressure. The stirred solution is cooled spontaneously to -30° during the process and the complex is filtered off, washed with EtOH (2 x 5ml at 0°) and dried *in vacuo*. It is identified by the characteristic IR bands (Nujol) at 907, 970, 980 and 1002 cm⁻¹; and the vinyl CH resonance in the ¹H NMR (CDCl₃) spectrum at δ 4.3. [Crabtree et al. *Synth React Inorg Met-Org Chem* **12** 407 1982, DOI:10.1080/00945718208063124.] [Fieser **5**, 113.] It is the metal complex precursor with allyamines for asymmetric allylic substitutions; for resting state and kinetic studies see Markovic and Hartwig [*J Am Chem Soc* **129** 11680 2007, DOI: 10.1021/ja074584h].

Iridium(I) (1,5-cyclooctadiene)-η⁵-(indenyl) {(1,5-cyclooctadiene)-η⁵-(indenyl)iridium (I), [(COD)(C₉H₇)Ir(I)], [(Ind)Ir(I) (COD)]} [102525-11-1] (C₉H₇)Ir(C₈H₁₂), **M 414.5, m 126-131°**. The complex is obtained by adding solid indenyllithium (0.73g, 5.98mmol) to a solution of the preceding dimer [Ir(COD)Cl]₂ (2.0g, 2.98mmol) in THF (50ml) and stirring at ~20° for 1 hour. The solvent is removed *in vacuo*, extracted with pentane (6 x 50ml) and the combined extracts are slowly evaporated to deposit pale yellow crystals (2.15g, 86%).

It has ^1H NMR (200MHz) with δ at 1.65-1.86 (m, 8H, COD CH_2), 3.99 (m, 4H, COD CH), 5.00 (d, 2H, indenyl H1/H3), 5.76 (t, 1H, indenyl H2), 6.98-7.16 (m, 4H, indenyl H4-H7); and ^{13}C NMR (50MHz) with δ at 34.36 (COD CH_2), 51.35 (COD CH), 72.80 (indenyl C1/C3), 85.15 (indenyl C2), 110.23 (indenyl C8/C9), 121.75 and 124.89 (indenyl C4-C7). [Merola & Kacmarcik *Organometallics* **8** 778 1989, DOI: 10.1021/om00105a031; Abad *Inorg Chim Acta* **121** 213 1986, DOI: 10.1016/S0020-1693(00)84522-3; Crabtree et al. *Synth React Inorg Met-Org Chem* **12** 407 1982, DOI: 10.1080/00945718208063124; Naderer et al. *Organomet Chem* **518** 181 1996, DOI:10.1016/0022-328X(96)06144-X.] [Uson et al. *Inorg Synth* **23** 126 1985, DOI: 10.1002/9780470132548.ch25.] **[(Ind)Ir(I) (COD)]** is a powerful C-H activation catalyst for preparing phenols from arenes in the presence of pinacolborane and bis(diphenylphosphino)ethane that borylate arenes to the intermediate pinacol arylboryl esters which are then converted to the respective phenols by reaction with *oxone* [$2\text{KHSO}_5 \cdot \text{KHSO}_4 \cdot \text{K}_2\text{SO}_4$] in Me_2CO at 25° within ~ 7 minutes in 51 to 88% yields depending on the substituents [Maleczka et al. *J Am Chem Soc* **125** 7792 2003, DOI: 10.1021/ja0349857.]

Iridium(I) μ -chloro-bis(cyclooctene) dimer {di- μ -chlorotetrakis(cyclooctene)diiridium(I), $[\text{Ir}(\text{COE})_2\text{Cl}]_2$ [12246-51-4] $\text{C}_{32}\text{H}_{56}\text{Ir}_2\text{Cl}_2$, M 896.1, m $150^\circ(\text{dec})$, $160\text{--}165^\circ(\text{dec})$. This solid yellow iridium complex is air sensitive and decomposes to a dark green solid that eventually turns black. However, it can be stored *in vacuo* in a desiccator for prolonged periods of time. It is preferable to store it in aliquots in sealed ampoules under an inert atmosphere or in a vacuum. It is soluble in Me_2CO , CHCl_3 and $^*\text{C}_6\text{H}_6$, but it oxidises more readily in solution than in the solid state, and the necessary precautions to strictly exclude air should be exercised.

It is prepared by stirring (stirrer bar) a mixture of $\text{IrCl}_3 \cdot 3\text{H}_2\text{O}$ (2.0g), *iso*-propanol (22ml), H_2O (8ml) and cyclooctene (40ml), while a slow stream of dry N_2 is bubbled through the solution, and refluxed at 78° for 3 hours. During this time the colour of the solution changes from dark red to orange-yellow and the complex separates from the solution which is cooled to $\sim 25^\circ$. It is collected rapidly by filtration, washed rapidly with ice-cold MeOH to remove excess of cyclohexene (this is done preferably in a dry box under N_2). Drying *in vacuo* at 25° for 4 hours gives pure di- μ -chlorotetrakis(cyclooctene)diiridium(I) (1.5g, 59%), which decomposes at 150° and has the correct elemental analysis for C, H, Cl and Ir. [Herde et al. *Inorg Synth* **15** 18 1974, DOI: 10.1002/9780470132463.ch5; Herd $\acute{\text{e}}$ & Senoff *Inorg Nucl Chem Lett* **7** 1029 1971, DOI: 10.1016/0020-1650(71)80023-5.] By using $(\text{NH}_4)_3\text{IrCl}_6$ (6g) suspended in an oxygen-free suspension of *iso*-PrOH (30ml) and H_2O (90ml), and cyclooctene (12ml), and refluxing with stirring for 3-4 hours under N_2 , followed by cooling, an orange oil which solidifies is formed and is collected, washed with cold EtOH, and allowed to crystallise from EtOH, collected, dried *in vacuo*, all under N_2 , to give the $[\text{Ir}(\text{COE})_2\text{Cl}]_2$ (80-92% yield) as yellow needles. It should be stored under N_2 at room temperature. [Van Der Ent et al. *Inorg Synth* **28** 90 1989, DOI: 10.1002/9780470132593.ch23.]

Although this compound is air sensitive, with the correct stoichiometry, it forms a less sensitive stable tris-boryl complex with pinacolborane (Bpin) and 4,4'-di-*tert*-butyl-2,2'-dipyridine (dtbpy) which has been crystallised, and its X-ray analysis shows that it has the composition $[\text{Ir}(\text{dtbpy})(\text{COE})(\text{Bpin})_3]^\ddagger$. These complexes are catalytically active in the C-H borylation of arenes. Thus dissolution of the latter complex in $^*\text{C}_6\text{D}_6$ (or $^*\text{C}_6\text{H}_6$) provided $^*\text{C}_6\text{D}_5$ -Bpin within minutes at room temperature and in 80% yield. Kinetic studies revealed a *kinetic isotope effect* $k_{\text{H}}/k_{\text{D}}$ of 3.8. The iridium complexes containing dtbpy are easily prepared and are air-stable. The reaction of bis(pinacolato)diboron (B_2pin_2) and $\frac{1}{2}[\text{IrCl}(\text{COE})_2]_2/\text{dtbpy}$ (5 mol% of Ir) in 60 equivalents of $^*\text{C}_6\text{H}_6$ at 25° for 4.5 hours gives *Ph-Bpin* in 83% yield; and with $\frac{1}{2}[\text{IrCl}(\text{COE})_2]_2/\text{dtbpy}$ (0.02 mol% of Ir) at 100° for 16 hours gives *Ph-Bpin* in 80% yield with 8000 turnovers. These catalysts provide a simple and direct route for the synthesis of arylboronates—which were previously obtained by transmetalation with aryl lithium or arylmagnesium reagents and trialkylborates. [Ishiyama et al. *J Am Chem Soc* **124** 390 2002, DOI: 10.1021/ja0173019; cf. Ishiyama et al. *Angew Chem Int Ed* **41** 3056 2002, DOI: 10.1002/1521-3773(20020816)41.]

$^\ddagger[\text{Ir}(\text{dtbpy})(\text{COE})(\text{Bpin})_3]$ is prepared in a dry box under N_2 : a glass flask containing a mixture of $[\text{Ir}(\text{COE})_2\text{Cl}]_2$ (350mg, 390mmol), dtbpy (209mg, 778mmol) and B_2pin_2 (494mg, 1.95mmol), to which is added mesitylene (or xylene) (50ml) and heated with very slow stirring at 50° for 5 hours. The solvent is evaporated off at $\sim 25^\circ$, the residue is dissolved in Et_2O and allowed to evaporate slowly at $\sim 25^\circ$ to afford red cubes (blocks) of the complex in 28% yield (52mg). Its ^1H NMR (400MHz, cyclohexane- d_{12}) has δ at 1.15 (s, 12H), 1.17 (s, 12H), 1.18 (s, 12H), 1.33 (m, 12H), 1.41 (s, 18H), 3.74 (d, 2H, $J = 10.8\text{Hz}$), 7.09 (dd, 2H, $J = 6.4, 2.0\text{Hz}$), 7.93, (s, 2H), 9.45 (d, 2H, $J = 6.4\text{Hz}$) from Me_4Si ; and its ^{11}B NMR (128MHz, CDCl_3) has δ at 37 from external BF_3OEt_2 . [Ishiyama et al. *J Am Chem Soc* **124** 390 2002, DOI: 10.1021/ja0173019.]

Iridium(I) (methoxy)(1,5-cyclooctadiene) dimer {bis(η^4 -1,5-cyclooctadiene) di- μ -methoxy diiridium (I) dimer, $[(\text{COD})(\text{OMe})\text{Ir}(\text{I})]_2$ [12148-71-9] $[(\text{C}_8\text{H}_{12})(\text{CH}_3\text{O})\text{Ir}]_2$, **M 662.9**, **m 154-179°** (dec). Deoxygenated solvents should be used and reactions should be carried out under N_2 or argon. The methoxylated catalyst is prepared in Schlenk equipment in an inert atmosphere by adding a suspension of $[\text{Ir}(\text{COD})\text{Cl}]_2$ (148mg, 222mmol, [12112-67-3]) in MeOH (10ml) to KOH (25mg, 445mmol) in MeOH (5ml), whereby the colour becomes orange-red and a yellow crystalline solid is formed. Stir for 30 minutes then add H_2O (40ml), collect the solid, wash it with H_2O (6 x 5ml), and dry it over P_2O_5 *in vacuo* to give $[\text{Ir}(\text{COD})(\text{OMe})]_2$ (124mg, 85%, decomposing at 145-155° becoming black at 155°) as a yellow air-stable solid. It is soluble in chlorinated solvents to give solutions that are air sensitive. It is soluble in MeOH, Me_2CO , hexane, $^*\text{C}_6\text{H}_6$ and Et_2O but insoluble in H_2O . For IR see references; the ^1H NMR (CDCl_3) has δ at 3.57 (8H, vinyl), 2.22 (8H, allylic H), 1.45 (8H, allylic H) and 3.28 (sharp singlet, 6H, MeO). [Uson et al. *Inorg Synth* **23** 126 1981, DOI: 10.1002/9780470132548.ch25.] *Alternatively*, $[\text{Ir}(\text{COD})(\text{OMe})]_2$ can be prepared by boiling $[\text{Ir}(\text{COD})\text{Cl}]_2$ (1g) in MeOH (40ml) under reflux with anhydrous Na_2CO_3 (0.6g) for 1 hour, filtering hot, cooling, and the yellow plates of the desired complex are filtered off, washed with MeOH and dried *in vacuo*. [cf. Rhodium analogue in Chatt & Venanzi *J Chem Soc* 4735 1957, DOI: 10.1039/JR9570004735.] It has been used successfully, by preparing it *in situ*, on adding NaOMe to the solution containing $\text{Ir}(\text{COD})\text{Cl}]_2$ (see below).

$[\text{Ir}(\text{I})(\text{OMe})(\text{COD})]_2$, like the preceding complex, is a powerful C-H activation catalyst for preparing phenols from arenes in the presence of pinacolborane and 3,3'-di-*tert*-butyl-1,1'-bipyridyl which borylates arenes to the intermediate pinacol arylboronates which are then converted to the respective phenols by reaction with *oxone* [$2\text{KHSO}_5 \cdot \text{KHSO}_4 \cdot \text{K}_2\text{SO}_4$] in Me_2CO at 25° within 7 minutes in ~70% yields depending on the substituents [Maleczka et al. *J Am Chem Soc* **125** 7792 2003, DOI: 10.1021/ja0349857.] The catalyst, prepared *in situ*, in the presence of bipyridines has been used as in the previous reference for borylation of arenes in high yields and should be useful for preparing arylboronates [Ishiyama et al. *Angew Chem Int Ed* **41** 3056 2002, DOI: 10.1002/1521-3773(20020816)41.]

Iridium(III) (1,1,1,5,5,5-trifluoroacetylacetonato)(bis-cyclooctene) [(hfacac)(COE) $_2$ Ir] [58616-58-3] $(\text{C}_{16}\text{H}_{28})(\text{C}_5\text{H}_2\text{F}_6\text{O}_2)\text{Ir}$, **M 620.7**, **m 96-97°**. It is prepared by adding $\text{Ti}(\text{hfacac})^\dagger$ (600mg) to a suspension of $[\text{Ir}(\text{COE})_2\text{Cl}]_2$ (920mg, see above) in pentane (100ml) at ~ 25°, and stirring for 3 hours then filtering. The red filtrate is concentrated to 10ml, kept at -30° for 24 hours and the red crystals of $[(\text{hfacac})(\text{COE})_2\text{Ir}]$ (786.5g, 70%) are collected and dried *in vacuo* to give *analytically pure complex* (C and H). It is a *monomeric* complex in $^*\text{C}_6\text{H}_6$ and for IR see references. The ^1H NMR (60MHz, CDCl_3 , TMS) has δ at 6.26 (s, 1H), 2.8-2.4 (br m, 4H), 2.3-1.8 (br m, 8H), 1.7-1.3 (br m, 16H).

$(\text{Acac})(\text{COE})_2\text{Ir}$ is similarly prepared from $\text{Ti}(\text{acac})$ {[25955-51-6], **M 303.5**, **m 134°** obtained as for $\text{Ti}(\text{hfacac})$ below} and $[\text{Ir}(\text{COE})_2\text{Cl}]_2$ in 86% yield forming orange-yellow crystals with **m 114-115°**, **M 512.6**, and for IR (KBr) see refs. The ^1H NMR (60MHz, CDCl_3 , TMS) has δ at 5.43 (s, 1H), 1.92 (s, 6H), 2.4-1.7 (br m, 8H), 1.7-1.0 (br m, 20H). [Diversi et al. *J Organomet Chem* **125** 253 1977, DOI: 10.1016/S0022-328X(00)89444-9.]

† **Thallium 1,1,1,5,5,5-hexafluoroacetylacetonate** $[\text{Ti}(\text{hfacac})]$ is obtained from a solution of thallos ethoxide (8g, prepared as in Fieser **2** 407; see [20398-06-5] in Chapter 4, 'Metal-Organic Compounds') in EtOH (60ml) reacted with hexafluoroacetylacetone (6.7g, [see 1522-22-1]) at ~ 25°/30 minutes; the solvent was removed *in vacuo*, the residue washed with pentane and dried. The $\text{Ti}(\text{hfacac})$ (11.8g, 95%) was thus obtained as a microcrystalline solid. [Ingrosso et al. *J Organomet Chem* **84** 75 1975, DOI: 10.1016/S0022-328X(00)88776-8.]

Iron(0) 2,2'-bipyridine [bpyFe(0)] [bpyFe(1⁺) 500295-31; bpyFe(2⁺) 73871-24-6; bpyFe(3⁺) 51232-88-3] $(\text{C}_5\text{H}_5\text{N})_2\text{Fe}$, **M 214.0**. This iron catalyst mediates *ene-carbocyclisations* such as formal [4+4] ene reactions of trienes [Takacs & Anderson *J Am Chem Soc* **109** 2200 1987, DOI: 10.1021/ja00241a059; Takacs et al. *Tetrahedron* **46** 5507 1990, DOI: 10.1016/S0040-4020(01)87749-8], regio- and chemo- selective diene to olefin cross coupling reactions [Takacs et al. *Organometallics* **5** 2395 1986, DOI: 10.1021/om00142a044], and stereoselective and regiocontrolled formation of substituted tetrahydropyrans [Takacs et al. *Tetrahedron Lett* **28** 5627 1987, DOI: 10.1016/S0040-4039(00)96797-2]. The 'active bpyFe(0) catalyst' is prepared *in situ* from $\text{Fe}(\text{acac})_2$ (see [14024-17-0]) generated by a minimum of 3.0 equivalents of Et_3Al and $\text{Fe}(\text{acac})_3$ (see [14024-18-1], in $^*\text{C}_6\text{H}_6$ at 0°), and 2,2'-bipyridine (1:1 with respect to Fe, sublimed at 65°/0.01mm) to which is added the olefin(s) at or near room temperature. [The $\text{Fe}(\text{acac})_3$ is recrystallised from EtOH or $^*\text{C}_6\text{H}_6$ /hexanes, dried at 25°/0.01mm, or sublimed at 100°/0.05mm.] Prior to the reduction of the iron, 1.1 equivalents of 2,2'-bipyridine and 10 equivalents of 'addend' (furan or 2-methylfuran) are added to the $\text{Fe}(\text{acac})_3$ in $^*\text{C}_6\text{H}_6$. After

reduction, the substrate(s) are added and the reaction is allowed to proceed at 25° and monitored. The ‘addend’ is added mainly to slow down the disproportionation of the active bpyFe(0) into inactive $\text{bpy}_2\text{Fe(0)}$ and Fe_{metal} [Takacs et al. *Tetrahedron* **46** 5507 1990, DOI: 10.1016/S0040-4020(01)87749-8]. **Note:** The catalytically active 1:1 complex bpyFe(0) has not been isolated, but is assumed to be in solution and is most probably stabilised in solution by coordinating with some entity such as ethylene (derived from Et_3Al), or perhaps even furan or methylfuran as these are required for its stability.

The reduction of FeCl_3 by 3.1 equivalents of *iso*-PrMgBr in the presence of 1.1 equivalents of 2,2'-bipyridine and 2,3-dimethylbutadiene yields an active catalyst by forming the ***bpyFe(0)2,3-butadiene complex*** [104714-94-5] in which the olefin may be involved in the reaction [Takacs et al. *Organometallics* **5** 2395 1986, DOI: 10.1021/om00142a044]. Carbocyclisations are also mediated by Pd, Ni, Co, and Rh complexes [bibliography : Takacs et al. *Tetrahedron* **46** 5507 1990, DOI: 10.1016/S0040-4020(01)87749-8]. The Fe(0) valence is **dubious**.

(2,3-O-Isopropylidene)-2,3-dihydroxy-1,4-bis-(diphenylphosphino)butane (DIOP) [4*R*,5*S*-(-)- 32305-98-9, 4*S*,5*R*-(+)- 37002-48-5] $\text{C}_{31}\text{H}_{32}\text{O}_2\text{P}_2$, *M* 498.5, *m* 88-89°, 88-90°, $[\alpha]_{\text{D}}^{19}$ (-) and (+) 26 (c 2.3, CHCl_3), also **4*R*,5*S*-** has $[\alpha]_{\text{D}}^{22}$ (-) 12.3 (c 4.6, $^*\text{C}_6\text{H}_6$), $\text{pK}_{\text{Est}} \sim 0.0$. These are quite stable in air and have been recrystallised from $^*\text{C}_6\text{H}_6$ /petroleum ether. After 2 recrystallisations from EtOH, they are generally pure by TLC on silica gel using Me_2CO /hexane as solvent. The optical purity can be determined by GC [Chiral Select 1000 column 15m x 0.25mm using He carrier gas (1ml/minute); or Chirasil-L-Val fused silica 25m x 0.25mm] or by HPLC using an (*S,S*)-Whelk-01 column (5.0µm, 25cm x 0.46cm). [Details of the general preparation of **DIOP** are described under **3,4-O-isopropylidene-(3*S*,4*S*)-dihydroxy-(2*R*,5*R*)-bis(diphenylphosphino)hexane (RSSR-dimeDIOP**, see [258873-45-9]) in Part 2. The rhodium(I) complexes described below catalyse the asymmetric hydrogenation of β -substituted α -acetamidoacrylic acids to the corresponding chiral α -amino acid derivatives with high stereoselectivity. [Kagan *J Am Chem Soc* **94** 6429 1972, DOI: 10.1021/ja00773a028.]

The DIOP derived catalysts used are **$\text{Rh(I)}^+(\text{DIOP})(\text{COD}) \text{X}^-$** where X^- is Cl^- , BF_4^- , SbF_6^- or PF_6^- and are prepared *in situ* by reaction of $\text{Rh(I)}^+(\text{COD})_2 \text{X}^-$ with the DIOP ligand in a solvent (CH_2Cl_2 , MeOH, PhCH_3 , THF or $^*\text{C}_6\text{H}_6$), using a DIOP/Rh ratio of <2 (preferably 1.1:1.0) to avoid the formation of $\text{Rh(I)}^+(\text{DIOP})_2$ which exhibits very little, if any, catalytic activity due to its inability to coordinate with the olefinic substrate. A **typical hydrogenation** is carried out in a Fisher-Porter tube which is filled (in a dry-box) with the enamide substrate (0.25mmol), the appropriate solvent (5ml), and pre-formed $\text{Rh(I)}^+(\text{DIOP})(\text{COD}) \text{X}^-$ [Rh precursor, $\text{Rh(I)}^+(\text{COD})_2 \text{X}^-$ (0.005mol) in the appropriate solvent (3ml), DIOP (0.055ml of 0.1M solution in, e.g. PhCH_3 , 0.0055mmol) and stir for 10 minutes]. After sealing, the tube is removed from the dry-box, placed behind a proper body shield, and *ca* five vacuum/refilling cycles of H_2 are performed. Hydrogenation is performed at room temperature, with the desired pressure of H_2 (1.1 to 50 bar, usually ~30psi), and is stirred vigorously for ~10 hours (but could be up to 60 hours). The reaction is completed by releasing the H_2 , the catalyst is filtered off through a short silica column, and the filtrate is worked up in the appropriate manner. [Kagan *J Am Chem Soc* **94** 6429 1972, DOI: 10.1021/ja00773a028; Li & Zhang *J Org Chem* **65** 5871 2000, DOI: 10.1021/jo0004613; Yan & RajanBabu *Org Lett* **26** 4137 2000, DOI: 10.1021/ol006591f.] A variety of DIOP derivatives have been synthesised and tested as ligands with pre-Rh catalyst salts having different X^- counter ions in several solvents. The combination which gave high yields and ‘ee’ excesses of ~98% was the all-equatorial **3,4-O-isopropylidene-(3*S*,4*S*)-dihydroxy-(2*R*,5*R*)-bis(diphenylphosphino)hexane**, i.e. **RSSR-dimeDIOP**, $\text{X}^- = \text{BF}_4^-$ and $\text{Rh(I)}^+(\text{COD})_2 \text{BF}_4^-$ [35138-22-8], in CH_2Cl_2 , at 20-40psi and 10 hours [Yan & RajanBabu *Org Lett* **26** 4137 2000, DOI: 10.1021/ol006591f]. Other examples of the application of Rh-catalysed reactions requiring the DIOP ligand include regioselective hydroformylation of methyl *N*-aminoacrylate [and involving $\text{RhH}(\text{CO})(\text{DOP})$] [Gladiali & Pinna *Tetrahedron: Asymmetry* **1** 693 1990, DOI: 10.1016/S0957-4166(00)82375-7], 1,4-disilylation of α , β -unsaturated ketones [Matsumoto et al. *Tetrahedron* **50** 335 1994, DOI: 10.1016/S0040-4020(01)80758-4], the Heck reaction with alkenyl iodides [Sato et al. *Tetrahedron* **50** 371 1994, DOI: 10.1016/S0040-4020(01)80761-4], as well as the asymmetric hydrogenation of enamides [Morimoto et al. *Tetrahedron: Asymmetry* **6** 23 1995, DOI: 10.1016/0957-4166(94)00342-9; Morimoto et al. *Tetrahedron: Asymmetry* **6** 75 1995, DOI: 10.1016/0957-4166(94)00355-F].

Lanthanide trifluoromethanesulfonate (triflate) salts. These behave mostly as water-tolerant **Lewis acids** which catalyse a variety of organic functional group transformations. The lanthanide triflates below have been prepared and purified by the general procedure described here unless otherwise stated. The salts are prepared by adding excess of lanthanide (III) or (IV) oxide (>99% purity, 30mmol) to an aqueous solution of trifluoro

methanesulfonic acid (50% v/v, 21.2ml) and heating to boiling for 30 minutes to 1 hour (*alternatively*, at 100° for 2 hours). The mixture is filtered to remove any unreacted oxide. The water is then removed under a vacuum. The resulting hydrated salt is dried by heating under vacuum at 180° to 200° for 48 hours (200°/0.5mm for 40 hours was also reported). These salts are quite **hygroscopic** and all manipulations of reactions using these salts should be carried out using Schlenk equipment or glove boxes under an inert atmosphere (N₂, He or argon) to prevent contamination with H₂O that will lead to the formation of insoluble lanthanide hydroxides, and to maximise the activity of the catalyst. [Forsberg et al. *J Org Chem* **52** 1017 1987, DOI: 10.1021/jo00382a009; Kobayashi & Hachiya *J Org Chem* **59** 3590 1994, DOI: 10.1021/jo00092a017.]

Cerium(IV) trifluoromethanesulfonate [cerium triflate, Ce(OSO₂CF₃)₄ · H₂O, Ce(OTf)₄ · H₂O] [698999-65-4] (CF₃SO₃)₄Ce · xH₂O, M 736.4 (anhydrous), slowly decomposes above 120° to give trivalent cerium species. Preparation of Ce(OTf)₄ from CeO₄ or Ce(OH)₄ and trifluoromethanesulfonic acid as above for lanthanide triflates is usually unsuccessful. It is best prepared by adding, with vigorous stirring, a solution of K₂CO₃ (17.3g, 120mmol) in H₂O (95ml) to a solution of cerium(IV) ammonium nitrate (27.4g, 50mmol) [16774-21-3] in H₂O (80ml), whereupon a pale yellow carbonate separates. This is filtered off and washed several times with H₂O. CF₃SO₃H (triflic acid, 17.7ml, 200mmol) is slowly added at 0° to this ‘wet’ cerium(IV) carbonate whereby it dissolves, and the resulting orange coloured solution is evaporated under reduced pressure, and the residue is dried *in vacuo* at 70° for 10 hours to give Ce(OTf)₄ as a yellow powder (36.2g). [Note that a ‘wet’ carbonate is important as very dry carbonate does not react readily with triflic acid at 0°.] The IR (KBr) has characteristic bands at ν_{S=O} 1230 and ν_{C-F} 1010 cm⁻¹ and it analyses as a **monohydrate** (by Karl Fischer). The salt is **hygroscopic** and should be stored under N₂ and preferably in aliquots in sealed containers. It is soluble in H₂O, EtOH, THF, 1,2-dimethoxyethane and dioxane, but almost insoluble in hexane, *C₆H₆ and CHCl₃. It has very good oxidising ability, thus converting benzylic alcohols to aldehydes or ketones, and benzylic type CH₂ to CO, e.g. substituted toluenes to their corresponding benzaldehydes, ethylbenzene to acetophenone and diphenylmethane to benzophenone in high yields. [Imamoto et al. *Chem Lett* 1445 1990, DOI: 10.1246/cl.1990.1445]. Ce(OTf)₄ is an efficient catalyst for the ring opening of epoxides with high regio and stereo selectivity. Ring opening, e.g. of styrene oxides (phenyloxiranes), yields essentially *trans* products with the OH, OR or OAc entering groups (when H₂O, alcohols or AcOH are used) attacking the ‘benzylic carbon’ atom, and with high optical purity when the reaction is carried out at ~-10° [e.g. *R*(+)-styrene oxide provides *S*(+)-PhCH(OMe)CH₂OH]. Ring opening of thiiranes with H₂O, alcohols or AcOH also yield *trans* products that dimerise to the corresponding dithianes. [Iranpoor et al. *Synth Commun* **28** 347 1998, DOI: 10.1080/00397919808005728; cf. Vougioukas & Kagan *Tetrahedron Lett* **28** 6065 1987, DOI: 10.1016/S0040-4039(00)96865-5.]

Gadolinium(III) trifluoromethanesulfonate [52093-29-5] (CF₃SO₃)₃Gd, M 604.5, decomposes > 120°. Gd(OTf)₃ is prepared by the above general method. If suspect, then add aqueous triflic acid (50% v/v) and proceed as above. It catalyses the aminolysis of epoxides in an extraordinarily efficient manner in aprotic solvents (e.g. toluene, CH₂Cl₂) with complete *trans* stereoselectivity and high regioselectivity [Chini et al. *Tetrahedron Lett* **35** 433 1994, DOI: 10.1016/0040-4039(94)85073-9]. It also catalyses the reactions between nitriles and amines to yield a variety of amidines which, depending on the amine, can be used to prepare cyclic amidines, pyrimidines and *s*-triazines [Forsberg et al. *J Org Chem* **52** 1017 1987, DOI: 10.1021/jo00382a009]. It is a water-tolerant **Lewis acid** used in aldol reactions of silyl enol-ethers and aldehydes in ~79-89% yields (see below) [Kobayashi & Hachiya *J Org Chem* **59** 3590 1994, DOI: 10.1021/jo00092a017].

Hafnium(IV) trifluoromethanesulfonate hydrate [161337-67-3] (CF₃SO₃)₄Hf · H₂O, M 774.8 (anhydrous), m >350°. Hf(OTf)₄ is prepared by the general method described above. If suspect, then add aqueous triflic acid (50% v/v) and proceed as in the general method above. It is an efficient catalyst in the Fries rearrangement of acyloxy benzene or naphthalene derivatives, and for the regioselective direct acylation of phenol and naphthol derivatives with acid chlorides [Kobayashi et al. *Tetrahedron Lett* **37** 2053 1996, DOI: 10.1016/0040-4039(96)00216-X]. It is an excellent and recyclable catalyst for mono-nitration of *o*-nitrotoluene [Waller et al. *Tetrahedron Lett* **39** 1641 1998, DOI: 10.1016/S0040-4039(97)10861-9].

Neodymium(III) trifluoromethanesulfonate [Nd(OTf)₃] [34622-08-7] (CF₃SO₃)₃Nd, M 698.3, decomposes > 120°. Nd(OTf)₃ is prepared by the general method described above. If suspect, then add aqueous triflic acid (50% v/v) and proceed as above. It catalyses the aminolysis of epoxides in an extraordinarily efficient manner in aprotic solvents (e.g. toluene, CH₂Cl₂) with complete *trans* stereoselectivity and high regioselectivity [Chini et al. *Tetrahedron Lett* **35** 433 1994, DOI: 10.1016/0040-4039(94)85073-9]. It also catalyses the reactions between nitriles and amines to yield a variety of amidines, which, depending on the amine, can be used to prepare cyclic amidines, pyrimidines and *s*-triazines [Forsberg et al. *J Org Chem* **52** 1017

1987, DOI: 10.1021/jo00382a009]. It is a **water-tolerant Lewis acid** used in aldol reactions of silyl enol-ethers and aldehydes in ~83-85% yields (see below) [Kobayashi & Hachiya *J Org Chem* **59** 3590 1994, DOI: 10.1021/jo00092a017].

Ytterbium(III) trifluoromethanesulfonate hydrate [$\text{Yb}(\text{OTf})_3 \cdot x \text{H}_2\text{O}$] [252976-51-5] (CF_3SO_3)₃Yb.
 $x\text{H}_2\text{O}$, M 620.3 (anhydr), decomposes > 120°. $\text{Yb}(\text{OTf})_3$ is prepared by the general method described above. If suspect, then add aqueous triflic acid (50% v/v) and proceed as above. It can be recrystallised from MeCN/ CH_2Cl_2 . It has IR (KBr) bands at 3650, 3350, 2300, 1650, 1300, 1040 cm^{-1} ; and ^{13}C NMR (270MHz, D_2O) at δ 122.4 (q, $J = 317\text{Hz}$) using sodium 2,2-dimethyl-2-silapentane-5-sulfonate) as internal standard. It catalyses the aminolysis of epoxides in an extraordinarily efficient manner in aprotic solvents (e.g. toluene, CH_2Cl_2) with complete *trans* stereoselectivity and high regioselectivity [Chini et al. *Tetrahedron Lett* **35** 433 1994, DOI: 10.1016/0040-4039(94)85073-9]. It also catalyses the *trans* addition of indole (at position 3) to epoxides (e.g. to phenoxymethyloxirane) in >50% yields at 60° (42 hours) under pressure (10 Kbar) and was successfully applied for an enantioselective synthesis of (+)-**diolmycin A2** [Kotsuki *Tetrahedron Lett* **37** 3727 1996, DOI: 10.1016/0040-4039(96)00670-3]. Of the ten lanthanide triflates, $\text{Yb}(\text{OTf})_3$ gave the highest yields (> 90%, see above)) of condensation products by catalytically activating formaldehyde, and a variety of aldehydes, in hydroformylations and aldol reactions, respectively, with trimethylsilyl enol-ethers in THF at room temperature. **All the lanthanide triflates can be recovered from these reactions for re-use.** [Kobayashi & Hachiya *J Org Chem* **59** 3590 1994, DOI: 10.1021/jo00092a017.]

Methyltrioxorhenium (MTO, trimethylrhenium(VII) trioxide, Me_3ReO_3) [70197-13-6] $\text{C}_3\text{H}_6\text{ReO}_3$, **M 249.2, m 110°, 111°, $\text{pK}_a^{25} 7.53$.** MTO is an **air-stable** carbon-rhenium oxide which is prepared from tetramethylrhenium oxide (Me_4ReO , [53022-70-1]) or trimethylrhenium dioxide (Me_3ReO_2 , [56090-01-8]) (100mg) in a 1L evacuated bulb, and dry air is admitted up to a pressure of ~760mm (atmospheric). After a few days long needles of Me_3ReO_3 are formed, and after 4 weeks in excess of 50% yields of the trioxide are obtained. These crystals can be **resublimed** to analytical purity *in vacuo* (at ~25°/1mm or 65°/0.001mm), have a sharp melting point (110°), are not decomposed in the gas phase <300°. **Alternatively on a larger scale**, silver tetraoxorhenate (14.0g, 39mmol, AgReO_4 [7784-00-1]) is dissolved in MeCN (150ml) followed by addition of Me_3SiCl (10.8ml, 85mmol, TMSCl [75-77-4]) whereby a white precipitate of AgCl separates and the solution becomes orange in colour. The suspension is treated with Me_4Sn (6.0ml, 43mmol, [594-27-4]), stirred for 24 hours, filtered into a Schlenk sublimation tube (~0.75L) and evaporated in a vacuum (20mbar), using a liquid N_2 trap (for volatile Sn compounds). A cold finger is then inserted into the tube and the Me_3SnCl [1066-45-1] is sublimed out at ~25°/20mbar; this compound is removed and the vacuum is increased to 0.01mbar when MTO sublimes out as a yellow-white crystalline solid. Traces of **toxic Sn compounds** may be present in this MTO (causing a pungent odour), but are removed by spreading it on filter paper in a fume hood for 1-3 hours, and the purity is checked by elemental analysis. MTO (7.8g) is thus obtained in 80% yield. Similar results are obtained on a 100g scale. The recovered Me_3SnCl can be converted to SnMe_4 by methylating with MeMgCl (23% in THF). High-purity MTO can be obtained by recrystallisation from CH_2Cl_2 /hexane (73%). The white solid turns grey on standing without alteration in its activity, but this can be minimised by storing it away from light under N_2 . **Note** that the Sn compounds are **HIGHLY TOXIC, and all work should be carried out in an efficient fume cupboard**. [Herrmann & Kratzer *Inorg Synth* **33** 111 2002, DOI: 10.1002/0471224502.ch2; Herrmann et al. *Angew Chem Int Ed* **36** 2652 1997, DOI: 10.1002/anie.199726521.]

MTO forms colourless needles that develop a greyish tint on prolonged storage in light. It differs from methylperrhenate in being stable to hydrolysis by H_2O . It is soluble in MeCN, $^*\text{C}_6\text{H}_6$, CHCl_3 , EtOH and H_2O , but sparingly soluble in CS_2 and hexane. Its MS has parent ion peaks of MeReO_3^+ at 248 and 250 mass units for the species containing ^{185}Re and ^{187}Re in a ratio consistent with the relative isotopic abundances. The UV/VIS (Et_2O) has λ_{max} (ϵ) at 260nm (1400, $\text{Lmol}^{-1}\text{cm}^{-1}$) and 232nm (1900, $\text{Lmol}^{-1}\text{cm}^{-1}$). ^1H , ^{13}C and ^{17}O NMR (CDCl_3 , 25°) spectroscopy shows signals with δ_{H} at 2.61 (sharp), δ_{C} at 19.03 ($^2J = 138\text{Hz}$, C-H) and δ_{O} at 829 respectively; and the IR (hexachlorobutadiene mull) has ν_{max} at 1360ms (*CH deformation*), 2895 and 2980 (*CH stretching*) cm^{-1} ; the IR in gaseous state (at 70°) has ν_{max} at 1003w(*ReO₃ sym str*), 985w, 975vs, 962vs(*ReO₃ antisym str*), 743mw(*CH₃-rock*), 574w(*Re-C*), 324w(*ReO₃ deformation*) cm^{-1} , and the IR (argon matrix) has ν_{max} at 1000ms(*ReO₃ sym str*), 970vs and 966m(*ReO₃ antisym str*), 566w(*Re-C*) cm^{-1} . Its Raman spectrum in CS_2 has ν_{max} at 999s(*ReO₃ sym str*) cm^{-1} , and in the solid state it has ν_{max} at 999s(*ReO₃ sym str*), 964m(*ReO₃ antisym str*), 530m(*Re-C*), 330m and 242m(*ReO₃ deformation*) cm^{-1} . [Beattie & Jones *Inorg Chem* **18** 2318 1979, DOI: 10.1021/ic50198a056.]

MTO is a *catalytic oxygen carrier* in conjunction with an oxidant, e.g. H_2O_2 , UHP (urea- H_2O_2 complex). For example 3.7mmol of 3,4,6-tri-*O*-acetyl-D-glucal in MeOH is oxidised by 7.4mmol of H_2O_2 in the presence of a catalytic amount of MTO (3.6×10^{-2} mmol) at $\sim 25^\circ$ in 2 hours to give the corresponding β -D-*gluco*- and α -D-*manno*- pyranosides (2:1) in 78% isolated yields *via* the intermediate epoxide. [Boyd et al. *Green Chem* **5** 679 2003.] **MTO** is a versatile oxidation catalyst and very effective in olefin epoxidation, olefin isomerisation, olefin metathesis, Baeyer-Villiger oxidation of ketones to lactones, aromatic oxidation to *p*-quinones, and in Diels-Alder reactions [Summarised in Herrmann *J Organomet Chem* **500** 149 1995, DOI: 10.1016/0022-328X(95)00518-U]. It is a catalytic oxidant for the conversion of imines to nitrones [Soldaini et al. *Org Lett* **9** 473 2007, DOI: 10.1021/ol062862w; Baldwin & Long *Org Lett* **6** 1653 2004, DOI: 10.1021/ol049505a], and catalyses the efficient and stereospecific desulfurisation of thiirane (episulfides) by Ph_3P at $\sim 25^\circ$, and more so if MTO is pre-treated with H_2S , probably due to the formation of Re(V)S species as the active catalyst [Jacob & Espenson *JCS Chem Commun* 1003 1999, DOI: 10.1039/A901708I].

Molybdenum dioxo bis(2,4-dipentandionato- $\kappa\text{O}_2,\kappa\text{O}_4$) [bis(acetylacetonato)dioxomolybdenum (IV), dioxobis(2,4-pentadionato)molybdenum (IV), $\text{MoO}_2(\text{acac})_2$ [17524-05-9] $\text{C}_{10}\text{H}_{14}\text{O}_6\text{Mo}$, M 326.2, m 179° , 184-185 $^\circ$, 184 $^\circ$ (dec). This complex is readily prepared by dissolving powdered ammonium *paramolybdate* $\{(\text{NH}_4)_6\text{Mo}_7\text{O}_{24} \cdot 4\text{H}_2\text{O}$, 3.0g, 2.4mmol, see [12054-85-2]) in 15% aqueous NH_3 (6.0ml, or 6.0ml of 58%) with stirring. The clear solution (but if some solid remains undissolved, it will dissolve when nitric acid is finally added) is treated with 2,4-pentanedione (7.0ml, 6.8mmol) with stirring and the colour turns yellow. Concentrated HNO_3 (5.0ml, d 1.42) is slowly added with stirring to the mixture which warms up and the colour turns greenish yellow. The yellow complex separates as the mixture is allowed to cool to room temperature. After ~ 0.5 hours the solid is filtered off washed well with H_2O , then with EtOH, and dried *in vacuo* to give a 72% yield (4.0g) of the complex which can be increased to 93% if the hot solution is kept in a freezer at -10° in 18 hours. It can be recrystallised by dissolving ~ 2.0 g in hot 2,4-pentanedione (6.0ml) at $\sim 90^\circ$ then keeping at -10° in 18 hours. The yellow crystals are filtered off with suction, washed with EtOH and dried *in vacuo* to give 1.8g of product. **Note** that if all the 2,4-pentanedione is not removed the crystals slowly develop a reddish tint. It decomposes slowly when stored in air or under fluorescent light, and turns blue. However, it is stable when stored *in vacuo* or under N_2 . It is insoluble in H_2O , but slightly soluble in EtOH, CHCl_3 , CH_2Cl_2 and soluble in MeCN. [Chakravorti et al. *Inorg Synth* **29** 129 1992, DOI: 10.1002/9780470132609.ch31; and from MoO_3 and 2,4-pentanedione see Fernelius et al. *Inorg Synth* **6** 147 1960, DOI: 10.1002/9780470132371.ch46.] Its IR (KBr) has ν_{max} at 1560, 1500, 1350, 1260, 1015, 930 ($\text{Mo}=\text{O}$ symm), 900 ($\text{Mo}=\text{O}$ antisymm), 795, 665, 570 and 445 cm^{-1} ; the ^1H NMR (CD_2Cl_2 , TMS) has δ at 2.12, 2.14 and 5.83; and for ^{13}C NMR see references.

$\text{MoO}_2(\text{acac})_2$ catalyses: (a) the oxidation of primary (e.g. cinnamyl alcohol) or secondary alcohols (e.g. 1-phenylpropan-1-ol) to their respective aldehydes or ketones in high yields using sodium percarbonate (see [15630-89-4]) as oxidant and Adogen 464 (0.2 equivalents, as phase transfer catalyst see [72749-59-8] below) in boiling (6-24 hours) CH_2Cl_2 [Maignien et al. *Synlett* 439 1996, DOI: 10.1055/s-1996-5449]; (b) the mild and efficient deprotection of acetals (e.g. benzaldehyde dimethyl acetal, dodecanal dimethyl acetal) in > 70 -95% yields using $\sim 10\%$ of catalyst in MeCN under N_2 for ~ 4 hours [Kantam et al. *Synth Commun* **25** 2529 1995, DOI:10.1080/00397919508011796], (c) the oxidation of 3- β -cholesteryl esters to the corresponding 5,6- β -epoxides in the presence of isobutyraldehyde and O_2 in CH_2Cl_2 in yields $> 75\%$ [Kantam et al. *Synth Commun* **24** 961 1994, DOI:10.1080/00397919408020771], and (d) the methoxymethylation of primary, secondary and tertiary alcohols (R-OH) to their corresponding methoxymethyl ethers ($\text{R-OCH}_2\text{OCH}_3$) in 75-95% yields using excess of methylal $\{\text{CH}_2(\text{OMe})_2$ see [109-87-5]\} in refluxing CHCl_3 [Kantam & Santhi *Synlett* 429 1993, DOI: 10.1055/s-1993-22483].

(2S)-3-exo-(Morpholino)isoborneol [(-)MIB] [287105-48-0] and (2R)-3-exo-(morpholino)iso-borneol [(+)MIB] (Nugent's reagent) $\text{C}_{14}\text{H}_{25}\text{NO}_2$, M 239.4, m 60 - 65° , 65 - 67° , $[\alpha]_{\text{D}}^{25}$ -6 and +6 (c 1, MeOH). The (2R)-*enantiomer* is prepared from (2R)(+)-*cis*-3-exo-(amino)isoborneol (4.53g, 26.8mmol; Chittenden & Cooper *J Chem Soc C* 49 1970, DOI: 10.1039/J39700000049) in DMSO (25ml) and Et_3N (10ml), to which is added dropwise, with stirring, a solution of di(2-bromoethyl) ether (8.07g, $\sim 90\%$ pure, 31.3mmol) in DMSO (20ml). After 72 hours the mixture is poured into H_2O (250ml), basified with aqueous M NaOH (60ml), extracted with Et_2O (3 x 100ml), and the extract is evaporated *in vacuo*. The residue is then dissolved in Et_2O , extracted into aqueous M HCl (50ml), basified with aqueous NaOH and re-extracted into Et_2O which is dried (MgSO_4), filtered, evaporated *in vacuo*, and the residue is dissolved in hot hexanes (4ml/g crude), filtered and

cooled to -30° to give *analytically pure* (C, H and N) (+)**MIB** (2.95g, 46%). It is a white crystalline solid, **m 65-67 $^{\circ}$** , that can be stored at ambient temperature for 3 months with no alteration in spectroscopic properties, or catalytic performance when compared with freshly prepared material. It has ^1H NMR ($^*\text{C}_6\text{D}_6$) with δ_{H} at 0.69 (s, 3H), 0.73 (m, 1H), 0.88 (m, 1H), 1.04 (s, 3H), 1.31 (td, 1H), 1.52 (m, 1H), 1.67 (d, 1H), 1.99 (d, 1H), 2.13 (br, 2H), 2.31 (br, 2H), 3.32-3.42 (m, 5H total), 3.92 (br d, 1H); and for ^{13}C NMR see reference. [Nugent *JCS Chem Commun* 1369 1999, DOI: 10.1039/A904042K.]

MIB acts as an efficient catalyst for the enantioselective addition of the ethoxy vinyl zinc reagent to aldehydes providing hydroxy vinyl ethers, which in turn, are easily converted to chiral hydroxy aldehydes [Jeon et al. *Org Lett* **7** 1729 2005]. It generally catalyses the addition of organo-zinc reagents (e.g. from transmetalation of allylic boranes with alkylZn compounds) to aldehydes with very high enantiomeric preference, *viz* (*R*)-**MIB** providing the (*R*)-alcohol from the corresponding aldehyde [Nugent *JCS Chem Commun* 1369 1999, DOI: 10.1039/A904042K; Jeon et al. *Org Lett* **7** 1729 2005, DOI: 10.1021/ol050255n], and this catalytic method has been adopted for the preparation of α -amino acids [Chen et al. *J Am Chem Soc* **124** 12225 2002, DOI: 10.1021/ja027271p], of γ -unsubstituted β -amino acids [Lurain & Walsh *J Am Chem Soc* **125** 10677 2003, DOI: 10.1021/ja035213d] and for epoxy-alcohols with up to three contiguous stereocentres [Lurain et al. *J Am Chem Soc* **126** 13608 2004, DOI: 10.1021/ja046750g; Lurain et al. *J Org Chem* **70** 1262 2005, DOI: 10.1021/jo048345d].

Nickel(II) acetylacetonate $[\text{Ni}(\text{acac})_2]$ [3264-82-2] ($\text{C}_5\text{H}_7\text{O}_2$) $_2\text{Ni}$, **M 256.9**, **m 229-230 $^{\circ}$** , **b 220-235 $^{\circ}$ /11mm**, **d 17 1.455**. It is obtained by adding a solution of acetylacetone (50g, 0.5mole) in MeOH (100ml) to $\text{NiCl}_2 \cdot 6\text{H}_2\text{O}$ (59.4g, 0.25mole) in H_2O (250ml) with stirring, followed by a solution of $\text{NaOAc} \cdot 3\text{H}_2\text{O}$ (68g, 0.5mole) in H_2O (150ml), heating briefly on a hot plate, cooling in a refrigerator for several hours, and filtering the crystals off. Wash the emerald green solid with H_2O , dry it in a vacuum desiccator and recrystallise it from MeOH. [Charles & Pawlikowski *J Phys Chem* **62** 440 1958, DOI: 10.1021/j150562a017.] The complex can be conveniently dehydrated by azeotropic distillation with toluene, and the crystals can be isolated by concentrating the toluene solution. [Wilkinson et al. *J Am Chem Soc* **76** 1970 1954, DOI: 10.1021/ja01636a080; *Beilstein* **1** IV 3677.] It is soluble in organic solvents such as EtOH, CHCl_3 , and $^*\text{C}_6\text{H}_6$ but insoluble in Et_2O and hexanes. Its UV spectrum has λ_{max} nm(log ϵ) at 265 (4.44) and 298 (4.34) (10^{-4} M in CHCl_3). It is a **trimer** in the solid state and a **monomer** in the vapour phase. When the metal in $\text{Ni}(\text{acac})_2$ is coordinated with the carbenes derived from $\text{IPr} \cdot \text{Cl}$, $\text{IPr} \cdot \text{BF}_4$, (see below) or related 1,3-dimesityl-imidazolium chloride [141556-45-8], or with tri-*tert*-butylphosphine, it efficiently catalyses the cross-coupling of aryl- and heteroaryl- halides with aryl Grignard reagents [Böhm et al. *Angew Chem Int Ed* **39** 1602 2000, DOI: 10.1002/(SICI)1521-3773(20000502)39:9].

Nickel(II) bis(triphenylphosphine) dichloride [bis(triphenylphosphine)nickel(II) dichloride] [14264-16-5] $[(\text{C}_6\text{H}_5)_3\text{P}]_2\text{NiCl}_2$, **M 654.2**, **m 247-250 $^{\circ}$ (dec)**, **250 $^{\circ}$ (dec)**. It is best prepared by adding $\text{NiCl}_2 \cdot 6\text{H}_2\text{O}$ (2.38g, 10mmol) in H_2O (2ml) to a solution of Ph_3P (5.25g, 20mmol) in glacial AcOH (25ml) when an olive-green precipitate separates, but changes to dark blue crystals after standing in the mother liquors for 24 hours. These are filtered off washed with glacial AcOH and dried in a vacuum desiccator (H_2SO_4 and KOH) until AcOH is removed to give 84% of dark blue crystals. [Venanzi *J Chem Soc* 719 1958, DOI: 10.1039/JR9580000719; Cotton et al. *J Am Chem Soc* **83** 344 1961, DOI: 10.1021/ja01463a021; Kocienski et al. *J Org Chem* **54** 1215 1989, DOI: 10.1021/jo00266a047; *Beilstein* **16** IV 953.]. With butyl chloride at 150-180 $^{\circ}$ in a sealed tube followed by crystallisation from BuOH, it provides blue crystals with **m 176 $^{\circ}$** of $[\text{Ni}(\text{BuCl})(\text{Ph}_3\text{P})_2\text{Cl}_2]$ [Yamamoto *Bull Chem Soc Jpn* **27** 501 1954, DOI: 10.1246/bcsj.27.501; *Beilstein* **16** IV 953, for applications see Fieser **2** 110, **9** 147, **14** 125, **16** 118.] In small amounts it catalyses the formation of terminal alkenes, e.g. from alkyl bromides or iodides in the presence of BuLi/DBU in THF at 25 $^{\circ}$ in 50-80% yields [Jeropoulos & Smith *JCS Chem Commun* 1621 1986, DOI: 10.1039/C39860001621]. It also catalyses the replacement of OH groups in allyl alcohols with the alkyl group of alkylmagnesium bromide thus forming a C-C bond, and also causes dehydrohalogenation of alkylhalides to form the corresponding terminal olefin [Chuit et al. *JCS Chem Commun* 1604 1968, DOI: 10.1039/C19680001604].

Nickel(0) bis(1,5-cyclooctadiene) [bis-(1,5-cyclooctadiene)nickel(0), $\text{Ni}(\text{COD})_2$] [1295-35-8] $\text{C}_{16}\text{H}_{24}\text{Ni}$, **M 275.1**, **m 60 $^{\circ}$ (dec)**, **142 $^{\circ}$ (dec)**. It is available in sealed ampoules under N_2 . All procedures should be carried out in a dry box and in an atmosphere of N_2 or argon in subdued light because the complex is light and oxygen sensitive, and flammable. The solid is washed with dry Et_2O (under argon) and separates from toluene as yellow

crystals. Filter this under argon gas pressure, place the crystals in a container and dry them under a vacuum of 0.01mm to remove adhered toluene, flush with argon and seal them under argon or N₂ in glass ampoules. It catalyses cycloaddition reactions of 1,3-dienes [Semmelhack *Org Reactions* **19** 115 1972, DOI: 10.1002/0471264180.or019.02; Bogdanović et al. *Justus Liebigs Ann Chem* **699** 1 1966, DOI: 10.1002/jlac.19666990102; Wender & Jenkins *J Am Chem Soc* **111** 6432 1989, DOI: 10.1021/ja00198a071; Fieser **4** 33, **16** 29, **17** 32]. It also catalyses the addition of allyl phenyl sulfide to alkynes leading to 1,4-dienes. The reaction with acetylenes affords high yields, and in the presence of chiral phosphine ligands the reaction proceeds with high stereoselectivity. The reaction tolerates a variety of functional groups [Hua et al. *Org Lett* **9** 263 2007, DOI: 10.1021/ol062686r]. **SUSPECTED CARCINOGEN.**

Oxime Palladacycle dimers (Nájera Catalysts, OPs) are Pd complexes of aromatic oximes, e.g. where palladium(II) is complexed with the oxime nitrogen atom and co-metalated with the *o*-position of the aromatic ring. They catalyse a variety of coupling reactions such as Suzuki-Miyaura, Heck, Stille, Sonogashira (all cross coupling) reactions and the Ullmann (homocoupling) reactions. *Note that the concentration of Pd in the products from Suzuki-Miyaura coupling reactions can be reduced from ~8000ppm to 100ppm by treating the reaction mixture with toluene and 20% aqueous NaHSO₃ at ~60° for ~1 hour* [Bullock, Mitchell and Toczko *Organic Process Research & Development* **12** 896 2008, DOI: 10.1021/op800064y].

Oximes are prepared by a common procedure: A solution of the required aryl-ketone (22mmol) in MeOH (10ml) is added to a solution of hydroxylamine hydrochloride (3.06g, 44mmol), anhydrous NaOAc (3.6g, 44mmol) in H₂O pre-heated at 60° for 1 hour. If some solid separates, add enough MeOH to obtain a clear solution, then stir at this temperature overnight. On cooling to ~25°, the oximes that separate in >90% yield are filtered off, washed with H₂O and recrystallised to ¹H NMR purity (>98%). **Acetophenone oxime** has **m** 59.1-59.7° (from MeOH then cyclohexane, Pearson & Ball *J Org Chem* **14** 118 1949, DOI: 10.1021/jo01153a018), **4,4'-dichlorobenzophenone oxime** has **m** 135.2-136.9° (from MeOH, Sieger & Klein *J Org Chem* **22** 951 1957, DOI: 10.1021/jo01359a026), **4,4'-dihydroxybenzophenone oxime** has **m** 266-267°dec (from EtOH, Zigeuner & Ziegler *Monatsh für Chemie* **89** 359 1949, DOI: 10.1007/BF00897769); **4-hydroxyacetophenone oxime** has **m** 144-145° (from aqueous MeOH), **4-methoxyacetophenone oxime** has **87°** (from petroleum ether, v. Auwers et al. *Chem Ber* **58** 36 1925, DOI: 10.1002/cber.19250580109), **4-methylacetophenone oxime** has **m** 87-88° (from MeOH then cyclohexane, Pearson & Ball *J Org Chem* **14** 118 1949, DOI: 10.1021/jo01153a018), and **9-fluorenone oxime** has **m** 195°, 198° (yellow crystals from *C₆H₆ or xylene, Anet et al. *Can J Chem* **35** 180 1957, DOI: 10.1139/v57-027; Wislicenus & Waldmuller *Chem Ber* **41** 3334 1908, DOI: 10.1002/cber.19080410309).

Oxime palladacycles are generally dimers that can be prepared by a general procedure. A solution of the oxime (10mmol) in methanol (20ml) containing NaOAc.3H₂O (1.63g, 10mmol) is added to a 0.5M solution of Li₂PdCl₄ (20ml, i.e.2.62g, 10mmol, see [123334-21-4] in this Chapter, 'Catalysts-Part 2') and stirred at ~25° for 2 to 3 days. Small quantities of precipitate may form and the colour of the reddish black solution turns to yellow. This is filtered and H₂O (10ml) is added to precipitate the palladacycles as yellow solids (~90% yields) that are filtered off, washed with a little MeOH and H₂O, and dried *in vacuo*. [Botella & Nájera *J Organomet Chem* **663** 46 2002, DOI: 10.1016/S0022-328X(02)01727-8; Onoue, Minami and Nakagawa *Bull Chem Soc Jpn* **43** 3480 1970, DOI: 10.1139/v57-027; for preparation *via* ligand exchange see Ryabov et al. *Inorg Chem* **31** 3083 1992, DOI: 10.1021/ic00040a018.] They can be used directly to catalyse reactions, their solubilities vary with the complex, but some can be crystallised from CH₂Cl₂-*n*-hexane, *C₆H₆/*n*-hexane or CHCl₃. They catalyse Heck couplings (in *N*-methylpyrrolidone with Et₃N as base), Heck couplings under Jeffrey's conditions (DMF with Bu₄NBr), Sonogashira reactions (with acetylenes in pyrrolidine and CuI), Stille coupling (in toluene), Suzuki coupling (in toluene with K₂CO₃ as base), and Ullmann-type homocoupling (in DMF, in the necessary presence hydroquinone), and typical reaction conditions have been described [Alonso, Nájera and Pacheco *Org Lett* **2** 1823 2000, DOI: 10.1021/ol0058644; Iyer & Ramesh *Tetrahedron Lett* **41** 8981 2000, DOI: 10.1016/S0040-4039(00)01594-X].

Di-chloro-bis[5-chloro-2-[(4-chlorophenyl)(hydroxyimino)methyl]phenyl-C]-di-palladium {Nájera Catalyst I, di-μ-chlorobis[5-chloro-2-[(4-chlorophenyl)(hydroxyimino-κN)methyl]phenyl-κC]-palladium (II) dimer} [287410-78-0] has C₂₆H₁₆Cl₆N₂O₂Pd₂, **M** 814.0 and **m** 208-210°. [Alonso, Nájera and Pacheco *Org Lett* **2** 1823 2000, DOI: 10.1021/ol0058644].

Di-chloro-bis[5-hydroxy-2-[1-(hydroxyimino)ethyl]phenyl-C]-di-palladium {Nájera Catalyst II, di-μ-chlorobis[5-chloro-2-[1-(hydroxyimino-κN)ethyl]phenyl-κC]-palladium(II) dimer} [419581-64-9] has

($\text{C}_8\text{H}_8\text{ClNO}_2$) $_2\text{Pd}_2$, **M 584.1** and **m >250°, 251-255°**. It has IR (KBr) with ν_{max} at 3425 (OH), 1662 (C=N) cm^{-1} ; the ^1H NMR (300MHz, DMF- d_7 , TMS) has δ_{H} at 9.96 (brs, 2H, 2 x OH), 9.71 (brs, 2H, 2 x OH), 7.20 (brs, 2H, ArH), 7.07 (d, $J = 8.5\text{Hz}$, 2H, ArH), 6.50 (dd, $J = 7.9, 2.4\text{ Hz}$, 2H, ArH), 2.24 (s, 6H, 2 x CH_3); and for ^{13}C NMR see references. Botella & Nájera [*J Organomet Chem* **663** 46 2002, DOI: 10.1016/S0022-328X(02)01727-8] obtained high turnover numbers and turnover frequencies in Suzuki-Miyaura cross-coupling reactions at room temperature conditions with this catalyst. Botella & Nájera [*J Org Chem* **70** 4360 2005, DOI: 10.1021/jo0502551] also studied this catalyst for Mirozoki-Heck couplings in aqueous *N,N*-dimethylacetamide in air using *N*-methyldicyclohexylamine (see [7560-83-0]) as base with or without Bu_4NBr .

Di-chloro-bis[2-[1-(hydroxyimino)ethyl]phenyl-C]-di-palladium {di- μ -chlorobis[2-[1-(hydroxyimino- κN)ethyl]phenyl- κC]-palladium(II) dimer} [32679-19-9] has ($\text{C}_8\text{H}_8\text{ClNO}$) $_2\text{Pd}_2$, **M 552.1** and **m 210°, 209-212°**. It has IR (KBr) with ν_{max} at 3426 (OH), 1640 (C=N) cm^{-1} . [Alonso, Nájera and Pacheco *Org Lett* **2** 1823 2000, DOI: 10.1021/ol0058644; Onoue, Minami and Nakagawa *Bull Chem Soc Jpn* **43** 3480 1970, DOI: org/10.1246/bcsj.43.3480 .]

Di-chloro-bis[5-hydroxy-2-[(4-hydroxyphenyl)(hydroxyimino)methyl]phenyl-C]-di-palladium, {di- μ -chlorobis[5-hydroxy-2-[(4-hydroxyphenyl)(hydroxyimino- κN)methyl]phenyl- κC]-palladium(II) dimer} [419581-64-9] has ($\text{C}_{13}\text{H}_{10}\text{ClNO}_3$) $_2\text{Pd}_2$, **M 740.74(777.0)** and **m >250°**. It has IR (KBr) with ν_{max} at 3405 (OH), 1612 (C=N) cm^{-1} ; the ^1H NMR (300MHz, DMF- d_7 , TMS) has δ_{H} at 10.50-9.40 (brs, 6H, 6 x OH), 7.41 (d, $J = 8.5\text{Hz}$, 4H, ArH), 7.26 (brs, 2H, 2 x OH), 7.20 (brs, 2H, ArH), 7.02 (d, $J = 8.5\text{Hz}$, 4H, ArH), 6.69 (d, $J = 8.5\text{Hz}$, 4H, ArH), 6.48 (dd, $J = 7.9, 2.7\text{ Hz}$, 2H, ArH); and its ^{13}C NMR (75MHz, DMF- d_7 , TMS) has δ_{C} at 167.8, 160.0, 157.5, 152.9, 134.3, 131.5, 129.4, 123.7, 121.0, 115.8, 111.5. [Botella & Nájera *J Organomet Chem* **663** 46 2002, DOI: 10.1016/S0022-328X(02)01727-8.]

Di-chloro-bis[5-chloro-2-[(4-phenyl)(hydroxyimino)methyl]phenyl-C]-di-palladium, {di- μ -chlorobis[5-chloro-2-[(4-phenyl)(hydroxyimino- κN)methyl]phenyl- κC]-palladium(II) dimer} [1145982-32-6; 30471-18-2 for stereoisomer] has ($\text{C}_{13}\text{H}_9\text{Cl}_2\text{NO}$) $_2\text{Pd}_2$, **M 745.6** and **m 139-141°**. It has IR (KBr) with ν_{max} at 3373 (OH), 1645 (C=N), 1569, 1435, 1340, 1024 cm^{-1} ; the ^1H NMR (300MHz, DMSO- d_6 , TMS) has δ_{H} at 6.69 (m, 4H), 7.09 (m, 4H), 7.35-8.10 (m, 10H). [Alonso, Nájera and Pacheco *Org Lett* **2** 1823 2000, DOI: 10.1021/ol0058644; Onoue, Minami and Nakagawa *Bull Chem Soc Jpn* **43** 3480 1970, DOI: org/10.1246/bcsj.43.3480.]

Di-chloro-bis[5-methoxy-2-[(4-methoxyphenyl)(hydroxyimino)methyl]phenyl-C]-di-palladium {di- μ -chlorobis[5-methoxy-2-[(4-methoxyphenyl)(hydroxyimino- κN)methyl]phenyl- κC]-palladium(II) dimer} [287410-79-1] has ($\text{C}_{15}\text{H}_{14}\text{ClNO}_3$) $_2\text{Pd}_2$, **M 796.9** and **m 135-137°**. It has IR (KBr) with ν_{max} at 3390 (OH), 1607 (C=N), 1579, 1559, 1253, 1233, 1177, 1026 cm^{-1} ; the ^1H NMR (300MHz, DMSO- d_6 , TMS) has δ_{H} at 3.71 (s, 6H), 3.83 (s, 6H), 6.71 (m, 4H), 7.08-7.25 (m with d at 7.10, $J = 8.6\text{Hz}$, 5H), 7.30-7.55 (m with d at 7.43, $J = 8.6\text{Hz}$, 5H). [Alonso, Nájera and Pacheco *Org Lett* **2** 1823 2000, DOI: 10.1021/ol0058644].

Palladium(II) acetate [Pd(OAc) $_2$] [3375-31-3] Pd(C $_2\text{H}_3\text{O}_2$) $_2$, M 244.5, m 205°(dec.), pK_1^{25} 1.0, pK_2^{25} 1.2 (for Pd $_2^{2+}$). It crystallises from CHCl_3 as purple crystals. It can be washed with AcOH and H_2O and dried in air. Large crystals are obtained by dissolving it in $^*\text{C}_6\text{H}_6$, adding half its volume of AcOH and allowing it to evaporate slowly at room temperature. It forms green adducts with nitrogen donors, it dissolves in KI solution to form solid PdI_2 and a red solution of PdI_4^{2-} , but is insoluble in aqueous saturated NaCl, and NaOAc. It dissolves in HCl to form PdCl_4^{2-} . It is soluble in CHCl_3 , CH_2Cl_2 , Me_2CO , MeCN, Et_2O , but it is insoluble in H_2O , and decomposes when warmed in alcohols in which it is also insoluble. [Morehouse et al. *Chem Ind (London)* 544 1964, Stephenson et al. *J Chem Soc* 3632 1965, DOI: 10.1039/JR9650003632; Skapski & Smart *J Chem Soc (D)* 658b 1970, DOI: 10.1039/C2970000658B; Heck *Acc Chem Res* **12** 146 1979, DOI: 10.1021/ar50136a006.] **Pd(OAc) $_2$** should be an orange/red solid. Sometimes **commercial** samples, or old samples, give lower yields in catalytic reactions and it is found that good results are obtained after recrystallisation from $^*\text{C}_6\text{H}_6$ [Anderson et al. *Org Synth* **84** 148 2007, DOI: 10.15227/orgsyn.084.0148]. It is useful as a catalyst for coupling reactions [Kang et al. *J Am Chem Soc* **128** 6194 2006, DOI: 10.1021/ja060185v], and is a precursor for preparing heterogeneous and homogeneous catalysts [Jujjuri et al. *J Catal* **239** (2) 486 2006, DOI: 10.1016/j.jcat.2006.02.022; Xu et al. *J Nano Res* **7** 449 2005, DOI: 10.1007/s11051-005-005-4273-3].

Palladium(II) acetylacetonate [Pd(acac) $_2$] [14024-61-4] Pd(C $_5\text{H}_7\text{O}_2$) $_2$, M 304.6, m 200-250°(dec). It can be recrystallised from $^*\text{C}_6\text{H}_6$ /petroleum ether and sublimes *in vacuo*. It is soluble in heptane, $^*\text{C}_6\text{H}_6$ (1.2% at 20°, 2.2 at 40°), toluene (0.56% at 20°, 1.4% at 40°) and acetylacetonate (1.2% at 20°, 0.05% at 40°). [West & Riley *J*

Inorg Nucl Chem **5** 295 1958, DOI: 10.1016/0022-1902(58)80007-X; Fernelius & Bryant *Inorg Synth* **5** 105 1957, DOI: 10.1002/9780470132364.ch29; *Beilstein* **1** IV 3676; for applications see Fieser **6** 45, **17** 269.] It is a soluble Pd source for the preparation of various soluble Pd catalysts by transferring the metal to a variety of phosphorus and other ligands for homogeneous catalysis [cf. *Handbook of Organopalladium for Organic Synthesis* Negishi ed. Wiley, Hoboken NJ 2002, ISBN 0-471-31506-0.]

A **complex consisting of Cu 1.10-phenanthroline** [which mediates decarboxylation of arylcarboxylic acids with formation of aryl Cu species] **and Pd(acac)₂** [for coupling] was made and used to catalyse the decarboxylative cross-coupling of the Cu species with aryl halides. This bimetallic system allows direct coupling of a variety of aryl, heteroaryl or vinyl carboxylic acids with aryl or heteroaryl bromides, chlorides or iodides at 160° in *N*-methylpyrrolidine in the presence of K₂CO₃. [Goossen et al. *J Am Chem Soc* **129** 4824 2007, DOI: 10.1021/ja068993+].

Palladium EnCat. Ley and co-workers [Ramarao et al. *JCS Chem Commun* 1132 2002, DOI: 10.1039/B200674J; Pears & Smith *Aldrichimica Acta* **38** 23 2005] have described a method of encapsulating Pd(OAc)₂ in the matrix of 20-250 μm microcapsules prepared from a dispersion of an aromatic polyfunctional isocyanate and Pd(OAc)₂ in CH₂Cl₂ into H₂O containing stabilisers and surfactants. At the point when the oily dispersion reaches the desired size, e.g. 20-250 μm, polymerisation is initiated (evolution of heat) whereby part of the isocyanate groups are hydrolyse to carbamate then to amino groups. The latter condense with the unhydrolysed isocyanate groups to form a **crosslinked polyurea matrix entrapping the metal catalyst**. After the necessary washing etc., polyurea microcapsules (MC average size ~150 μm) are formed which are hard, porous and highly crosslinked spheres. These beads are catalytically active, robust and recyclable and have been used in conventional and supercritical media (e.g. liquid CO₂). They catalyse **MC-[Pd]** mediated Heck coupling (*p*-aryl and 4-heteroaryl nitro, methoxy and fluoro compounds with acrylic esters), carbonylation of 4-substituted arenes or heterocycles, Suzuki-type (ArB(OH)₂ + Ar to form crossed biaryls) and Stille couplings without requiring supplementary ligands [Ley et al. *JCS Chem Commun* 1134 2002, DOI: 10.1039/B200677B]. For **EnPd(OAc)₂** [**Pd EnCatTM**] and **Pd⁽⁰⁾ EnCatTM** see entries in Chapter 7. Encapsulated Pd catalysts such as **Pd EnCatTM** are available commercially. These particles are defined by their matrix content e.g. 30% or 40%, the latter having the smaller pore size. [See also Bremeyer et al. *Synlett* 1843 2002, DOI: 10.1055/s-2002-34862; Yu et al. *JCS Chem Commun* 678 2003, DOI: 10.1039/B300074P; Vickerstaffe et al. *Org Biomol Chem* **1** 2419 2003, DOI: 10.1039/B305713E.]

Palladium(II) trifluoroacetate [42196-31-6] **Pd(C₂F₃O₂)₂**, **M 332.4**, **m ~210°(dec)**. Suspend it in trifluoroacetic acid and evaporate it on a steam bath a couple of times. The residue is then dried in vacuum (40-80°) to give a brown powder. It is **hygroscopic** and should be stored in a dry atmosphere, preferably aliquoted in sealed vials [Stephenson et al. *J Chem Soc* 3632 1965, DOI: 10.1039/JR9650003632; Trost & Metzner *J Am Chem Soc* **102** 3572 1980, DOI: 10.1021/ja00530a042.]

Pd(CF₃CO)₂ catalyses the decarboxylation of electron-rich aromatic acids (e.g. with OMe groups) in DMSO/DMF at 70-90° (1 to 24 hours) in high yields and is not affected by steric hinderance [Dickstein et al. *Org Lett* **9** 2441 2007, DOI: 10.1021/ol070749f], and (in the presence of Cu(OAc)₂ with Cesium pivalate + 3-nitropyridine as additives at 110-140° in a microwave) it catalysed direct cross-coupling between unactivated arenes and *N*-acetylindoles with coupling mostly at C3 of indoles but with no homo-coupling [Stuart & Fangou *Science* **316** 1172 2007, DOI: 10.1126/science.1141956]. In the presence of AcOH, benzoquinone (as oxidant) and *o*-methoxyacetophenone or Ph₃P as ligands, Pd(tfa)₂ catalyses selective allylic oxidation of olefins into their allyl acetates [McMurry & Kočotovský *Tetrahedron Lett* **25** 4187 1984, DOI: 10.1016/S0040-4039(01)81391-5]. [Fieser **10** 302, **12** 373, **13** 236, **14** 253.]

Phosferrox ligands and SK-Naud catalysts. The ligands are (diphenylphosphinoferrocenyl)oxazolines and are complexes where one of the cyclopentadienyl rings has two different substituents. These molecules have '**planar chirality**', i.e. are asymmetric, and exist in two enantiomeric forms. The oxazolines are 4,5-dihydro-oxazoles which for this use have a 4-sustituent, usually alkyl or aryl, thus introducing a chiral centre at C-4 of this heterocyclic ring. Phosferrox ligands are prepared by lithiation of chiral 4-alkyloxazolin-2-ylferrocene in which lithiation is directed predominantly to one of the *ipso* positions of the cyclopentadienyl ring, generating '**planar asymmetry**' in the ferrocene moiety. It is highly diastereoselective with *ortho*-lithiation yields of ~84~99% de, producing the *S*-stereochemistry at the ferrocene moiety as shown by X-ray analysis and CD spectra of the products. Reaction of the lithiated ferrocene with Ph₂PCl furnishes the desired chiral phosferrox

ligand. The stereochemistry of the reaction has been studied in detail [Sammakia et al. *J Org Chem* **60** 10 1995, DOI: 10.1021/jo00106a005; Sammakia & Latham *J Org Chem* **60** 6002 1995, DOI: 10.1021/jo00124a003; Richards & Mulvaney *Tetrahedron: Asymmetry* **7** 1419 1996, DOI: 10.1016/0957-4166(96)00159-0; Nishibayashi et al. *J Organomet Chem* **545-546** 381 1997, DOI: 10.1016/S0022-328X(97)00368-9.] Advantage is taken of the diastereoselectivity of the lithiation reaction in order to obtain the **enantiomeric R-ferrocene**. Thus, after lithiation, the lithium is displaced by a trimethylsilyl group (by reaction with Me₃SiCl), the *S*-TMS derivative is lithiated again, but at the other *ipso* position of the same cyclopentadienyl ring, followed by reaction with Ph₂PCl to form the **1-diphenylphosphino-3-TMS-2-(oxazolin-2-yl)ferrocene**. Finally, removal of the TMS group, e.g. with *tetra-n-butylammonium fluoride* (TBAF, see [429-41-4; 3 H₂O 87749-50-6]), provides the phosferrox where the stereochemistry at the ferrocene moiety is now **R**. Chiral phosferrox ligands react with RuCl₂(Ph₃P)₃ [15529-49-4] to form **Nauk catalysts** which are **phosferrox-RuCl₂(Ph₃P)** complexes that reduce aryl ketones in the presence of *i*-PrOH/*i*-PrOK to form the respective alcohols with high stereoselectivity [Sammakia & Stangeland *J Org Chem* **62** 6104 1997, DOI: 10.1021/jo9711044; Nishibayashi et al. *Organometallics* **18** 2291 1999, DOI: 10.1021/om990210o].

***S*-2-[(*S*)-2-(Diphenylphosphino)ferrocenyl]-4-(1-methylethyl)oxazoline (S,S-*i*-Pr-Phosferrox)** [163169-29-7] C₂₈H₂₈FeNOP, M 481.4, m 157-158°, [α]_D²⁴ +112 (c 0.1, EtOH). This phosferrox is prepared in a Schlenk tube at -78° under N₂, by adding dropwise *n*-BuLi (0.38ml, 0.7mmol) to a yellow-orange stirred solution of *S*-2-ferrocenyl-4-(1-methylethyl)oxazoline (0.158g, 0.53mmol) and TMEDA (0.10ml, 0.7mmol) in Et₂O (6ml) which had formed a yellow precipitate, and is stirred for 2 hours; the tube containing the orange non-homogeneous mixture is transferred to an ice-bath and stirred for 5 minutes further. To this now orange-red homogeneous solution is added Ph₂PCl (0.12ml, 0.7mmol, see [1079-66-9]), the mixture is allowed to warm to ~25°, and after 15 minutes it is quenched with saturated aqueous NaHCO₃ (10ml), the layers are separated, the aqueous layer is extracted with Et₂O (10ml), the combined Et₂O solutions are dried (Mg SO₄), filtered, and evaporated to give an orange crystalline solid. This is purified by column chromatography (pre-adsorbed on silica, eluting with 10% EtOAc/petroleum ether) to give a yellow-orange crystalline solid (0.163g, 64%) that provided **S,S-*i*-Pr-Phosferrox** as an analytically pure single diastereomer upon recrystallisation from hexane. It has CD (CHCl₃) λ_{max} (Δε) 456 (+2.20), 368 (+0.49), 342 (-1.00), 315 (+216) nm; the IR has ν_{max} (nujol) at 1652 (C=N) cm⁻¹; the ¹H NMR (360Mz, CDCl₃) has δ_H at 0.68 (3H, d, *J* = 7 Hz, -CH₃), 0.82 (3H, d, *J* = 7 Hz, -CH₃), 1.61-1.69 (1H, m, CH(CH₃)₂), 3.61 (1H, brs, Fc), 3.67 (1H, t, *J* = 8 Hz, -OCHH), 3.83-3.90 (1H, m, -NCH-), 4.22 (5H, s, C₅H₅), 4.22-4.30 (1H, m, -OCHH-), 4.37 (1H, brs, Fc), 4.99 (1H, brs, Fc), 7.18-7.24 (5H, m, Ph), 7.36-7.37 (3H, m, Ph), 7.46-7.51 (2H, m, Ph); the ³¹P NMR (CDCl₃) has a single peak with δ_P at -16.92, and for ¹³C NMR and MS (EI) see references. [Richards & Mulvaney *Tetrahedron: Asymmetry* **7** 1419 1996, DOI: 10.1016/0957-4166(96)00159-0].

***R,R*-*i*-Pr-Phosferrox** [541540-70-9] C₂₈H₂₈FeNOP, M 481.4, [α]_D²⁴ -112 (c 0.1, EtOH) is the mirror image of the preceding *phosferrox*.

***S*-2-[(*R*)-2-(Diphenylphosphino)ferrocenyl]-4-(1-methylethyl)oxazoline [S-(*R*-*i*-Pr-Phosferrox)]** [163169-10-6] C₂₈H₂₈FeNOP, M 481.4, has m 132-132.5°, [α]_D²⁰ -53 (c 0.15, EtOH). This diastereomer is obtained by de-trimethylsilylation of *S*-2-[*R*-2-(diphenylphosphino)-5-(trimethylsilyl)ferrocenyl]-4-(1-methylethyl)oxazoline (0.14g 0.25mmol) with a yellow solution of 1M TBAF (see [429-41-4; 3 H₂O 87749-50-6], in 'Catalysts-Part 2') in THF (10ml) containing ca 5% H₂O by boiling for 4 hours, evaporating *in vacuo* to a small volume, shaking with Et₂O (10ml) and H₂O (10ml), separating, the aqueous layer is extracted with Et₂O, the ethereal layers are combined, dried (Mg SO₄), filtered, evaporated *in vacuo*, and the residue is chromatographed in a silica column (eluted with 10% EtOAc/petroleum ether b 40-60°) to give **S-(*R*-*i*-Pr-Phosferrox)** as a yellow crystalline solid (0.090, 75%). Its CD (CHCl₃) has λ_{max} (Δε) 492 (+0.45), 434 (-0.43), 361 (-0.31), 344 (+0.35), 314 (-2.38) nm; the IR has ν_{max} (nujol) at 1660 (C=N) cm⁻¹; the ¹H NMR (360Mz, CDCl₃) has δ_H at 0.63 (3H, d, *J* = 7 Hz, -CH₃), 0.65 (3H, d, *J* = 7 Hz, -CH₃), 1.50-1.58 (1H, m, CH(CH₃)₂), 3.62 (1H, brs, Fc), 3.89-4.08 (3H, m, -OCH₂CH-), 4.22 (5H, s, C₅H₅), 4.36 (1H, brs, Fc), 4.94 (1H, brs, Fc), 7.19-7.23 (5H, m, Ph), 7.35-7.37 (3H, m, Ph), 7.48-7.53 (2H, m, Ph); the ³¹P NMR (CDCl₃) has a single peak at δ_P -18.03; for ¹³C NMR and MS (EI) see refs. [Richards & Mulvaney *Tetrahedron: Asymmetry* **7** 1419 1996, DOI: 10.1016/0957-4166(96)00159-0].

Other phosferrox ligands prepared are **S,S-4'-*Me*-Phosferrox**, **S,R-4'-*Me*-Phosferrox**, **S,S-4'-*Et*-Phosferrox**, **S,S-4'-*n*-Bu-Phosferrox**, **S,S-4'-*iso*-Bu-Phosferrox**, **S,S-4'-*t*-Bu-Phosferrox**, and **S,S-4'-*Ph*-Phosferrox**, **S,S-4'-*benzyl*-Phosferrox** [Sammakia & Stangeland *J Org Chem* **62** 6104 1997, DOI: 10.1021/jo9711044; Nishibayashi et al. *J Organomet Chem* **545-546** 381 1997, DOI: 10.1016/S0022-328X(97)00368-9].

***S*-2-[*S*_P-2-(Diphenylphosphino)ferrocenyl]-4-*iso*-propyl-2-oxazoline triphenylphosphine ruthenium**

(II) dichloride complex [*S,S*-*i*-Pr-Phosferrox-Ru(II) Ph_3PCl_2 complex, SK-Naud Catalyst-N003-2z] [212133-11-4] $\text{RuCl}_2[(\text{C}_6\text{H}_5)_3\text{P}](\text{C}_{28}\text{H}_{28}\text{FeNOP})$, **M 915.6**, $[\alpha]_{\text{D}}^{20}$ -1127 (c 0.1, CHCl_3). This catalyst can be prepared *in situ* from the phosferrox and $\text{RuCl}_2(\text{PhP}_3)_3$, but it can also be isolated. Thus a mixture of $\text{RuCl}_2(\text{PhP}_3)_3$ (480mg, 0.50mmol) and *S,S*-*i*-Pr-Phosferrox (0.50mmol) and toluene (15ml) are stirred under N_2 at $\sim 25^\circ$ for 20 hours when the original purple solution changes to a red suspension. Addition of *n*-hexane caused the crystalline catalyst to separate, and the red crystals are recrystallised to analytical purity from $\text{CH}_2\text{Cl}_2/n$ -hexane ($\sim 81\%$ yield). The ^1H NMR (270Mz, CDCl_3) has δ_{H} at 0.57 (3H, d, $J = 7$ Hz, CH_3), 0.97 (3H, d, $J = 7$ Hz, CH_3), 2.12 (1H, dd, $J = 8$ and 8Hz), 3.21 (1H, m), 3.28 (1H, m), 3.80 (1H, dd, $J = 3$ and 8Hz), 4.02 (5H, s), 4.59 (1H, m), 4.68 (1H, m), 4.84 (1H, m), 6.5-8.4 (25H, m); and the ^{31}P NMR (107Mz, CDCl_3) indicated a diastereomerically pure complex with δ_{P} 40.1 (d, $J = 45\text{Hz}$) and 77.0 (d, $J = 45\text{Hz}$). It causes a high conversion with high stereoselectivity (with 88- $<99\%$ 'ee') in the Ru-catalysed asymmetric transfer hydrogenation of ketones using *i*-PrOH as H source, as well as catalysing the asymmetric oxidation of *sec*-alcohols to ketones [Nishibayashi et al. *Organometallics* **18** 2291 1999, DOI: 10.1021/om990210o].

R-2-[*R*_p-2-(Diphenylphosphino)ferrocenyl]-4-iso-propyl-2-oxazoline triphenylphosphine ruthenium(II) dichloride complex (SK-Naud Catalyst-N003-1z) [849921-25-1] $\text{RuCl}_2[(\text{C}_6\text{H}_5)_3\text{P}](\text{C}_{28}\text{H}_{28}\text{FeNOP})$, **M 915.6**, $[\alpha]_{\text{D}}^{20}$ +1127 (c 0.1, CHCl_3) can be prepared and purified as in the preceding entry.

S-2-[*S*_p-2-(Diphenylphosphino)ferrocenyl]-4-phenyl-2-oxazoline triphenylphosphine ruthenium (II) dichloride complex [*S,S*-*Ph*-Phosferrox-Ru(II) $\text{Ph}_3\text{P Cl}_2$ complex] [212210-73-6] $\text{RuCl}_2[(\text{C}_6\text{H}_5)_3\text{P}](\text{C}_{31}\text{H}_{26}\text{FeNOP})$, **M 949.6**. As above (SK-Naud catalyst-N003-2z), the catalyst can be prepared *in situ*, and can also be isolated in pure crystalline form, and its structure has been confirmed by X-ray crystallography. A mixture of *S,S*-4'*Ph*-Phosferrox [257mg, 0.50mol, red solid purified by flash chromatography with silica gel and eluting with hexanes the 10:1 hexanes/EtOAc, and recrystallisation from hexane, **m 184-185° dec**, $[\alpha]_{\text{D}}^{23}$ +25.6 (c 0.67, CHCl_3)] and $\text{RuCl}_2(\text{PhP}_3)_3$ (480mg, 0.50mmol) in toluene (15ml) are stirred under N_2 at $\sim 25^\circ$ for 20 hours when the original purple solution changes to a red suspension. Addition of *n*-hexane causes the crystalline catalyst to separate, and the red crystals are recrystallised to analytical purity from $\text{CH}_2\text{Cl}_2/n$ -hexane to provide the catalyst (with 1 mol of CH_2Cl_2) in 80% yield. The ^1H NMR (270Mz, CDCl_3) has δ_{H} at 2.68 (1H, dd $J = 8$ and 9Hz), 3.29 (1H, dd, $J = 1$ and 9Hz), 4.12 (5H, s), 4.23 (1H, dd, $J = 1$ and 8Hz), 4.71 (1H, m), 4.80 (1H, m), 5.04 (1H, m), 6.6-8.2 (30H, m); and the ^{31}P NMR (107Mz, CDCl_3) indicated almost diastereomerically pure complex with δ_{P} 40.8 (d, $J = 45\text{Hz}$) and 75.7 (d, $J = 45\text{Hz}$). It has similar catalytic activity as the SK-Naud catalysts [Nishibayashi et al. *Organometallics* **18** 2291 1999, DOI: 10.1021/om990210o; Sammakia & Stangeland *J Org Chem* **62** 6104 1997, DOI: 10.1021/jo9711044].

Platinum(0) bis(1,5-cyclooctadiene) [$\text{Pt}(\text{COD})_2$, Pt bis(1,2,5,6- η -1,5-octadiene)] [12130-66-4] $(\text{C}_8\text{H}_{12})_2\text{Pt}$, **M 411.6**, **m** $> \sim 200^\circ$. $\text{Pt}(\text{COD})_2$ is prepared in a dry, O_2 -free, N_2 atmosphere by adding dropwise an ethereal solution of $\text{Li}_2(\text{C}_8\text{H}_8)$ [ca 40ml of a 0.24mol dm^{-3} of dilithium cyclooctatetraene [40698-91-7] in Et_2O (Katz *J Am Chem Soc* **82** 3784 1960, see also below)] to a finely powdered suspension of $\text{Pt}(\text{COD})\text{Cl}_2$ (3.7g, 10mmol, see above entry) in 1,5-cyclooctadiene at -30° , stirring for 30 minutes as the temperature rises to -10° ; and then the solvent is evaporated *in vacuo* to dryness. Extraction of the residue with toluene (6 x 50ml) gives a brown solution which is filtered through an Al_2O_3 column (8 x 2.5cm, Brockman Activity III), the filtrate is evaporated to $\sim 15\text{ml}$ and the supernatant is decanted from the off-white product which is washed with several small volumes of cold toluene until the washings are almost colourless. This Pt catalyst (1.6-2.4g, 40-60%) is useful for most purposes. Analytically pure (C and H) white crystals can be obtained by dissolving it in petroleum ether (b 40° - 60°) ($\sim 80\text{ml}$ for each mmol of catalyst), filtering through an Al_2O_3 column (6 x 2.5cm) and cooling to -78° . The ^1H NMR (100MHz, $^*\text{C}_6\text{D}_6$) has τ at 5.80 (m, 8H, $\text{CH}=\text{CH}$, $J_{\text{Pt-H}} = 55\text{Hz}$) and 7.81 (m, 16H, CH_2); and the ^{13}C NMR (25.15MHz, $^*\text{C}_6\text{D}_6$ - ^1H decoupled) has ^{13}C shifts with δ measured relative to SiMe_4 (positive values to high frequency) 73.3 ($\text{C}=\text{C}$, $J_{\text{Pt-C}} = 143\text{Hz}$) and 33.2 (CH_2 , $J_{\text{Pt-C}} = 15\text{Hz}$), and for IR see references. The white crystalline complex is *stable in air* and can be handled without difficulty. [Spencer et al. *Inorg Synth* **19** 213 1979, DOI: 10.1002/9780470132500.ch49; Crascall et al. *Inorg Synth* **28** 126 1990, DOI: 10.1002/9780470132593.ch34.] *Note that attempts to prepare the corresponding Pd complex [Pd(COD)₂] in a similar manner failed; however, it was obtained by the reaction of [Pd(COD)Cl₂] with $\text{Li}_2[\text{C}_8\text{H}_8]$ in the presence of excess of propene as a white crystalline solid which was stable below -20° , but decomposed rapidly to Pd and 1,5-cyclooctadiene at ambient temperatures.* [Green et al. *JCS Dalton Trans* 271 1977, DOI: 10.1039/DT9770000271; Crascall et al. *Inorg Synth* **28** 126 1990, DOI: 10.1002/9780470132593.ch34.] $\text{Pt}(\text{COD})_2$ catalyses the *cis*-diborylation of olefins in $>80\%$ yields under ambient cond-

itions with e.g. CatB-BCat (see below). [Iverson & Smith *Organometallics* **16** 2757 1997, DOI: 10.1021/om970199x; Müller & Göser *Angew Chem Int Ed* **6** 364 1967, DOI: 10.1002/anie.196703642.]

Platinum(0) bis(dibenzylideneacetone) [Pt(dba)₂, bis(C³,O³-η-1,5-diphenyl-1,4-pentadien-3-one)-platinum(0)] [33677-56-4] (C₃₄H₂₈O₂)Pt, M 663.8, m >170° (to Pt and dba). Pt(dba)₂ is a deep purple air stable complex which is prepared in 41% yield from K₂PtCl₄ with 3 mol equivalents of dba in refluxing EtOH under N₂ in the presence of NaOAc. It crystallises from Me₂CO and is soluble in MeOH and CHCl₃. It is useful for preparing complexes with (PPh₃)₂, (AsPh₃)₂, (PEt₃)₂, (AsEt₃)₂, as well as with other ligands. The *tris complex Pt(dba)₃* is formed as a yellow microcrystalline solid in 40% yield by carrying out the above reaction in refluxing aqueous MeOH *in the presence of air or oxygen*. It is stable in the solid form, and is soluble in organic solvents but gives green solutions that rapidly turns purple, from which Pt(dba)₂ and dba (1:1) can be isolated. Physical properties showed that complete dissociation of Pt(dba)₃ had occurred. Both complexes decompose >170° on heating to Pt and dba, on treatment with CO, or when solutions are set aside for long periods. The IR (CHCl₃) of Pt(dba)₂ has ν_{max} at 1656w(CO), 1613vs, br(C=C), 1579m(C=C aromatic), 1544m, br(CO) cm⁻¹; and complex Pt(dba)₃ has ν_{max} at 1652w(CO), 1624vs(C=C), 1593s, br(C=C aromatic), 1579m(C=C aromatic), 1527m br(CO) cm⁻¹. The UV-VIS (CHCl₃) of Pt(dba)₂ has λ_{max} nm (logε) at 235 (4.48), 336 (4.54), 378sh(4.27) 538 (3.92, metal→ ligand: d→ π*); and Pt(dba)₃ has λ_{max} nm (logε) at 235 (4.27), 336 (4.48), 568 (3.51, metal→ ligand: d→ π*); and dba has λ_{max} nm (logε) at 233 (3.90), 325 (4.38). The ¹H NMR (100MHz, CDCl₃) of Pt(dba)₂ and dba are surprisingly similar [see however: Müller & Gröser *Angew Chem Int Ed* **6** 364 1967, DOI: 10.1002/anie.196703642]. It is a useful catalyst for the oligomerisation of acetylenes [Moseley & Maitlis *JCS Chem Commun* 982 1971, DOI: 10.1039/C29710000982; Cherwinski et al. *JCS Dalton Trans* 1405 1974, DOI: 10.1039/DT9740001405], and is used in the Pt(dba)₂-catalysed selective *cis*-addition of bis(pinacolato)borane to terminal alkenes and cyclic alkenes with internal strain to form bis(boryl)alkanes in 76-86% yields at 50° [Ishiyama et al. *JCS Chem Commun* 689 1997, DOI: 10.1039/A700878C] among other uses in catalysis.

Rhodium(II) acetate dimer (2H₂O) [dirhodium tetraacetate, tetrakis(acetato)dirhodium(II)] [15956-28-2] [Rh(C₂H₃O₂)₂]₂·2H₂O, M 478.0. Dissolve 5g of the dimeric salt in boiling MeOH (ca 600ml) and filter. Concentrate the filtrate to 400ml and chill overnight at ca 0° to obtain dark green crystals of the MeOH adduct. Concentration of the mother liquors gives a further crop of [Rh(OAc)₂]₂·2MeOH. The adduct is then heated at 45° in a vacuum for 2 hours (all MeOH is lost) to leave the emerald green crystals of the acetate. [Legzdins et al. *J Chem Soc (A)* 3322 1970, DOI: 10.1039/J19700003322; Rempel et al. *Inorg Synth* **13** 90 1972, DOI: 10.1002/9780470132449.ch16.] Alternatively, dissolve the acetate in glacial AcOH and reflux for a few hours to give an emerald green solution. Evaporate most of the AcOH on a steam bath, then heat the residue at 120°/1hour. Extract the residue with boiling Me₂CO. Filter, concentrate to half its volume and keep at 0°/18hours. Collect the crystals, wash them with ice cold Me₂CO and dry them at 110°. It is moderately soluble H₂O, MeOH, Me₂CO (see above), and in many organic solvents to give green solutions. It forms adducts with MeOH, Me₃N and Me₂S, and gives solutions with different colours varying from green to orange and red depending on solvent and dilution. The IR (Nujol) has ν_{max} at 1580s, 1425s, and 1350m cm⁻¹, and the IR (hexachlorobutadiene) has ν_{max} at 1445s, 1415s and 1350m cm⁻¹ among other bands. [UV: Johnson et al. *Inorg Chem* **2** 960 1963, DOI: 10.1021/ic50009a020; Beilstein **1** H 124, **2** 124; for applications see Fieser **5** 571, **8** 434, **13** 266, **15** 278, **16** 289, **17** 298.]

It is a homogeneous catalyst [Black *Aldrichimica Acta* **15** 13 1982], is used in an efficient synthesis for β-hydroxy-α-acrylates involving the decomposition of diazoester intermediates with concomitant 1,2-aryl migration [Xiao et al. *Tetrahedron Lett* **48** 1147 2007, DOI: 10.1016/j.tetlet.2006.12.062], and is an effective catalyst for the formation of allylsulfonium ylides (e.g. from 3,3-dimethylallyl methyl sulfide and trimethylsilyldiazomethane) which undergo [2.3] sigmatropic rearrangements (e.g. to 2,2-dimethyl-1-methylthio-1-trimethylsilylbut-3,4-ene) [Carter & Van Vranken *Tetrahedron Lett* **40** 1617 1999, DOI: 10.1039/J19700003322]. The reactions of alkyl diazoacetates with carbodiimides (to form 2-imino-4-oxazolines) [Drapier et al. *Tetrahedron Lett* 559 1979, DOI: 10.1016/S0040-4039(01)86000-7], with acetylenes (to form cyclopropenes) [Petiniot et al. *Tetrahedron Lett* 1239 1978, DOI: 10.1016/S0040-4039(01)94511-3], and with ROH, HOH and R'COOH (replacing H to form the respective O-CH₂CO₂Alkyl) [Paulissen et al. *Tetrahedron Lett* 2233 1973, DOI: 10.1016/S0040-4039(01)87603-6] are all catalysed by [Rh(OAc)₂]₂ at about room temperature.

Rhodium(III) acetylacetonate [2,4-pentanedione rhodium(III), Rh(acac)₃] [14284-92-5] Rh(C₅H₇O₂)₃, M 400.2, m 263-264°. It is prepared from Rh(NO₃)₃ solution (0.1g in Rh) in 0.2N HNO₃ (10ml) which is neutralised with aqueous NaHCO₃ (10%) to pH 4, whereby the light yellow hydroxide (or basic nitrate) begins to separate. Acetylacetone (5ml) is added, the mixture is refluxed, and after a few minutes orange-yellow crystals begin to separate while the pH of the solution decreases. After 30 minutes' reflux, the pH of the solution is re-adjusted and reflux is continued for a further 15 minutes. The orange-yellow complex is collected and forms monoclinic plates (0.3g, 75%, m 260°) upon recrystallisation from aqueous MeOH, and sublimates at 240°/1.0mm. It decomposes above 280° depositing a rhodium mirror. It is insoluble in H₂O, slightly soluble in EtOH and petroleum ether, but freely soluble in *C₆H₆ and CHCl₃. It is stable in boiling dilute acids and 10% aqueous NaOH. Molecular weight determination (~490, by Rast in camphor) indicates that it is *monomeric*. [Dwyer & Sargeson *J Am Chem Soc* **75** 984 1953, DOI: 10.1021/ja01100a503; *Beilstein* **1** IV 3677.] Rh(acac)₃ is a very effective catalyst (~90% yields) for the hydrogenation of monocarboxylic or ω-dicarboxylic acids to the respective alcohols in DME at 100 atmospheres (16 hours at 160°) when combined with Re(CO)₁₀ or Mo(CO)₆ [He et al. *Tetrahedron Lett* **36** 1059 1995, DOI: 10.1016/0040-4039(94)02453-I].

Rhodium(I) bis(1,5-cyclooctadiene) tetrafluoroborate complex [bis(1,5-cyclooctadiene)rhodium(I) tetrafluoroborate, Rh(COD)₂⁺.BF₄⁻] [35138-22-8] RhC₁₆H₂₄BF₄Rh, M 406.1, m ~190° (dec), the xHydrate [207124-65-0] has m 165° (dec). This complex is prepared from the dimer [Rh(COD)Cl]₂ (1.47g, 2.98mmol, cf. [12092-47-6] in Catalysts-Part 2) in CH₂Cl₂ (20ml), to which is added COD (1.1ml, 8.97mmol) followed by AgBF₄ (1.33g, 6.83mmol) in Me₂CO (10ml) which results in a deep red solution containing a white precipitate. The mixture is stirred for 20 minutes, filtered through Celite, THF (20ml) is then added and the volume is reduced on a Rotovap at 25° down to 10ml. The deep red crystals are filtered off, washed with THF (2 x 5ml), Et₂O and dried in air (2.35g, 97%). [Schenck et al. *Inorg Chem* **24** 2334 1985, DOI: 10.1021/ic00209a003.] A general purification procedure involves dissolving the complex in the minimum volume of CH₂Cl₂, adding an equal volume of EtOH and completing the crystallisation by dropwise addition of Et₂O, filtration, and drying the solid *in vacuo*. It is used for preparing *cationic COD-Rh complexes* with phosphine ligands for enantioselective [2+2+2] cycloaddition of unsymmetrical diynes with styrene and norbornene derivatives to yield bi- and tetra-cyclic products with good (~50%) to very good (>90%) enantiomeric enrichment [Shibata et al. *Tetrahedron* **63** 12853 2007, DOI: 10.1016/j.tet.2007.10.053], for hydrogenation [Nagel et al. *Chem Ber* **119** 3326 1986, DOI: 10.1002/cber.19861191112; Ojima et al. *Tetrahedron* **45** 6901 1989, DOI: 10.1016/S0040-4020(01)89159-6; Sawamura et al. *J Am Chem Soc* **117** 9602 1995, DOI: 10.1021/ja00142a044], and for hydrosilylation [Takeuchi et al. *J Org Chem* **60** 3045 1995, DOI: 10.1021/jo00115a020].

[Rhodium(*S,S*-Chiraphos)(COD)] ClO₄.THF {[1,2,5,6-η)-1,5-cyclooctadiene][1,2-dimethyl-1,2-ethanediyl-(di-phenylphosphine)-*P,P'*]rhodium(1⁺) perchlorate. THF, η⁴-1,5-cyclooctadiene[(2*S*,3*S*)-2,3-bis(diphenylphosphino)butane]rhodium(I) perchlorate.THF} [61886-03-1, 61886-02-0 THF-free] C₃₆H₄₀P₂Rh⁺ ClO₄⁻ C₄H₈O, M 809.1, 737.0 (THF-free). The *S,S*-catalyst is obtained by adding 70% perchloric acid (0.080g, 1 equivalent) in pure THF (1ml) under N₂ to a mixture of *S,S*-chiraphos (0.244g, see [64896-28-2]) and *[Rh(COD)(acac)] (0.180) in THF (4ml); and the red mixture is allowed to stand at 25° for 14 hours. The bright orange block crystals are collected, washed with cold THF and dried in air to give analytically pure (C, H, P and Cl analysis) catalyst perchlorate THF (0.40g). The presence of solvent is confirmed by NMR. A single crystal X-ray (absolute) structural determination of this THF-pre-catalyst shows that it is as predicted, i.e. the chiral centres are *S* and the methyl groups are equatorial to give a δ-chelate ring [Ball & Payne *Inorg Chem* **16** 1187 1977, DOI: 10.1021/ic50171a042]. The configuration in solution is assumed to be the same, being consistent with the stereochemistry of the catalytically produced products.

The above two complexes are *efficient homogeneous catalysts* for the hydrogenation of α-*N*-acylamioacrylic acids at room temperature and pressure in THF, dioxane or *C₆H₆, EtOH or aqueous EtOH to provide the amino acid derivatives in very high optical purity [e.g. of alanine (91%), DOPA (83%) and tyrosine (92%)], and in almost quantitative chemical yields with turnover numbers ranging from 3 x 10⁻² to 6 x 10⁻⁴ sec⁻¹, from which essentially completely *optically pure 'non-natural' R-α-amino acids* can be obtained by recrystallisation. [Fryzuk & Bosnich *J Am Chem Soc* **99** 6262 1977, DOI: 10.1021/ja00461a014]. An ingenious application of this *S,S*-chiraphos catalyst using hydrogen, deuterium and tritium has been adopted for the asymmetric synthesis of chiral lactic acid in which the methyl group is chiral by virtue of its having a hydrogen, a deuterium and a

tritium atom on the methyl carbon atom. [Fryzuk & Bosnich *J Am Chem Soc* **101** 3043 1979, DOI: 10.1021/ja00505a035]. In the hydrogenation of selected Z-olefins with this pre-catalyst, addition of Et₃N appears to give higher optical yields of products [Onuma et al. *Bull Chem Soc, Jpn* **53** 2012 1980, DOI: org/10.1246/bcsj.53.2012].

* **[Rh(COD)(acac)]** [12245-39-5] **C₁₃H₁₉O₂Rh**, **M 310.2**, **m 138-140°**, is prepared from [Rh(COD)Cl]₂ (4.4g, 9 mmol, see [12092-47-6], cf. Catalysts-Part 2) and acetylacetone (acacH, 0.9ml, 9 mmol) in CH₂Cl₂ (50ml) chilled to -80° by treating dropwise with a solution of Na₂CO₃ (5.7g) in H₂O (100ml) and shaking; then warming to 0° while shaking and more CH₂Cl₂ (50ml) is added. The layers are separated, and the organic layer is evaporated *in vacuo* until the complex crystallises out. [Fryzuk & Bosnich *J Am Chem Soc* **99** 6262 1977, DOI: 10.1021/ja00461a014].

Rhodium(S,S-Chiraphos)(NBD)] ClO₄ {(bicyclo[2.2.1]heptadiene)[(2*S*,3*S*)-2,3-bis(diphenylphosphino)butane]-rhodium (I) perchlorate, [(1,2,5,6-η)bicyclo[2.2.1]hepta-2,5-diene][*S,S*-1,2-dimethyl-1,2-ethanediyl)-bis(diphenylphosphine)-*P,P'*]rhodium(1⁺) perchlorate} [65012-74-0; 65012-73-9 for cation; *Rh(S,S-chiraphos)(MeOH)₂⁺* has 71264-71-6] (**C₂₈H₂₈P₂**)(**C₇H₈Rh⁺**) **ClO₄⁻**, **M 720.97**. This catalyst can be prepared by two methods. In the *first*, 7% HClO₄ (0.173g, 1 equivalent) in THF (6ml) is added to a solution of the complex *[Rh(NBD)(acac)] (0.356g) and *S,S*-chiraphos (0.516g, see [64896-28-2]) in dry, freshly distilled, THF (7.5ml) under N₂, and the deep red coloured solution is allowed to stand at 25° for 24 hours. The orange-red crystals that separate are collected, washed with cold THF, dried at 40° *in vacuo* to give the catalyst *perchlorate* (0.6g), which should be stored at 0° under N₂ in a sealed container. In the *second*, *S,S*-chiraphos (0.308g) and the complex **[Rh(NBD)₂]ClO₄ (0.290g) are dissolved in CH₂Cl₂ (5ml) and THF (5ml) under N₂, followed by addition of hexane (6ml), allowing to stand at 25° for 1 hour, then at 5° for 2 hours. The orange-red needles of the catalyst *perchlorate* (0.43g) are collected, dried and stored as in the first method. *Note* that it loses catalytic activity if stored in air. [Fryzuk & Bosnich *J Am Chem Soc* **99** 6262 1977, DOI: 10.1021/ja00461a014.] Its ³¹P NMR (CDCl₃, with external H₃PO₄ as reference) has δ at +56.9 (d, *J*_{Rh-P} = 153Hz), i.e. downfield from H₃PO₄ [Slack et al. *Inorg Chem* **18** 3125 1979, DOI: 10.1021/ic50201a034]. The salt, (**C₂₈H₂₈P₂**)(**C₇H₈Rh⁺**) **BF₄⁻** [79790-89-9] **M 708.3** forms orange-red crystals from CH₂Cl₂/Et₂O.

*[*Rh(NBD)(acac)*] [32354-50-0] (**C₇H₈Rh(C₅H₇O₂)**), **M 294.2**, **m 176-177°**, is prepared from the dimer [Rh(NBD)Cl]₂ [12257-42-0] and Tl(acetylacetonate) ([25955-51-6]) as described for [(hfacac)(COE)₂Ir] [58616-58-3] [see this chapter and Diversi et al. *J Organomet Chem* **125** 253 1977, DOI: 10.1016/S0022-328X(00)89444-9].

** [*Rh(NBD)₂*]⁺**ClO₄⁻** [60576-59-1] (**C₇H₈**)₂**Rh⁺** **ClO₄⁻**, **M 386.6**, is also prepared from the dimer [Rh(NBD)Cl]₂ (0.35g, [12257-42-0]) and NBD (0.14g) in CH₂Cl₂ (15ml) under N₂, adding AgClO₄ (0.315g), stirring for 1 hour, filtering and adding THF (15ml). When the CH₂Cl₂ in the mixture is evaporated off under a vacuum, [Rh(NBD)₂]ClO₄ separates as orange needles which are collected, washed with cold THF and dried *in vacuo* to give now orange-brown crystals (0.5g). *Alternatively*, add Et₂O instead of THF (15ml), filter, wash with Et₂O, and dry *in vacuo* (yield 90%). Its ¹H NMR in (CD₃)₂CO has δ at 5.23 (br, olefin H), 4.09 (bridgehead CH) and 1.50 (CH₂). [Fryzuk & Bosnich *J Am Chem Soc* **99** 6262 1977, DOI: 10.1021/ja00461a014; cf. Cramer *J Am Chem Soc* **86** 217 1964, DOI: 10.1021/ja01056a022.] [*Rh(NBD)₂*]⁺**BF₄⁻** has [36620-11-8] (**C₇H₈**)₂**Rh⁺** **BF₄⁻**, **M 373.99**, **m 157-159°**, and is prepared as for the perchlorate, but replacing AgClO₄ by an equivalent of AgBF₄, and recrystallising from CH₂Cl₂/Et₂O. The cation [*Rh(NBD)₂*]⁺ has [35015-46-4] (**C₇H₈**)₂**Rh⁺**, **M 287.2**. [Green et al. *J Chem Soc (A)* 2334 1971, DOI: 10.1039/J19710002334; Osborn & Schrock *J Am Chem Soc* **93** 3089 1971, DOI: 10.1021/ja00741a069; Green & Kuc *JCS Dalton Trans* 832 1972, DOI: 10.1039/DT9720000832.]

[Rhodium(R,R-Chiraphos)(NBD)]ClO₄ {(bicyclo[2.2.1]heptadiene)[(2*R*,3*R*)-2,3-bis(diphenylphosphino)butane]-rhodium (I) perchlorate} [74892-62-9] (**C₂₈H₂₅P₂**)(**C₇H₈Rh⁺**) **ClO₄⁻**, **M 720.97**. This pre-catalyst is prepared and purified as its enantiomer above in 77% yield in orange-red crystals after recrystallising from CH₂Cl₂/hexane. Its catalytic properties are very similar to those of its enantiomer above except that optically pure 'natural' *S*-α-amino acids are produced. [Köttner & Gerber *Chem Ber* **113** 2323 1980, DOI: 10.1002/cber.19801130627.] The mechanism of the above asymmetric hydrogenations has been studied in some detail [Brown & Chaloner *J Am Chem Soc* **102** 3040 1980, DOI: 10.1021/ja00529a029; Brown et al. *J Organomet Chem* **216** 263 1981, DOI: 10.1016/S0022-328X(00)85766-6].

[Rhodium (*R*-Prophos)(NBD)] ClO₄. 0.5CH₂Cl₂ {(bicyclo[2.2.1]heptadiene)[*R*(+)-1,2-bis(diphenylphosphino)-propane]-rhodium(I) perchlorate. 0.5CH₂Cl₂, [(2,3,5,6-η)bicyclo[2.2.1]hepta-2,5-diene][(1-methyl-1,2-ethanediyl)bis(diphenylphosphine)-*P,P'*]rhodium(I⁺) perchlorate. 0.5CH₂Cl₂} [67881-59-8; 67884-58-7 for the cation; *Rh*(*R*-prophos)(MeOH)₂⁺ has 71264-72-7] (C₂₇H₂₈P₂)(C₇H₈)Rh⁺ ClO₄⁻, **M 706.9**, decomposes on heating. The precatalyst is made from freshly recrystallised [Rh(NBD)₂]ClO₄ (0.388g, 1mmol, [12257-42-0]) and *R*-prophos (0.437g, 1.106mmol) in a mixture of CH₂Cl₂ (4ml) and pure THF (4ml) under N₂; and to this orange red solution is added hexane (4ml) dropwise, and the mixture is then allowed to stand at 25° for 5 hours then at 5° for 12 hours. The orange-red solid is filtered off quickly, washed with ice-cold THF then hexane, and dried under a stream of dry N₂ to give analytically pure [Rh(*R*-Prophos)(NBD)]ClO₄ (with 0.5CH₂Cl₂ by NMR). It remains catalytically active indefinitely if kept at 0° under N₂. Its ³¹P NMR (CDCl₃, with external H₃PO₄ as reference) has δ at +60.5 (d, *J*_{Rh-P} = 172Hz), +41.8 (q, *J*_{Rh-P} = 139Hz, *J*_{P-P} = 34Hz) i.e. downfield [Slack et al. *Inorg Chem* **18** 3125 1979, DOI: 10.1021/ic50201a034]. Its absolute crystal X-ray structure has been determined and is the one predicted, i.e. the chelate ring is λ, the methyl group is equatorially disposed and the absolute configuration of the diphosphine is *R*. The configuration in solution is assumed to be the same, being consistent with the stereochemistry of the catalytically produced products.

Like the above rhodium complexes of *S,S*-chiraphos, this *R*-prophos rhodium complex is an efficient homogeneous catalyst for the production of α-amino acids, and the optical yields appear to be insensitive to the nature of the substituents on the substrates which provide the ‘natural’ *S*-amino acids in 90±3% optical yields and high chemical yields (<87%) [compare with the *S,S*-chiraphos rhodium precatalysts above which give the ‘non-natural’ *R*-α-amino acids]. Furthermore, the catalyst *R*-prophos rhodium complex can breed its own chirality so that large quantities of *R*-prophos can be made from the catalytic hydrogenation of ethyl acetoxyacrylate, via ethyl *S*(-)-O-acetylactate, by the *R*-prophos catalyst itself. This procedure has been used to produce *S*-prophos if the pre-catalyst used is [Rh(*S,S*-chiraphos)(NBD)]ClO₄. [Fryzuk & Bosnich *J Am Chem Soc* **100** 5491 1978, DOI: 10.1021/ja00485a037.] An ingenious application of this *R*-prophos rhodium precatalyst using hydrogen, deuterium and tritium has been adopted for the asymmetric synthesis of chiral lactic acid in which the methyl group is chiral by virtue of its having a hydrogen, a deuterium and a tritium atom on the methyl carbon atom. The orientations here being opposite to those obtained with the *S,S*-chiraphos rhodium precatalysts. [Fryzuk & Bosnich *J Am Chem Soc* **101** 3043 1979, DOI: 10.1021/ja00505a035.]

Note that these rhodium catalysts are in fact *pre-catalysts* because in the presence of hydrogen their {e.g. [Rh(*S,S*-chiraphos)(NBD)]⁺} strong deep red-orange colours are reduced to a light straw yellow colour of the *true catalyst* {e.g. [Rh(*S,S*-chiraphos)(H)₂]⁺}. The mechanism of these asymmetric hydrogenations has been studied in some detail [Brown & Chaloner *J Am Chem Soc* **102** 3040 1980, DOI: 10.1021/ja00529a029; Brown et al. *J Organomet Chem* **216** 263 1981, DOI: 10.1016/S0022-328X(00)85766-6].

Rhodium(III) chloride [RhCl₃ ± x H₂O] [10049-07-7; 20765-98-4 x H₂O] RhCl₃, **M 209.3** (anhydrous), **m 100°** (hydrate, dec). The anhydrous form is crystalline and is *hydrophobic*. It is a red powder insoluble in H₂O, but soluble in aqueous NaOH or cyanide solutions. The *hydrate*, however, is soluble in H₂O. [cf. preparation of the *trihydrate* by Anderson, Basolo and Onstott [*Inorg Chem* **7** 214 1963, DOI: 10.1002/9780470132388.ch57].

It catalyses the direct conversion of methane to AcOH [Lin & Sen et al. *Nature* **368** 613 1994, DOI: 10.1038/368613a0], and is a catalyst for conjugate reduction of cinnamaldehydes followed by cross-coupling with arylboronic acids [Wang et al. *JCS Chem Commun* 1192 2004, DOI: 10.1039/B401511H]. It also catalyses the deuteration of saturated hydrocarbons [Takahashi et al. *J Am Chem Soc* **97** 7489 1975, DOI: 10.1021/ja00859a019], and in the presence of Ph₃P (*Wilkinson's catalyst*) the trimethylsilylation of methacrylic esters to 1-alkoxy-1-trimethylsilyl oxyethylenes with rearrangement of a methyl group in the presence of Me₃SiH in THF [see Revis & Hilty *J Org Chem* **55** 2972 1990, DOI: 10.1021/jo00296a080.]

Rhodium(II) pentafluorobenzoate dimer [tetrakis(pentafluorobenzoato)dirhodium(II)] [75863-37-5] C₂₈F₂₀O₈Rh₂, **M 1050.82**. This dimer is obtained from a solution of perfluorobenzoic acid (4.5g) and hydrated RhCl₃ (1g) in EtOH (80ml), to which is added NaOH (0.8g), and heated under reflux for 3 hours, cooled and the solid is filtered off. The solid is refluxed for a further hour with fresh EtOH (80ml) and is filtered again. The combined EtOH solutions are evaporated *in vacuo*, the solid green residue is extracted with several volumes of Et₂O until the extracts are colourless; the Et₂O is concentrated and chromatographed on SiO₂ (eluting with toluene/Et₂O 9:1). Some of the complex (1g) is collected in one of the fractions, and dried *in vacuo* for 3 hours

[150° at 10⁻² mm] and is analytically pure. The IR(KBr) has ν_{\max} at 1655m, 1597s, 1525m, 1500s, 1433s, 1405s, 1297w, 1118m, 997s, 942w, 768m cm⁻¹. Like the **triflate salt** (following entry) it catalyses the addition of carbenes (derived from alkyl diazoesters) to aromatic compounds; its catalytic efficiency is unaffected by the bulk of the diazo ester alkoxy group [Anciaux et al. *J Org Chem* **46** 873 1981, DOI: 10.1021/jo00318a010].

Rhodium(II) trifluoroacetate dimer [dirhodium tetra-trifluoroacetate, tetrakis(trifluoroacetato)-dirhodium(II), Rh₂(TfI)₄] [31126-95-1] [(CF₃CO₂)₂Rh]₂, M 657.9. It can be purified in much the same way as the preceding acetate dimer (see [15956-28-2]) and is recrystallised from *C₆H₆ before use. It is a homogeneous catalyst [Black *Aldrichimica Acta* **15** 13 1982]. Tetrakis(perfluorocarboxylato)dirhodium(II) catalyses the addition of carbenes (generated from diazo esters) to aromatic compounds at room temperature to generate 1-carbalkoxycyclohepta-2,4,6-trienes (with high kinetic selectivity for the *nonconjugated* isomers) in very good yields. A competitive reduction of the catalyst occurs simultaneously causing the reaction to slow down, and should be taken into account [Anciaux et al. *J Org Chem* **46** 873 1981 DOI: 10.1021/jo00318a010].

Ruthenium(III) acetylacetonate [Ru(acac)₃] [14284-93-6] (C₅H₇O₂)₃Ru, M 398.4, m 240°(dec). Purify the complex by recrystallisation from *benzene. [Wilkinson *J Am Chem Soc* **74** 6146 1952, DOI: 10.1021/ja01143a538; *Beilstein* **1** IV 3677.] It catalyses the hydrogenation of dimethyloxalate to ethylene glycol under mild conditions [70bar H₂ pressure at 100°] and the best ligand is MeC(CH₂PPh₂)₃ with a trace of Zn (0.07% of oxalate) in MeOH with yields of >84% and with turnover numbers of 875 (turnover frequency/hour of 53.5) [Teunissen & Elsevier *JCS Chem Commun* 667 1997, DOI: 10.1039/A700862G].

Ruthenium (benzylidene)dichloro-bis-(tricyclohexylphosphine) [phenylmethylene-bis-(tricyclohexylphosphine) dichlororuthenium (Grubbs catalyst—first generation) [172222-30-9] C₄₃H₇₂Cl₂P₂Ru, M 823.0, m 153° (dec). Wash it repeatedly with Me₂CO and MeOH and dry it in a vacuum. *Alternatively*, dissolve it in warm CH₂Cl₂, concentrate it to half its volume, filter, add MeOH to precipitate it as purple microcrystals. Filter these off, wash several times with Me₂CO and MeOH and dry them in a vacuum for several hours. [Schwab et al. *J Am Chem Soc* **118** 100 1996, DOI: 10.1021/ja952676d; Miller et al. *J Am Chem Soc* **118** 9606 1996, DOI: 10.1021/ja961626l; Furstner & Langemann *J Am Chem Soc* **119** 9130 1997, DOI: 10.1021/ja9719945.] It is used to catalyse ring-closing metathesis [Schrodi & Pederson *Aldrichimica Acta* **40** 45 2007, Schmidt *Angew Chem Int Ed* **42** 4996 2003, DOI: 10.1002/anie.200301688], and promotes olefin metathesis with ruthenium based catalysts [Grubbs *Tetrahedron* **60** 7117 2004, DOI: 10.1016/j.tet.2004.05.124]. § A polymer supported version is also commercially available [Schwab et al. *Angew Chem Int Ed* **34** 2039 1995, DOI: 10.1002/anie.199520391].

Ruthenium(III) chloride (RuCl₃) [3H₂O 13815-94-6; xH₂O 14898-67-0; Anhydrous 10049-08-8] RuCl₃·xH₂O, M 207.4 (anhydrous), 261.5 (3H₂O), d²⁰ 3.11. The *anhydrous salt* exists in two forms. The **α -form** is produced by the slow reaction of Cl₂ with Ru metal in siliceous containers at >600° to give black lustrous hexagonal crystals which are *antiferromagnetic* and insoluble in H₂O or EtOH [Hill & Beamish *J Am Chem Soc* **72** 4855 1950, DOI: 10.1021/ja01167a002]. The second **β -form** is prepared by heating Ru metal in a stream of CO and Cl₂ at 340°, to avoid the formation of carbonyl compounds such as Ru(CO)₂Cl₂ and is free from the metal or α -RuCl₃. It is formed in dark brown fluffy hexagonal crystals that are soluble in EtOH. It is the metastable form because at or about the transition temperature of 450° the **β -form** is slowly *converted* to the **α -form** (irreversibly, with $t_{0.5}$ ~1 hour), and is the best way to prepare the **α -form**. [Fletcher et al. *Nature* **199** 1089 1963, DOI: 10.1038/1991089a0.]

Hydrated RuCl₃ is one of the *most useful* inorganic ruthenium compounds, and particularly for the preparation of Ru coordinated compounds. It is prepared by evaporating RuO₄ in concentrated hydrochloric acid in a stream of HCl gas. Unlike the anhydrous form, the hydrate is soluble in H₂O, but a fresh aqueous solution (brown to brown-green colour) does not precipitate AgCl with AgNO₃ solution because the halogen atoms (as well as one molecule of H₂O) are coordinated to the metal. However, on warming in H₂O the halogen atoms are displaced by H₂O molecules and the Cl⁻ ions can be precipitated (or titrated) with AgNO₃ [Connick & Fine *J Am Chem Soc* **83** 3414 1961, DOI: 10.1021/ja01477a014]. It is a *soluble catalyst* used for the oxidative cyclisation of 1,7-dienes to oxepane diols [Piccialli et al. *Tetrahedron Lett* **48** 5131 2007, DOI: 10.1016/j.tetlet.2007.05.078], and promotes a site-specific hydroxylation of tertiary carbon-hydrogen bonds of cyclic ethers in the presence of periodate or bromate [Lee et al. *J Org Chem* **72** 5820 2007, DOI: 10.1021/jo070382s].

It also catalyses oxidation reactions with an oxidant, e.g. the methylene group in cyclopropylmethyl-compounds to a carbonyl group in the presence of *meta*periodate [Hasegawa et al. *Chem Lett (Jpn)* 1385 1985, DOI: 10.1246/cl.1985.1385; Carlsen et al. *J Org Chem* **46** 3936 1981, DOI: 10.1021/jo00332a045; cf. Review Gore *Platinum Metals Rev* **27** 111 1983]. It catalyses the synthesis of 2-ethyl-3-methylquinolines from primary aromatic amines and triallylamine [Cho et al. *Tetrahedron Lett* **40** 1499 1999, DOI: 10.1016/S0040-4039(98)02661-6], and has been used for the selective hydrogenation of unsaturated aldehydes [Fujita et al. *J Catal* **255** 95 2004, DOI: 10.1016/j.jcat.2004.03.037]. [For further applications see Fieser **4** 421, **8** 437, **10** 343, **11** 462, **13** 268, **15** 280.]

Ruthenium [1,3-bis-(2,4,6-trimethylphenyl)-2-imidazolidinylidene] dichloro (benzylidene) (tri-cyclohexylphosphine) {Benzylidene-[1,3-bis(2,4,6-trimethylphenyl)-4,5-dihydro-2-imidazolinyldene]dichloro-(tri-cyclohexylphosphine)-ruthenium, [(1,3-Bis(2,4,6-trimethylphenyl)-2-imidazolinyldene) dichloro (phenyl-methylene)(tricyclophosphine)ruthenium, (Grubbs catalyst—second generation)} [246047-72-3] **C₄₆H₆₅Cl₂N₂PRu, M 849.0, m 143.5-148.5°**. This catalyst is prepared by adding a solution of *tert*-BuOK (1.25g, 10.2mmol, 1.4 equiv) in dry THF (200ml) slowly to a suspension of 1,3-dimesityl-4,5-dihydroimidazolium tetrafluoroborate (4.2g, 10.2mmol, 1.4 equivalent) in dry THF (100ml) in a flame dried Schlenk flask (500ml) under N₂ (magnetic stirring) at ~20°. The BF₄⁻ salt dissolves immediately, the cloudy yellow mixture is stirred at ~20° for 1 hour, and then transferred (cannula) to a larger Schlenk flask (twice size) under argon. To this is added RuCl₂(=CHPh)(PCy₃)₂ (7.29, 7.29mmol, 1.0 equivalents) in dry *C₆H₆ (400ml) and heated at 80° for 30 minutes when the reaction is complete (by ¹H NMR). The volatiles are removed *in vacuo* and the residue is washed with dry MeOH, or pentane (4 x 100ml) to provide the active catalyst as a pink-brown microcrystalline powder (4.64g, 75%). It has ¹H NMR (400MHz, CD₂Cl₂) with δ at 19.16 (s, 1H), 7.37-7.05 (m, 9H), 3.88 (s, 4H), 2.56–0.15 (m, 51H); ³¹P NMR (161.9MHz, CD₂Cl₂) with δ at 31.41 and HRMS (FAB) at 848.3306 for M⁺ [Scholl et al. *Org Lett* **1** 953 1999, DOI: 10.1021/ol990909q].

It is used in ruthenium catalysed *ring closure metathesis (RCM)* and olefin metathesis [Kulkarni & Diver *J Am Chem Soc* **126** 8110 2004, DOI: 10.1021/ja0476922; Schmidt *Angew Chem Int Ed* **42** 4996 2003, DOI: 10.1002/anie.200301688; Scholl et al. *Org Lett* **1** 953 1999, DOI: 10.1021/ol990909q], and to generate unsaturated sultones (e.g. 2,7-3*H*-dihydro-[1,2]oxathiepine-2,2-dioxide) from olefinic sulfonates (e.g. but-3-enyl allyl-sulfonate) *via* ring closure metathesis [Le Flohic et al. *Tetrahedron* **62** 9017 2006, DOI: 10.1016/j.tet.2006.07.010].

Ruthenium [1,5-Cyclooctadiene][1,3,5-cyclooctatetraene] (Ru[cod][cot]) [127382-91-6] **C₁₆H₂₂Ru, M 315.4, m 88-94°, 92-94°**. For preparation see Chapter 4, ‘Metal-Organic Compounds’. Purify the ruthenium complex (~0.3g) by dissolving it in *n*-pentane (~70ml) and filtering the solution through a column of alumina (Merck, Brockman activity II–III, 20cm). Collect the yellow band and reduce its volume to ~5ml then cool it at -70°, preferably under N₂ or argon. After ~6 hours collect the yellow solid (~0.2g) and dry it *in vacuo*. Recrystallisation from *n*-pentane results in a 75-85% recovery. [Itoh et al. *J Organomet Chem* **272** 179 1984, DOI: 10.1016/0022-328X(84)80465-9; Petrici et al. *JCS Dalton Trans* 1961 1980, DOI: 10.1039/DT9800001961]. It is a highly selective catalyst for amine alkylations in which, unlike other ruthenium catalysts, alkylates the amino group of e.g., 2-, 3-, or 4- aminopyridines as well as 2-aminopyrimidine to give the respective monoethylamino derivatives by using EtOH in high yields with very little, if any, of diethylamino derivatives. Thus it stops almost *exclusively at monoalkylation*. [Watanabe et al. *J Org Chem* **61** 4214 1996, DOI: 10.1021/jo9516289]. It also catalyses the dimerisation of 2,5-norbornadiene *via* [2+2] cycloaddition reactions and eyne generation in the presence of dimethylfumarate in THF at 40° for 1 hour to form the ‘*cup-shaped*’ molecule *pentacyclo[6.6.0.0^{2,6}.0^{3,13}.0^{10,14}]tetradeca-4,11-diene* in 96% yield. An alicyclic compound which would otherwise require many steps to synthesise. [Mitsudo et al. *J Am Chem Soc* **121** 1839 1999, DOI: 10.1021/ja9835741.]

Ruthenium [(R-P-Phos)(acac)₂] {Ruthenium [R-(+)-2,2',6,6'-tetramethoxy-4,4'-bis(diphenylphosphino)-3,3'-bipyridine]-bis(acetylacetonate)} is prepared by mixing equimolar amounts of *R*-P-Phos (see *R*-(+)-[221012-82-4] in ‘Part-2’) with Ru(acac)₃ [14284-93-6] in the presence of a reducing agent (Zn powder) in refluxing EtOH to give the solid catalyst in 97% yield which was characterised by ¹H, ³¹P NMR, elemental analysis and X-ray crystallographic structure determination. *Ru[(S-P-Phos)Cl₂(DMF)_n]* is prepared in a similar manner with the appropriate ingredients (using *S*-(*-*)-*P*-Phos [362524-23-0]) in DMF. [Pai et al. *J Am Chem Soc*

122 11513 2000, DOI: 10.1021/ja000163n.] These (see also below) *atropisomeric* bi-heteroaromatic diphosphines (P-Phos and its variants) which complex with transition metals are effective in catalysing asymmetric reactions including asymmetric hydrogenation of 2-arylacrylates, β -ketoesters, arylketones, hydrosilylation and C-C bond formation with high stereospecificity. [Wu & Chan *Acc Chem Res* **39** 711 2006, DOI: 10.1021/ar0680015; Au-Yeung & Chan *Coordination Chemistry Reviews* **248** 2151 2004, DOI: 10.1016/j.ccr.2004.08.026; Pai et al. *J Am Chem Soc* **122** 11513 2000, DOI: 10.1021/ja000163n.]

Ruthenium [(*R*-Xylyl-P-Phos)(C₆H₆)]Cl₂ {**Ruthenium** *R*-(+)-2,2',6,6'-tetramethoxy-4,4'-bis[di(3,5-xylyl)phosphino]-3,3'-bipyridine(benzene) dichloride] is prepared in 88% yield by mixing equimolar amounts of [RuCl₂(C₆H₆)₂] [37366-09-9] with *R*-Xylyl-P-Phos (see *R*-(+)- [442905-33-1] in 'Part-2') in EtOH/*C₆H₆ (8:1) and heating at 50-60° for 1 hour. This ruthenium complex is a highly active, enantioselective, air stable catalyst for the asymmetric hydrogenation of β -ketoesters (P_{H2} = 200-350psi at 70-90° for 1-15 hours) to optically active β -hydroxyesters with ~100% conversion and >91% enantiomeric enrichment. It is *very stable* in the solid state in air; and a stirred solution for 10 hours under air showed no loss of activity or stereoselectivity; and its ³¹P NMR (200MHz, CDCl₃) is not different from that obtained for the original sample under N₂ which has an AB set of signals with δ_P at 33.49 (d, J_{PA-PB} = 62.9Hz) and 39.96 (d, J_{PA-PB} = 62.5Hz). The coordination of ruthenium to benzene makes the phosphorus atoms *non-equivalent*. The resonance observed in the ¹H NMR (500MHz, CDCl₃) at 5.65ppm is assignable to the 6H resonances of the η^6 -coordinated benzene ring [Wu et al. *Tetrahedron Lett* **43** 1539 2002, DOI: 10.1016/S0040-4039(02)00062-X]. Other P-Phos related ligands, such as the chiral 4-bis(tolylphosphino) (*Tol-P-Phos*) and chiral 4-bis(cyclohexylphosphino) (*Cy-P-Phos*) ligands, were prepared and purified in a similar, or slightly modified, manner using the appropriate phosphines. They also form ruthenium complexes that have related catalytic activities [Wu & Chan *Acc Chem Res* **39** 711 2006, DOI: 10.1021/ar0680015; Au-Yeung & Chan *Coordination Chemistry Reviews* **248** 2151 2004, DOI: 10.1016/j.ccr.2004.08.026].

Ruthenium(II)-tris(triphenylphosphine) dichloride [tris(triphenylphosphine)ruthenium(II) dichloride] [15529-49-4] [(C₆H₅)₃P]₃RuCl₂, **M 958.8; 433 (in Me₂CO/N₂), m 132-134°**. This catalyst (compare with Wilkinson's Rhodium catalyst above) is obtained by heating under reflux RuCl₃·3H₂O (0.26g, 1mmol) and Ph₃P (1.57g, 6mmol) in MeOH (65ml) under N₂ for 3 hours, when the reddish-brown tris-complex separates. It is washed well with MeOH, then Et₂O and dried *in vacuo* at 60° (yield 75%). It is soluble in CHCl₃, Me₂CO, *C₆H₆, EtOAc and hot *i*-PrOH, but insoluble in H₂O, Et₂O and sparingly soluble in alcohol. When less Ph₃P is used in e.g. MeOH, complexes such as *Ru(III)-di(Ph₃P)(MeOH) Cl₂* are formed. *Note that if the above mixture is shaken, not refluxed, Ru(II)-tetrakis(triphenylphosphine) dichloride is formed which turns green on exposure to air.* [Stephenson & Wilkinson *J Inorg Nucl Chem* **28** 945 1966, DOI: 10.1016/0022-1902(66)80191-4; Sammakia & Stangeland *J Org Chem* **62** 6104 (footnote) 1997, DOI: 10.1021/jo9711044.] The *tris-complex* catalyses a variety of reactions including hydrogen transfer [Sasson & Blum *Tetrahedron Lett* 2167 1971, DOI: 10.1016/S0040-4039(01)96811-X; Regen & Whitesides *J Org Chem* **37** 1832 1972, DOI: 10.1021/jo00976a038], insertion of *t*-BuOO- into the carbon atom α to the N-atom of, e.g. *N*-methoxycarbonyl pyrrolidine and 1,2,3,4-tetrahydroisoquinoline with *t*-BuOOH [Kondo et al. *J Org Chem* **56** 487 1991, DOI: 10.1021/jo00002a003], intramolecular cyclisation of unsaturated α,α -dichloroesters or amides [Hayes et al. *J Org Chem* **51** 5501 1986, DOI: 10.1021/jo00376a109], the reduction of -CHO by HCOOH/Et₃N/THF without affecting NO₂, RCOR, ester or *tert*-amide groups [Kai & Arcelli *Tetrahedron Lett* **26** 3365 1985, DOI: 10.1016/S0040-4039(00)98299-6], in the synthesis of furans from allenyl sulfides [Peng et al. *Angew Chem Int Ed* **46** 1905 2007, DOI: 10.1002/anie.200604299], and for other examples of catalysis by this Ru-complex see Fieser **4** 564, **5** 740, **6** 654, **10** 141, **12** 179, **13** 107, **14** 130, **16** 126. When complexed further with chiral (phosphinoferrocenyl)oxazolines, hydrogen transfer is not only considerably enhanced but is also highly stereospecific [see further in this chapter].

Samarium (II) iodide (SmI₂) **Samarium (II) iodide (SmI₂)** see [32248-43-4]; **Silver acetate (AgOAc)** see [563-63-3], and **Silver tetrafluoroborate [AgBF₄]** see [14104-20-2], all in Chapter 4, 'Inorganic Compounds'.

Tetrakis(triphenylphosphino)palladium(0) [palladium-tetrakis(triphenylphosphine)] [14221-01-3] [(C₆H₅)₃P]₄Pd, **M 1155.6, m 100-105°, 115°, ~116° (dec, sealed tube under N₂)**, (**m** is unreliable and is not

a criterion of purity because it varies). This catalyst is prepared from PdCl_2 (5.9g, 0.1mol), Ph_3P (43.7g, 0.5ml) in Me_2SO (400ml) under a vacuum N_2 system with pressure release, and the yellow mixture is heated (oil bath) with stirring until dissolution (140°). The bath is removed, the mixture is stirred rapidly for 15 minutes and hydrazine hydrate (6.7g 0.4mol) is injected within 1 minute. A vigorous evolution of N_2 occurs and the dark solution is rapidly cooled with a cool water bath and crystallisation takes place ($\sim 125^\circ$). When the temperature reaches $\sim 20^\circ$ the solid is filtered off (coarse sintered glass) under N_2 , washed with EtOH (2 x 50ml), Et_2O (2 x 50ml) and the yellow crystalline solid is dried by a slow stream of N_2 through the funnel overnight (37.4g 97%). **Alternatively**, it can be prepared freshly by mixing $\text{Pd}(\text{NO}_3)_2$ (2mmols) and PPh_3 (2mmols) in hot $^*\text{C}_6\text{H}_6$ when vigorous evolution of nitric oxide occurs (fume cupboard) and a solid mass separates. This is collected and crystallised from EtOH. It should not be heated excessively as it dissociates to $\text{Pd}(\text{PPh}_3)_3$ and PPh_3 , and then further to $\text{Pd}(\text{PPh}_3)_2$ and PPh_3 . It is also air sensitive as PPh_3 is oxidised to PPh_3O . It is stable only for short periods because on exposure to heat or air it turns from yellow to orange and dissociates in solution, so the solutions should be used directly. Its **cryoscopic constant** in $^*\text{C}_6\text{H}_6$ (at 0.601g/20ml) corresponds to **M 1156** [Malatesta & Angoletti *J Chem Soc* 1186 1957, DOI: 10.1039/JR9570001186]. Molecular weight determination in $^*\text{C}_6\text{H}_6$ indicates considerable dissociation, and the solution absorbs O_2 rapidly to give an insoluble green oxygen complex [Nyman et al. *J Chem Soc A* 561 1968, DOI: 10.1039/J19680000561]. It is moderately soluble in $^*\text{C}_6\text{H}_6$ (5g/100ml), CH_2Cl_2 and CHCl_3 but less soluble in Me_2CO , THF and MeCN. $\text{Pd}(\text{Ph}_3\text{P})_4$ may be handled in air but it is best stored under N_2 . [Coulson et al. *Inorg Synth* **28** 107 1990, DOI: 10.1002/9780470132593.ch28; Malatesta & Angoletta *J Chem Soc* 1186 1990, DOI: 10.1039/JR9570001186; Beilstein **16** IV 954, Fieser **6** 571, **7** 357, **8** 472, **9** 451, **10** 384, **11** 503, **12** 468, **13** 289, **14** 295, **15** 300, **16** 317, **17** 327.]

$\text{Pd}(\text{Ph}_3\text{P})_4$ is a very **versatile catalyst** for promoting the dimerisation of butadiene to 1,3,7-octatriene [Takahashi et al. *Bull Chem Soc Jpn* **41** 454 1968, DOI: org/10.1246/bcsj.41.454], catalysing various coupling reactions, without homo-coupling occurring [Brocato et al. *Tetrahedron Lett* **33** 7433 1992, DOI: 10.1016/S0040-4039(00)60208-3; Arcadi et al. *Tetrahedron Lett* **34** 2813 1993, DOI: 10.1016/S0040-4039(00)73569-6; McClure & Danishefsky *J Am Chem Soc* **115** 6094 1993, DOI: 10.1021/ja00067a026; Paquette & Astles *J Org Chem* **58** 165 1993, DOI: 10.1021/jo00053a031; Sahoo et al. *Org Lett* **8** 4141 2006, DOI: 10.1021/ol061763f], including **Suzuki** coupling [Trost *Tetrahedron* **33** 2615 1977, DOI: 10.1016/0040-4020(77)80284-6]. [Beilstein **16** IV 954.] This palladium catalyst bound to a polymer support ($\sim 0.06\text{mmol/g}$) is also commercially available [cf. Fenger & Le Drain *Tetrahedron Lett* **39** 4287 1998, DOI: 10.1016/S0040-4039(98)00757-6].

Tetra-*n*-propylammonium perruthenate (TPAP, tetrapropyl tetraoxoruthenate) [114615-82-6] (C_3H_7)₄ NRuO_4 , **M 351.4**, **m 160° (dec)**. This stable dark green solid is a useful catalyst for a variety of oxidation reactions at about room temperature. It is used at a concentration of about 5 mol% with an equivalent of ***N*-methylmorpholine *N*-oxide (NMO)** which provides the required oxygen with typical turnovers of about 250. The reactions are carried out in CH_2Cl_2 and/or MeCN, although the latter assists the reactions in CH_2Cl_2 considerably. **Care** should be taken to add TPAP slowly as the reaction can be vigorous. Addition of finely ground **4Å molecular sieves moderate** the rate and efficiency of the reaction. The reactions usually proceed to completion within 5 minutes to one hour. Cooling and slow addition of TPAP is necessary with reactions on a large scale. Workup is simple, and is carried out by passing the solution through a short silica-gel column and eluting with EtOAc followed by evaporation and crystallisation or distillation of the product. When MeCN is the solvent, or is present in the solvent, it is advisable to evaporate the solvent prior to passage through the silica-gel column. [Ley et al. *Synthesis* 639 1994, DOI: 10.1055/s-1994-25538; see entry in the ‘Metal-Organic Compounds’ chapter 4, for further information].

Triphenylphosphinegold(I) bis(trifluoromethanesulfonyl)imide ($[\text{Ph}_3\text{PAu(I)}]^+ [\text{NTf}_2]^-$) [866395-16-6] ($[\text{C}_6\text{H}_5]_3\text{PAu}^+$ (CF_3SO_2)₂ N^- , **M 739.4**, **m $>230^\circ$** . AgNTf_2 is first prepared by mixing Ag_2CO_3 (276mg, 1.0mmol) and HNTf_2 (562mg, 2.0mmol, 2 equivalents [82113-65-3]) in H_2O (10ml), and refluxing for 3 hours (to eliminate CO_2); then evaporating to dryness *in vacuo*. It can be used as a solid or dissolved in dry CH_2Cl_2 (5ml). [Li et al. *J Org Chem* **73** 4323 2008, DOI: 10.1021/jo8003875.] The versatile catalyst is then prepared by dissolving Ph_3PAuCl (2mmol, [14243-64-2]) in CH_2Cl_2 (5ml) and adding to it solid AgNTf_2 (786mg, 2mmol) or the preceding CH_2Cl_2 (5ml) solution. Immediate separation of AgCl occurs. After stirring for 15 minutes the ^{31}PMR reveals a single peak of the Au salt. The AgCl is removed by filtration through Celite, the pale col-

oured solution is evaporated *in vacuo*, and the residue is dried *in vacuo* to give a **quantitative yield** of the catalyst. It forms small white crystals which are air stable, but should be preferably stored in a dry inert atmosphere. It is soluble in Me₂CO, CH₂Cl₂, and CHCl₃ but insoluble in petroleum ethers. Its IR (CCl₄) has ν_{\max} at 1482, 1437, 1405, 1384, 1216, 1196, 1133, 1103 and 960, cm⁻¹; the ¹H NMR (400MHz, CDCl₃) has δ_{H} at 7.47-7.55 (m, 12H, aromatic H) and 7.56-7.62 (m, 3H, aromatic H close to Au); the ³¹P NMR (121.5MHz, CD₂Cl₂) has δ_{P} at 30.7, and for ¹³C NMR see references. [Mézaillies et al. *Org Lett* **7** 4133 2005, DOI: 10.1021/ol0515917.] NTf₂⁻ is a weakly coordinating counter ion that confers stability to the complex. ³¹P NMR spectra of pre-formed solutions of the complexes Ph₃PAuBF₄, Ph₃PAuPF₆ and Ph₃PAuSbF₆, using the respective Ag salts, showed that they were not very stable. Attempts to isolate the complexes were unsuccessful. [Mézaillies et al. *Org Lett* **7** 4133 2005, DOI: 10.1021/ol0515917.]

At concentrations of 1-2mol% in CH₂Cl₂ or Me₂CO, [Ph₃PAu(I)]⁺ [NTf₂]⁻ catalyses the stereoselective isomerisation of butynediol monobenzoates into functionalised 2,5-dihydrofurans (in a sequence of two steps) in high yields at room temperature and in 15 to ~30 minutes [Buzas et al. *Org Lett* **8** 1957 2006, DOI: 10.1021/ol0606839]. It is **exceedingly active** in catalysing a wide variety of *eyne* cycloisomerisations [Mézaillies et al. *Org Lett* **7** 4133 2005, DOI: 10.1021/ol0515917]. Also under similar mild conditions, in the presence of electrophilic bromine (e.g. from *N*-bromosuccinimide) or iodine (e.g. from *N*-iodosuccinimide), it catalyses the formation of linear α -halo-enones from propargyl acetates [Yu et al. *Tetrahedron* **65** 1846 2009, DOI: 10.1016/j.tet.2008.11.107], and 4-haloalkylidene-1,3-dioxalan-2-ones with propargyl *tert*-butylcarbonates [Buzas et al. *Tetrahedron* **65** 1889 2009, DOI: 10.1016/j.tet.2008.11.108], products which are suitable substrates for Pd-catalysed cross coupling reactions. Like the following OTf⁻ Au catalyst, it promotes the glycosylation of 1,2-anhydrosugars as donors using protected sugars with one free sterically unhindered OH group [Li et al. *J Org Chem* **73** 4323 2008, DOI: 10.1021/jo8003875].

Triphenylphosphinegold(I) trifluoromethanesulfonate ([Ph₃PAu(I)]⁺ [OTf]⁻) [156397-47-6] [(C₆H₅)₃PAu]⁺ CF₃SO₃⁻, **M 608.2, m >230°**. Ph₃PAu(I)⁺ [OTf]⁻ differs from the preceding catalyst in the counter anion, and is **much less air and moisture sensitive**. Due precautions need to be taken in use and storage however. It is prepared by dissolving Ph₃PAuCl (0.1mmol, [14243-64-2]) and silver trifluoromethanesulfonate (0.1mol, [2923-28-6]) in dry CH₂Cl₂ (1ml) and stirred for 5 minutes. Filtration from AgCl provides a 0.1M solution of catalyst in CH₂Cl₂. Generally the AgCl does not interfere in the catalytic process and so the catalyst can be prepared *in situ*. Colourless Ph₃PAu(I)⁺ [OTf]⁻ can be obtained by evaporating the CH₂Cl₂ solution and stored appropriately. It is a superior catalyst to anhydrous ZnCl₂ for the well established glycolsylation reactions with 1,2-anhydrosugars as donors [Li et al. *J Org Chem* **73** 4323 2008, DOI: 10.1021/jo8003875], and it catalyses the intramolecular hydroamination of terminal alkenes in high yields with 1-5mol% of catalyst by heating in toluene at 100° for 12-48 hours, or the intra- and inter- molecular hydroamination in ClCH₂CH₂Cl by microwave radiation as heat source in ~30 minutes in high yield [Liu et al. *Org Lett* **8** 2707 2006, DOI: 10.1021/ol060719x]. In a useful application, the catalyst [Ph₃PAu(I)]⁺ [OTf]⁻ promotes a cascade cyclisation/oxidative cleavage of a carbon-carbon triple bond in *Z*-enynols, e.g. 5-Ph- or 5-*n*-butyl- pent-2-ene-4-yne-1-ols, in the presence of molecular oxygen to give high yields of the corresponding 2,5-dihydrofuran-2-ones and releasing C-5 with its substituent as the aldehyde or acid. The reaction involves free radicals as it is completely suppressed in the presence of the radical scavenger 4-hydroxy-TEMPO [Liu et al. *J Am Chem Soc* **128** 11332 2006, DOI: 10.1021/ja062610q].

Tris(dibenzylideneacetone)dipalladium(0) chloroform adduct [Pd₂(dba)₃(CDCl₃)] [52522-40-4] (C₁₇H₁₄O)₃Pd₂, **M 1035.1, m 122-124°, 131-135°; solvent free tablet form (1μmol Pd per MgAlsilicate tablet) Pd₂(dba)₃ [51364-51-3] M 915.7, m 152-155°**. When PdCl₂ (1.05g, 5.92mmol) is added to a solution of dibenzylidene acetone (dba, 4.60g, 19.6mmol) and NaOAc (3.9g, 47.5mmol) in MeOH (150ml at 50°), mixed at 40° for 4 hours then cooled, a reddish-purple precipitate separates. This is collected, washed with H₂O then Me₂CO and dried *in vacuo*. The precipitate (3.39g) is purified by dissolving it in hot CDCl₃ (120ml), filtering, and to the deep violet filtrate is added slowly Et₂O (170ml) when the **chloroform adduct** separates as deep purple needles which are collected and dried *in vacuo* (80% yield, **m 122-124°**). Recrystallisation from *C₆H₆ instead of CHCl₃ gives deep-violet needles of **tris(dibenzylideneacetone)-dipalladium(0) benzene adduct** (62.5% yield, **m 141-142°**). By using toluene instead of CHCl₃, deep-violet needles of **tris(dibenzylideneacetone) dipalladium(0) toluene adduct** (36% yield, **m 140-141°**) are obtained. The solvates can be exchanged with each other (i.e. CDCl₃, *C₆H₆ or toluene) without affecting the coordinating reactivity of the complex with other ligands such as Ph₃P, Bipy, *o*-phen, olefins, *o*-quinones etc. [Ukai et al. *J Organomet Chem* **65** 253 1974,

DOI: 10.1016/S0022-328X(00)91277-4; for applications see Fustero *Org Lett* **8** 4129 2006, DOI: 10.1021/ol061733c.] If the procedure is used with bis(*p*-methoxybenzylidene)acetone and the complex crystallised from CDCl₃/Et₂O, **Pd₂(*p,p'*-methoxy-dba)₃** is obtained as deep violet needles (72.3% yield, m 141-143°) which do not complex with the solvent. [cf. **Pd(dba)₂** above.] [Ukai et al. *J Organomet Chem* **65** 253 1974, DOI: 10.1016/S0022-328X(00)91277-4.]

Tris(dibenzylideneacetone)palladium [Pd(dba)₃]: When a fourfold excess of dba is added to solution of Pd(dba)₂ or Pd₂(dba)₃ in *C₆H₆, heated and reduced in volume, the dark red colour of Pd(dba)₂ turns to brown, and on complete evaporation, orange brown crystals of **Pd(dba)₃** contaminated with yellow crystals of dba are obtained. Washing the crystals with *C₆H₆ leads to crystal decomposition. The IR of the complex has ν_{\max} at 1651 (C=O) and olefine bands at 1598, 1580 and 1531 cm⁻¹. X-ray analysis of the Pd(dba)₃—*C₆H₆ crystals showed that each pentadienone ligand is coordinated through one olefin group, and the Pd atom is trigonal with ~C₃ symmetry [Mazza & Pierpont *Inorg Chem* **12** 2955 1973, DOI: 10.1021/ic50130a043; cf. *Handbook of Organopalladium for Organic Synthesis* Negishi ed. Wiley, NJ 2002, ISBN 0-471-31506-0].

Tris(triphenylphosphinegold(I))oxonium tetrafluoroborate ([Ph₃PAu(I)]₃O⁺ [BF₄]⁻) [53317-87-6] **C₅₄H₄₅P₃OAu₃BF₄**, **M 1480.6, m 207° (dec), 207-208° (dec), 220-221° (dec)**. The catalytic oxonium salt is prepared by adding a solution of AgBF₄ (0.2g, 1.03mmol) in MeOH (5ml) to a solution of Ph₃PAuCl (0.5g, 1.01mmol, [14243-64-2]) in THF (20ml) and the AgCl that precipitated is filtered off. A solution of KOH (0.1g, 1.78mmol) and NaBF₄ (0.5g, 4.55mmol) in MeOH (100ml) is added to the filtrate, stirred for 1 hour, the solvent is evaporated off *in vacuo*, the residue is extracted with CHCl₃ (2 x 30ml). The combined extract is filtered, and hexane (100ml) is added to precipitate the oxonium salt which gives analytically pure [Ph₃PAu(I)]₃O⁺ [BF₄]⁻ (0.42g, 84%), **m 220-221° (dec)**, after recrystallisation from a saturated CHCl₃ solution on adding ~1.5-fold volume of Me₂CO. **Alternatively**, Ag₂O freshly prepared from AgNO₃ (5.0g, 29.5mmol), and NaBF₄ (5.0g, 45.5mmol, finely powdered) are added to a solution of Ph₃PAuCl (4.0g, 8.08mmol) in Me₂CO (600ml), stirred vigorously for 1 hour, and the solvent evaporated *in vacuo*. The residue is extracted with *C₆H₆ to removed unreacted Ph₃PAuCl, and the oxonium salt is extracted with CHCl₃ (3 x 40ml), filtered, and hexane (400ml) is added to crystallise [Ph₃PAu(I)]₃O⁺ [BF₄]⁻ (3.6g, 90%) out. It is soluble in CHCl₃ and CH₂Cl₂ but insoluble in hexane and Et₂O. Its IR (nujol mull) has ν_{\max} at 1050-1070 (br, BF₄⁻) cm⁻¹. It is a versatile air and moisture tolerant catalyst that has been used in a variety of reactions (see below). X-ray crystallography showed that in the crystals the oxonium ions are **dimeric**, with the two pyramidal monomeric (Ph₃PAu)₃(μ₃-O)⁺ fragments interacting *via* Au-Au' bonds (~3.16Å) involving two of the three Au atoms in each unit. The pyramidal structure has the O⁺ atom centrally above the Au₃ triangular plane, and the resulting six-membered Au₂OAu₂O heterocycle has a chair conformation. The **oxonium trifluoroacetate m 209-210° (dec)** from CHCl₃/*C₆H₆, and the **oxonium permanganate m 131-131.5° (dec)** from CHCl₃/hexane (1.2:4.0 v/v) were prepared similarly. [Nesmeyanov et al. *J Organomet Chem* **201** 343 1980, DOI: 10.1016/S0022-328X(00)92589-0; cf. Bruce et al. *Inorg Synth* **26** 324 1988, DOI: 10.1002/9780470132579.ch59.] Oxonium complexes with other phosphine ligands have been similarly prepared and if H₂¹⁷O is used in the preparations, then it is incorporated into the oxygen of the cation. **[Ph₃PAu(I)]₃¹⁷O⁺ [BF₄]⁻** has ³¹P NMR (36MHz, CD₂Cl₂, external 85% H₃PO₄) with δ_P at 24.0, and ¹⁷O NMR (CD₂Cl₂ and external H₂O) with δ_{17O} at +19.7 (br s, w/2 = 152Hz). [Yang et al. *Inorg Chem* **32** 1946 1993, DOI: 10.1021/ic00062a012.]

[Ph₃PAu(I)]₃O⁺ [BF₄]⁻ catalyses the Claisen rearrangement of propargyl vinyl ethers, e.g. 1-phenylhept-2-yne-1-yl vinyl ether, to respective aldehydes which are usually reduced *in situ* with NaBH₄ to give the homoallenlic alcohol, i.e. 1-phenyl-3-*n*-butyl-3-(2-hydroxyethyl)allene, in high yield with 1mole% of catalyst in CH₂Cl₂ at room temperature. Chirality is efficiently transferred in the rearrangement. [Sherry & Toste *J Am Chem Soc* **126** 15978 2004, DOI: 10.1021/ja044602k]. In the presence of sterically hindered secondary amines, e.g. (iso-Pr)₂NH or (iso-Pr)CyNH, it catalyses the **5-*exo-dig* cyclisation** of formyl alkynes, e.g. 6-formyl-4,4-bis(methoxycarbonyl)hex-1-yne, to 1,1-(bismethoxycarbonyl)-3-formyl-4-exomethylenecyclopentane in CDCl₃, 70°/3-24 hours, in ~70% yields [Binder et al. *Org Lett* **10** 1025 2008, DOI: 10.1021/ol800092p]. In the presence of O₂ in THF at ~50° [Ph₃PAu(I)]₃O⁺ [BF₄]⁻ also catalyses the cascade cyclisation/oxidative cleavage of a carbon-carbon triple bond in Z-enynols efficiently [see Ph₃PAu(I)]⁺ [OTf]⁻ above, Liu et al. *J Am Chem Soc* **128** 11332 2006, DOI: 10.1021/ja062610q].

(Xantphos)Rh(H)(CO)(PPh₃) is prepared by stirring a solution of (PPh₃)Rh(H)(CO) (100mg, 0.1mmol) and

Xantphos (63.6mg, 0.11mmol, see [161265-03-8] in ‘Catalysts—Part 2’) in $^*\text{C}_6\text{H}_6$ (10ml) at 30° for 4 hours, evaporating the solvent *in vacuo*, washing the residue with MeOH (1ml) and drying *in vacuo* to give the **analytically pure** complex with the formula $\text{C}_{58}\text{H}_{48}\text{O}_2\text{P}_3\text{Rh}$. The complex has IR (CHCl_3) with ν_{max} at 1996.9vs, 1909.6m cm^{-1} ; the ^1H NMR (300MHz, C_6D_6) has δ at 7.82 (apparent q, 4H, $J = 4.8\text{Hz}$, ar), 7.66 (m, 6H, ar), 7.53 (apparent q, 4H, $J = 4.9\text{Hz}$, ar), 7.11 (dd, 2H, $J = 7.3, 1.3\text{Hz}$, CHCHCC), 7.0-6.9 (ar), 6.79 (‘d’, 4H), 1.48 (s, 3H, CCH_3), 1.38 (s, 3H, CCH_3), -9.14 ($J_{\text{H-P}} = 12.2\text{Hz}$, $J_{\text{H-P}'} = 18.2\text{Hz}$, $J_{\text{H-Rh}} = 1.7\text{Hz}$); and the $^{31}\text{P}\{^1\text{H}\}$ NMR (121.5MHz, C_6D_6 , referenced to external 85% H_3PO_4) has δ at 42.67 ($J_{\text{P-Rh}} = 151.1\text{Hz}$, $J_{\text{P-P}} = 119.1\text{Hz}$, PPh_3), 25.65 ($J_{\text{P-Rh}} = 127.9\text{Hz}$, $J_{\text{P-P}} = 119.1\text{Hz}$, Xantphos-P); and MS has m/z at 961 (M-CO), 726 (M- PPh_3 -2H), 698 (M- PPh_3 -CO-2H); HR-MS found M is 578.1916 (calc for $\text{C}_{58}\text{H}_{48}\text{O}_2\text{P}_3$ is 578.1928), and for $^{13}\text{C}\{^1\text{H}\}$ NMR see references. [Kranenburg et al. *Organometallics* **14** 3081 1995, DOI: 10.1021/om00006a057.]

Xantphos-Ru complex formed *in situ* from equimolar amounts of Xantphos and $\text{Ru}(\text{PPh}_3)(\text{CO})\text{H}_2$ in refluxing toluene containing piperidinium acetate is a good catalyst for alkylating active methylene compounds, e.g. *t*-butyl cyanoacetate, and hydroxy compounds, i.e. PhCH_2OH , to provide α -substituted cyanoacetates, or *t*-butyl 2-cyano-3-phenylpropionate [Slatford et al. *Tetrahedron Lett* **47** 6787 2006, DOI: 10.1016/j.tetlet.2006.07.069].

Xantphos-Pd complexes formed *in situ* from xantphos and $\text{Pd}(\text{OAc})_2$ or $\text{Pd}_2(\text{dba})_3$ catalyse the cross-coupling reactions between the amide nitrogen and aryl and heteroaryl halides [Manley & Bilodeau *Org Lett* **6** 2433 2004, DOI: 10.1021/ol049165t] or *meso*-brominated porphyrins [Gao, Chen and Zhang *Org Lett* **6** 1837 2004, DOI: 10.1021/ol049440b] to form the corresponding aryl-N-CO- and *meso*-porphyrin-NCO- in high yields respectively in the presence of a base e.g. CsCO_3 . Similar cross-coupling catalysis with *in situ* Xantphos and $\text{Pd}(\text{OAc})_2$ between thenylbromides and 2-aminopyridines or aminoquinolines have been achieved in high yields [Begouin et al. *Synthesis* 2794 2006, DOI: 10.1055/s-2006-942510]. It should be **noted** that a detailed study of the mixing of Xantphos and $\text{Pd}_2(\text{dba})_3$ identified the complexes (**Xantphos**) $\text{Pd}(\text{dba})$ and $\text{Pd}(\text{Xantphos})_2$. The former is a very active catalyst whereas the latter bis(xanthane based phosphine) $_2\text{Pd}(0)$ is less soluble and inherently less active. It is therefore **important**, when forming the pre-catalyst, to keep the ratio of xantphos to Pd source less than 1.5:1 for a more effective catalyst involving cross-coupling between a nitrogen atom and an organic halide. [Klingensmith et al. *Organometallics* **25** 82 2006, DOI: 10.1021/om050715g.]

(**Xantphos**) $_2$ **Pd** is prepared in flame dried Schlenk equipment, evacuated/backfilled with argon three times, by stirring xantphos (579mg, 1.00mmol) and $\text{Pd}_2(\text{dba})_3$ (229mg, 0.25mmol) in toluene (300ml) under argon for 4 hours. This is filtered *via* a cannula into a separate dry flask under argon, concentrated somewhat overnight and any palladium black which may have settled is filtered off and the filtrate is evaporated to dryness. The yellow residue is stirred overnight with toluene (100ml) to remove excess (or unused) dibenzylidene acetone and unreacted Xantphos. The remaining yellow $\text{Pd}(\text{Xantphos})_2$ has **m 164° (dec.)**, is sparingly soluble in most common organic solvents, and is characterised by the correct elemental analyses for $\text{C}_{78}\text{H}_{64}\text{O}_2\text{P}_4\text{Pd}$. Its IR (KBr) has ν_{max} at 2924, 2854, 1461, 1398, 1377, 1222 cm^{-1} ; and MALDI-MS: calcd for $\text{C}_{78}\text{H}_{64}\text{O}_2\text{P}_4\text{Pd}$: *theoretical*: 1260.2894 (22.9%), 1262.2909 (63.4%), 1262.2911 (100.0%), 1263.2907 64.6%), 1264.2914 (77.2%), 1265.2933 (47.5%), 1266.2931 (35.5%), 1267.2949 (25.1%); *Found*: 1260.3405 (24.0%), 1261.3285 (67.4%), 1262.3166 (100.0%), 1263.3162 (73.2%), 1264.3300 (79.3%), 1265.3424 (47.5%), 1266.3491 (35.5%), 1267.3104 (25.1%). [Klingensmith et al. *Organometallics* **25** 82 2006, DOI: 10.1021/om050715g.]

Zirconocene chloride hydride (bis[cyclopentadienyl]zirconium(IV) hydride chloride, Cp_2ZrClH (Schwartz’ reagent) [37342-97-5] $\text{C}_{10}\text{H}_{11}\text{ClZr}$, M 257.9. It is moisture and light sensitive. Determine its purity by reaction with a slight excess of Me_2CO whereby the active H reacts to produce $\text{Cp}_2\text{ZrClOPr}^i$ and the integrals of the residual Me_2CO in the ^1H NMR will show its purity. The presence of Cp_2ZrH_2 can be determined because it forms $\text{Cp}_2\text{Zr}(\text{OPr}^i)_2$. For a very active compound, it is best to prepare it freshly from the **dichloride** {see below by reduction with Vitride [$\text{LiAl}(\text{OCH}_2\text{CH}_2\text{OH})_2\text{H}_2$]}, the white precipitate is filtered off, washed with tetrahydrofuran, then Et_2O and dried in a vacuum. Store it dry in the dark. [Carr & Schwartz *J Am Chem Soc* **101** 3521 1979, DOI: 10.1021/ja00507a017; Negishi & Takahashi *Aldrichimica Acta* **18** 31 1985, Buchwald et al. *Tetrahedron Lett* **28** 3895 1987, DOI: 10.1016/S0040-4039(00)96413-X; Negishi & Takahashi *Synthesis* 1 1988, DOI: 10.1055/s-1988-27453; for applications see Fieser **6** 175, **7** 101, **8** 84, **9** 104, **14** 81, **15** 80, **16** 72, **17** 70.] It has been used for functionalising olefins and alkynes [Sun et al. *Org Synth* **71** 83 1992, DOI: 10.15227/orgsyn.071.0083; Negishi & Takahashi *Aldrichimica Acta* **18** 31 1985, Ganem & Franke *J Org Chem* **72** 3981 2007, DOI: 10.1021/jo070129s]. It was used also for mild and selective **hydrozirconation** of amides to aldehydes [Spletstoser et al. *J Am Chem Soc* **129** 3408 2007, DOI: 10.1021/ja066362+].

Zirconocene dichloride (bis[cyclopentadienyl]zirconium dichloride, Cp_2ZrCl_2) [1291-32-3] $\text{C}_{10}\text{H}_{10}\text{Cl}_2\text{Zr}$, **M 292.3, m 242-245°, 248°**. Recrystallise the dichloride from CHCl_3 or xylene and dry it in a vacuum. ^1H NMR (CDCl_3) has δ at 6.52 from Me_4Si . Store it dry in the dark under N_2 . [Reid et al. *Aust J Chem* **18** 173 1965, DOI: 10.1071/CH9650173; *Beilstein* **16** IV 1770; for applications see Fieser **10** 131, **14** 122, **15** 120, **16** 121, **17** 106.] Together with Zn and CH_2Cl_2 it is used for methylenation of carbonyl compounds [Tour et al. *Tetrahedron Lett* **30** 3927 1989], and has been useful for the synthesis of a wide range of early transition-metal complexes and organometallic compounds [Negishi & Takahashi *Aldrichimica Acta* **18** 31 1985].

NANO METAL CATALYSTS: see ‘Nanomaterials and Nanotechnology’ in Chapter 7.

ORGANOCATALYSIS

An important area of chemical research is in the development of efficient catalysts for carrying out organic transformations. Traditionally this field is dominated by metals, although increasingly *organocatalysis* and biocatalysis are gaining prominence. **Organocatalysts** are low molecular weight (<1000g/mol) metal-free organic molecules, or metal-containing compounds where the metal is *not* involved in the catalytic process. They efficiently and selectively catalyse a large variety of chemical reactions when used in small non-stoichiometric amounts. The potential advantages that organocatalysts can offer over metal catalysts include the ease of handling, good stability, low costs and their environmentally friendly nature. Their major disadvantage is the limited particular substrate and transformation scope. It should be noted that the field of organocatalysis is in its relative infancy but stunning progress has been made in the last ten years or so. For example, readily available organic compounds have been found to be capable of catalyzing a range of reactions. These include carbon-carbon bond forming reactions (Aldol, Mannich, Diels-Alder, alkylations, cyclopropanations, etc), epoxidation reactions, desymmetrisation reactions and so on [for excellent books, see Berkessel and Gröger ‘Asymmetric Organocatalysis: From Biomimetic Concepts to Applications in Asymmetric Synthesis’, Wiley-VCH 2005, ISBN-10:3-527-30517-3, 13:978-3-527-30517-9, and Dalko (Ed.) ‘Enantioselective Organocatalysis: Reactions and Experimental Procedures,’ Wiley-VCH, 2007. ISBN-10: 3527315225, ISBN-13: 978-3527315222.]. Classes of compounds that have demonstrated applications as organocatalysts include α -amino acids [Jarvo & Miller *Tetrahedron* **58** 2481 2002, DOI: 10.1016/S0040-4020(02)00122-9], N-heterocyclic carbenes [Marion et al. *Angew Chem Int Ed* **46** 2988 2007, DOI: 10.1002/anie.200603380], thioureas [Connon *Chemistry-A Eur J* **12** 5418 2006, DOI: 10.1002/chem.200501076], cinchona alkaloids [see below, Dalaigh *Synlett* 875 2005, DOI: 10.1055/s-2005-864796] and so on. One of the earliest practical demonstrations of the use of organocatalysts is exemplified by the Hajos-Parrish-Eder-Sauer-Wiechert reaction which was patented as early as 1971 [Eder, Sauer & Wiechert *Angew Chem Int Ed* **10** 496 1971, DOI: 10.1002/anie.197104961; Hajos & Parrish *J Org Chem* **39** 1612 1974, DOI: 10.1021/jo00925a002]. The Robinson aldol annulation reaction utilising a proline molecule as the organocatalyst is another, and is an early example of a highly enantioselective carbon-carbon bond forming reaction. Recent advances in organocatalysis have progressed to the level of sophistication that has enabled the synthesis of relatively complex organic molecules with multiple stereocenters in high chemical and optical yields [see Bertelsen & Jørgensen ‘Organocatalysis-after the gold rush’ *Chem Soc Rev* **38** 2178 2009, DOI: 10.1039/B903816G; Ting et al. ‘Bronsted base catalysts’ *Topics in Current Chemistry* (Ojima Ed) **291** 145 2010, Sereda et al. ‘Lewis acid organocatalysts’ *Topics in Current Chemistry* (Ojima Ed) **291** 349 2010; Kampen et al. ‘Chiral Bronsted acids for asymmetric organocatalysis’ *Topics in Current Chemistry* (Ojima Ed) **291** 395 2010; Lelais & MacMillan ‘Modern Strategies in Organic Catalysis: the Advent and Development of Iminium Activation’ *Aldrichimica Acta* **39** 79 2006; List ‘The Ying and Yang of Organic Asymmetric Aminocatalysis’ *JCS Chem Commun* 819 2006, DOI: 10.1039/B514296M; List, Yamamoto & Gong ‘Organocatalysis: a web Collection’ *JCS Chem Commun* **48** 10703 2012, DOI: 10.1039/C2CC90327J, Editorial; Zhong & X. Shi ‘When organocatalysts meets transition metal catalysis’ *Eur J Org Chem* 2999 2010, DOI: 10.1002/ejoc.201000004; also see ChemFiles, Vol 6, no 4 from Sigma-Aldrich for a compilation of some commercially available organocatalysts]. For the definition of Lewis and Brønsted/Lowry acids and Bases see at the end of ‘Catalysts—Part 2’ in this Chapter.

This section deals with some individual examples that have proved useful. Several are commercially available and the purification of some are described here. The preparations in some entries are also mentioned as these

assist in devising methods of purification. The purification of a number of organocatalysts is also described in other chapters. Specific examples include (–)-**quinine** [see 130-95-0] and (+)-**quinidine** [see 56-54-2] used to catalyse α -halogenation of carbonyl compounds, intramolecular Michael additions, cyclopropanation of enones, enolates etc, β -lactam synthesis from imines and ketenes, β -lactone synthesis from aldehydes and ketenes, Morita-Baylis-Hillman reactions, hydrophosphonylation of aldehydes, Diels-Alder reactions and desymmetrisation of *meso*-anhydrides; (–)-**cinchonidine** [see 405-71-2] and (+)-**cinchonine** [see 118-10-5] which catalyse additions to prochiral ketenes, desymmetrisation of *meso*-diols and of *meso*-epoxides; and their respective *N*-benzyl salts [see 69257-04-1, and 69221-14-3] are efficient chiral **phase transfer catalysts** (see below); **L-proline** [see 147-85-3] which catalyses intramolecular Michael addition, Mannich reactions, inter- and intra- molecular aldol reactions, addition to N=O (α -aminooxylation/hydroxylation of C=O compounds), addition to nitrones and to N=N double bonds (α -amination of C=O compounds); **L-phenylalanine** [see 63-91-2] which catalyses intramolecular aldol reactions; and the **metal-free Jacobsen's ligand** *N,N'*-bis(3,5-di-*tert*-butylsalicylidene)-1,2-*trans*-cyclohexanediamine [see *R,R'*-135616-40-9 or *S,S'*-135616-36-3] which catalyses hydrocyanation (with TMS-CN) of aldehydes. It should be noted that the **Phase Transfer Catalysts** described in a later section of this Chapter are **organocatalysts**.

RS-(±)-1,1'-Bi(2-naphthol) (± 1,1'-binaphthyl-2,2'-diol) [602-09-5] $C_{20}H_{14}O_2$, M 286.3, m 214-217°, 216-218°, 217-218°, 218°, $pK_{Est(1)}^{20} \sim 7.5$, $pK_{Est(2)}^{20} \sim 11.8$; **R-(+)-1,1'-bi(2-naphthol) (R-(+)-BINOL)** [18531-94-7] m 208-210°, 207.5-208.5°, $[\alpha]_D^{20} +34$ (c 1, THF), $[\alpha]_{546}^{25} +50.9$ (c 1, THF); **S-(-)-1,1'-bi(2-naphthol) (S-(-)-BINOL)** [18531-99-2] m 207-208°, 208-210°, $[\alpha]_D^{22} -34$ (c 1, THF), $[\alpha]_{546}^{25} +51.3$ (c 1, THF). 1,1'-Bi(2-naphthol) is easily prepared by slowly adding a solution of $FeCl_3 \cdot 6H_2O$ (28g, 104mmol, see [10025-77-1]) in H_2O (60ml) to a vigorously stirred and refluxing suspension of 2-naphthol (14.4g, 100mmol, see [135-19-3]) in H_2O (600ml) and boiled until the oily drops of 2-naphthol disappear and the **binaphthol** separates out in flakes. Boiling is continued for 10 minutes, and the solid is filtered off, washed with boiling H_2O and dried first in air then *in vacuo*, to give (±)-**BINOL** in almost quantitative yield. Recrystallisation from hot EtOH, toluene or $*C_6H_6$ (~150ml, using charcoal) gives colourless glistening plates m 218°. The (±)-**dimethyl ether** [75640-87-8, 75685-01-7] has m 190° (from AcOH). [A.I.Vogel *Practical Organic Chemistry* Longmans, Green & Co, London, p 639 1948, Pummerer et al. *Ber* 59 2159 1926, DOI: 10.1002/cber.19260590881; Walder *Ber* 15 2166 1882, DOI: 10.1002/cber.188201502162] The FT-IR (Nujol) has ν_{max} at 3486.6, 1617.9, 1508.8, 1322.7, 1252.4, 1177.2, 1126.4, 827.3 and 751.3 cm^{-1} (note below how it is different from its enantiomers). [Beilstein 6 I 519, 6 II 1026, 6 III 5877, 6 IV 7020; for applications see Fieser 12 190, 13 113, 15 135, 16 132, 17 116.]

R- and S- BINOL have been obtained by a variety of ways which attest to their usefulness. They have been prepared by resolution of the (±)-BNP-acid (see below), the (±)-BINOL-boron ester (*via* the quinine salt), the (±)-BINOL dibenzoate (*via* pancreas acetone powder), or by oxidative coupling of 2-naphthol in the presence of chiral amines [see Zimmer & Suhrbier *J Prakt Chem* 339 758 1997, DOI: 10.1002/prac.199733901138]. The preparations involving reduction of the (±)-BNP acid directly with $LiAlH_4$, or *via* the methyl phosphate derivative with RedAl [sodium bis(2-methoxyethoxy)aluminium hydride] in 3.4M toluene solution [Truesdale *Org Synth Coll Vol* 8 46 1993, DOI: 10.15227/orgsyn.067.0013] appear to be most convenient, and do **not** involve racemisation. The former is described here as it does not require an esterification step. Pure **R-(+)-BNP** acid (115.4g, 331mmol, below) in dry THF at 0° under N_2 , is treated with $LiAlH_4$ (31.4g, 830mmol) in small portions (exert extreme care here because of the large quantities used — fire hazard) during 1 hour with stirring. A further volume of dry THF (400ml) is added and the mixture is stirred at 25° under N_2 , for 17 hours, cooled to 0° under N_2 , and cold 6N aqueous HCl (250ml) is added very carefully. The upper phase is then decanted off, and the lower phase is mixed with 6N aqueous HCl (150ml) and THF (150ml), stirred, and is allowed to settle. The phases are separated and the lower phase is extracted with Et_2O . The combined organic phases are washed with brine, decolorised with charcoal and evaporated *in vacuo*. The residual oil is dissolved in $*C_6H_6$ (1L, toluene may be used) and evaporated until crystallisation sets in. The solid is collected (90.6g, 96%, m 202-207°), which upon crystallisation from $*C_6H_6$ or toluene gives optically pure BINOL (84.5g, 89%, m 207-208°) with (+) or (–) $[\alpha]_{589}^{25}$ 34.3, $[\alpha]_{578}^{25}$ 37.8, $[\alpha]_{546}^{25}$ 51.3, $[\alpha]_{436}^{25}$ 228 (c 1.1, THF). The FT-IR (Nujol) of the BINOL enantiomers are identical, though different from that of the racemate (see above), and have ν_{max} at 3509.9, 1617.9, 1511.5, 1319.1, 1220.9, 1148.9, 815.6, 749.6 and 566.0 cm^{-1} ; and the 1H NMR (1:1 $CDCl_3/DMSO-d_6$, TMS) has δ at 7.04 (d, 2H, $J = 8.8Hz$), 7.20-7.35 (m, 4H), 7.40 (d, 2H, $J = 8.8Hz$), 7.92 (d, 2H, $J = 8.8Hz$) and 9.21 (s, 2H). **Racemisation studies** showed that: BINOL (0.1g) is optically stable in an H_2O (10ml)/dioxane (12ml) mixture at 100° under N_2 during 26 hours; loses 44% and 66% of optical activity in

BuOH (10ml) containing KOH (0.71mmol) per BINOL (0.33mmol) at 118° under N₂ in 13 and 23 hours respectively; and loses 37% and 72% of optical activity of BINOL (100mg) in a 20% aqueous HCl (10ml)/dioxane (12ml) mixture at 100° under N₂ during 7 and 26 hours respectively. [Kyba et al. *J Org Chem* **42** 4173 1977, DOI: 10.1021/jo00862a001.] BINOL and 3,3'-diaryl derivatives act as organocatalysts for **Mirata-Baylis-Hillman reactions** involving C-H activation, e.g. condensation of a variety of aldehyde with cyclohex-2-en-1-one to provide chiral 2-(1-substituted-1-hydroxymethyl)cyclohex-2-en-1-ones in 40 to 80% yields and 'ee' values of ~90 in the presence of Et₃P [McDougal & Schaus *J Am Chem Soc* **125** 12094 2003, DOI: 10.1021/ja037705w], they catalyse hydrocyanation (with TMS-CN) of aldehydes with high 'ee' at -78° [Holmes & Kagan *Tetrahedron Lett* **41** 7453 DOI: 10.1016/S0040-4039(00)01275-2; and 7457 2000, DOI: 10.1016/S0040-4039(00)01276-4], and the **titanium complexes** catalyse the stereoselective reduction [with (MeO)₃SiH] of carbonyl compounds [Schiffers & Kaga *Synlett* 1175 1997, DOI: 10.1055/s-1997-982], as well as C-C forming reactions [Zimmer & Suhrbier *J Prakt Chem* **339** 758 1997, DOI: 10.1002/prac.199733901138] and Diels-Alder reactions [Mikami et al. *Tetrahedron: Asymmetry* **2** 643 1991, DOI: 10.1016/S0957-4166(00)86118-2]. The rare earth **La-S-BINOL complex** realises an asymmetric nitroaldol reaction with nitromethane in high product yield and enantiomeric enrichment [Sasai et al. *J Am Chem Soc* **115** 10372 1993, DOI: 10.1021/ja00075a068].

RS-(±)-1,1'-Binaphthyl-2,2'-diylhydrogen phosphate (1,1'-binaphthyl-2,2'-phosphoric acid, BNP) [36193-63-6] C₂₀H₁₃O₄P, M 348.3, m 217°, pK²⁰ 0.74; **R-(-)-1,1'-binaphthyl-2,2'-diylhydrogen phosphate** [39648-67-4] m 217°, [α]_D²⁰ -608 (c 1, MeOH), [α]₅₄₆²² -562.7 (c 0.97, CHCl₃), [α]₅₄₆²² -113 (c 0.95, CHCl₃); **S-(+)-1,1'-binaphthyl-2,2'-diylhydrogen phosphate** [35193-64-7] m 217°, [α]_D²⁰ +608 (c 1, MeOH). Recrystallise it from EtOH. Reflux for 3 hours in N NaOH is required to hydrolyse the cyclic phosphate. The **R-enantiomer** has been prepared by a general procedure in which **R-(+)-1,1'-binaphthyl-2,2'-ol** (1.90mmol, see *R-BINOL* [18531-94-7]) in pyridine (7.8ml) is treated with POCl₃ (250μl, 2.68mmol) and stirred at ~25° for 3 hours. The mixture is then quenched with H₂O (154μl) at 0°, stirred for 1 hour at ~25°, evaporated *in vacuo*, the residue is acidified with 6N HCl (20ml) at 0° then refluxed for 2 hours. After cooling to 0°, the **phosphoric acid** separates, is filtered off, washed with H₂O and dried *in vacuo*. Recrystallisation by dissolving in CH₂Cl₂ and pouring it into hexane provides pure crystals of the **R-(-)-acid** (1.58mmol, 83%). [Yamanaka et al. *J Am Chem Soc* **129** 6756 2007, DOI: 10.1021/ja0684803; for general procedure see Jacques & Fouquey *Org Synth* **67** 1 1989, DOI: 10.1522/orgsyn.067.0001.] Its ¹H NMR (400MHz, CDCl₃, TMS) has δ at 8.01 (d, 2H, J = 8.8 Hz), 7.94 (d, 2H, J = 8.2 Hz), 7.55 (d, 2H, J = 8.8 Hz), 7.49-7.44 (m, 2H), 7.38 (d, 2H, J = 8.2 Hz), 7.32-7.27 (m, 2H) and 3.22 (s, 1H); and for IR and ¹³C NMR see references. [Jacques et al. *Tetrahedron Lett* **48** 4617 1971, DOI: 10.1016/S0040-4039(01)97544-6; Arnold et al. *Tetrahedron* **24**, 343 1983, DOI: 10.1016/S0040-4039(00)81402-1; Beilstein **6** II 1027.] **Alternatively**, (and more useful as it can be carried out on a 100g scale), the **R-(-) and S-(+) BNP acids** are obtained by optical resolution of the diastereoisomeric **(+)-cinchonine salts** of the **(±)-BNP acid** via recrystallisation from hot MeOH. The crude **(+)-acid (+)-base salt** [which contains 91% of (+)-acid (+)-base and 9% of (-)-acid (+)-base] has [α]₅₄₆²⁵ +424 (c 0.99, MeOH) and the more soluble crude **(-)-acid (+)-base salt** obtained by evaporation of the mother liquors [which contains 81% of (-)-acid (+)-base and 19% of (+)-acid (+)-base] has [α]₅₄₆²⁵ -113 (c 0.95, MeOH). Pure **(+)-acid (+)-base salt** [α]₅₄₆²⁵ +492 (c 1, MeOH) and pure **(-)-acid (+)-base salt**, [α]₅₄₆²⁵ -256 (c 1, MeOH) as obtained from the pure enantiomeric BNP acids and (+)-cinchonine are recrystallised from MeOH/EtOAc and MeOH/Me₂CO/EtOAc respectively. The BNP acids are obtained from the respective salts (e.g. 77g) in boiling EtOH (e.g. 500ml) and adding 6N aqueous HCl (e.g. 570ml) with vigorous stirring during 30 minutes while keeping the temperature between 75° and 80°. After cooling the BNP acid is collected, washed with H₂O (e.g. 5 x 90ml), and air dried to give (~50 yield of ~99% 'ee') of almost pure acid (by NMR and HPLC). BNP acids from the filtrates can be purified by recrystallisation from hot MeOH or EtOH and are sparingly soluble in H₂O. The solubilities at 25° in g/100ml are: 5.7±0.2 for enantiomers (EtOH) and 10.3±0.5 for racemate (EtOH); and 2.1±0.1 for enantiomers (MeOH) and 2.5±0.1 for racemate (MeOH). Two recrystallisations from EtOH of the BNP enantiomeric acids obtained did not alter the rotations at 25° which are (+) or (-): 595° ±7 (589nm), 624° ±7 (578nm), 720° ±8 (546nm), 1328 ±15 (436nm), 2050° ±25 (365nm) in MeOH; 574° ±16 (589nm), 602° ±17 (578nm), 694° ±20 (546nm), 1267 ±25 (436nm), 1828° ±40 (365nm) in EtOH. [Jacques & Fouquey *Org Synth Coll Vol* **8** 50 1993, DOI: 10.1522/orgsyn.067.0001; see also Kyba et al. *J Org Chem* **42** 4173 1977, DOI: 10.1021/jo00862a001].

The BNP acids, and their 3,3'-aryl derivatives, are useful asymmetric organocatalysts in being efficient *Bronsted acid catalysts* for enantioselective Mannich reactions. [Yamanaka et al. *J Am Chem Soc* **129** 6756 2007, DOI: 10.1021/ja0684803.] [For applications see Fieser **12** 49, **16** 25.]

2,2'-Bis(diphenylphosphino)-1,1'-binaphthyl (BINAP) [*RS* see 98327-87-8] [(C₆H₅)₂PC₁₀H₆—]₂, **M 622.7, m 283-286°**, [*R*-(+)-see 76189-55-4] **m 241-242°**, [*S*-(-)-see 76189-56-5] **m 241-242°**. In addition to being ligands for Pd, Rh and Ru catalysts, *S*-BINAP also acts as an organocatalyst for Mirata-Baylis-Hillman reactions involving C-H activation, e.g. in the condensation of 5-formylpyrimidine and derivatives with methyl acrylate to form methyl β-hydroxy-β-(pyrimidin-5-yl)-α-methylenepropionates with moderate enantiomeric excess at the chiral hydroxy centre [Hayase et al. *Chem Commun* 1271 1998, DOI: 10.1039/A802594K].

Bode-Rovis-Enders NHC precatalysts

These are a group of 3,4-fused-2-aryl substituted 1,2,4-triazolium salt pre-catalysts, which when converted into the corresponding NHCs (N-heterocyclic carbenes) where C-5 of the heterocyclic ring becomes a carbene carbon atom, are *organocatalysts* that promote a wide variety of organic reactions. The carbene is usually generated *in situ* on addition of diisopropylamine or BTU. Among the reactions that these NHCs catalyse are cyclocondensation reactions, enantioselective azadiene Diels-Alder reactions, aza-Claisen reactions, annulation reactions, formation of trisubstituted indanes *via* catalysed cascade annelations, desymmetrisation of cyclohexadienones, benzoin condensations, Stetter reaction, a³ to d³ Umpolung, transesterification, 1,2-additions, etc. [Bode *Speciality Chemicals Magazine* **27**(6) 28 pages 2007, Moor & Rovis *Top Curr Chem* **291** 77-144 2010, DOI: 10.1007/978-3-642-02815-1_18, PMID: 21494949; Enders, Niemeier and Henseler *Chem Rev* **107** 5606-5655 2007, DOI: 10.1021/cr068372z.] Described below are such four chiral organoprecatalysts whose carbenes catalyse highly stereospecific reactions in good to very good yields and with very high enantiomeric excess(ee), and two related non-chiral salts which also catalyse non-chiral reactions in high yields.

(4aR,9aS)-4,4a,9,9a-Tetrahydro-1-oxa-4-azafluoren-3-one, **(4aR,9aS)-indeno[2,1-b]-1,4-oxazin-3(2H)-one** [862095-79-2] C₁₁H₁₁NO₂, **M 189.2, m 180-183°(dec)**, [α]_D²³ **-17.3 (c 1.08, MeOH)**. This is the key chiral intermediate *fluorene-3-one* that uses (+)-*cis*-(1*R*,2*S*)-1-aminoindan-2-ol (see [136030-00-7]) for the preparation of chiral *indeno-triazolo-oxazinium* pre-catalysts. Details of its synthesis are not given here because they have been reported in great detail by Rovis and coworkers [Vora et al. *Org Synth* **87** 350 2010; DOI: 10.15227/orgsyn.087.0350], but the physical and spectroscopic properties, and final purification procedures are provided in order to assess the purity of the commercial product and to purify it further if necessary. Thus the aminoindanol is converted to the -ONa salt with NaH/THF that is reacted with ClCH₂CO₂Et to form the *fluorenone*. The final purification is carried out by stirring vigorously the crude light brown solid with hexanes under a reflux condenser in an oil bath at 70° for 2 hours, cooling to ~25°, filtering the solid off and drying it at 70° *in vacuo* (2mm Hg) for 1 hour to give the *fluorenone* as a white solid in 80-86% yield of almost analytical (C, H and N) purity. The ¹H NMR (400MHz, Me₂CO-*d*₆) with δ at 2.97 (d, *J* = 16.8Hz, 1H), 3.23 (dd, *J* = 16.6, 4.9Hz, 1H), 3.89 (d, *J* = 16.2Hz, 1H), 4.05 (d, *J* = 16.2Hz, 1H), 4.57 (t, *J* = 4.6Hz, 1H), 4.82 (t, *J* = 4.0Hz, 1H), 7.23-7.29 (m, 3H), 7.46-7.49 (m, 1H), 8.21 (br s, 1H); and for IR, ¹³C NMR and HRMS (APCI⁺) see references. The enantiomeric **(4aS,9aR)-4,4a,9,9a-1-oxa-4-azafluorenone** can be similarly prepared by starting from (-)-*cis*-(1*S*,2*R*)-1-aminoindan-2-ol (see [126456-43-7]) and differs only in the sign of the specific rotation.

(5aR,10bS)-5a,10b-Dihydro-2-(2,4,6-trimethylphenyl)-4H,6H-indeno[2,1-b]-1,2,4-triazolo[4,3-d]-1,4-oxazinium chloride monohydrate [903571-02-8 for anhydrous] C₂₁H₂₂ClN₃O. H₂O, **M 385.9, m 217-219°, 226-230° (anhydrous ?)**, [α]_D²⁰ **-133.5° (c 1.00, EtOH)**, [α]_D²⁰ **-156.0 (c 1.00, CHCl₃)**. Here also details of its synthesis are not given because they have been reported in great detail by Bode and coworkers [see Struble & Bode *Org Synth* **87** 362 2010, DOI: 10.15227/orgsyn.087.0362], but the physical and spectroscopic properties, and final purification procedures are provided in order to assess the purity of the commercial products and to purify further if necessary. The preceding **(4aR,9aS)-fluorenone** is treated with Me₃OBf₄/CH₂Cl₂ (~25°/16 hours) then NaHCO₃, and dried in a high vacuum (12 hours) to give a 92-93% yield (95% purity) of **(4aR,9aS)-3-methoxy-2,4a,9,9a-tetrahydroindeno[2,1-b][1,4]oxazine** as a pale brown solid with **m 73-75°, [α]_D²⁰ -15.7 (c 1.26, EtOH)**, and consistent IR, NMR and MS. This *methoxy-oxazine* is then added to a solution containing one equivalent of **2,4,6-trimethylphenylhydrazinium chloride*** in MeOH and a catalytic amount of 4M HCl/dioxane (obligatory) and heated at 60° under a reflux condenser and N₂ atmosphere for 48 hours. The solvent is removed *in vacuo* and the crude orange solid residue is suspended in EtOAc (must be absolutely dry for high yields; dry by passing through an Al₂O₃ drying column in an argon atmosphere) with vigorous stirring

at 90° (under N₂) for 30 minutes, cooled to 0°, and the light yellow solid is filtered off, washed with EtOAc and dried at ~25° (<1 mbar, 12 hours) to provide an 81-82% yield of **(Z)-2-mesityl-1-[(4aR,9aS)-4,4a,9,9a-tetrahydroindeno-[2,1b][1,4]-oxazin-3-(2H)ylidene)hydrazinium chloride** as a light yellow powder which has **m** 200-201°(dec), $[\alpha]_D^{20} +66.7$ (c 1.42, EtOH), and consistent IR, NMR and MS. The precatalyst is then obtained by cyclising the hydrazinium chloride with triethylorthoformate (8 equivalents) in chlorobenzene (dried over 4Å molecular sieves) and anhydrous HCl (4M in dioxane, 1 equivalent) by stirring under N₂ at 120° for 1 hour, and the clear brown solution is evaporated *in vacuo*. The brown residue is treated with toluene, heated at 120° for 5 minutes, cooled in ice/water, the white solid is filtered off and the filtrate is evaporated at 60° *in vacuo* and dried at high vacuum (<0.75mm). The brown residue dissolves in toluene at 60° within 3-5 minutes then a white precipitate is formed rapidly. The cooled solution is filtered, the white solid is washed with a little toluene and dried at high vacuum (<1 mbar) to constant weight (>24 hours) to give a 60-64% yield of the desired **(5aR,10bS)-oxazinium chloride** pre-catalyst as an analytically pure (C, H and N) white powder with **m** 217-219°. The ¹H NMR (400MHz, DMSO-*d*₆) has δ at 212 (s, 6H), 2.37 (s, 3H), 3.16 (d, *J* = 17.0Hz, 1H), 3.50 (dd, *J* = 16.9, 4.8Hz, 1H), 4.99 (t, *J* = 4.4Hz, 1H), 5.08 (d, *J* = 16.0Hz, 1H), 5.26 (d, *J* = 16.0Hz, 1H), 6.12 (d, *J* = 4.0Hz, 1H), 7.21 (s, 2H), 7.33-7.45 (m, 3H), 7.65 (d, *J* = 7.2Hz, 1H), 11.34 (s, 1H); and for IR, ¹³C NMR and HRMS (ESI) see references. The enantiomeric **(5aS,10bR)-5a,10b-Dihydro-2-(2,4,6-trimethylphenyl)-4H,6H-indeno[2,1-b]-1,2,4-tri-azolo[4,3-d]-1,4-oxazinium chloride monohydrate** C₂₁H₂₂ClN₃O · H₂O, **M** 385.9, **m** 212-216°, $[\alpha]_D^{20} +158.0$ (c 1.00, CHCl₃) can be prepared by starting from the enantiomeric **(4aS,9aR)-4,4a,9,9a-tetrahydro-1-oxa-4-azafluoren-3-one** and is purified in the same way.

*The purity of **2,4,6-mesitylhydrazinium chloride** [76195-82-9] **M** 186.7 is critical for obtaining pure mesityl-pre-catalysts. The preparation from 2,4,6-trimethylaniline is described in great detail by Struble and Bode [*Org Synth* **87** 362 2010, DOI: 10.15227/orgsyn.087.0362] who emphasise that it is imperative for diazotisation and reduction to be strictly monitored. It is paramount that NaNO₂ and SnCl₂ solutions be added at such a rate that the internal temperature is carefully monitored at <5°. The final purification is by suspending the salt (~7.5g) in absolute EtOH/Et₂O (1:1, 15ml), and sonicating (85W) for 5 minutes while maintaining the temperature below 30°. The pale orange solid is filtered off using a fine porosity filter, washed with EtOH/Et₂O (1:1, 3 x 5ml), and dried under high vacuum (~25°, <0.75mm) for 12 hours to give analytically pure (C, H and N) **2,4,6-mesitylhydrazinium chloride** (36-40% yield). *Note* that the material may be very fine to filter in a short period, in which case it should be dissolved in dry MeOH and concentrated by evaporation to give a more crystalline product that can be filtered more easily. *Alternatively*, it should be collected by centrifugation, decantation of the supernatant, washed with EtOH/Et₂O (1:1, 3 x 5ml) also by decantation and sonication in the same way in between washes. It has **m** 195-197°(dec) (167-173° also reported); the ¹H NMR (400MHz, DMSO-*d*₆) has δ at 2.20 (s, 3H), 2.35 (s, 6H), 6.60 (br s, 1H), 6.86 (s, 2H), 9.27 (variable br s, 3H, NH, NH₂); and for IR, ¹³C NMR and HRMS (ESI) see references. [Struble and Bode *Org Synth* **87** 362 2010, DOI: 10.15227/orgsyn.087.0362].

(5aR,10bS)-5a,10b-Dihydro-2-(pentafluorophenyl)-4H,6H-indeno[2,1-b]-1,2,4-triazolo[4,3-d]-1,4-oxazinium tetrafluoroborate [740816-14-2] C₁₈H₁₁BF₉N₃O, **M** 467.1, **m** 223-226°, 235°(dec), $[\alpha]_D^{23} -130.8$ (c 1.28, MeCN). Details of its synthesis are not given here because they have been reported in great detail by Rovis and coworkers [Vora et al. *Org Synth* **87** 350 2010, DOI: 10.15227/orgsyn.087.0350], but the physical and spectroscopic properties, and final purification procedures are provided in order to assess the purity of the commercial product and to purify it further if necessary. The common intermediate **(4aR,9aS)-4,4a,9,9a-1-oxa-4-azafluorenone** (1.0 equivalent see [862095-79-2] above) is added with stirring, and under anhydrous conditions in an argon atmosphere, to a mixture of Me₃OBf₄ (1.0 equivalent *and no more*, care **Toxic**) in CH₂Cl₂ (containing <20ppm of H₂O, by filtering through activated Al₂O₃ under argon) and stirred at ~25° until the homogeneous reaction is complete (1-2 hours by ¹H NMR). Pentafluorophenylhydrazine (1.0 equivalent) is then added to the mixture under argon, stirred until the reaction is complete (~4 hours, when ¹H NMR signals of activated amidate disappear), the mixture is evaporated *in vacuo*, dried at 100°/2mm; then chlorobenzene (as solvent) followed by triethyl orthoformate (2.5 equivalents) are added and heated under reflux to 130° with stirring (open to atmosphere) for 24 hours. A second portion of (EtO)₂O (2.5 equivalents, syringe) is added and stirred for a further 24 hours, and this process is repeated once more. Excess of toluene is added, the slurry is filtered, and the solid washed with toluene and hexanes. The solid is triturated with EtOAc and MeOH (4:1), filtered through a medium frit funnel, washed with cold EtOAc to give the **oxazinium tetrafluoroborate** pre-catalyst which is dried (100°/2mm) to constant weight (~1 hour, 61-64% yield), gave an off white solid that analysed as the ~0.5 hydrate. The ¹H NMR (400MHz, Me₂CO-*d*₆) has δ at 3.28 (d, *J* = 17.2Hz, 1H), 3.55 (dd, *J* = 17.1, 4.9Hz, 1H), 5.19 (t, *J* = 4.5Hz, 1H), 5.25 (d, *J* = 16.4Hz, 1H), 5.39 (d, *J* = 16.4Hz, 1H), 6.33 (d, *J* = 16.4

Hz, 1H), 6.33 (d, $J = 4.0$ Hz, 1H), 7.34 (t, $J = 7.3$ Hz, 1H), 7.43 (q, $J = 7.4$ Hz, 2H), 7.63 (d, $J = 7.6$ Hz, 1H), 11.09 (br s, 1H); for IR, ^{13}C NMR and HRMS (APCI⁺) see references. [see Vora et al. above].

The enantiomeric (*5aS,10bR*)-*5a,10b-Dihydro-2-(pentafluorophenyl)-4H,6H-indeno[2,1-b]-1,2,4-triazolo-[4,3-d]-1,4-oxazinium tetrafluoroborate* can be prepared and purified in a similar way but starting with the enantiomeric (*4aS,9aR*)-*4,4a,9,9a-1-oxa-4-azafluorenone*

2-Pentafluorophenyl-6,7-dihydro-5H-pyrrolo[2,1-c][1,2,4]triazol-2-ium tetrafluoroborate (*2-pentafluoro-2,5,6,7-tetrahydro[2,1-c][1.2.4]-tetrazol-4-ium tetrafluoroborate*) [862095-91-8] $\text{C}_{11}\text{H}_7\text{BF}_9\text{N}_3$, **M 363.0**, **m 242-245°**, **245°**. Details of its synthesis are not given here because they have been reported in great detail by Rovis and coworkers [Vora et al. *Org Synth* **87** 350 2010, DOI: 10.15227/orgsyn.087.0350], but the physical and spectroscopic properties, and final purification procedures are provided in order to assess the purity of the commercial product and to purify it further if necessary. Starting with 2-pyrrolidinone (1.0 equivalent, [616-45-5]) which is converted into 2-methoxy-3,4-dihydro-5H-pyrrolidinium tetrafluoroborate by reaction with trimethyloxonium tetrafluoroborate (1.0 equivalent) in CH_2Cl_2 under argon with stirring at $\sim 25^\circ$ until homogeneous, then pentafluorophenylhydrazine (1.0 equivalent, [828-73-9]) is added with stirring for 4 hours (when amidate signals disappear in the ^1H NMR spectrum), the mixture is evaporated *in vacuo*, chlorobenzene and triethyl orthoformate (2.0 equivalents, syringe) are added and heated at 130° for 24 hours under a reflux condenser open to the atmosphere. Addition of triethyl orthoformate (2.0 equivalents, syringe) and heating is repeated for 24 hours, followed by evaporation to dryness *in vacuo*, excess of toluene is added to the residue and the slurry is filtered, and the solid is washed with toluene and hexanes, then with EtOAc. Final purification is by washing well with cold EtOAc, and the off-white solid is thoroughly dried at 100° *in vacuo* (2mm) to constant weight (~ 1 hour) to give the *pre-catalyst tetrafluoroborate* in 74-76% yield. The ^1H NMR (400MHz, $\text{Me}_2\text{CO}-d_6$) has δ at 3.00 (ddd, $J = 15.0, 7.7, 7.7$ Hz, 2H), 3.42 (t, $J = 8.0$ Hz, 2H), 4.76 (t, $J = 8.0$ Hz, 2H), 10.19 (s, 1H); and for IR, ^{13}C NMR and HRMS (APCI⁺) see references. [Vora et al. *Org Synth* **87** 350 2010, DOI: 10.15227/orgsyn.087.0350].

2-Mesityl-6,7-dihydro-5H-pyrrolo[2,1-c][1,2,4]triazol-2-ium chloride (*2-mesityl-2,5,6,7-tetrahydropyrrolo[2,1-c][1.2.4]-tetrazol-4-ium chloride*) [Bode Catalyst 3] $\text{C}_{14}\text{H}_{18}\text{N}_3$, **M 263.8**, **m 202-205°**. This is prepared and purified in much the same way as the preceding salt except for using trimethyloxonium chloride for preparing the ethoxy pyrrolidinium chloride followed by reaction with mesitylhydrazinium chloride instead of perfluorophenyl-hydrazine. This is a highly selective, *N-heterocyclic carbene (NHC) Bode catalyst* that promotes aza-Diels-Alder reactions with *exceptional* diastereo- and enantio- selectivities ($>99\%$ 'ee') [He et al. *J Am Chem Soc* **128** 8418 2006, DOI: 10.1021/ja062707c].

Boron trifluoride diethyl etherate [109-63-7] (C_2H_5)₂O. BF_3 , **M 141.9**, **m -58°** , **b $46^\circ/10\text{mm}$** , **126-129°/atm**, **d_4^{25} 1.15**, **n_D^{20} 1.344**. If the liquid has darkened due to oxidation by air it can be redistilled to give a colourless liquid. It is recommended that excess dry Et_2O (10ml) be added to the etherate (500ml) and distilled from CaH_2 (2g) in an all glass still at (b $46^\circ/10\text{mm}$) which removes volatile acids and avoids bumping [Zwiefel & Brown *Organic Reactions* **13** 28 1963, DOI: 10.1002/0471264180.or013.01]. It is a fuming liquid which hydrolyses in moist air forming highly toxic fluorides on decomposition. Store it under N_2 or argon and work in an efficient fumecupboard. It is a versatile *Lewis acid* for catalysis, acetylation, alkylation, dehydration, condensations, Beckmann rearrangements and polymerisation reactions. It catalyses the synthesis of cyclopentyl- and cycloheptyl[b]indoles from aryl cyclopropyl ketones *via* [3+2] cycloaddition [Venkatesh et al. *Eur J Org Chem* 5378 2006, DOI: 10.1002/ejoc.200600342], and is a mild and efficient catalyst for Beckmann rearrangement in anhydrous MeCN [An et al. *Chin J Chem* **29** 947 2011, DOI: 10.1002/cjoc.201190193]. [Beilstein **1** IV 1321; for applications see Fieser in vols **1** to **17** 52.] It is available as a $\text{BF}_3\cdot\text{EtNH}_2$ complex [75-23-0] **M 112.9**, **m $85-89^\circ$** [Beilstein **4** II 588]; and as a 50% w/w $\text{BF}_3\cdot\text{MeOH}$ solution [373-23-0] **M 99.9** in MeOH, is used for making germinal bis-peroxides from ketals and enol ethers with *t*-BuOOH [Terent'ev et al. *Synthesis* 2215 2005, DOI: 10.1055/s-2005-872093], and used for the mild deacetylation of amines [Miltsov et al. *Tetrahedron Lett* **44** 2301 2003, DOI: 10.1016/S0040-4039(03)00271-5.] [Beilstein **1** IV 1220; Fieser **1** 73, **9** 64.]

CBS (Corey-Bakshi-Shibata) Catalysts:

R-(+)-2-Methyl-CBS-oxazaborolidine (*R-1-methyl-3,3-diphenyltetrahydro-1H,3H-pyrrolo[1,2-c][1,3,2]-oxazaborole*, α,α -diphenyl-D-prolinolmethylboronic acid cyclamide) [112022-83-0] $\text{C}_{18}\text{H}_{20}\text{BNO}$, **M 277.2**, **m $74-87^\circ$** , **$77-82^\circ$** , **$85-95^\circ$** , [α]_D²⁰ **+76.8** (c 1, toluene) and *S-(-)-2-methyl-CBS-oxazaborolidine* [112022-81-8] $\text{C}_{18}\text{H}_{20}\text{BNO}$, **M 277.2**, **m $85-95^\circ$** . These CBS (Corey-Bakshi-Shibata) catalysts effect the asym-

metric reduction of *pro*-chiral ketones, the stereoselective synthesis of α -amino acids and α -hydroxy acids, the preparation of ferrocenyl diols and propargyl alcohols. The catalyst is usually best prepared *in situ* from the respective α,α -diphenyl-(*R* or *S*)-prolinol [22348-32-9 or 79815-20-6] and a slight excess of trimethylboroxine [823-96-1]. Their suitability are determined by capillary GC [DB-1. 200°, retention time 4.9 minutes] and ^1H NMR, where possible impurities have signals at δ (CDCl_3) for starting prolinol at $\sim 4.3(\text{t})$, trimethylboroxine at 0.45(s), the B—OH intermediate at 0.35 to -0.50 multiple B-Me singlet's, and/or the ring-B-O-B(Me)-OH intermediate at -0.25 B-Me brs. An **analytical sample** is obtained by dissolving it in toluene, filtering, then evaporating at 50°/0.001mm to give the catalyst as a white solid. Store it under dry N_2 or argon. The ^1H NMR (0.2M in CDCl_3) has δ at 7.65-7.15 (m, 10 H, ArH), 4.4 (dd, $J = 5.8, 10.0$ Hz, 1 H, C3a-H), 3.45-3.30 (m, 1 H, C6-H), 3.15-3.00 (m, 1 H, C6-H), 1.90-1.55 (m, 3 H, C4-H, C5-H₂), 0.95-0.75 (m, 1 H, C4-H), 0.40 (s, 3 H, B-CH₃); and for IR see references. They are also available commercially as 1M solutions in toluene or tetrahydrofuran. [Mathre et al. *J Org Chem* **56** 751 1991, DOI: 10.1021/jo00002a049; Corey et al. *J Am Chem Soc* **124** 3808 2002, DOI: 10.1021/ja025848x; Ryu & Corey *J Am Chem Soc* **126** 8106 2004, DOI: 10.1021/ja0475959; Ryu et al. *J Am Chem Soc* **124** 9992 2002, DOI: 10.1021/ja027468h; Hu et al. *J Am Chem Soc* **126** 13708 2004, DOI: 10.1021/ja046154m; Quallich & Woodall *Tetrahedron Lett* **34** 785 1993, DOI: 10.1016/0040-4039(93)89012-F.] They are excellent catalysts for asymmetric reductions [Corey & Helal *Angew Chem Int Ed* **37** 1986 1998, DOI: 10.1002/(SICI)1521-3773(19980817)37:15<1986::AID-ANIE1986>3.0.CO;2-Z; Baar et al. *J Am Chem Soc* **126** 8216 2004, DOI: 10.1021/ja040021j].

***R*-(+)- and *S*-(-)- 2-*o*-Tolyl-CBS-oxazaborolidine {*R*- and *S*- 3,3-diphenyl-1-*o*-tolyltetrahydro-1H,3H-pyrrolo[1,2-*c*] [1,3,2]-oxazaborole}** [*R*-865812-10-8; *S* 463941-07-3] $\text{C}_{24}\text{H}_{24}\text{BNO}$, **M 353.3**, $[\alpha]_{\text{D}}^{20}$ **+18 and -18 (c 1, toluene)**. These Corey-Bakshi-Shibata (CBS) oxazaborolidines are available commercially as 0.5M solutions in toluene (d_4^{25} 0.9009g/ml). They produce chiral **Lewis acids** when protonated with trifluoromethanesulfonamide and are used in enantioselective Diels-Alder reactions. They are prepared and purified as for the 2-methyl-CBS-oxazaborolidines above except that they are obtained as liquids on evaporation of the toluene solutions. [For applications see Ryu & Corey *J Am Chem Soc* **125** 6388 2003, DOI: 10.1021/ja035393r; Stemmler *Synlett* 997 2007, DOI: 10.1055/s-2007-973876; Corey et al. *J Am Chem Soc* **109** 7925 1987, DOI: 10.1021/ja00259a075; Cho & Kim *Tetrahedron: Asymmetry* **12** 2043 2001, DOI: 10.1016/S0957-4166(01)00359-7; Cho et al. *JCS Perkin Trans I* 1204 2001, DOI: 10.1039/B010155I; Lebsack et al. *J Am Chem Soc* **123** 4851 2001, DOI: 10.1021/ja015802o; and references therein.]

2-Methyl-CBS-oxazaborolidines derived from 1*S*,2*R*-(*-*) [126456-43-7] **and 1*R*,2*S*-(*+*) -*cis*-1-Amino-2-hydroxyindane** [136030-00-7] are also best prepared *in situ* by reaction of *cis*-1-aminoindan-2-ol (1*S*,2*R* see [126456-43-7]; 1*R*,2*S* see [136030-00-7]) with trimethylboroxine (see [823-96-1]) in a solvent (usually toluene), and they are **sensitive to moisture and air**. They catalyse the reduction of ketones and α -haloketones to their respective alcohols by BH_3 or some of its complexes, e.g. $\text{BH}_3\text{-DMS}$, $\text{BH}_3\text{-THF}$, with high stereospecificities ('ee' above 80%) at room temperature or below because of the steric effects of the methyl and indane groups around the heterocyclic boron atom. In a **typical preparation**, trimethylboroxine (0.19ml, 1.34mmol) is added to a solution of pure 1*R*,2*S*-1-aminoindan-2-ol (2.01mmol) in dry toluene (10ml), stirred at $\sim 25^\circ$ for 30 minutes, more toluene (10ml) is added, and the solution is concentrated by distillation to *ca* 2ml. The procedure is repeated twice then the toluene is removed *in vacuo* to give the **CBS-catalyst** as a white solid. Dry THF ($\sim 2\text{ml}$) is added to produce an approximately 1.0M solution of the catalyst in THF for later reactions. The solution should be stored under dry N_2 or argon and is stable for about 48 hours. For reduction reactions, the borane hydride (1.83mmol) is added to a solution of the chiral B-methylaminoindanol (0.17ml, 0.166mmol) in the desired solvent (e.g. toluene) (2ml) stirred under N_2 at $\sim 25^\circ$ for 30 minutes, cooled to 0° and the ketone (1.67mmol) in the desired solvent (e.g. toluene, 1ml) is added *via* a cannula, stirred at $\sim 25^\circ$ for 30 minutes, quenched with MeOH (5ml), H_2O (20ml) is added, the MeOH is removed *in vacuo*, and the resulting optically active alcohol is isolated from the aqueous solution. [Gilmore & Jones *Tetrahedron: Asymmetry* **14** 2115 2003, DOI: 10.1016/S0957-4166(01)00359-7; Hong et al. *Tetrahedron Lett* **35** 6631 1994, DOI: 10.1016/S0040-4039(00)73453-8.]

4-Dimethylaminopyridine (DMAP) [1122-58-3] $\text{C}_7\text{H}_{10}\text{N}_2$, **M 122.2**, **m 108-109°, 112-113°, b 191°/atm (under N_2)**, **pK^{5.4} 10.14, pK²⁰ 9.70, pK²⁵ 9.61, pK³⁵ 9.33 (in H_2O)**. Recrystallise DMAP from toluene [Sadownik et al. *J Am Chem Soc* **108** 7789 1986, DOI: 10.1021/ja00284a050], or from EtOH or $^*\text{C}_6\text{H}_6$ and dried *in vacuo*. It is prepared easily from 4-chloropyridine and Me_2NH in EtOH (2 hours/100°, or in a sealed tube for 12 hours/115-130°, Wibaut & Broekman *Recl Trav Chim Pays-Bas* **80** 309 1961, DOI: 10.1002/recl.

19610800309), or in very high yield from 4-trimethylsilyloxypyridine with $\text{Me}_2\text{NH}/\text{HgCl}_2$ (48 hours at 120° , Vorbrüggen *Angew Chem Int Ed* **11** 305 1972, DOI: 10.1002/anie.197203041). This **organocatalyst** is a strong base (see pKa) and promotes a large variety of acylation reactions, e.g. of *m*-chloroaniline and of benzyl alcohol at relative rates that are 3.14×10^6 and 3.45×10^8 compared with those of the uncatalysed reactions. It is useful industrially. The acylating agent is an acid chloride or an acid anhydride. Typically, catalytic amounts of DMAP are used together with one equivalent of a tertiary base, e.g. Me_3N , to neutralise the anion formed, i.e. Cl^- from the acid chloride or AcO^- from the anhydride, and to avoid protonating the catalyst. [Scriven *Chem Soc Rev* **12** 129 1983, DOI: 10.1039/CS9831200129.] Its **picrate** has **m 108°** (EtOH), and its **tetrafluoroborate salt** has **m 126°** (from MeCN/Et₂O). The UV has λ_{max} (ϵ) at 261nm (18,300 cm².mole⁻¹) for **free base** at pH >12.0, and 280.5nm (19,600 cm².mole⁻¹) for the **mono-cation** at pH <7.0 in H₂O (Essery & Schofield *J Chem Soc* 3939 1961, DOI: 10.1039/JR9610003939); and the IR (CHCl₃) has ν_{max} (ϵ_{A}) at 2840 (sh, 45), 1448 (95, Me bend), 1380 (155, CN str), 1346 (35), 1179 (35, Me₂ rock), 1108 (30, Me₂ rock), 1063 (35, NMe₂, C-N-C stretch) and 951 (55, NMe₂, C-N-C stretch) cm⁻¹ (Katritzky & Jones *J Chem Soc* 3674 1959, DOI: 10.1039/JR9590003674). [Beilstein **22** III/IV 4101, **22/9** V 112; for applications see Fieser vols **3** to **16**.]

DMAP is an efficient **desymmetrisation** catalyst. [Willis *JCS Perkin 1* 1765 1999, DOI: 10.1039/A906269B.] Thus it allows the use of an achiral or *meso* molecule, e.g. 3-substituted glutaric anhydride to react with chiral alcohols e.g. *R*-1-(1'-naphthyl)ethanol to produce (1'*R*,3*R*)-1(1'-naphthyl)ethyl methyl 3-substituted-pentanoate di-esters (after treatment with diazomethane), which are now chiral at C-3. The achiral carbon atom at C-3 in the glutaric anhydride becomes chiral in the pentanoate (mono-ester of glutarate) produced. [Theisen & Heathcock *J Org Chem* **53** 2374 1988, DOI: 10.1021/jo00245a051; Theisen & Heathcock *J Org Chem* **58** 142 1993, DOI: 10.1021/jo00053a027.] § It is commercially available in 0.04mmol impregnated tablets, and a polystyrene supported version (PS-DMAP) with a loading of ~3.0mmol/g is also available for recyclable acylation purposes.

***N,N'*-Dimethylethylenediamine [DMEDA, bis(methylamino)ethane] [110-70-3] C₄H₁₂N₂, M 88.2, 110-112°/750mm, 119°/atm, d²⁵ 0.819 g/ml, n_D²⁰ 1.431, pK₁⁰ 8.30, pK₂⁰ 10.89, pK₁²⁵ 6.80 (7.01), pK₂²⁵ 9.88 (10.03).** It has been prepared in several ways, but most commonly when ethylene dibromide (1mol), MeNH₂ (5mols) and H₂O (*ca* 3mols) are boiled under reflux for 15 hours. A 32% aqueous solution of NaOH is added, excess MeNH₂ is distilled out, and the rest of the liquid is distilled off. The diamine in the distillate is salted out with solid NaOH, dried (Na₂CO₃, or KOH), filtered and fractionally distilled (in *ca* 50% yield) until the NMR indicates that it is free from impurities. Store it in sealed containers under N₂ or argon as it is a strong base (see pKa). The ***N,N'*-bis(*p*-toluenesulfonamide)** has **m 164°** (from AcOH), ***N,N'*-bis(benzenesulfonamide)** has **m 129-131°** (from EtOH and AcOH, or EtOAc), the ***N,N'*-bis(benzamide)** has **m 177-178°** (from *C₆H₆), the **monopicrate** has **m 140°** (plates from EtOH or Me₂CO) and the **dihydrochloride** decomposes at **m 235°**. [Kermack & Wight *J Chem Soc* 1421 1935, DOI: 10.1039/JR9350001421; Boon *J Chem Soc* 307 1947, DOI: 10.1039/JR9470000307; Woodburn & O'gee *J Org Chem* **17** 1235 1953, DOI: 10.1021/jo50009a008; Bauer *J Am Chem Soc* **78** 1945 1956, DOI: 10.1021/ja01590a049.]

In the presence of resublimed *tert*-BuOK (1.5mmol), DMEDA (0.1mmol) **catalyses** cross-coupling between aryl and substituted aryl halides (0.5mmol) with arenes, e.g. *benzene, naphthalene (0.5mmol) at 80°/~4 hours, to provide the corresponding biaryls in high yields (~70 to >90%). *Ortho* substitution leads to decreased yields, and replacing KOBu^t with other bases (e.g. KOH, Na₂CO₃, KOAc, *tert*-BuONa, or *tert*-BuOLi) results in no reaction or extremely low yields. Of the amines studied, DMEDA is the **most effective catalyst** and evidence is presented that a **radical reaction** is involved. **It is the first example of a cross-coupling reaction that does not require a metal.** [Liu et al. *J Am Chem Soc* **132** 16737 2010, DOI: 10.1021/ja103050x; Beilstein **4** H 250, **4** I 425, **4** II 689, **4** III 512, **4** IV 1171.]

***R*-(+)-Indoline-2-carboxylic acid [98167-06 -7] C₉H₉NO₂, M 163.2, m 177°(dec), [α]_D²⁰ +11.0 (c 0.3, MeOH). Purify it as for the *S*-enantiomer below. The ***R*-(+)-hydrochloride** has [172152-19-1]. It is a proline based organocatalyst which promotes the enantioselective formation of cyclopropanes by reaction between 2-(dimethyl- λ^4 -sulfanylidene)-1-phenyl-ethanone (for stable sulfonium ylides see Ratts & Yao *J Org Chem* **31** 1185 1966, DOI: 10.1021/jo01342a047) and but-2-enals in high yields and very high stereoselectivity involving **Direct Electrostatic Activation (DEA)** [Kunz & MacMillan *J Am Chem Soc* **127** 3240 2005, DOI: 10.1021/ja042774b.]**

S*-(-)-Indoline-2-carboxylic acid [79815-20-6] C₉H₉NO₂, M 163.2, m 177°(dec), [α]_D²⁰ -114 (c 1, N HCl). It is purified as the racemate (see [78348-24-0] in 'Heterocyclic Compounds', Chapter 3). The ***S*-(-)-hydro-*

chloride [82923-76-0] **M 199.7, m 133^o(dec), [α]_D²⁰ -70 (c 1, EtOH)** crystallises from Et₂O/propan-2-ol. It is a proline based organocatalyst which promotes the enantioselective formation of cyclopropanes by reaction between 2-(dimethyl- λ^4 -sulfanylidene)-1-phenyl-ethanone (for stable sulfonium ylides see Ratts & Yao *J Org Chem* **31** 1185 1966, DOI: 10.1021/jo01342a047) and but-2-enals in high yields and very high stereoselectivity involving Direct Electrostatic Activation (DEA) [Kunz & MacMillan *J Am Chem Soc* **127** 3240 2005, DOI: 10.1021/ja042774b.]

α -Methyl-L-proline (S-2-methylproline) [42856-71-3] **C₆H₁₁NO₂, M 129.2, m 248-252^o(dec), [α]_D²⁵ -75 (c 2.0, MeOH), and (R-2-methylproline)** [58123-62-9]. This derivative catalyses reactions similar to those of proline. The general three-step synthesis from proline devised by Seebach and co-workers provides enantiomerically pure compound [*Helv Chim Acta* **64** 2704 1981, DOI: 10.1002/hlca.19810640829; *J Am Chem Soc* **105** 5390 1983, DOI: 10.1021/ja00354a034; Beck et al. *Org Synth* **72** 62 1993, DOI: 10.15227/orgsyn.072.0062]. It has been prepared on a 20-40g scale starting from S- or R- proline which is converted into **2S,5S- (or 2R,5S)- 2-tert-butyl-1-aza-3-oxabicyclo[3.3.0]octan-4-one** (*tert*-BuCHO, CF₃CO₂H as catalyst, in pentane and reflux for 114 hours with removal of H₂O) in 67-74% yield, which is methylated to **2S,5S- (or 2R,5S)- 2-tert-butyl-5-methyl-1-aza-3-oxabicyclo[3.3.0]octan-4-one** [86046-11-9] (MeI, *iso*-Pr₂NLi/THF/-78^o) with **complete retention of configuration** at C-5 in 93-95% yield; and hydrolysed (1.3M HCl/reflux for 1 hour, or 15% aqueous HBr/~25^o for 8 hours) to give the respective **2-methyl-proline hydrochloride** which is purified through a Dowex 50W x 8 (H⁺ form) column by washing with H₂O, then eluting with 3N aqueous NH₃ and evaporating to give the free **2-methylproline** in 85-90% yield. The last step is apparently quite tedious to carry out and is time consuming. The following procedure is more convenient and gives better than 90% yields of **pure 2-methylproline**. The above butyl-methyl-azaoxa-bicyclooctanone (13.4mmol) in MeOH/H₂O (6:1, 35ml) is treated with Silica gel 200-400 mesh, 60Å (3.0g) and stirred overnight. The silica gel is filtered off, washed with MeOH, the combined washings and filtrate are evaporated *in vacuo*, the pale yellow residue is dissolved in CHCl₃/MeOH (20:1), filtered through a 0.45µ filter disk (Acrodisc® CR PTFE), evaporated to dryness and the residue is triturated with Et₂O and dried *in vacuo*. Although it may be yellow in colour, it is almost spectroscopically pure, and can be purified further by dissolving in MeOH, treating with activated charcoal (5% w/w), filtering the whole suspension through Celite and evaporating. It can be crystallised by slow evaporation of the CHCl₃/MeOH solution, or from an MeOH/Et₂O mixture to give a first crop (50-70% yield) of white crystals leaving remaining **2-methylproline** and traces of yellow impurities in the mother liquors. Additional product crystallisation can be achieved by vapour diffusion of the supernatant against Et₂O in a closed desiccator. It also crystallises nicely from MeOH/EtOAc in colourless plates, unlike many α -amino acids. [Genin et al. *Tetrahedron Lett* **35** 4967 1994, DOI: 10.1016/S0040-4039(00)73294-1.]

S-2-Methylproline with [α]_D²⁰ -72.1 (c 1.0, MeOH) was shown to have an enantiomeric excess 'ee' of 99.0 ± 0.5% by chiral capillary gas chromatography. For this purpose the amino acid (5mg) is heated with *iso*-propyl isocyanate (0.3ml) in CH₂Cl₂ (0.3ml) at 100^o for 15 minutes, cooled, excess of volatiles are removed by a strong stream of dry air, the residue is dissolved in Et₂O (1ml), filtered through cotton wool and 10µl applied to the chiral capillary column. A Chirasil-Val fused-silica capillary column of Machery-Nagel (25m, 0.4mm) in a Carlo-Erba-Fraetovap 4160 HR GC unit has been used. The column was heated at 160^o for 15 minutes then raised by 2^o/min up to 200^o (argon as carrier gas ?). The amino acid has ¹H NMR (200MHz, D₂O, HDO at d 4.80) with δ at 1.52 (s, 3H, CH₃), 1.75-2.40 (m, 4H, 3-CH₂ + 4-CH₂), and 3.20-3.45 (m, 2H, 5-CH₂); and its ¹³C{¹H decoupled}NMR (50MHz, D₂O) has δ at 179.87 (CO₂H), 73.2 (C-2), 48.06 (CH₃-2), 38.27 (N-C-5), 25-85 (C-3) and 23.99 (C-4). [Beck et al. *Org Synth* **72** 62 1993, DOI: 10.15227/orgsyn.072.0062]. **S-2-Methylproline hydrobromide** [42856-71-3], prepared, by hydrolysis with 15% aqueous HBr (see above) evaporating and triturating with dry Et₂O, as colourless crystals (80%), has **m 136-137^o(dec), [α]_D²⁰ -28.9 (c 1.0, MeOH)** and its ¹H NMR (200MHz, D₂O, HDO at d 4.70) has δ at 1.60 (s, 3H, CH₃), 1.80-2.45 (m, 4H, 3-CH₂ + 4-CH₂), and 3.36 (t, 2H, *J* = 7 Hz, 5-CH₂) [Seebach et al. *J Am Chem Soc* **105** 5390 1983, DOI: 10.1021/ja00354a034; Overberger & Jon *J Polym Sci, Polym Sci Ed* **15** 1413 1977, DOI: 10.1002/pol.1977.170150612].

α -Methyl-DL-proline (RS-2-methylproline) [68078-09-1] **C₆H₁₁NO₂, M 129.2, m 263-264.5^o**. The racemic 2-methylproline was prepared by hydrolysing 5-(3-chloropropyl)-5-methylhydantoin (2g, 10mmol) with Ba(OH)₂ (6.3g, 20mmol) and H₂O (50ml) in a pressure bomb at 160^o for 30 minutes, cooling, adjusting the pH to 6 with 6N H₂SO₄, filtering off the BaSO₄, applying the filtrate through a column of Amberlite IRC 120 (H⁺ form), washing with H₂O, eluting with 4N NH₄OH, and evaporating the eluate to give a white solid m 252-258^o

(1.2g, 90%). Recrystallisation of the solid from MeOH/Et₂O gave *analytically pure* *RS*-2-methylproline **m 263-264.5°**. It has IR (KBr) with ν_{\max} at 3450 and 3200 (OH and NH), and 1600 (C=O) cm⁻¹; and its ¹H NMR (60MHz, CD₃OD, TMS) has δ at 1.6 (s, 3H, CH₃), 1.9 (m, 4H, 3-CH₂ + 4-CH₂), and 3.3 (m, 2H, 5-CH₂). [Ellington & Honigberg *J Org Chem* **39** 104 1974, DOI: 10.1021/jo00915a026]. It should be possible to prepare it by Seebach's procedure (see preceding entry) and starting with *RS*-proline.

Tetra-*n*-butylammonium fluoride (TBAF) [anhydrous, 1 M in THF 429-41-4; 3H₂O 87749-50-6; hydrate 22206-57-1] C₁₆H₃₆N⁺ F⁻, **M 261.5** (anhydrous), **315.5** (3 H₂O), **m liquid** (anhydrous), **62-63°** (3 H₂O), **37°** (18 H₂O). It is a powerful catalyst for the regioselective ring opening of epoxides with thiols [Albanese et al, *Synthesis* 34 1994, DOI: 10.1055/s-1994-25399], the addition reactions of trimethylallylsilane [Majetich et al. *J Org Chem* **51** 1745 1986, DOI: 10.1021/jo00360a021], Michael, Knoevenagel, aldol condensations, and by acting as a base in organic synthesis [Clark *Chem Rev* **80** 429 1980, DOI: 10.1021/cr60327a004]. It is a mild and highly efficient *source of nucleophilic (naked) fluoride ions*, causing substitution of halogen or tosyl groups by fluorine. TBAF is also used for *cleaving* silyl ethers and silyl protecting groups [cf. P.G.M. Wuts and T.W. Greene in *Green's Protective Groups in Organic Synthesis* 4t edn, J. Wiley, NY 2006, ISBN: 978-0-471-69754-1]. When an aqueous solution of tetra-*n*-butylammonium hydroxide is titrated with hydrofluoric acid, a homogeneously melting *octadecahydrate* (**m 37°**) is obtained [Fowler et al. *J Am Chem Soc* **62** 1140 1940, DOI: 10.1021/ja01862a039]. When the octadeca- or tri- hydrates are heated under high vacuum at temperatures of above 80°, decomposition occurs giving tetrabutylammonium bifluoride, tributylamine and 1-butene; with loss of a lot of its useful properties. When the *trihydrate* is warmed at 40° under high vacuum, almost '*anhydrous*' **TBAF** is obtained. This salt contained 0.1-0.3 molar equivalents of H₂O (by ¹H NMR). It is an oil at room temperature [Cox et al. *J Org Chem* **49** 3216 1984, DOI: 10.1021/jo00191a035]. **Virtually anhydrous TBAF in DMF** can be prepared without heating by using aliquots (~10 to 30mg) of the trihydrate and placing them under high vacuum (<0.1mm Hg) for 0.5 hour at ~25°, the flask is flushed with N₂, and dry DMF (2ml) is added. The solution is transferred (syringe) to a round-bottomed flask (under N₂), containing 4Å Molecular sieves (which had been activated by flame-drying for 5 minutes under high vacuum), and stirred with a micro stirrer bar for 30 minutes [Majetich et al. *J Org Chem* **51** 1745 1986, DOI: 10.1021/jo00360a021]. This is then used for catalytic or other reactions in anhydrous media. [Beilstein **4** III 292.]

TBAF is also available commercially as a 1 M solution in THF, 75% solution in H₂O, on alumina support 15wt% loading [429-41-4], on silica gel at ~1.5mmol/g F⁻ as a non-hygroscopic support for use as a source of '*naked*' *fluoride ions* [Ricci et al. *Tetrahedron Lett* **23** 577 1982, DOI: 10.1016/S0040-4039(00)86894-X; Gambacort et al. *Synth Commun* **19** 2441 1989, DOI: 10.1080/00397918908052645]. [For applications see Fieser **4** 477, **5** 645, **7** 353, **8** 467, **9** 444, **10** 379, **11** 499, **12** 458, **13** 286, **14** 293, **15** 298, **17** 324.]

PHASE TRANSFER CATALYSTS

Phase transfer catalysts (PTCs) are agents that are soluble in both lipophilic and hydrophilic solutions, i.e. soluble in aqueous solutions and in non-water soluble organic solvents, in varying degrees. They are mostly cationic species that assist the transfer of anions. e.g. CN⁻, from the aqueous phase into the non-aqueous phase, e.g. hexane, CHCl₃, which would contain the reactive compound, e.g. 1-octyl chloride, to form the product, i.e. 1-octylnitrile, in the non-aqueous phase in high yields. However, the catalysts can also be essentially neutral as with *crown ethers*, *cryptands* (tricyclic ethers containing nitrogen atoms at the bridgehead allowing formation of three rings) and *ionophores* that are useful for transferring metal ions through the bilayer interface. Some PTCs are polymer-bound making it easy for their removal and recycling. Other reactions that are phase transfer catalysed include: β -eliminations, *N*-alkylations, nucleophilic aromatic substitutions, metal-organic reactions, oxidation and reduction reactions. The reactions may proceed in either phase and/or at the interphase of the phases. They invariably occur in the organic phase, but in *Reverse Phase Transfer* reactions they occur in the aqueous phase. This section provides a selection of such agents, some of which are chiral and are capable of promoting stereoselective, or stereospecific reactions. [Starks *J Am Chem Soc* **93** 195 1971, DOI: 10.1021/ja00730a033; Brändström *Adv Phys Org Chem* **15** 267 1978, DOI: 10.1016/S0065-3160(08)60120-3; G.W. Gokel *Crown Ethers and Cryptands*, Royal Society of Chemistry, Cambridge, 1991, ISBN 0-85186-996-3; S. R. Cooper (Ed.), *Crown Compound: Towards Future Applications*, VCH, Weinheim, 1992, ISBN 1-56081-024-6, 3-527-28073-1; B. Deitrich, P. Viout and J-M, Lehn, *Macrocyclic Chemistry: Aspects of Organic and Inorganic Supramolecular Chemistry*, VCH, Weinheim, 1993, ISBN 3-527-28330-7; M. Hiraoka (Ed.), *Crown Ethers and Analogous Compounds*, Elsevier, Amsterdam, 1992, ISBN 0-444-88191-3; Ooi & Maruoka *Aldri-*

chimica Acta **40** 77 2007; K. Maruoka (Ed.), *Asymmetric Phase Transfer Catalysis*, Wiley-VCH, Weinheim, 2008, ISBN 978-3-527-31842-1; for *The Enantioselective Synthesis of α -Aminoacids by Phase-Transfer Catalysis with Chiral Schiff Base Esters* see O'Donnell [*Acc Chem Res* **37** 506 2004, DOI: 10.1021/ar0300625.]

Adogen 464 [methyl trialkyl(C8—C10)ammonium chloride] [72749-59-8] **M 416.2, d²⁵ 0.898, n_D²⁰ 1.4665**. This is purified in the same way as Aliquat 336 in the following entry. [US Patent 3 992 432].

Aliquat 336 (methyltricaprylammonium chloride, tri-*n*-octylmethylammonium chloride) [5137-55-3, a replacement product, Aliquat 128, has 63393-96-4] [$\text{CH}_3(\text{CH}_2)_6\text{CH}_2\text{N}^+\text{CH}_3 \text{Cl}^-$, **M 404.2, d²⁵ 0.884, n_D²⁰ 1.4665**. A 30% (v/v) of Aliquat 336 solution in *benzene is washed twice with an equal volume of 1.5M HBr. [Petrov & Allen, *Anal Chem* **33** 1303 1961, DOI: 10.1021/ac60178a004.] It is purified by dissolving 50g in CHCl_3 (100ml) and shaking with 20% NaOH solution (200ml) for 10 minutes, and followed by 20% NaCl (200ml) for 10 minutes. It is then washed with a small volume of H_2O , and filtered through a dry filter paper [Starks *J Am Chem Soc* **93** 195 1971, DOI: 10.1021/ja00730a033; Adam & Pribil *Talanta* **18** 733 1971, DOI: 10.1016/0039-9140(71)80114-5.]

***N*-Anthracenylmethyl)cinchonidinium chloride** [199588-80-2] $\text{C}_{34}\text{H}_{33}\text{ClN}_2\text{O}$, **M 521.1, m ~158-161°**. The salt can be purified by recrystallisation from $\text{CH}_2\text{Cl}_2/\text{Et}_2\text{O}$ and dried *in vacuo*. This *chiral phase-transfer-catalyst* (PTC) assists in the allyl alkylation of the α -carbon of glycine imino ester to yield finally α -allyl-substituted α -amino acids. Thus with a *ratio of imino ester to allyl compound*: i.e. achiral phosphines (e.g. 1,2-bis[diphenyl-phosphino]ethane DIPHOS, DPPE [1663-45-2], its oxide, triphenoxyphosphine or its oxide): base (KOH): [Pd(p-allyl)Cl]₂: PTC of 100:200:150:3.5:10 produced the α -allyl imino ester in varying yields and stereoselectivity. However, the derivative ***N*-(9-anthracenylmethyl)-*O*(9)-methyl-cinchonidinium iodide** (prepared from the chloride with MeI/50% aqueous NaOH/20°/4 hours) was a *more effective PTC*, providing better yields and with up to ~94% enantiomeric excess. The orange *iodide*, which is soluble in CH_2Cl_2 , is crystallised from $\text{CH}_2\text{Cl}_2/\text{Et}_2\text{O}$ and has [α]_D¹⁹ -288 (c 1, CH_2Cl_2); the ¹H NMR (500MHz, CDCl_3) has δ_{H} at 1.38-1.55 (m, 2H), 1.95-2.03 (m, 1H), 2.12-2.21 (m, 1H), 2.35-2.43 (m, 1H), 2.64-2.72 (m, 1H), 3.09 (dd, 1H, *J* = 5.6, 5.6Hz), 3.83 (s, 3H), 4.42-4.51 (m, 1H), 4.84 (d, *J* = 4.8Hz, 1H), 5.07 (d, *J* = 5.2Hz, 1H), 5.19 (d, *J* = 8.5Hz, 1H), 5.25 (bs, 1H), 5.87-6.02 (m, 2H), 6.52 (bs, 1H), 6.97 (d, *J* = 6.9Hz, 1H), 7.54-7.60 (m, 3H), 7.69-7.73 (m, 2H), 7.82-7.89 (m, 2H), 7.97 (brs, 1H), 8.05 (d, *J* = 4.3Hz, 1H), 8.14 (d, *J* = 4.1Hz, 1H), 8.20 (d, *J* = 4.3Hz, 1H), 8.32 (dd, *J* = 4.6Hz, 1H), 8.65 (s, 1H), 9.06 (d, *J* = 2.3Hz, 2H), 9.65 (d, *J* = 4.4Hz, 1H); and for IR and ¹³C NMR (125MHz, CD_3OD) see references. This *iodide* in CH_2Cl_2 was converted to another useful PTC ***N*-(9-anthracenylmethyl)-*O*(9)-methyl-cinchonidinium hexafluorophosphate** [by addition of NH_4PF_6 in H_2O at 25°/1 hour, dilution with CH_2Cl_2 and H_2O , drying the organic layer (MgSO_4) and evaporating] which gave yellow crystals from $\text{CH}_2\text{Cl}_2/\text{CHCl}_3$ with **m 163-164°**, [α]_D²³ -314 (c 1.1, CH_2Cl_2), with IR and NMR quite similar to those of the iodide. [Nakoji et al *J Org Chem* **67** 7418 2002, DOI: 10.1021/jo0260645.]

8*S*,9*R*-(*-*)-*N*-Benzylcinchonidinium chloride (BCDC) [69257-04-1] $\text{C}_{26}\text{H}_{29}\text{ClN}_2\text{O}$, **M 421.0, m 212-213° (dec), [α]_D²⁰ -180, -183 (c 5, H_2O), p*K*_{Est} ~5**. Dissolve the chloride in the minimum volume of H_2O and add absolute Me_2CO . Filter it off and dry it in a vacuum. It can also be recrystallised from hot EtOH or EtOH/ Et_2O . It is a good *chiral phase transfer catalyst* producing the *opposite enantiomer* to the one using *N*-benzylcinchoninium chloride as catalyst — see below. [Colonna et al. *JCS Perkin Trans 1* 547 1981, DOI: 10.1039/P19810000547; Imperiali & Fisher *J Org Chem* **57** 757 1992, DOI: 10.1021/jo00028a068.] [*Beilstein* **23** H 446.] See cinchonidine [485-71-2].

8*R*,9*S*-(*+*)-*N*-Benzylcinchoninium chloride (BCNC, *N*-benzyl-9*S*-hydroxycinchoninium chloride) [69221-14-3] $\text{C}_{26}\text{H}_{29}\text{ClN}_2\text{O}$, **M 421.0, m 265° (dec), [α]_D²⁰ +169 (c 0.4, H_2O), p*K*_{Est} ~ 5**. Recrystallise the chloride from isoPrOH, toluene or small volumes of H_2O . It is a good *chiral phase transfer catalyst* producing the *opposite enantiomer* to the one using the above *N*-benzylcinchonidinium chloride as catalyst — see above. [Juliá et al. *JCS Perkin Trans 1* 574 1981, DOI: 10.1039/P19810000574; Dolling et al. *J Am Chem Soc* **106** 446 1984, DOI: 10.1021/ja00314a045; Hughes et al. *J Org Chem* **52** 4745 1987, DOI: 10.1021/jo00230a017; *Beilstein* **23** IV 2832; Fieser **12** 380.] See cinchonine [118-10-5].

***Trans*-4-*n*-Butyl-4'-hydroxyethoxy-(ethoxy)₂-azobenzene (C-4-Azo-PEG1, $\text{C}_4\text{AzoOC}_2\text{E}_2$)** $\text{C}_{22}\text{H}_{30}\text{N}_2\text{O}_4$,

M 386.5, m 60°. This *micellar phase transfer catalyst* is a photochromic ionic surfactant with a photochromic core (azobenzene), a hydrophobic C4 butyl chain and a hydrophilic polyethylene glycol hydrophilic head group. It is prepared by the diazo-coupling of 4-*n*-butylbenzenediazonium ion with phenol and the resulting 4-*n*-butyl-4'-hydroxyazobenzene is condensed with trimethyleneglycol monotosylate. It is purified by silica gel chromatography (EtOAc/cyclohexane 7:3), the yellow eluate is evaporated and the residual yellow solid is recrystallised from pentane. The ^{13}C NMR (100MHz, CDCl_3 , TMS) has δ 160.9, 150.9, 147.2, 145.8, 129.0(2x), 124.5(x2), 122.5(2x), 114.7(2x), 72.4, 70.8, 70.3, 69.6, 67.6, 61.7, 35.5, 33.4, 22.3, 13.9 (the ^1H NMR is also reported). The surfactant undergoes reversible photoisomerisation when irradiated alternately with 365 and 254nm 500W lamp causing reversible conversion of *trans* to *cis* to *trans* with *photo-organising* and *de-organising* in the medium. UV-analysis showed that the CMCs (critical micellar concentrations) of the *trans*-form and *cis*-form in water are 4.1 μM and 8 μM respectively. Experimentation using Pd catalysed Tsuji-Trost reactions (cross-coupling for constructing C-C, C-N, C-S and C-O bonds [Trost & Van Vranken *Chem Rev* **96** 395 1996, DOI: 10.1021/cr9409804; Trost & Crawley *Chem Rev* **103** 2921 2003, DOI: 10.1021/cr020027w; Poli et al. *Top Organomet Chem* **38** 1 2012, DOI 10.1007/978-3-642-22749-3, ISBN 978-3-642-22748-6]) revealed that it is superior to other surfactants (such as SDS, Tween 20 and CTAB) when used at 3CMC (between the two isomers), when irradiated at 365nm during ~30 minutes before extraction of products, using microwave and heating. This *photochromic surfactant* was efficient (shorter time, high yields and high chemo-, regio- and stereoselectivities), recyclable and versatile for Pd-catalysed coupling reactions. [Billamboz et al. *J Org Chem.* **79** 493 2014, DOI: 10.1021/jo401737t; Shang et al. *Langmuir* **19** 10763 2003, DOI: 10.1021/la0350958.]

(11bR)-(-)- and (11bS)-(+)- 4,4-Dibutyl-4,5-dihydro-2,6-bis(3,4,5-trifluorophenyl)-3H-dinaphth[2,1-c:1',2'-e]azepinium bromide [*R*- 887938-70-7; *S*- 851942-89-7] $\text{C}_{42}\text{H}_{36}\text{BrF}_6\text{N}$, **M 748.6, m 223-228°, 224-229°, $[\alpha]_{\text{D}}^{29}$ - and + 11.7 (c 1.01, CHCl_3).** The precursor *chiral 2,2'-bisbromomethyl-1,1'-binaphthyl* (280mg, 0.4mmol), di-*n*-butylamine (140 μl , 0.8mmol) and K_2CO_3 (82mg, 0.6mmol) in MeCN (5ml) is refluxed with stirring for 10 hours. The mixture is poured into H_2O , extracted with CH_2Cl_2 , the extract is dried (Na_2SO_4), filtered and evaporated to dryness. The residue is purified by column chromatography on silical gel (eluting with MeOH/ CH_2Cl_2 , 1:20) to give the *bromide salt* (275mg, 92%) which may be recrystallised from MeOH/ Et_2O . The *S-enantiomer* has ^1H NMR (400MHz, CDCl_3) with δ_{H} at 7.97-7.95 (4H, m, Ar-H), 7.55-7.51 (2H, m, Ar-H), 7.27-7.23 (8H, m, Ar-H), 4.99 (2H, d, $J = 14.1\text{Hz}$, Ar- CH_2), 3.74 (2H, d, $J = 14.1\text{Hz}$, Ar- CH_2), 3.32 (2H, t, $J = 12.5\text{Hz}$, N- CH_2 - CH_2), 2.56 (2H, t, $J = 12.3\text{Hz}$, N- CH_2 - CH_2), 1.06-0.97 (6H, m, CH_2), 0.71 (6H, t, $J = 6.9\text{Hz}$, CH_3), 0.23 (2H, bs, CH_2); and for IR and ^{13}C NMR see references. [Kitamura et al. *Angew Chem Int Ed* **44** 1549 2005, DOI: 10.1002/anie.200462257; Ooi et al. *J Org Chem* **68** 4576 2003, DOI: 10.1021/jo030032f.]

These are potent phase transfer organocatalysts for asymmetric α -alkylation of *N*-arylidene-glycine *tert*-butyl ester derivatives for the synthesis of chiral α -substituted α -amino acids at extremely low concentrations of catalyst [Ooi et al. *Tetrahedron: Asymmetry* **17** 603 2006, DOI: 10.1016/j.tetasy.2006.01.019].

Cetyltrimethylammonium bromide (cetrionium bromide, CTAB) [57-09-0] $\text{CH}_3(\text{CH}_2)_{15}\text{N}^+(\text{CH}_3)_3 \text{Br}^-$, **M 364.5, m 227-235°(dec).** Crystallise it from EtOH, EtOH/*benzene or from wet acetone after extracting twice with petroleum ether. Shake it with anhydrous diethyl ether, filter and dissolve it in a little hot MeOH. After cooling in the refrigerator, the precipitate is filtered off at room temperature and re-dissolved in MeOH. Anhydrous ether is added and, after warming to obtain a clear solution, it is cooled and the crystalline material is collected. [Dearden & Woolley *J Phys Chem* **91** 2404 1987, DOI: 10.1021/j100293a040; Hakemi et al. *J Phys Chem* **91** 120 1987, DOI: 10.1021/j100285a028; Beilstein **4** IV 819; Fieser **15** 77.] It is a cationic surfactant with a micellar average weight of 62,000 Daltons, aggregation number of 170 and a CMC (critical micellar concentration) of 1mM (~25°). **Cetyltrimethylammonium chloride** [112-02-7] $\text{CH}_3(\text{CH}_2)_{15}\text{N}^+(\text{CH}_3)_3 \text{Cl}^-$, **M 320.0.** Crystallise the chloride from acetone/ether mixture, EtOH/ether, or from MeOH. [Moss et al. *J Am Chem Soc* **109** 4363 1987, DOI: 10.1021/ja00248a038; Beilstein **4** IV 819.]

Decyltrimethylammonium bromide [2082-84-0] $\text{CH}_3(\text{CH}_2)_9\text{N}^+(\text{CH}_3)_3 \text{Br}^-$, **M 280.3, m 239-242°.** Crystallise the salt from 50% (v/v) EtOH/ Et_2O , or from acetone and wash with ether. Dry it under vacuum at 60°. Also recrystallise it from EtOH and dry it over silica gel. [McDowell & Kraus *J Am Chem Soc* **73** 2170 1952, DOI: 10.1021/ja01149a074; Dearden & Woolley *J Phys Chem* **91** 2404 1987, DOI: 10.1021/j100293a040]

Didodecyldimethylammonium bromide [3282-73-3] $[\text{CH}_3(\text{CH}_2)_{11}]_2\text{N}^+(\text{CH}_3)_2 \text{Br}^-$, **M 463.6, m 157-162°.**

Recrystallise the salt from acetone, acetone/ether mixture, then from ethyl acetate, wash with ether and dry it in a vacuum oven at 60° [Chen et al. *J Phys Chem* **88** 1631 1984, DOI: 10.1021/j150652a038; Rupert et al. *J Am Chem Soc* **107** 2628 1985, DOI: 10.1021/ja00295a012; Rupert et al. *J Am Chem Soc* **108** 3920 1986, DOI: 10.1021/ja00274a011; Allen et al. *J Phys Chem* **91** 2320 1987, DOI: 10.1021/j100293a022].

Diocetadecyldimethylammonium bromide [3700-67-2] $[\text{CH}_3(\text{CH}_2)_{17}]_2\text{N}^+(\text{CH}_3)_2 \text{Br}^-$, **M 630.9, m 161-163°**. Crystallise the bromide from acetone, then MeOH [Lukac *J Am Chem Soc* **106** 4386 1984, DOI: 10.1021/ja00328a016]. Also purify it by chromatography on alumina by washing with $^*\text{C}_6\text{H}_6$ and eluting with Me_2CO , evaporating and crystallising from MeCN [Swain & Kreevoy *J Am Chem Soc* **77** 1122 1955, DOI: 10.1021/ja01610a012]. [*Beilstein* **4** IV 829.]

N,N-Dioctadecyl methylamine (hydrogen ionophore III) [4088-22-6] $[\text{CH}_3(\text{CH}_2)_{17}]_2\text{NCH}_3$, **M 536.0, m 40°, 44-46°, 48-49°, b 252-259°, $\text{pK}_{\text{Est}} \sim 10$** . It can be distilled at high vacuum, but by dissolving in $^*\text{C}_6\text{H}_6$, filtering and evaporating, a waxy solid suitable for electrode use can be obtained. It recrystallises from Me_2CO or MeCN. It is a **neutral ionophore** for polymeric membrane electrodes in proton sensitive solvents. [Hoerr et al. *J Org Chem* **9** 201 1944, DOI: 10.1021/jo01184a011; Wu & Yu *Talanta* **34** 577 1987, DOI: 10.1016/0039-9140(87)80193-5; *Beilstein* **4** III 435.]

Dodecyltrimethylammonium chloride [112-00-5] $\text{CH}_3(\text{CH}_2)_{11}\text{N}^+(\text{CH}_3)_3 \text{Cl}^-$, **M 263.9, m 246°(dec)**. Dissolve the chloride in MeOH, treat with active charcoal, filter and dry it *in vacuo* [Walderhaug *J Phys Chem* **88** 1655 1984, DOI: 10.1021/j150652a043], or recrystallise it several times from 10% EtOH in acetone. It has also been repeatedly crystallised from EtOH/ether or MeOH, and its solubility in H_2O is 50mg/ml at ~25°. [Cella et al. *J Am Chem Soc* **74** 2061 1952, DOI: 10.1021/ja01128a061; *Beilstein* **4** IV 79.]

Hexyltrimethylammonium bromide [2650-53-5] $\text{CH}_3(\text{CH}_2)_5\text{N}^+(\text{CH}_3)_3 \text{Br}^-$, **M 224.3, m 186°**. Recrystallise it from acetone. It is extremely *hygroscopic*. [McDowell and Kraus *J Am Chem Soc* **73** 2170 1951, DOI: 10.1021/ja01149a074; *Beilstein* **4** IV 710.]

Octadecyl trimethylammonium bromide (stearyl trimethylammonium bromide) [1120-02-1] $\text{CH}_3(\text{CH}_2)_{17}\text{N}^+(\text{CH}_3)_3 \text{Br}^-$, **M 392.5, m ~250°dec, 230-240°(dec)**. Crystallise it from EtOH or H_2O (solubility is 1 in 1000parts). It is very soluble in Me_2CO . It is also a bactericide. [Shelton et al. *J Am Chem Soc* **68** 753 1946, DOI: 10.1021/ja01209a012; Grieger & Kraus *J Am Chem Soc* **70** 3803 1948, DOI: 10.1021/ja01191a078; *Beilstein* **4** IV 827.]

Tetra-n-amylammonium bromide (tetra-n-pentylammonium bromide) [866-97-7] $[\text{CH}_3(\text{CH}_2)_4]_4\text{N}^+ \text{Br}^-$, **M 378.5, m 100-101°**. Crystallise it from petroleum ether, $^*\text{benzene}$ or acetone/ether mixtures and dry it *in vacuo* at 40-50° for 2 days. It is used in ion-paired chromatography [Sagara et al. *J Chromatogr* **328** 289 1985, DOI: 10.1016/S0021-9673(01)87399-4]. [*Beilstein* **4** IV 677.]

Tetra-n-amylammonium iodide [2498-20-6] $[\text{CH}_3(\text{CH}_2)_4]_4\text{N}^+ \text{I}^-$, **M 425.5, m 135-137°**. Crystallise the iodide from EtOH and dry it at 35° under a vacuum. It has also been purified by dissolving in acetone and precipitating by adding diethyl ether, and drying at 50° for 2 days. [*Beilstein* **4** IV 677.]

Tetradecyltrimethylammonium bromide (myristyl trimethylammonium bromide) [1119-97-7] $\text{CH}_3(\text{CH}_2)_{13}\text{N}^+(\text{CH}_3)_3 \text{Br}^-$, **M 336.4, m 244-245°, 244-249°, 245-250°**. Crystallise the bromide from acetone or a mixture of Me_2CO and >5% MeOH, or $\text{Me}_2\text{CO}/\text{EtOH}$. Wash it with diethyl ether and dry it in a vacuum oven at 60°. It is a **cationic detergent** with a micellar average weight of 27,000 Daltons, aggregation number 80 and CMC (critical micellar concentration) of 4-5mM ~25°. Its solubility is 1g/5g H_2O . [Dearden & Woolley *J Phys Chem* **91** 2404 1987, DOI: 10.1021/j100293a040; Shelton et al. *J Am Chem Soc* **68** 753 1946, DOI: 10.1021/ja01209a012; *Beilstein* **4** III 419, **4** IV 813.]

Tetra-n-heptylammonium iodide [3535-83-9] $[\text{CH}_3(\text{CH}_2)_6]_4\text{N}^+ \text{I}^-$, **M 537.7, m 102-103°**. Crystallise the iodide from EtOH or aqueous EtOH. [Eriksen et al. *J Org Chem* **25** 849 1960, DOI: 10.1021/jo01075a616; *Beilstein* **4** IV 736 for triheptylamine.]

Tetra-n-hexylammonium bromide [4328-13-6] $[\text{CH}_3(\text{CH}_2)_5]_4\text{N}^+ \text{Br}^-$, **M 434.6, m 99-100°**. Wash the bromide with ether, and dry it in a vacuum at room temperature for 3 days. [For application see Fieser **10** 383.]

Tetra-n-hexylammonium chloride [5922-92-9] $[\text{CH}_3(\text{CH}_2)_5]_4\text{N}^+ \text{Cl}^-$, **M 390.1, m 111-113°**. Crystallise the chloride from EtOH.

Tetra-n-hexylammonium iodide [2138-24-1] $[\text{CH}_3(\text{CH}_2)_5]_4\text{N}^+ \text{I}^-$, **M 481.6, m 99-101°, 102-103°**. Wash the iodide with diethyl ether and dry it at room temperature *in vacuo* for 3 days. It is soluble in CH_2Cl_2 . [Eriksen et al. *J Org Chem* **25** 849 1960, DOI: 10.1021/jo01075a616; *Beilstein* **4** IV 711 for trihexylamine.]

Tetrahexylammonium perchlorate [4656-81-9] $[\text{CH}_3(\text{CH}_2)_5]_4\text{N}^+ \text{ClO}_4^-$, **M 454.1, m 104-106°**. Crystallise the salt from acetone and dry it *in vacuo* at 80° for 24 hours.

Tetra-*n*-propylammonium bromide [1941-30-6] $[\text{CH}_3\text{CH}_2\text{CH}_2]_4\text{N}^+ \text{Br}^-$, **M 266.3, m >280°(dec)**. Crystallise it from ethyl acetate/EtOH (9:1), acetone or MeOH. Dry it at 110° under reduced pressure. [Beilstein 4 IV 471.]

Tetra-*n*-propylammonium iodide [631-40-3] $[\text{CH}_3\text{CH}_2\text{CH}_2]_4\text{N}^+ \text{I}^-$, **M 313.3, m >280°(dec)**. Purify the iodide by crystallising it from EtOH, EtOH/diethyl ether (1:1), EtOH/water or aqueous acetone. Dry it at 50° under a vacuum and store it over P_2O_5 in a vacuum desiccator. Store it away from light. [Beilstein 4 IV 472.]

Tri-*n*-butylhexadecylphosphonium bromide (TBHDTB, hexadecyltributylphosphonium bromide) [14937-45-2] $\text{CH}_3(\text{CH}_2)_{15}\text{P}^+[(\text{CH}_2)_3\text{CH}_3]_3 \text{Br}^-$, **M 507.7, m 54°, 56-58°**. It is made by heating 1-bromohexadecane (1 mol, 112-82-3) and tri-*n*-butylphosphine (1 mol, 998-40-3) at 65° for 3 days. The mixture solidifies on cooling, and the solid is recrystallised from hexane and dried *in vacuo* to give 68% of the phosphonium salt. [Starks *J Am Chem Soc* **93** 195 1971, DOI: 10.1021/ja00730a033; for preparing fluorides see Landini et al. *Synthesis* 37 1974, DOI: 10.1055/s-1974-23231; Landini et al. *Synthesis* 428 1974, DOI: 10.1055/s-1974-23333; for making disulfides see Landini & Rolla *Synthesis* 565 1974, DOI: 10.1055/s-1974-23371; and Fieser **5** 322, **6** 271.]

Tri-*n*-dodecylamine (Hydrogen ionophore I) [102-87-4] $[\text{CH}_3(\text{CH}_2)_{11}]_3\text{N}$, **M 522.0, m 15.7°, b 220-228°/0.03mm, d_4^{25} 0.833, n_D^{20} 1.4577, $\text{pK}_{\text{Est}} \sim 11.0$** . Distil tridodecylamine under high vacuum and N_2 , and store it in the absence of CO_2 . It can be recrystallised from 95%EtOH/* C_6H_6 at low temperature under vacuum. The **hydrochloride** has **m 78-79°**. [Ralston et al. *J Org Chem* **9** 259 1944, DOI: 10.1021/jo01185a009; Beilstein 4 III 413, 4 IV 801.]

Tri-*n*-dodecylammonium nitrate [2305-34-2] $[\text{CH}_3(\text{CH}_2)_{11}]_3\text{N}^+ \text{NO}_3^-$, **M 585.0**. Crystallise the salt from *n*-hexane/acetone (95:5) and keep it in a desiccator over P_2O_5 under vacuum. [Beilstein 4 IV 801 for tridodecylamine.]

Tri-*n*-dodecylammonium perchlorate [5838-82-4] $[\text{CH}_3(\text{CH}_2)_{11}]_3\text{N}^+ \text{ClO}_4^-$, **M 622.4**. Recrystallise the salt from *n*-hexane or acetone and keep it in a desiccator over P_2O_5 . (Potentially explosive.)

Tri-*n*-octylammonium chloride [1188-95-0] $[\text{CH}_3(\text{CH}_2)_7]_3\text{N}^+ \text{Cl}^-$, **M 384.2, m 78-79°, pK^{25} 8.35 (in 70% aqueous EtOH)**. Crystallise it from Et_2O , then *n*-hexane (see above). [Burrows et al. *J Chem Soc* 197 1947, DOI: 10.1039/JR9470000197; Beilstein 4 H 196.]

Tri-*n*-octylammonium perchlorate [2861-99-6] $[\text{CH}_3(\text{CH}_2)_7]_3\text{N}^+ \text{ClO}_4^-$, **M 454.2, m >300°(dec)**. Crystallise the perchlorate from *n*-hexane. (Possibly explosive.) [Beilstein 4 IV 754.]

IMIDAZOLINIUM IONIC LIQUID CRYSTAL CATALYSTS

—A general discussion on ionic liquid crystals has already been made (see Chapter 2), and this section includes some commercially available liquid crystal 1,3-dialkylimidazolium salts that participate in the catalytic process, i.e. by coordination with the metal component of catalysts, as well as assisting in other ways such as acting as a solvent, affecting the dielectric, and in phase transference of reagents. Depending on the *N*-alkyl substituents and the counter anion the salts may be more, or less, soluble in water or organic solvents. They are non-volatile (i.e. environmentally friendly), stable to water and air, and many may be heated to temperatures as high as 300°, making it possible to distil off some products from them. They can be recovered for re-use. [See P. Wasserscheid and T. Welton, *Ionic Liquids in Synthesis* (2 volume set), Wiley VCH, Weinheim, 2008. ISBN-10: 3527312390, ISBN-13: 978-3527312399.]

Preparation: Alkylation of *N*-1 of imidazole can be carried out using alkyl halide and alkoxide as for 1-ethylimidazole [1072-62-4], and the second alkylation onto *N*-3 is performed by further reaction with the desired alkylbromide (slightly less than a molar equivalent) in a solvent, e.g. 1,1,1-trichloroethane, at reflux for 2 hours. The molten bromide salt is separated from the solvent, washed with trichloroethane, dried on a Rotavap at 70°/0.1mm to give the bromide salt as a white solid or a liquid in high yield (~>90%) and high state of purity if the starting reagents are pure. Check purity by ^1H NMR and elemental analysis. The imidazolium bromide salts are most useful as they can be converted into other salts, e.g. AcO^- with AgOAc , TfO^- with AgOTf , bis(trifluoromethylsulfonyl)imide $[\text{NTf}_2^-]$ with NTf_2Li , nonafluorobutanesulfonate $[\text{NfO}^-]$ with NfOK , etc. Generally the imidazolium bromide and the metal salt of the required anion are dissolved in the least volume of H_2O at 70°, stirred for 1 hour (if silver halide precipitates it is filtered off), and the imidazolium salt can be extracted into an organic solvent, e.g. CH_2Cl_2 or 1,1,1-trichloroethane, and the extract is then evaporated *in vacuo* and dried at 50°/0.1mm for 2 hours or until there is no further loss in weight. **Alternatively**, where possible, the 1-alkylimidazole is alkylated with e.g. alkyl tosyl ester to provide the 1,3-dialkylimidazolium toluene-*p*-sulfonate directly. [Bonhôte et al. *Inorg Chem* **35** 1168 1996, DOI: 10.1021/ic951325x].

The following liquid crystal salts can be readily prepared in small or large quantities:

1-Butyl-3-methylimidazolium Salts (BMIM⁺ X⁻, see Park & Kazlauskas *J Org Chem* **66** 8395 2001, DOI: 10.1021/jo015761e; for improved preparation, purification and use at ambient temperature of some of the following ionic liquids in lipase-catalysed enantio- and regio- selective acylation reactions):

BMIM⁺ AcO⁻ [284049-75-8] **C₁₀H₁₈N₂O₂**, **M 198.6**; **BMIM⁺ NTf₂⁻** [174899-83-3] **C₁₀H₁₅F₆N₃O₄S₂**, **M 419.4**, **n_D²⁰ 1.428**, [good synthetic activity of soluble *Candida antarctica* lipase B in this ionic liquid was obtained towards the enantioselective and operational stability for butyl butyrate synthesis and kinetic resolution of 1-phenylethanol in supercritical CO₂, Lozano et al. *JCS Chem Commun* 692 2002, DOI: 10.1039/B200055E]; **BMIM⁺ Br⁻** [85100-77-2] **C₈H₁₅BrN₂**, **M 219.1**, [used in Heck reactions with Pd(OAc)₂ where Pd-Imetal bond is formed, Xu et al. *Organometallics* **19** 1123 2000, DOI: 10.1021/om990956m; **BMIM⁺ Cl⁻** [79917-90-1] **C₈H₁₅ClN₂**, **M 174.7**, **m ~70°**, [used with AlCl₃ in Friedel-Crafts alkylation reactions, Chauvin et al. *J Mol Catal* **92** 155 1994, DOI: 10.1016/0304-5102(94)00065-4]; **BMIM⁺ (n-BuO)₂P(O)O⁻** [663199-28-8] **C₁₆H₃₃N₂O₄P**, **M 348.4**, **n_D²⁰ 1.472**; **BMIM⁺ (NC)₂N⁻** [448245-52-1] **C₁₀H₁₅N₅**, **M 205.3**; **BMIM⁺ SbF₆⁻** [174645-81-9] **C₈H₁₅F₆N₂Sb**, **M 375.0**; and **BMIM⁺ PF₆⁻** [174501-64-5] **C₈H₁₅F₆N₂P**, **M 284.2**, **n_D²⁰ 1.411**, [used in allylation of RCHO with (Allyl)Sn, Gordon & McCluskey *JCS Chem Commun* 1431 1999, DOI: 10.1039/A903661J; in coupling of aryl halides by (Ph₃)_nNi(0), Howarth et al. *Tetrahedron Lett* **41** 10319 2000, DOI: 10.1016/S0040-4039(00)01854-2; and catalyse the addition of CN (from TMSCN) to arylimines (from ArCHO + RNH₂) to form α-aminonitriles, Yadav et al. *New J Chem* **27** 462 2003, DOI: 10.1039/B208844B]; **BMIM⁺ HCO₃⁻** [366491-15-8] **C₉H₁₆N₂O₃**, **M 200.2**, as 50% in 2:3 MeOH:H₂O; **BMIM⁺ HSO₄⁻** (as **BASIONIC[®] AC 28**) [262297-13-2] **C₈H₁₆N₂O₄S**, **M 236.3**; **BMIM⁺ MeSO₃⁻** (as **BASIONIC[®] ST 78**) [342789-81-5] **C₉H₁₉N₂O₃S**, **M 234.3**; and **BMIM⁺ MeSO₄⁻** [401788-98-5] **C₉H₁₉N₂O₄S**, **M 250.3**, **n_D²⁰ 1.478**, [a halogen-free ionic liquid that catalyses the addition of CN (from TMSCN) to arylimines (from ArCHO + RNH₂) to form α-aminonitriles, Yadav et al. *New J Chem* **27** 462 2003, DOI: 10.1039/B208844B; Itoh et al. *Chem Lett* (Jpn) **32** 654 2003, DOI: 10.1246/cl.2003.654]; **BMIM⁺ BF₄⁻** [174501-65-6] **C₈H₁₅BF₄N₂**, **M 226.0**, {specifically catalyses the *Biginelli reaction* (formation of 3,4-dihydropyrimidin-2(1H)-ones from aldehydes + urea + MeCOCH₂COR), Peng & Deng *Tetrahedron Lett* **42** 5917 2001, DOI: 10.1016/S0040-4039(01)01139-X; it is an *ionic liquid* used in many reactions such as hydrogenations [Dyson et al. *JCS Chem Commun* 25 1999, DOI: 10.1039/A807447J], asymmetric hydrogenations proceeding in higher enantioselectivity than in homogenous phases [Monteiro et al. *Tetrahedron: Asymmetry* **8** 177 1997, DOI: 10.1016/S0957-4166(96)00485-5], and in Suzuki cross-coupling at room temperature [Mathews et al. *JCS Chem Commun* 1249 2000, DOI: 10.1039/B002755N]; **BMIM⁺ PF₆⁻** [174501-64-5] **C₈H₁₅F₆N₂P**, **M 284.2**, [catalyses the addition of CN (from TMSCN) to arylimines (from ArCHO + RNH₂) to form α-aminonitriles, Yadav et al. *New J Chem* **27** 462 2003, DOI: 10.1039/B208844B; assists in the bi-phasic hydrogenation of arenes at room temperature using a ruthenium cluster catalyst which coordinates with it, Dyson et al. *JCS Chem Commun* 25 1999, DOI: 10.1039/A807447J; and the molten salt catalyses the asymmetric hydrogenation of 2-arylacrylic acids by immobilised the Ru-BINAP complex, Montiero et al. *Tetrahedron: Asymmetry* **8** 177 1997, DOI: 10.1016/S0957-4166(96)00485-5; and is used in Palladium [Pd(Ph₃)₄] catalysed Suzuki cross-coupling (aryl halides and arylboronic acids) at ambient temperature, Mathews et al. *JCS Chem Commun* 1249 2000, DOI: 10.1039/B002755N; and it specifically catalyses the *Biginelli reaction* (formation of 3,4-dihydropyrimidin-2(1H)-ones from aldehydes + urea + MeCOCH₂COR), Peng & Deng *Tetrahedron Lett* **42** 5917 2001, DOI: 10.1016/S0040-4039(01)01139-X}; and **BMIM⁺ CF₃SO₃⁻** [174899-66-2] **C₉H₁₅F₃N₂O₃S**, **M 288.3**, **n_D²⁰ 1.434**; **BMIM⁺ octyloSO₃⁻** [445473-58-5] **C₁₆H₃₂N₂O₄S**, **M 348.5**, [this ionic liquid inceases the yield and enzyme stability of β-galactosidase in enzyme-catalysed syntheses, Kaftzik et al. *Org Process Res Dev* **6** 553 2002, DOI: 10.1021/op0255231].

1-Ethyl-3-methylimidazolium Salts (EMIM⁺ X⁻, see Park & Kazlauskas *J Org Chem* **66** 8395 2001, DOI: 10.1021/jo015761e; for improved preparation and use of ambient temperature of some of the following ionic liquids in lipase-catalysed enantio- and regio- selective acylation reactions):

EMIM⁺ Br⁻ [65039-08-9] **C₆H₁₁BrN₂**, **M 191.1**, [it is *hydrophobic* and a highly conductive salt melting at ~25°, Bonhôte et al. *Inorg Chem* **35** 1168 1996, DOI: 10.1021/ic951325x]; **EMIM⁺ Cl⁻** [65039-09-0] **C₆H₁₁ClN₂**, **M 146.6**, **m 77-79°**; **EMIM⁺ (n-BuO)₂P(O)O⁻** [869858-84-4] **C₁₄H₂₉N₂O₄P**, **M 320.4**, **n_D²⁰ 1.469**; **EMIM⁺ AcO⁻** [143314-17-4] **C₈H₁₄N₂O₂**, **M 170.2**, **n_D²⁰ 1.502**; **EMIM⁺ (CF₃CF₂-SO₂)₂N⁻** [216299-76-2] **C₁₀H₁₁F₁₀N₃O₄S**, **M 491.3**, **m ≤1°**, [very stable fluorinated ionic liquids that are *extremely*

hydrophobic]. **EMIM⁺ NTf₂⁻** [174899-82-2] **C₈H₁₁F₆N₃O₄S, M 391.3, m ≥15°**, is a useful medium for the enantioselective cyclopropanation of styrene with ethyldiazoacetate, promoted by two different Cu-bis(oxazoline) complexes and its recovery, Fraile et al. *Tetrahedron: Asymmetry* **12** 1891 2001, DOI: 10.1016/S0957-4166(01)00315-9. Good synthetic activity of soluble *Candida antarctica* lipase B in this ionic liquid was obtained towards the enantioselective and operational stability for butyl butyrate synthesis and kinetic resolution of 1-phenyl ethanol in supercritical CO₂, Lozano et al. *JCS Chem Commun* 692 2002, DOI: 10.1039/B200055E]; **EMIM⁺ (CN)₂N⁻** [370865-89-7] **C₈H₁₁N₅, M 177.2**, [its conductivity is ~26,000 μS/cm, and its electrochemical window is -2.4 to +3.3 V, cf. Bonhôte et al. *Inorg Chem* **35** 1168 1996, DOI: 10.1021/ic951325x; **EMIM⁺ PF₆⁻** [155371-19-0] **C₆H₁₁F₆N₂P, M 256.1, m 58-62°**, [it is prepared by mixing EMIM⁺ Cl⁻ (29.3g, 200mmol) and 60% aqueous HPF₆ (9.g, 200mmol) in H₂O (300ml), the resulting mixture of white solid and liquid are cooled in an ice bath for 2 hours and the EMIM⁺ PF₆⁻ (31.8g 62%) is dried *in vacuo*. Recrystallisation from MeOH provides crystals for X-ray structural analysis. [Fuller et al. *JCS Chem Commun* 299 1994, DOI: 10.1039/C39940000299]; and **EMIM⁺ BF₄⁻** [143314-16-3] **C₆H₁₁BF₄N₂, M 198.0, m 15°, b >350°, d₄²⁵ 1.294, n_D²⁰ 1.413**, [its conductivity is ~11,500 μS/cm, and its electrochemical window is -2.2 to +3.5 V, cf. Bonhôte et al. *Inorg Chem* **35** 1168 1996, DOI: 10.1021/ic951325x]; the salt is prepared by stirring Ag₂O (23.2g, 100mmol) with 48% aqueous HBF₄ (36.9g, 200mmol) in H₂O (300ml) until the Ag₂O has reacted completely to give a clear solution, then EMIM⁺ Cl⁻ (29.2g, 200mmol) dissolved in H₂O is added and the mixture is stirred for 2 hours, the AgCl is filtered off, the filtrate is evaporated *in vacuo* and the colourless residue is dried in a vacuum oven at 60° to give the **BF₄⁻ salt** (33.6g, 85%). [Fuller et al. *JCS Chem Commun* 299 1994, DOI: 10.1039/C39940000299].

Enzymes as catalysts: see ‘Introduction’ and ‘Proteins and Enzymes’ in Chapter 6.

CATALYSTS—Part 2

ORGANIC COMPOUNDS USED FOR MAKING LIGANDS THAT ASSIST CATALYSIS

Ligands and reagents in this section are mainly ones that are used within this and the former section Catalysts-Part 1. Other such ligands and related reagents not included here will be found scattered in Chapters 3, 4 and 6, and can be located by their CAS Registry Numbers (CASRNs) in the CASRN Index, in the General Index (as abbreviations) or from their commonly used names in the relevant chapters.

(η^3 -Allyl)(η^5 -cyclopentadienyl)palladium(II) [(allyl)(cyclopentadienyl)palladium(II)] [1271-03-0] $\text{C}_8\text{H}_{10}\text{Pd}$, **M 212.5, m 61° (dec), 63-63.5°** This complex is volatile and should be handled in an efficient fume hood. Using Schlenk equipment under N_2 or argon and strictly dry conditions, a clear yellow solution of bis(η^3 -allyl)di- μ -chloro-dipalladium(II) (9.9g, 27mmol, see [12012-95-2]) in THF (100ml) and $^*\text{C}_6\text{H}_6$ (100ml) is prepared, and cooled in an ice-NaCl bath to -20° . A solution of sodium cyclopentadienyl (54mmol in 28ml of THF, [4984-82-1]) in a N_2 flushed syringe is added dropwise to the yellow solution at -20° with stirring, whereby the colour changes to dark red. The ice bath is removed after stirring for 1 hour, the temperature is allowed to rise to $\sim 25^\circ$, stirring is continued for 30 minutes and the solvents are removed by evaporation *in vacuo* (30-60 torr; no higher than 20° because the Pd complex will begin to sublime at $\sim 25^\circ$) to yield a dark red solid. The residue is extracted with hexane (80ml), the extract is filtered under N_2 (use fluted filter paper as a glass frit is likely to become clogged). The filtrate is evaporated as before (*in vacuo* at 30-60 torr) to give red needles of the **palladium(II) complex** (9.2g, 80%). **Note** that by using mechanical stirring the yield can be improved to 98%. This product is satisfactory for most preparations of Pd(0) complexes, but an analytical sample is readily obtained as red needle-like crystals by recrystallisation from light petroleum (b 40 - 60°), or by subliming it at 40° (bath temperature)/30mm. It has an **unpleasant odour**, is relatively stable in air at $\sim 25^\circ$ for a few days, unchanged in a refrigerator for several weeks, but decomposes gradually in air at $\sim 25^\circ$ to form a black solid which is not soluble in hexane. Best to stored below -20° under N_2 or argon in sealed glass tubes. It is **diamagnetic** and has a low dipole moment ($< 1.5\text{D}$) consistent with both the cyclopentadienyl and the allyl groups as sandwich ligands. The ^1H NMR ($^*\text{C}_6\text{D}_6$) has δ at 2.14 (d, $J = 11\text{Hz}$, 2H), 3.11 (d, $J = 6\text{Hz}$, 6H) and 4.63 (m, 1H) complex for allyl protons and 8.1 (s, 5H, cyclopentadienyl protons). [Shaw *Proc Chem Soc (London)* 247 1960, **notes start on p 233**, DOI: 10.1039/PS9600000233; Tatsuno et al. *Inorg Synth* **19** 220 1979, DOI: 10.1002/9780470132500.ch51;] The complex is useful for preparing Pd(0) complexes by reaction with hindered alkyl phosphines [Otsuka et al. *J Am Chem Soc* **98** 5850 1976, DOI: 10.1021/ja00435a017], reacts with isonitriles to form $\text{Pd}(\text{CNR})_2$ complexes [Fischer & Werner *Chem Ber* **95** 703 1962, DOI: 10.1002/cber.19620950321; Otsuka et al. *J Am Chem Soc* **91** 6994 1969, DOI: 10.1021/ja01053a017], and has been used to prepare BINAP complexes such as $\text{Pd}[(R)\text{-BINAP}]_2$ for asymmetric catalysis [Ozawa et al. *Organomet* **12** 4188 1993, DOI: 10.1021/om00034a064].

Ammonium perrhenate (NH_4ReO_4) [13598-65-7] NH_4ReO_4 , **M 268.2, m 365° (dec.), d 25 3.97g/ml, pK 25 - 1.25 (for HReO_4)**. The higher solubility of the ammonium salt in H_2O (17g/L at 0° , and 162g/L at $\sim 50^\circ$) compared to that of the potassium salt (see [10466-65-6]) has made this salt preferable for use in the preparation of a variety of rhenium compounds. It is prepared from perrhenic acid (see [13768-11-1]) which is obtained from KReO_4 (10.0g, 0.036mol) using the procedure of Watt et al. (*Inorg Synth* **7** 187 1963, DOI: 10.1002/9780470132388.ch51) whereby a solution in H_2O (100ml, solubility is 14% at 100°) at 90° is passed through a column of cation exchange resin (Dowex 50-Wx2, but not x8) held at 90° (preferably coated with 'Instatherm', Ace Glass Co., Inc Vineland, NJ, USA, to withstand the temperature), washed with H_2O , and the combined aqueous eluates are concentrated *in vacuo* down to the critical volume of 5ml. If it is concentrated further and the colour darkens, then one drop of 30% aqueous H_2O_2 should be added to decolourise it. The solution is cooled to $\sim 0^\circ$, and while being stirred, a chilled mixture of 2-propanol (15ml, saturated with gaseous NH_3) and Et_2O (50ml) is added. The cold mixture is allowed to stand for 1 hour, and the white hexagonal plates ($\sim 90\%$ yield) are filtered onto a sintered-glass funnel, washed three times with 2-propanol/ Et_2O (1:9), dried *in vacuo*, then at 110° to give NH_4ReO_4 (8.9g, 96% yield) that is **99.4-99.95% pure** by microanalysis, and by passage through a cation-exchange resin in H_2O and titrating with standard base. [Thompson et al. *Inorg*

Synth **8** 171 1966, DOI: 10.1002/9780470132395.ch44; see also Smith & Long *J Am Chem Soc* **70** 354 1948, DOI: 10.1021/ja01181a110.]

Benzotriazol-1-yloxy-tris(dimethylamino)phosphonium hexafluorophosphate (BOP.PF₆, Castro's reagent) [56602-33-6] C₁₂H₂₂F₆N₆OP₂, M 442.3, m >130°, 147-148°. 130°. Castro's reagent is prepared by adding dropwise hexamethylphosphoric triamide (180g, 1 mol, see [680-31-9]) to a vigorously stirred solution of phosgene in toluene (180ml, 20%) during 4 hours (use a bubbler to control the evolution of CO₂). After stirring for 16 hours at ~25°, excess of COCl₂ is removed under reduced pressure, CH₂Cl₂ (300ml) is added to ensure a homogeneous solution, and solid hydroxybenzotriazole monohydrate (153g, see [123333-53-9]) is added. The mixture is cooled to -5° (Me₂CO/Dry-ice) and Et₂NH (100ml, 1mol) is added during 15 minutes, and stirring is continued at 0° for 4 hours. The precipitated Et₂NH.HCl is filtered off (sintered glass funnel), the solvent is evaporated *in vacuo*, H₂O (500ml) is added and the aqueous solution is extracted with Et₂O (3 x 100ml). This aqueous solution of BOP chloride is added to a filtered solution of KPF₆ (200g) in H₂O (2L) and the precipitated BOP.PF₆ is extracted into CH₂Cl₂. The organic layer is dried (MgSO₄), filtered and concentrated under reduced pressure. Addition of Et₂O causes the separation of the first crop of BOP.PF₆ which is collected, and further crops are obtained by addition of CH₂Cl₂ to the mother liquors, concentrating and adding Et₂O. The combined crops are washed with Et₂O and dried *in vacuo* to give the salt (354g, m 147-149°, 80%) which may be recrystallised further from Me₂CO/ Et₂O. The reagent is usually pure enough for efficient coupling of peptides without this recrystallisation. Its IR(KBr) has ν_{\max} at 1010 (P-N), 840, 770, and 560 (PF₆⁻) cm⁻¹; the ¹H NMR (acetone-*d*₆) has δ_{H} at 3.0 (d, 18H, N(CH₃)₂, $J_{\text{H-P}}$ = 10Hz), 7.9 (m, 4H_{arom}); and the ³¹P NMR (CH₂Cl₂/D₃PO₄) has δ at -43.7 (s, P⁺), +144.2 (septet, PF₆⁻). [Castro et al. *Synthesis* 751 1976, DOI: 10.1055/s-1976-24189.]

Benzotriazol-1-yloxy(tripyrrolidino)phosphonium hexafluorophosphate (PyBOP) [128625-52-5] C₁₈H₂₈N₆OP. BF₆, M 520.4, m 154-156°, 156-157°. It can be prepared and purified in a similar manner as for BOP except that pyrrolidine replaced Et₂NH [cf. Castro's reagent above, see [56602-33-6] and Dormoy & Castro *Tetrahedron Lett* 3321 1979, DOI: 10.1016/S0040-4039(01)95397-3] by recrystallisation from Me₂CO/ Et₂O and drying *in vacuo*. It is non-hygroscopic and can be stored at ~25°. It has ³¹P NMR (CDCl₃) at δ : 31.8 (s) and -143.7 (heptet, J = 713Hz). It is an analogue of the BOP and a very good coupling agent which does not form the carcinogenic HMPA as byproduct [Coste et al. *Tetrahedron Lett* **31** 205 1990, DOI: 10.1016/S0040-4039(00)94371-5; Seebach et al. *Helv Chim Acta* **77** 1313 1994, DOI: 10.1002/hlca.19940770513].

(2-Biphenyl)-di-*tert*-butylphosphine [JohnPhos, 2-(di-*tert*-butylphosphino)biphenyl] [224311-51-7] C₆H₄C₆H₄P [C(CH₃)₃]₂, M 298.4, m 86-88°. This *Buchwald ligand* is prepared in Schlenk-type equipment under argon containing Mg turnings (617mg, 25.4mmol) and a small crystal of I₂ (to activate the metal) at ~25°, which is treated with a solution of 2-bromobiphenyl (5.38g, 23.1mol, see [2052-07-5]) in THF (40ml), the mixture is refluxed for 2 hours and allowed to cool to 25°. Anhydrous Cu(I) Cl (2.40g, 24.2mmol) is added, the flask is capped with a septum, purged with argon for 2 minutes and di-*tert*-butylchlorophosphine (5.0g, 24.2mmol; see [13716-10-4]) is injected *via* a syringe, and the mixture is refluxed for 8 hours. The reaction mixture is cooled to ~25°, diluted with 1:1 hexanes/Et₂O (200ml) and the suspension is filtered, the solids are washed with hexanes (60ml), and the solid material is added to 1:1 hexanes/EtOAc (150ml) followed by H₂O (100ml) and 30% aqueous NH₄OH (60ml). This slurry is stirred at ~25° for 5 minutes, the layers are separated and the organic layer is washed with brine (100ml), dried (Na₂SO₄), filtered, evaporated *in vacuo* and the residue is recrystallised from MeOH (2 crops are collected) to provide *JohnPhos* as a white solid (4.46g, 67%) with m 86-86.5°. It has ¹H NMR (300MHz, CDCl₃) with δ_{H} at 7.95-7.85 (m, 1H), 7.40-7.21 (m, 8H), 1.15 (d, J = 11.6Hz, 18H); the ³¹P NMR (121MHz, CDCl₃) has δ_{P} at -18.7; and for IR and ¹³C NMR see references.

This ligand with Pd(OAc)₂ catalyses Suzuki coupling at ~25° between aryl halides (Br and Cl) with arylboronic acids with 0.5-1.0mol%Pd in high yields [Wolfe et al. *J Am Chem Soc* **121** 9550 1999, DOI: 10.1021/ja992130h; see cyclohexyl-JohnPhos below]. The ligand was used for the amination of aryl halides and aryltriflates [Wolfe et al. *J Org Chem* **65** 1158 2000, DOI: 10.1021/jo991699y]. This bulky biarylphosphine ligand was also used in the Pd-catalysed Stille cross-coupling reaction [Artamkina et al. *Synlett* **2** 235 2006, DOI: 10.1055/s-2005-923596], and in the Pd-catalysed 2,3-diarylation of α,α -disubstituted-3-thiophenemethanols *via* cleavage of C-H and C-C bonds [Nakano et al. *J Org Chem* **71** 8309 2006, DOI: 10.1021/jo061412e].

(2-Biphenyl)-dicyclohexylphosphine [cyclohexyl-JohnPhos, 2-(dicyclohexylphosphino)biphenyl]

[247940-06-3] **C₂₄H₃₁P, M 350.5, m 102-106°**. This *Buchwald ligand* is prepared in a way similar to the preceding JohnPhos except that 2-bromobiphenyl in THF is converted to the 2-lithiobiphenyl with *n*-BuLi at -78°/45 minutes, followed by reaction with dicyclohexylchlorophosphine at -78°/15 minutes. The desired product is recrystallised from MeOH to give *cyclohexyl-JohnPhos* (71%) as white crystals **m 103°**. The ¹H NMR (300MHz, CDCl₃) has δ_H at 7.62-7.51 (m, 1H), 7.40-7.10 (m, 8H), 1.95-1.45 (m, 13H), 1.35-0.95 (m, 9H); and the ³¹P NMR (121MHz, CDCl₃) has δ_P at -12.7; and for IR and ¹³C NMR see references.

This ligand with Pd(OAc)₂ allows Suzuki coupling at low catalyst loadings (0.000001-0.02mol%Pd) between aryl halides (Br and Cl) with arylboronic acids in high yields, and tolerates a wide range of functional groups and substrate combinations including sterically hindered substrates. It was the most active catalyst system in terms of temperature of reaction, turnover numbers and steric tolerance in 1999 [Wolfe et al. *J Am Chem Soc* **121** 9550 1999, DOI: 10.1021/ja992130h]. This ligand was also used for the amination of aryl halides and aryltriflates [Ali & Buchwald *J Org Chem* **66** 2560 2001, DOI: 10.1021/jo0008486], and was employed in the Pd-catalysed synthesis of 1,3,5-tris(2'-aminophenyl)-benzene from *o*-aminophenylboronic acid and 1,3,5-triiodobenzene, which may be used as a three-directional core building block for potential ionic receptors [Piatek & Slomiany *Synlett* 2027 2006, DOI: 10.1055/s-2006-948199].

2,2'-Bipyridine (2,2'-dipyridyl, α,α'-dipyridyl) [366-18-7] **C₁₀H₈N₂, M 156.2, m 69.7°, b 272-273°/atm, pK_a 4.50**. The reaction of pyridine with Na produces a mixture of bipyridyls which can be separated by fractional distillation where the 2,2'-isomer distils at 272.5°/atm and solidifies on cooling (**m 69.5°**). The distilled oil can be purified further by dissolving in Et₂O, adding an equal volume of petroleum ether then the 2,3'- and 3,3'- isomers are washed out with several portions of H₂O. The organic layer is evaporated and the oily residue is recrystallised from aqueous EtOH. 2,2'-Bipyridyl also sublimates at 65°/0.01mm. Its solubility in H₂O is 0.5%, but it is very soluble in organic solvents. Its UV spectrum has λ_{max} nm(ε) 233 (10,200) and 280 (13,300) for the neutral species in H₂O. Unlike the other isomers it complexes with metals, e.g. it gives an intense red colour with ferrous salts. The *picrate* has **m 69.7°** (from aqueous EtOH). It is a metalloprotease inhibitor with high affinity for Fe²⁺ containing enzymes at 10⁻⁸ M. [Smith *J Am Chem Soc* **46** 414 1924, DOI: 10.1021/ja01667a016; UV Krumholz *J Am Chem Soc* **73** 3487 1951, DOI: 10.1021/ja01151a146; *Beilstein* **23/8** IV 28, **23/16** V 8.] [TOXIC]

1,3-Bis(1-adamantyl)-1,3-dihydro-2H-imidazol-2-ylidene (IAd) [131042-77-8] **C₂₃H₃₂N₂, M 336.3, m 240-241°**. This is a stable *N*-heterocyclic carbene (NHC) which has been prepared in 96% yield from IAdCl by deprotonation in THF with catalytic amounts of the dismyl anion [CH₃S(O)CH₂⁻] in the presence of 1 equivalent of NaH, or with *tert*-BuOK. In this reaction H₂ is liberated and NaCl is precipitated. The carbene is stable in the absence of oxygen and moisture, and recrystallises from toluene to give clear, colourless rectangular prisms with a sharp melting point that is unaltered by melting and re-solidifying. The ¹H NMR (*C₆H₆-*d*₆) has δ_H at 1.58 (s, Ad₄, 6', 10', 12H), 2.01 (s, Ad₃, 5', 7', 6H), 2.29 (s, Ad₂, 8', 9', 12H), 6.91 (s, 3,4-CH, 2H); and for IR, ¹³C NMR and EI-MS see references. The X-ray crystal structure has been determined and showed a small N-C-N angle at the carbene centre. [Arduengo et al. *J Am Chem Soc* **113** 361 1991, DOI: 10.1021/ja00001a054; see also Arduengo et al. *J Am Chem Soc* 114 5530 1992, DOI: 10.1021/ja00040a007.] It is as effective, if not better in some cases, in many of the metal mediated catalytic reactions as other NHCs (Nitrogen Heterocyclic Carbenes). For further detail see the entry on IPr.Cl [250285-32-6]. [Arduengo USPatent 5 077 414 1991, *Chem Abstr* **116** 106289 1002, Kantchev, O'Brien & Organ *Aldrichimica Acta* **39** 97 2006, Phillips, Chan & Scheidt *Aldrichimica Acta* **42** 55 2009].

1,3-Bis(1-adamantyl)imidazolium tetrafluoroborate (IAd.BF₄) [286014-42-4] **C₂₃H₃₃N₂. BF₄, M 424.3, m 277-282°**. This NHC (N-Heterocyclic Carbene) precursor is prepared by established procedures from 2 mols of amine, 1 mol of glyoxal and one mol of formaldehyde in toluene/H₂O in the presence of HBF₄. It is as effective, if not better in some cases, in many of the metal mediated catalytic reactions as other NHCs. For further detail see the entry on IPr.Cl [250285-32-6]. [Arduengo USPatent 5 077 414 1991, *Chem Abstr* **116** 106289 1002, Kantchev, O'Brien & Organ *Aldrichimica Acta* **39** 97 2006, Phillips, Chan & Scheidt *Aldrichimica Acta* **42** 55 2009].

2,2'-Bis(1,3,2-benzodioxaborole) [bis(catcholato)diboron, Cat-BB-Cat] [13826-27-2] **C₁₂H₈BO₄, M 237.8, m 189-196°, 195-198°**. This borole is prepared by distilling B₂Cl₄ (3.95mmol) [Wartik et al. *J Am Chem Soc* **71**

3265 1949, DOI: 10.1021/ja01177a538; Urry et al. *J Am Chem Soc* **76** 5293 1954, DOI: 10.1021/ja01650a010] and CH_2Cl_2 (30ml) into a reaction vessel containing catechol (8.03mmol) at -196° , and then the reaction is allowed to proceed over the temperature range ~ 78 to 25° for 15 hours to give a clear solution (slightly less than 16mmol of HCl is liberated). CH_2Cl_2 and HCl are distilled off *in vacuo* from the reaction vessel leaving a white residue which is sublimed at $120\text{--}130^\circ/10^{-4}\text{mm}$ to give the **pure borole** in $\sim 70\%$ yield. A similar result is obtained by using $\text{B}_2[\text{NMe}_2]_4$ instead of B_2Cl_4 . Cat-BB-Cat has one peak at -30.7ppm (from BF_3OEt_2 external reference) in the ^{11}B NMR (19MHz, CH_2Cl_2), and one peak for the aromatic protons at 2.69ppm (from solvent reference) in the ^1H NMR (60MHz, CH_2Cl_2); and for IR see references. Molecular weight determination in $^*\text{C}_6\text{H}_6$ indicated that it is a **monomer**, and alkaline hydrolyses provided 0.96 mole of hydrogen per mole of B-B compound. [Welch & Shore *Inorg Chem* **7** 225 1968, DOI: 10.1021/ic50060a011.] Cat-BB-Cat is efficient in olefin diboration catalysed by base-free Pt complexes such as $\text{Pt}(\text{COD})_2$ and $\text{Pt}(\text{norbornene})_2$ [Iverson & Smith *Organometallics* **16** 2757 1997, DOI: 10.1021/om970199x], consequently producing useful synthons for cross-coupling reactions and related conversions to other functional groups [Miyaura & Suzuki *Chem Rev* **95** 2457 1995, DOI: 10.1021/cr00039a007].

Bis[(*R,R,S*)-diazaphos-SPE] {2,2',2'',2'''-(1,2-phenylenebis(1*R*,3*R*)-tetrahydro-5,8-dioxo-1*H*-[1,2,4]-diazaphospholo[1,2-*a*]pyridazine-2,1,3(3*H*)-triyl)tetrakis(*N*-[(1*S*)-1-phenethyl]benzamide} [851609-33-1] $\text{C}_{78}\text{H}_{72}\text{N}_8\text{O}_8\text{P}_2$, **M 1311.40**, **m 183-195°**, $[\alpha]_{\text{D}}^{20} -82$ (c 1, THF), and the diastereomeric **bis[(*S,S,S*)-Diazaphos-SPE] {2,2',2'',2'''-(1,2-phenylenebis(1*S*,3*S*)-tetrahydro-5,8-dioxo-1*H*-[1,2,4]-diazaphospholo[1,2-*a*]pyridazine-2,1,3(3*H*)-triyl)tetrakis(*N*-[(1*S*)-1-phenethyl]benzamide}** [851770-14-4] $\text{C}_{78}\text{H}_{72}\text{N}_8\text{O}_8\text{P}_2$, **M 1311.40**, **m 289-299°**, $[\alpha]_{\text{D}}^{20} +15$ (c 1, THF). The central diazaphospholane tetracarboxylic acid in the two molecules are enantiomeric and these are converted to the phenethylamide using the same *S*-phenethylamine. Thus the two substances are diastereomeric. They are prepared by mixing the diazaphospholane tetracarboxylic acid (0.34mmol) with 5 equivalents of PyBOP {(benzotriazol-1-yloxy)tripyrrolidinophosphonium hexafluorophosphate, m 154-156° cf. [128625-52-5]} under N_2 , adding degassed CH_2Cl_2 (100ml) followed by 5 equivalents of *N,N*-diisopropylethylamine and 5 equivalents of (α *S*)- α -methylbenzylmethanamine (*S*-2-phenylethylamine), and stirring overnight. The solution is exposed to the atmosphere and washed with saturated NaHCO_3 (50ml), 2M HCl (50ml), saturated NaHCO_3 (50ml) again, then H_2O (50ml). The organic layer is dried (MgSO_4), filtered, and evaporated *in vacuo*. The residue is purified by flash chromatography (Al_2O_3 , eluted with 2:1 $\text{CH}_2\text{Cl}_2/\text{EtOAc}$), and the separation of diastereoisomers is accomplished by liquid chromatography using a Zorbax Rx-Sil column (4.6 x 250 column). The ***R,R,S*-diastereomer** (21% yield) has ^1H NMR (500MHz, d-THF) with δ_{H} at 1.37 (d, 6H, $J = 6.7\text{Hz}$, CH_3), 1.56 (d, 6H, $J = 6.7\text{Hz}$, CH_3), 2.2-2.7 (m, 6H, CH_2CH_2), 2.65-3.0 (m, 2H, CH_2CH_2), 5.06 (dq, 2H, $J = 7.5$, 6.9Hz, CHCH_3), 5.41 (dq, 2H, $J = 7.7$, 6.9Hz, CHCH_3), 6.39 (d, 2H, $J = 7.3\text{Hz}$), 6.5-6.9 (m, 10H), 7.06-7.36 (m, 26H), 7.51 (m, 2H), 7.60 (d, 4H, $J = 7.4\text{Hz}$), 7.97 (d, 2H, $J = 8.3\text{Hz}$, NHCH CH_3), 9.23 (d, 2H, $J = 7.8\text{Hz}$, NHCH CH_3); the for ^{13}C NMR see references. The NMR spectra of the ***S,S,S*-diastereomer** is similar but is **not** identical. [Clark et al. *J Am Chem Soc* **127** 5040 2005, DOI: 10.1021/ja050148o; Clark & Landis *J Am Chem Soc* **125** 11792 2003, DOI: 10.1021/ja036359f.]

It is a diazaphospholane ligand which displays high conversion and selectivity in Rh catalysed asymmetric hydroformylation reactions [Axtel et al. *Angew Chem Int Ed* **44** 5834 2005, DOI: 10.1002/anie.200501478; Clark et al. *J Am Chem Soc* **127** 5040 2005, DOI: 10.1021/ja050148o; US Pat 7.071.357B.]

***R,R*(-)- and *S,S*(+)- *N,N'*-Bis(3,5-di-*tert*-butylsalicylidene)-1,2-*trans*-cyclohexanediamine (Jacobsen's ligand)** [*R,R*- 135616-40-9, *S,S*- 135616-36-3] $\text{C}_{36}\text{H}_{54}\text{N}_2\text{O}_2$, **M 546.8**, **m 203-206°**, **205-208°**, **205-208°**, $[\alpha]_{\text{D}}^{20}$ (+) and (-) **310** (c 1, CH_2Cl_2). If this general ligand is to be purified, dissolve it in boiling EtOH and while cooling add H_2O to the yellow solution until a yellow solid separates. This is filtered off, washed with a little of 95% aqueous EtOH and dried *in vacuo*. It is the ligand for preparing Jacobsen's Catalysts [Jacobsen et al. *J Am Chem Soc* **113** 7063 1991, DOI: 10.1021/ja00018a068; Deng & Jacobsen *J Org Chem* **57** 4320 1992, DOI: 10.1021/jo00041a054].

On a large scale, the *R,R*-ligand ($\sim 60\text{g}$) is dissolved in CH_2Cl_2 (500ml), washed with H_2O (2 x 300ml) and brine (100ml), dried (Na_2SO_4), and the solvent is removed to yield a yellow powder with **m 200-203°**, $[\alpha]_{\text{D}}^{20}$ (+) and (-) **315** (c 1, CH_2Cl_2), the required ^1H and ^{13}C NMR, and IR with ν_{max} (KBr) at 1595, 1631, 2869, 2960 cm^{-1} . However, if it is felt that the product is of insufficient purity then it should be crystallised in two crops from Me_2CO (1:20 w/v with typical 86-93% recovery) [Larrow et al. *J Org Chem* **59** 1939 1994, DOI: 10.1021/

jo00086a062.]

A **dimeric Jacobsen ligand**, formed by joining two units with a methylene bridge has been prepared and crystallised from CH_2Cl_2 /pentane. Its **bis-Mn catalyst** exhibited improved retention in a poly-dimethylsiloxane membrane for the asymmetric epoxidation of olefins [Janssen et al. *Tetrahedron Asymmetry* **8** 3481 1997, DOI: 10.1016/S0957-4166(97)00465-5].

***1,5-Bis(3',5'-dimethoxyphenyl)penta-1E,4E-dien-3-one (dm-dba)** [39777-58-7] $\text{C}_{21}\text{H}_{22}\text{O}_5$, **M 354.1, m 132-134°**. To a cooled (ice-water bath) solution of NaOH (1.8g, 2.5equivalents) in H_2O (18ml) diluted with EtOH (15ml) is added slowly 3',5'-dimethoxybenzaldehyde (3g, 1equivalent) and analytical grade acetone (0.52g, 0.5 equivalent) during 15 minutes, then it is stirred at 25° for 1 hour. A yellow solid separates, and after a further hour it is filtered off, washed with Et_2O (3 x 50ml) and dried *in vacuo*. Purification by flash chromatography (*ca* 150 mesh Al_2O_3 deactivated with 6% v/w H_2O prior to use) and elution with petroleum ether (b 40°-60°)/EtOAc (4/1 v/v) gives **dm-dba** as a yellow solid (2.2g, 69%) which is recrystallised by layering a concentrated solution of CD_2Cl_2 with Et_2O ($\text{CD}_2\text{Cl}_2/\text{Et}_2\text{O}$, 1/3). The IR (CH_2Cl_2) has ν_{max} at 1652m (C=O), 1621vs (C=C), 1596w (C=C aromatic), 1572w (C=C aromatic) and 989m (CH trans) cm^{-1} ; the UV has λ_{max} (THF) at 239 (π - π^*) and 371 (n- π^*) nm; the ^1H NMR (400MHz, CDCl_3) has δ_{H} at 6.48 (d, 1H, $^3J = 15.7\text{Hz}$, H-2), 7.01 (1H, d, $^3J = 15.7\text{Hz}$, H-1), 6.71 (2H, t, $^3J = 6.7\text{Hz}$, 7.9Hz, H-6'), 7.62 (1H, d, $^3J = 8.2\text{Hz}$, H-4'), 3.70 (s, 12H); and for the ^{13}C NMR see references. [Fairlamb et al. *Org Lett* **6** 4435 2004, DOI: 10.1021/ol048413i; Beilstein **7** IV 1747.]

2,2'-Bis(diphenylphosphino)-1,1'-binaphthyl (BINAP) [RS 98327-87-8] $[(\text{C}_6\text{H}_5)_2\text{PC}_{10}\text{H}_6-]_2$, **M 622.7, m 283-286°**, [R-(+)-76189-55-4] **m 241-242°**, [S-(-)-76189-56-5] **m 241-242°**, $[\alpha]_{\text{D}}^{20}$ **R (+) and S (-) 233 (c 0.3, toluene)**. It has been prepared from (\pm)-BINOL which is converted to (\pm)-2,2'-dibromo-1,1'-binaphthyl (with Ph_3PBr_2) then to (\pm)-2,2'-bis(diphenylphosphinyl)-1,1'-binaphthyl (**BINAPO**) [with $\text{Mg}/\text{Ph}_2\text{P}(\text{O})\text{Cl}$] which is resolved into **S-(-)- and R-(+)- BINAPO** *via* separation of the diastereoisomeric (-)-2,3-di-O-benzoyl-L-tartrate salts in $\text{CHCl}_3/\text{EtOAc}$ followed by treatment with 0.75N NaOH, extraction into CHCl_3 , and isolation. The chiral BINAPO enantiomers are reduced to the respective **S-(-)- and R-(+)- BINAP** with excess of Cl_3SiH [see 10025-78-1] and Et_3N in toluene (100-120°) followed by treatment with 30% aqueous NaOH, extraction into toluene, isolation and purification as follows. Dissolve the enantiomer in toluene, wash it with 30% aqueous NaOH, three times with H_2O , dry (Na_2SO_4), evaporate to ~15% of its volume and add an equal volume of degassed MeOH. Collect the solid, wash it with MeOH and dry it at 80°/0.005mm for 6 hours. Recrystallise it from a 1:1 mixture of toluene/EtOH to **optical purity (m 241-242°)**. **Pure (99%) (S-(-)-BINAP** has ^1H NMR (300MHz, CDCl_3 , TMS) with δ_{H} at 6.81 (d), 6.90 (t), 7.10 (m), 7.23 (s), 7.33 (t), 7.43 (dd), 7.82 (d) and 7.89 (d) where multiplicities are not necessarily coupled signals; and for the ^{13}C NMR see reference. The purity is determined by GLC analysis using an OV-101 5m capillary column at 200-280° (argon carrier gas ?). TLC analysis (Kieselgel 60 PF₂₅₄, with MeOH/ CHCl_3 1:19) is used to identify BINAP (R_F 0.83), BINAPO (R_F 0.42) and BINAP-monooxide (R_F 0.67). (\pm)-**BINAPO** [86632-33-9] crystallises from hot toluene (solubility is 3.6g/100ml at 110°) or from hexane/toluene, and analytically pure crystals have **m 304-306°** (299-300° also reported). **S-(-)-BINAPO** [94041-18-6] crystallises from a hexane/toluene mixture and has **m 261-262°**, $[\alpha]_{\text{D}}^{24}$ **-396 (c 0.47, C_6H_6), $[\alpha]_{\text{D}}^{24}$ -168 (c 0.5, EtOH)**; and **R-(+)-BINAPO** [94041-16-4], similarly recrystallised, has **m 262-263°**, $[\alpha]_{\text{D}}^{24}$ **+396 (c 0.5, C_6H_6)**. [Takaya et al. *Org Synth Coll Vol* **8** 57 1993, DOI: 10.15227/orgsyn.067.0020]. [Noyori & Takaya *Acc Chem Res* **23** 345 1990, DOI: 10.1021/ar00178a005; Kitamura et al. *Org Synth* **71** 1 1993, DOI: 10.15227/orgsyn.071.0001; Takaya et al. *Org Synth* **72** 74 1995, DOI: 10.15227/orgsyn.072.0074; Kitamura et al. *J Org Chem* **57** 4053 1992, DOI: 10.1021/jo00040a068.]

§ A polymer supported version of BINAP is available.

2R,3R-(+)-2,3-Bis(diphenylphosphino)butane (R,R-CHIRAPHOS) [74839-84-2], **2S,3S-(-)-2,3-bis(diphenylphosphino)butane (S,S-CHIRAPHOS)** [64896-28-2] $\text{C}_{28}\text{H}_{28}\text{P}_2$, **M 426.5, m 108-109°**, **R- $[\alpha]_{\text{D}}^{20}$ (+) and S- $[\alpha]_{\text{D}}^{20}$ (-) 200 (c 1.5, CHCl_3)**. For **S,S-CHIRAPHOS**: Ph_2PLi [prepared *in situ* from Ph_3P (95g) and Li (5g) in THF (300ml) under N_2 at ~55°/1 hour then at 25°/2 hours, and the PhLi formed is decomposed with *tert*-BuCl (33g) for 45 minutes; the clear orange solution is boiled for 5 minutes then cooled to -4°] solution is treated with **(+)-(2R,3R)-butanediol bis(tosylate)** (35g, [64896-27-1]) in dry THF (100ml) over 1 hour with stirring, the temperature is allowed to rise to 25° and stirring is continued for 30 minutes. N_2 purged H_2O (100ml) is added, the THF layer is separated and evaporated *in vacuo* to give impure **S,S-chiraphos (note**

inversion of configuration) as a colourless oil. The oil is extracted under N_2 with Et_2O (2 x 150ml), dried (Na_2SO_4), and filtered (under N_2) into a solution of $Ni(ClO_4)_2 \cdot 6H_2O$ (15g) in $EtOH$. The Na_2SO_4 is washed with Et_2O which is added to the Ni solution. The oily red-brown deposit of the *Ni-chiraphos* sometimes contains some yellow solid of the *Ni-perchlorate complex* which is discarded (*pyrophoric?*). To the brown-oily (or part crystalline) mixture is added $NaCNS$ (15g) in hot $EtOH$ and stirred vigorously until a homogenous yellow-brown crystalline $[Ni(S,S\text{-chiraphos})_2NCS]NCS$ is obtained. It is collected, washed thoroughly with $EtOH$ then finally Et_2O .

This *Ni-NCS complex* (15g), suspended in 95% $EtOH$ (150ml) under N_2 is stirred, brought to the boil, and $NaCN$ (4g) in H_2O (20ml) is added rapidly when the Ni complex slowly dissolves to give a clear blood-red solution which then turns cloudy beige in colour. The hot solution is stirred until all the complex which had dissolved is converted to a yellow slurry. After cooling in ice-water, the solid is collected, washed with H_2O (2 x 25ml) then rapidly with ice-cold $EtOH$ (25ml) to give impure beige coloured *S,S-chiraphos*. This solid is purified by drying at 25° , dissolving in boiling absolute $EtOH$ (~125ml), filtering through a frit under N_2 and allowing to stand at $\sim 25^\circ$ for 12 hours, whereby *S,S-chiraphos* deposits as lustrous colourless plates. A second crystallisation from absolute $EtOH$ (60ml) provides *optically pure (-)-(2S,3S)-bis(diphenylphosphino)butane (S,S-CHIRAPHOS)* (5.5g), **m 108-109°** (sealed tube under N_2) as colourless plates with $[\alpha]_D^{27} -211$ (c 1.5, $CHCl_3$), which is unchanged on further recrystallisation [Fryzuk & Bosnich *J Am Chem Soc* **99** 6262 1977, DOI: 10.1021/ja00461a014]. Its ^{31}P NMR ($CDCl_3$, with external H_3PO_4 as reference) has δ at -10.7, i.e. upfield from H_3PO_4 [Slack et al. *Inorg Chem* **18** 3125 1979, DOI: 10.1021/ic50201a034].

For R,R-CHIRAPHOS: This enantiomeric ligand is obtained essentially by the same synthesis as its enantiomer above. The *(-)-(2S,3S)-butanediol bis(tosylate) [74839-83-1]* is prepared in 94% yield, and has **m 63-64°**, $[\alpha]_D^{25} -36.8$ (c 2.1, $CHCl_3$), and 1H NMR ($CDCl_3$) with δ at 1.20 (d, $J = 6Hz$; 6H, CH_3), 2.50 (s; 6H, tosyl- CH_3), 4.70 (q, $J \sim 7Hz$; 2H, CH), 7.70 (m; 8H, aromatic-H). It is converted into **bis[(2R,3R)-2,3-bis(diphenylphosphino)butan]thiocyanato-nickel(II) thiocyanate** in 56% yield (*note* inversion of configuration) after washing with $EtOH$ and Et_2O , and decomposed with $NaCN$ to give *R,R-CHIRAPHOS* in 32% yield with **m 106-107°**, $[\alpha]_D^{27} +197$ (c 1.5, $CHCl_3$), and 1H NMR ($CDCl_3$) with δ at 1.10 (d, $J = 7Hz$; 3H, CH_3), 1.40 (d, $J = 6Hz$; 6H, CH_3), 2.60 (q, $J \sim 7Hz$; 2H, CH), 7.50 (m; 20H, aromatic-H). [Köttner & Gerber *Chem Ber* **113** 2323 1980, DOI: 10.1002/cber.19801130627.]

1,4-Bis-(diphenylphosphino)butane (dppb) [7688-25-7] (C_6H_5)₂P(CH₂)₄P(C_6H_5)₂, **M 426.5, m 135-136°, 136-137°**. Recrystallise it from $EtOH$ [Trippett *J Chem Soc* 4263 1961, DOI: 10.1039/JR9610004247]. [King *J Coord Chem* **1** 67 1971, DOI: 10.1080/00958977108070745; Tolman *Chem Rev* **77** 313 1977, DOI: 10.1021/cr60307a002.]

1,2-Bis-(diphenylphosphino)ethane (DIPHOS, ethylene bis(diphenylphosphine)) [1663-45-2] (C_6H_5)₂P(CH₂)₂P(C_6H_5)₂, **M 398.4, m 139-140°, 140-142°, 143-144°, pK_{Est} ~4.5**. Recrystallise it from aqueous $EtOH$ or $*C_6H_6$. The *dimethiodide*, when recrystallised from $MeOH$ has **m 305-307°**, and the *dioxide* when recrystallised from toluene or DMF (needles), or $*C_6H_6$ (plates) has **m 252-254° (276-278°)** [Isslieb et al. *Chem Ber* **92** 3175 1959, DOI: 10.1002/cber.19590921221; NMR: Aguiar & Beiser *J Org Chem* **29** 1660 1964, DOI: 10.1021/jo01029a524; Baekvall et al. *J Org Chem* **52** 5430 1987, DOI: 10.1021/jo00233a023]. [Beilstein **16** IV 958.]

R-(+)-1,2-Bis(diphenylphosphino)propane (R-PROPHOS) [67884-32-6] (C_6H_5)₂PCH(CH₃)CH₂—P(C_6H_5)₂, **M 412.4, m 68.5° (sealed tube under N_2), 71-73°, $[\alpha]_D^{26} R +186.0$ (c 1.0, Me_3CO)**. *S-(-)-Propane-1,2-diol bis(p-toluenesulfonate)* (51.9g, [60434-71-1]) in dry THF (75ml) is reacted with Ph_2PLi to give the *R-diol* (*note* change in absolute configuration) which is converted to its *Ni-perchlorate* salt and then converted into its *NiNCS* complex (45-55g) essentially as described for *chiraphos* above. This *NCS* complex (22g) is decomposed with $NaCN$ essentially as described for *chiraphos* above, and finally the crude oily *R-prophos* is dissolved in absolute $EtOH$ at 50° under N_2 , allowed to cool to 25° , then held at 5° for 24 hours and the *diphosphine* (10g) that separated is collected and recrystallised from absolute $EtOH$ (100ml) to give small colourless prisms of *analytically pure R-prophos* (7.5g) whose optical rotation is unchanged by further recrystallisation. [Fryzuk & Bosnich *J Am Chem Soc* **101** 3043 1979, DOI: 10.1021/ja00505a035]. Its ^{31}P NMR ($CDCl_3$, with external H_3PO_4 as reference) has δ at -20.6 (d, $J_{P-P} = 20.6Hz$), i.e. upfield, and +1.7 ppm, i.e.

downfield from H_3PO_4 [Slack et al. *Inorg Chem* **18** 3125 1979, DOI: 10.1021/ic50201a034]. Pure *S-prophos* may be prepared in a similar way.

Bis(2-hydroxyethyl)sulfide (2,2'-thiodiglycol, thiodiglycol) [111-48-8] $(\text{HOCH}_2\text{CH}_2)_2\text{S}$, M 112.2, m -16° , b $121\text{--}121.8^\circ/0.01\text{mm}$, $130^\circ/2\text{mm}$, $136\text{--}137^\circ/3\text{mm}$, $148^\circ/4\text{mm}$, $165^\circ/20\text{mm}$, $168^\circ/20\text{mm}$, $185.5^\circ/40\text{mm}$, $194^\circ/50\text{mm}$, $282^\circ/760\text{mm}$, $d_4^{25} 1.1973$, $d_4^{25} 1.1793$, $n_D^{20} 1.5203$, $n_D^{26} 1.5146$. The sulfide has been prepared on large scales for the manufacture of 'Mustard Gas'. The thioglycol has been prepared from ethylene oxide and H_2S , and the crude compound prepared from chlorohydrin contains considerable quantities of dithiane and polymeric impurities. These can be removed by distillation at $150^\circ/8\text{mm}$ and the polymeric material breaks down at $\sim 160^\circ$ [Masson *J Chem Soc* **49** 233 1886, DOI: 10.1039/CT8864900233]. The distillate is then diluted with H_2O until its boiling point at atmospheric pressure is reduced to 165° , and superheated steam is passed through it. After evaporating off the H_2O , the thiodiglycol is distilled at $147^\circ/6\text{mm}$. It is soluble in H_2O , lower alcohols, CHCl_3 , EtOAc ; and at 25° its solubility (w/w) in $^*\text{C}_6\text{H}_6$ is 1.07%, in absolute Et_2O it is 7.09 and in ligroin it is 0.06%. In organic acids, or alone, it is stable at 180° for many hours, but when heated at 100° with 2.5 parts of 0.1N aqueous NaOH for 30 minutes sulfide ions are formed, much more so (50%) with 1N NaOH at $140^\circ/10$ hours. $\text{Pb}(\text{OAc})_2$, and $\text{Cu}(\text{NO}_3)_2$ decompose it at 100° , but it is stable with BaO , CaO and Al_2O_3 even at $180^\circ/10$ hours, in unsuccessful attempts to dehydrate it to vinyl sulfide. Of the several esters of aliphatic alcohols that were reported, the *diacetate* had **b $139.5^\circ/8\text{mm}$** , the *dibutyrate* had **b $172^\circ/8\text{mm}$** , and the *dicaproate* had **b $207^\circ/7\text{mm}$** [Clayton & Reid *J Am Chem Soc* **64** 908 1942, DOI: 10.1021/ja01256a048]. The *bis-3,4-diphenylcarbamoyl derivative* had **m $141.4\text{--}142.5^\circ$** (from EtOH) [Beaver et al. *J Am Chem Soc* **79** 1236 1957, DOI: 10.1021/ja01562a053], and the *bis-4-nitrobenzoate ester* had **m 107.7°** (from EtOH) [Major *Bull Soc Chim Fr* **41** 634 1927]. [Nenitzescu & Scărlătescu *Chem Ber* **68** 587 1935, DOI: 10.1002/cber.19350680406; *Beilstein* **1 H** 470, **1 I** 244, **1 II** 525, **1 III** 2122, **1 IV** 2437.]

1,2-Bis[(4*S*)-4-isopropyl-2-oxazolin-2-yl]benzene [(4*S*,4'*S*)-2,2'-(phen-1,2-diyl)bis(4-isopropyl-4,5-dihydro-oxazole)] [131380-80-8] $\text{C}_{18}\text{H}_{24}\text{N}_2\text{O}_2$, M 300.4, b $175^\circ/10^{-2}$ mbar, $[\alpha]_{589}^{25} -98.8$ (c 2.13, CH_2Cl_2), $[\alpha]_{546}^{25} -119.4$ (c 2.13, CH_2Cl_2), $[\alpha]_{365}^{25} -415.2$ (c 2.13, CH_2Cl_2). This ligand is prepared by melting a small amount of anhydrous ZnCl_2 (68mg, 0.50mmol, m 293°) in high vacuum under argon, cooling to $\sim 25^\circ$, adding chlorobenzene followed by *o*-phthalonitrile (1.28g, 10mmol) and *S*-valinol (3.09g, 30mmol), and boiling under reflux for 24 hours. The cooled solution is extracted with H_2O (2 x 20ml), and the aqueous phase is extracted with CH_2Cl_2 (30ml). The combined organic extracts are dried (Na_2SO_4), filtered, evaporated *in vacuo*, and the residual oil is purified by flash chromatography and MPLC (pentane/ EtOAc 4:1); and distilled at high vacuum to provide optically (by NMR and chiral HPLC on CHIRACEL OD) and analytically pure (C, H and N) colourless *bis-oxazoline* (2.68g, 89%). It has IR(film) with ν_{max} at 1655 (C=N) and 1250 (C-O) cm^{-1} ; the ^1H NMR (300MHz, CDCl_3) has δ at 0.95 [d, $J = 6.8\text{Hz}$, 6H, $\text{CH}(\text{CH}_3)_2$], 1.04 [d, $J = 6.8\text{Hz}$, 6H, $\text{CH}(\text{CH}_3)_2$], 1.88 [sept, $J = 6.8\text{Hz}$, 2H, $\text{CH}(\text{CH}_3)_2$], 4.04–4.13 (m, 4H, OCH_2), 4.34–4.42 (m, 2H, CHN), 7.45–7.50 (m, 2H, 4- and 5-H), 7.74–7.77 (m, 2H, 3- and 6-H) from TMS; and for ^{13}C NMR see reference. [Bolm et al. *Chem Ber* **124** 1173 1991, DOI: 10.1002/cber.19911240532.] Like many chiral bis(oxazolin-2-yl) ligands it complexes with metals such as Zn (see its crystalline Zn complex [131380-93-3] in Part 1) and Cu, being involved in a variety of metal catalysed asymmetric synthesis such as allylation, aziridination, cyclopropanation, Diels-Alder and retro Diels-Alder, Mukaiyama aldol condensation and hydrosilylation [see reviews by Ghosh et al. *Tetrahedron: Asymmetry* **9** 1 1998, DOI: 10.1016/S0957-4166(97)00593-4; Pflatz *Acta Chem Scand* **50** 189 1996, DOI: 10.3891/acta.chem.scand.50-0189; Johnson & Evans *Acc Chem Res* **33** 325 2000, DOI: 10.1021/ar960062n; Jørgensen et al. *Acc Chem Res* **32** 605 1999, DOI: 10.1021/ar970347f.]

1,2-Bis[(4*S*)-4-isopropyl-2-oxazolin-2-yl]ethane [(4*S*,4'*S*)-2,2'-(ethane-1,2-diyl)bis(4-isopropyl-4,5-dihydro-oxazole)] [131380-80-8] $\text{C}_{14}\text{H}_{24}\text{N}_2\text{O}_2$, M 252.4, b $150^\circ/2.10^{-2}$ mbar, $[\alpha]_{589}^{25} -93.6$ (c 2.28, CH_2Cl_2), $[\alpha]_{546}^{25} -112.1$ (c 2.28, CH_2Cl_2), $[\alpha]_{365}^{25} -335.5$ (c 2.28, CH_2Cl_2). This colourless bis-oxazolinylethane (1.49g, 59% yield) was prepared by the general method described in the preceding entry from anhydrous ZnCl_2 (68mg, 0.50mmol), 1,2-dicyanoethane (0.80g, 10mmol), and *S*-valinol (3.09g, 30mmol) in optical and analytical purity. It has IR(film) with ν_{max} at 1645 (C=N) cm^{-1} ; the ^1H NMR (300MHz, CDCl_3 , TMS) has δ at 0.87 [d, $J = 6.7\text{Hz}$, 6H, $\text{CH}(\text{CH}_3)_2$], 0.95 [d, $J = 6.7\text{Hz}$, 6H, $\text{CH}(\text{CH}_3)_2$], 1.74 [sept, $J = 6.7\text{Hz}$, 2H, $\text{CH}(\text{CH}_3)_2$], 2.63 [s, 4H, CH_2], 3.84–3.98 (m, 4H, OCH_2), 4.22 (dd, $J = 8.5, 8.5\text{Hz}$, 2H, CHN); and for ^{13}C NMR see Bolm et al. [Bolm et al. *Chem Ber* **124** 1173 1991, DOI: 10.1002/cber.19911240532.] Like many chiral bis(oxazolin-2-yl) ligands it

complexes with metals such as Zn (see the crystalline Zn complex in Part 1) and Cu, being involved in a variety of metal catalysed asymmetric synthesis such as allylation, aziridination, cyclopropanation, Diels-Alder and retro Diels-Alder, Mukaiyama aldol condensation and hydrosilylation [see reviews by Ghosh et al. *Tetrahedron: Asymmetry* **9** 1 1998, DOI: 10.1016/S0957-4166(97)00593-4; Pflatz *Acta Chem Scand* **50** 189 1996, DOI: 10.3891/acta.chem.scand.50-0189; Johnson & Evans *Acc Chem Res* **33** 325 2000, DOI: 10.1021/ar960062n; Jørgensen et al. *Acc Chem Res* **32** 605 1999, DOI: 10.1021/ar970347f.]

1,3-Bis[(4*S*)-4-isopropyl-2-oxazolin-2-yl]propane [(4*S*,4'*S*)-2,2'-(propane-1,3-diyl)bis(4-isopropyl-4,5-dihydrooxazole)] [*131380-90-0*] $C_{15}H_{26}N_2O_2$, **M 266.4**, **b 120°/2.10⁻² mbar**, $[\alpha]_{D}^{25}$ **-57.4** (c 3.05, CH₂Cl₂), $[\alpha]_{D}^{25}$ **-68.4** (c 3.05, CH₂Cl₂), $[\alpha]_{D}^{25}$ **-204.0** (c 3.05, CH₂Cl₂). This colourless bis-oxazolinylpropane (1.12g, 42% yield) was prepared by the general method described in the preceding entries from anhydrous ZnCl₂ (68mg, 0.50mmol), 1,3-dicyanopropane (0.94g, 10mmol), and *S*-valinol (3.09g, 30mmol) in optical and analytical purity. It has IR(film) with ν_{max} at 1645 (C=N) cm⁻¹; the ¹H NMR (300MHz, CDCl₃) has δ at 0.88 [d, *J* = 6.7Hz, 6H, CH(CH₃)₂], 0.95 [d, *J* = 6.7Hz, 6H, CH(CH₃)₂], 1.74 [sept, *J* = 6.7Hz, 2H, CH(CH₃)₂], 2.33-2.61 [m, 6H, CH₂]₃, 3.85-3.98 (m, 4H, OCH₂CHN), 4.18-4.26 (m, 2H, NCHCH₂O) ppm from TMS; and for ¹³C NMR see Bolm et al. [Bolm et al. *Chem Ber* **124** 1173 1991, DOI: 10.1002/cber.19911240532.] Like many chiral bis(oxazolin-2-yl) ligands it complexes with metals such as Zn (see the crystalline Zn complex in the 'Catalyst' section Part 1) and Cu, being involved in a variety of metal catalysed asymmetric synthesis such as allylation, aziridination, cyclopropanation, Diels-Alder and retro Diels-Alder, Mukaiyama aldol condensation and hydrosilylation [see reviews by Ghosh et al. *Tetrahedron: Asymmetry* **9** 1 1998, DOI: 10.1016/S0957-4166(97)00593-4; Pflatz *Acta Chem Scand* **50** 189 1996, DOI: 10.3891/acta.chem.scand.50-0189; Johnson & Evans *Acc Chem Res* **33** 325 2000, DOI: 10.1021/ar960062n; Jørgensen et al. *Acc Chem Res* **32** 605 1999, DOI: 10.1021/ar970347f.]

2,2-Bis[(4*S*)-4-isopropyl-2-oxazolin-2-yl]propane {4,5-dihydro-2-[2(4,5-dihydro-(4*S*)-4-isopropoxyloxazol-2-yl)propan-2-yl]-(4*S*)-4-isopropoxyloxazole, (4*S*,4'*S*)-2,2'-(propane-2,2-diyl)bis(4-isopropyl-4,5-dihydro-oxazole)} [*relative stereochemistry* 929270-13-3; *absolute stereochemistry* 797757-81-4] $C_{15}H_{26}N_2O_2$, **M 266.4**, **b 95-100°/0.5 mm**, d_4^{25} **0.9864**, n_D^{20} **1.4665**, $[\alpha]_D^{24}$ **-107.5** (c 1, CH₂Cl₂). The propane ligand for metal assisted asymmetric catalysis is prepared in two steps. The *first* step is the synthesis of the bis-oxazoline precursor (-)-(*S,S*)-*N,N'*-bis(1-hydroxymethyl-2-methylpropyl)-2,2-dimethylmalonamide which is prepared in a flask under nitrogen purge at 0° (ice bath) containing a mixture of *S*-valinol (5.13g, 50mmol) and dry EtNH₂ (17.4ml, 124mmol, distilled from CaH₂) to which 2,2-dimethylmalondioyl chloride (3.3ml, 250mmol; Evans et al. *J Org Chem* **63** 4541 1998, DOI: 10.1021/jo980296f) is added dropwise during 25 minutes (temperature rising from 0° to 10°), then allowing to warm to ~25°. After stirring for 45 minutes the colourless precipitate in the mixture is dissolved by adding CH₂Cl₂ (120ml). Aqueous N HCl (30ml) is then added, the mixture is extracted with CH₂Cl₂ (3 x 13ml), the combined organic layers are washed with saturated aqueous NaHCO₃ (30ml), brine (30ml), dried (MgSO₄), filtered and evaporated *in vacuo* to provide a yellow solid which is crystallised from EtOAc (~40ml) to give the *malonamide* (6.4g, 84% in three crops). The amide has **m 98-99°**, $[\alpha]_D^{24}$ **-6.0** (c 0.5, CH₂Cl₂); the ¹H NMR (300MHz, CDCl₃) has δ at 0.92 (d, *J* = 6.8Hz, 6H), 0.96 (d, *J* = 6.8Hz, 6H), 1.50 (s, 6H), 1.82 (oct, *J* = 6.8Hz, 2H), 2.66 (br s, 2H), 3.52 (m, 2H), 3.69-2.86 (m, 4H), 6.41 (d, *J* = 8.6Hz, 2H); correct elemental analysis for C, H and N, and for IR and ¹³C NMR see references.

The *second* step is carried out, with stirring, under N₂ purge in a flask containing the preceding malondiamide (5.5g, 18.4mmol), 4-dimethylaminopyridine (204mg, 1.67mmol) in dry CH₂Cl₂ (130ml, filtered through activated Al₂O₃) at ~25°, and dry EtNH₂ (10.25ml, 73.4mmol, distilled from CaH₂) is added slowly (*via* syringe) followed by tosyl chloride (7.10g, 37mmol, 2 equivalents) dissolved in dry CH₂Cl₂ (15ml) dropwise during 30 minutes *via* a funnel which is rinsed with CH₂Cl₂ (2.5ml), and the mixture is stirred for 27 hours at ~25°. The mixture is then treated with saturated aqueous NH₄Cl (70ml), H₂O (40ml), the aqueous layer is separated, extracted with CH₂Cl₂ (3 x 55ml), the combined organic layers are dried (MgSO₄), filtered and evaporated. The oily residue is mixed with hot pentane (40ml), stirred for 5 minutes, the supernatant liquid is decanted, and the procedure repeated three times. The pentane layers are combined and evaporated *in vacuo* and the oily residue (4.05g 83%) is distilled (Kügelrohr) to give the *analytically pure bis-oxazole*; its ¹H NMR (300MHz, CDCl₃, TMS) has δ at 0.95 (d, *J* = 6.8Hz, 6H), 0.91 (d, *J* = 6.8Hz, 6H), 1.51 (s, 6H), 1.88-1.73 (m, 2H), 4.06-3.93 (m, 4H), 4.26-4.15 (m, 2H); and for IR and ¹³C NMR see Evans et al. [*Org Synth* **83** 97 2006, DOI: 10.15227/orgsyn.083.0097]. Like many chiral bis(oxazolin-2-yl) ligands it complexes with metals such as

Zn (see the crystalline Zn complex [131380-93-3] in the ‘Catalysts’ section, Part 1) Cu, Ir, Pd, W, being involved (with or without further ligands) in a variety of metal catalysed asymmetric synthesis such as allylation, aziridination, cyclopropanation, Diels-Alder and retro Diels-Alder, Mukaiyama aldol condensation and hydrosilylation [see reviews by Ghosh et al. *Tetrahedron: Asymmetry* **9** 1 1998, DOI: 10.1016/S0957-4166(97)00593-4; Pflatz *Acta Chem Scand* **50** 189 1996, DOI: 10.3891/acta.chem.scand.50-0189; Johnson & Evans *Acc Chem Res* **33** 325 2000, DOI: 10.1021/ar960062n; Jørgensen et al. *Acc Chem Res* **32** 605 1999, DOI: 10.1021/ar970347f.]

1,3-Bis(2,6-isopropylphenyl)imidazolium chloride (IPr.Cl) [250285-32-6] $C_{27}H_{37}N_2^+ Cl^-$, **M 425.1, m 278°(dec)**. IPr.Cl is a ligand precursor of an NHC (N-heterocyclic carbene— Kantchev, O’Brien & Organ *Aldrichimica Acta* **39** 97 2006, Phillips, Chan & Scheidt *Aldrichimica Acta* **42** 55 2009) which coordinates with transition metals to form soluble catalysts that promote a variety of reactions. IPr.Cl can be prepared in two steps. *1,4-Bis(2,6-diisopropylphenyl)diazabutadiene* [1,2-bis(2,6-diisopropylphenylimino)-ethane] is first prepared by dissolving 2,6-diisopropylaniline (100g, 560mmol) and glyoxal (31.5ml, 280mmol, in 40% H₂O) in absolute EtOH (500ml), and adding a few drops of formic acid as catalyst. The yellow coloured solution produces a yellow precipitate after a few hours, the mixture is stirred for 2 days, and the yellow solid is collected, washed with cold MeOH, and dried *in vacuo* to give **analytically pure diazabutadiene-ethane** (81.7g, 77.5%). It has ¹H NMR (400MHz, CDCl₃) with δ at 1.28 (d, *J* = 7.6 Hz, 24H, CH(CH₃)₂), 3.03 (sep, *J* = 6.4Hz, 4H, CH(CH₃)₂), 7.27 (m, 6H, CH(CH₃)₂-C₆H₃), 8.19 (s, 2H, NCH). In the second step the **diazabutadiene-ethane** (25g, 66mmol) in toluene (500ml) is treated with solid paraformaldehyde (2.0g, 66mmol) with stirring under N₂, heated to 100° until clear, then cooled to 40° and HCl (16.5ml, 66mmol, 4M in dioxane) is introduced with a syringe. The colour of the mixture turns to brown and a white precipitate separates within a few hours, but stirring is continued at ~25° for 36 hours. The solid is filtered off, washed with THF, and dried *in vacuo* to give off-white IPr.Cl (13.1g, 47%). It has ¹H NMR (400MHz, CD₂Cl₂) with δ at 1.24 (d, *J* = 7.2 Hz, 12H, CH(CH₃)₂), 1.27 (d, *J* = 7.2 Hz, 12H, CH(CH₃)₂), 2.42 (sep, *J* = 6.8Hz, 4H, CH(CH₃)₂), 7.18 (t, *J* = 7.2Hz, 2H, *p*-C₆H₃), 7.4 (m, 4H, *m*-C₆H₃), 7.80 (s, 2H, NCH), 11.0 (s, 1H, NC(HCl)); and its ¹³C NMR (100MHz, CD₂Cl₂) has δ_C{H} at 23.9 (CH(CH₃)₂), 26.1 (CH(CH₃)₂), 86.4 (NCHCHN), 125.2 (q C, phenyl), 125.8 (CH, phenyl), 128.7 (quaternary C, phenyl), 129.5 (CH, phenyl), 145.7 (CHCl).

IPr.Cl has been used, in dioxane under argon, as a ligand with Pd₂(dba)₃ (see above) to catalyse cross-coupling reactions between arylhalides and arylmagnesium bromides in a Kumada reaction to form diaryls efficiently [Huang & Nolan *J Am Chem Soc* **121** 9889 1999, DOI: 10.1021/ja991703n]; and in the presence of a base (*tert*-BuOK, or Cs₂CO₃) and Pd(OAc)₂ {or PdCl₂, [Pd(allyl)Cl]₂, PdCl₂(PCy₃)₂} and CO under pressure it catalyses the carbonylative cross-coupling of bromopyridines with arylboronic acids to provide high yields of arylpyridylketones [Maerten et al. *Tetrahedron* **63** 682 2007, DOI: 10.1016/j.tet.2006.11.008].

When IPr.Cl is treated with a base (e.g. Bu^tOK, Bu^tONa, K₂CO₃ or Cs₂CO₃), the carbene **1,3-bis(2,6-diisopropylphenyl)-1,3-dihydro-2H-imidazol-2-ylidene** [244187-81-3] $C_{27}H_{36}N_2$, **M 388.6, m 213-217°**, is formed and is stable enough to be isolated, stored and is available commercially. The carbene-carbon atom at C2 coordinates with metals; and catalyses the formation of C-C bonds, by C-H insertion, with acetylene, MeCN, HCCl₃, PhSOCH₃, and the structures of some of the products have been confirmed by X-ray analysis [Arduengo et al. *Helv Chim Acta* **82** 2348 1999, DOI: 10.1002/(SICI)1522-2675(19991215)82:12<2348>. The reduced **4,5-dihydro-1,3-bis(2,6-diisopropylphenyl)dihydroimidazolium tetrafluoroborate (SIPr-HBF₄)** has $C_{27}H_{39}N_2 \cdot BF_4$, **M 478.4, and m >300°**.

Bis(pinacolato)diboron [bis(pinacolyborane), **4, 4, 4', 4', 5, 5, 5', 5'-octamethyl-2,2'-bi-1,2,3-dioxaborolane, B₂pin₂**] [73183-34-3] $C_{12}H_{24}B_2O_4$, **M 253.9, m 138°, 137-140°**. This versatile borylating agent is prepared in four steps* from BBr₃ by conversion to tris(dimethylamino)borane {(Me₂N)₃B, m -16°, b 45°/12mm, 147-148°/atm [4375-83-1]} upon treatment with Me₂NH/pentane, followed by further reaction with BBr₃/pentane to give **bromobis(dimethylamino)borane** {(Me₂N)₂BBr, b 20-28°/0.5mm, 56-58°/12mm [6990-27-8]}, which on reaction with Na in toluene under reflux for ~3 hours provides **tetrakis(dimethylamino)diboron** [(Me₂N)₂B-B(Me₂N)₂, b 55-57°/2.5mm, 92°/12mm in 72% yield [Brotheron et al. *J Am Chem Soc* **82** 6242, 6245 1962, DOI: 10.1021/ja01509a009]. Then a mixture of (Me₂N)₂B-B(Me₂N)₂ [53.7g, 271mmol, one signal in the ¹H NMR at δ 2.67 (s, 24H) in CDCl₃] in dry toluene (510ml) and pinacol (64.4g, 545mmol) in dry toluene (340ml) under dry N₂ is stirred in an ice-water bath, and 5.4M HCl in Et₂O (203ml, 1.10 mol) is added dropwise during 2 hours. A white precipitate of Me₂NH.HCl separates immediately but the mixture is stirred further for 4

hours at $\sim 25^\circ$; the precipitate is filtered off and the filtrate is evaporated to dryness. The white residue is dissolved in pentane (~ 700 ml), washed with H_2O (3 x 500ml), dried (MgSO_4), filtered and evaporated to ~ 150 ml. This is warmed to dissolve any solid and chilled in a freezer (-30°). The first crop of **B₂pin₂** is collected and washed with cold pentane (2 x 30ml). The mother liquor is concentrated to give further crops of **B₂pin₂** which are combined with the first crop, and dried for 16 hours at $\sim 25^\circ/0.1$ mm to give colourless plates, **m 138^o**, in 79-91% yield. The diboron is air stable and is stored in a capped container. Its IR (KBr) has ν_{max} at 2978, 2930, 1372, 1289, 1189, 1177, 1127, 960, 850, 744, 660, 547 cm^{-1} ; the ^1H NMR (300MHz, CDCl_3) has δ_{H} at 1.25 (s, 24H); the ^{13}C NMR (100MHz, CDCl_3) has δ_{C} at 83.4, 24.9; and the ^{11}B NMR (128.3MHz, toluene) has δ_{B} at 30.61 ($\text{BF}_3 \cdot \text{Et}_2\text{O}$ as external reference with δ 0.00). [Ishiyama et al. *Org Synth* **77** 176 2000, DOI: 10.15227/orgsyn.077.0176; Ishiyama et al. *Org Synth* **82** 126 2005, DOI: 10.15227/orgsyn.082.0126, and DOI: 10.1002/0471264229.os082.20.] It borates ethylenes, acetylenes, allenes, conjugated olefins, replaces the halogen atom or a triflate group in arenes with a $-\text{B}(\text{OR})_2$ group effectively in the presence of catalysts such as $\text{Pt}(\text{PPh}_3)_4$, $\text{Pt}(\text{dba})_2$, or $\text{PdCl}_2(\text{dppf})$, $\text{Pd}(\text{dba})_2$ [Ishiyama et al. *J Am Chem Soc* **115** 11018 1993, DOI: 10.1021/ja00076a081; Lesley *Organometallics* **15** 5137 1996, DOI: 10.1021/om950918c; Ishiyama et al. *JCS Chem Commun* 689 1997, DOI: 10.1039/A700878C; Iverson & Smith *Organometallics* **16** 2757 1997, DOI: 10.1021/om970199x; Ishiyama et al. *JCS Chem Commun* 2073 1996, DOI: 10.1039/CC9960002073; Ishiyama et al. *Tetrahedron Lett* **39** 2357 1998, DOI: 10.1016/S0040-4039(98)00199-3; Ishiyama et al. *J Org Chem* **60** 7508 1995, DOI: 10.1021/jo00128a024; Ishiyama et al. *Tetrahedron Lett* **38** 3447 1997, DOI: 10.1016/S0040-4039(97)00642-4].

* **Caution:** All operations should be carried out under N_2 in a well ventilated fume cupboard because bromoborane derivatives fume in air by hydrolysing rapidly with evolution of heat.

Bis(trifluoromethanesulfonyl)amine [**NTf₂**, bis(trifluoromethanesulfonyl)amide, $(\text{CF}_3\text{SO}_2)_2\text{NH}$] [82113-65-3] ($\text{CF}_3\text{SO}_2)_2\text{NH}$, **M 281.2**, **m 49-50^o**, **46-57^o**, **b 90-91^o/atm**, has a **pK of a superacid** [$\Delta G_{\text{acid}} = 291.8$ kcal/mol]. **NTf₂** is a very strong neutral **Brønsted acid** which is to be compared with $\text{CHF}_2\text{CO}_2\text{H}$ ($\Delta G_{\text{acid}} = 323.8$) < HBr ($\Delta G_{\text{acid}} = 318.3$) < $\text{CF}_3\text{CO}_2\text{H}$ ($\Delta G_{\text{acid}} = 316.3$) and < $\text{CF}_3\text{SO}_3\text{H}$ ($\Delta G_{\text{acid}} = 299.5$) [Koppel et al. *J Am Chem Soc* **116** 3047 1994, DOI: 10.1021/ja00086a038; see pK values in AcOH below]. For purification see at the end of its synthesis below. **NTf₂** is a white crystalline solid that should be handled in closed systems (Schlenk equipment), or under very efficient ventilation as it is quite volatile, fumes in moist air and is very corrosive. However, it is soluble in H_2O , and it is stable in aqueous solutions where it can be titrated with NaOH as a typical acid with a pKa of ~ 1.7 . In glacial acetic acid, the pK value, by measuring $\delta(\text{OH})$ chemical shifts in the ^1H NMR spectra, is 7.8 for **NTf₂**, as compared with $\text{CF}_3\text{CO}_2\text{H}$ (11.4), HNO_3 (10.1), H_2SO_4 (7.0), HI (5.8), HClO_4 (4.9) and $\text{CF}_3\text{SO}_3\text{H}$ (4.2) measured in the same way. If these comparisons are valid then **NTf₂** is a **remarkably strong acid**. [Foropoulos & DesMarteau *Inorg Chem* **23** 3270 1984, DOI: 10.1021/ic00191a011; see also Nie et al. *J Fluorine Chem* **87** 45 1989, DOI: 10.1016/S0022-1139(97)00103-6.] The preparation of **NTf₂** in several steps from $\text{CH}_3\text{SO}_2\text{Cl}$ [Foropoulos & Desmarteau *Inorg Chem* **23** 3720 1984, DOI: 10.1021/ic00191a011;] has been improved from 48% to 80% overall yield [Desmarteau & Witz *J Fluorine Chem* **52** 7 1991, DOI: 10.1016/S0022-1139(00)80317-6], and is described here.

Methanesulfonyl chloride (286g, [124-63-0], **b 60^o/21mm**, 161^o/atm, redistilled from P_2O_5 , [Hearst & Noller *Org Synth Coll Vol* **4** 571 1963, DOI: 10.15227/orgsyn.030.0058]) is added slowly to solid KF (170g) and stirred for 1 hour at 25° , then $\text{CH}_3\text{SO}_2\text{F}$ is distilled off, fraction **b 123-124^o** is collected and redistilled from P_2O_5 to give pure **acid fluoride** (208g). Electrochemical fluorination of the methanesulfonyl fluoride in anhydrous HF at 4-5Volts and 7-9Amps continuously for 24 hours (as described by Gramstad & Haszeldine *J Chem Soc* 173 1956, DOI: 10.1039/JR9560000173) gave **CF₃SO₂F** (~ 180 g, [335-05-7]) and was isolated *via* condensation at -78° and redistillation at -21.7° .

CF₃SO₂F (76g) is bubbled into semi-frozen NH_3 (600ml, at $\sim -77^\circ$) with dry N_2 , during 0.5 hours while stirring with external cooling at -78° . Excess of NH_3 is allowed to evaporate under N_2 flow into a fume cupboard; NaOMe (54g) in MeOH (500ml) is then added to the residual slush and the mixture is heated to 60° for a few minutes, the NaF is filtered off, the filtrate is evaporated, the residue is dried in a high vacuum to give **CF₃SO₂NHNa** (81g, 95%, [91742-21-1]). This Na salt (81g) is refluxed with hexamethyldisilazane (645ml, 500g, HMDS see [999-97-3]) under N_2 flow (oil bath temperature $< 145^\circ$ to avoid excessive darkening) with strong mechanical stirring as the mixture thickens progressively. After release of NH_3 is complete (~ 12 hours), HMDS is distilled off (while increasing the vacuum), and the residual moisture sensitive solid is dried under high vacuum (in the same flask) to give **CF₃SO₂NNa(SiMe₃)** (106g, 92%, [91742-20-0]). This salt (106g) in dry

THF (370ml) is transferred to a stirrable stainless steel autoclave (~600ml) and gaseous $\text{CF}_3\text{SO}_2\text{F}$ (67g, ~26% excess) is transferred under N_2 pressure *via* a metal vacuum system, as it is not usually possible to cool the autoclave to below -50° . The autoclave is sealed and the mixture is stirred at 100° overnight, the volatile products are vented into an efficient hood, the autoclave is washed out with H_2O , the combined $\text{H}_2\text{O}/\text{THF}$ mixture is washed with CH_2Cl_2 , evaporated in a rotavap, and the residual solid is dried under high vacuum at 110° to give $(\text{CF}_3\text{SO}_2)_2\text{NNa}$ (129g, 98%, [91742-21-1]). This Na salt (129g) and 96% H_2SO_4 (150ml) are heated in a single necked flask at $60\text{--}90^\circ/2 \times 0.01\text{mm}$ and the $(\text{CF}_3\text{SO}_2)_2\text{NH}$ (NTf_2) (111g, 93%) is collected in an ascending tube cooled at -22° . If necessary (cf. spectra below), it is sublimed twice at 60° and/or recrystallised from CFCI_3 at -50° . The **analytically pure crystalline white solid** has **m 49-50°** and can be stored in a sealed container in a drybox.

NTf_2 has IR (gas in equilibrium with solid at 25°) with ν_{max} at 3395 (m), 3320 (br), 1463 (m), 1440 (m), 1300 (w), 1240 (s), 1224 (s), 1138 (s), 860 (m), 643 (vw), 614 (m), 570 (vw) and 505 (w) cm^{-1} ; Raman (solid -180°) has ν_{max} at 3205 (w), 1464 (vw), 1458 (w), 1450 (w), 1343 (w), 1263 (s), 1142 (m), 839 (w), 778 (s), 646 (w), 591 (w), 566 (w), 537 (w), 510 (w), 392 (m), 386 (m), 346 (s), 311 (s), 276 (s), 212 (w), 195 (w) and 128 (m) cm^{-1} ; the ^1H NMR (60MHz, external TMS) has peaks at 10.42 (in $\text{Me}_2\text{CO } d_6$), 7.92 (in CFCI_3) ppm; the ^{19}F NMR (20MHz, in CFCI_3 as solvent and internal standard) has a peak at -75.97 (s) ppm, i.e. is at higher field than that of CFCI_3 ; the MS[EI] has m/z at 281 (M^+), 211 ($\text{CF}_3\text{SO}_2\text{NSO}_2^+$), 147 ($\text{CF}_3\text{SO}_2\text{N}^+$), 133 (CF_3SO_2^+) and 69 (CF_3^+); and the MS[CI] has major m/z at 282 (MH^+), 150 ($\text{CF}_3\text{SO}_3\text{H}^+$ or $\text{CF}_3\text{SO}_2\text{NH}_3^+$) and 115 ($\text{CF}_2\text{SO}_2\text{H}^+$ or $\text{CF}_2\text{SONH}_2^+$). [Foropoulos & DesMarteau *Inorg Chem* **23** 3720 1984, DOI: 10.1021/ic00191a011.]

NTf_2 forms several derivatives including $(\text{CF}_3\text{SO}_2)_2\text{NCl}$ (viscous acid liquid, **m -96° to -93°**, [91742-17-5]), $(\text{CF}_3\text{SO}_2)_2\text{NNO}$ (white solid, **m ~118°**, [91742-19-7]), $(\text{CF}_3\text{SO}_2)_2\text{NNO}_2$ (white crystals **dec ~107°**, [91742-18-6]), $(\text{CF}_3\text{SO}_2)_2\text{Si}(\text{Me})_3$ (low-volatile colourless liquid, [82113-18-5]), and with CsF in MeCN it gives the cesium salt $(\text{CF}_3\text{SO}_2)_2\text{NCs}$ (white crystals **m 115°**, [91742-16-4]) [Foropoulos & DesMarteau *Inorg Chem* **23** 3720 1984, DOI: 10.1021/ic00191a011]. It forms an *N-trimethylsilyl derivative* (TMSNTf_2 prepared from allylTMS and NTf_2 , see above) which catalyses Diels-Alder reactions between methyl acrylate and various dienes [Mathieu & Ghosez *Tetrahedron Lett* **38** 5497 1997, DOI: 10.1016/S0040-4039(97)01208-2], as well as Friedel-Crafts acylations of anisole; and allylation and bis-allylation of carbonyl derivatives [Oshii et al. *Synlett* 1145 1997]. It readily forms the silver salt, AgNTf_2 , e.g. with Ag_2CO_3 .

1,3-Bis(2,4,6-trimethylphenyl)imidazolium chloride (IMes.Cl) [141556-45-8] $\text{C}_{21}\text{H}_{25}\text{N}_2^+ \text{Cl}^-$, **M 340.9, m >300°**. This NHC (N-Heterocyclic Carbene) precursor is prepared by established procedures from 2 mols of amine, 1 mol of glyoxal and one mol of formaldehyde [Arduengo USPatent 5 077 414 1991, *Chem Abstr* **116** 106289 1002, cf. Herrmann et al. *Chem Eur J* **2** 1627 1996, DOI: 10.1002/chem.19960021222] and washed with THF before use. It is useful for making IMesCuCl which in turn is used for making IMesAuCl and $[\text{IMesPtCl}]_2$ [Furst & Cazin *Chem Commun* **46** 6924 2010, DOI: 10.1039/C0CC02308F].

1,3-Bis(2,4,6-trimethylphenyl)imidazol-2-ylidene (IMes) [141556-42-5] $\text{C}_{21}\text{H}_{24}\text{N}_2$, **M 304.4, m 150-155°(dec.)**. By using Schlenk equipment and techniques under dry N_2 , this NHC is prepared from the preceding IMes.Cl (10.0g, 29.3mmol) suspended in dry THF (80ml) and stirred for 15 minutes, then solid *tert*-BuOK (3.5g, 31.2mmol) is added at $\sim 25^\circ$ in one portion. The dark grey solution is stirred for 20 minutes, and the volatiles are removed *in vacuo*. The residue is extracted into warm toluene (2 x 50ml), filtered through Celite and the solvent is removed *in vacuo* to provide small crystals of the carbene (7.55g, 84%). The crystals are recrystallised from hexane to yield colourless crystals, **m 150-155°(dec)** which are analytically pure and suitable for X-ray structural analysis. Its ^1H NMR (300MHz, THF- d_8 , TMS) has δ_{H} at 2.02 (s, 2',6'- CH_3 , 12H), 2.30 (s, 4'- CH_3 , 6H), 6.94 (s, ArH, 4H), 7.04 (s, 4,5-CH, 2H); and its ^{13}C NMR (75MHz, THF- d_8 , TMS) has δ_{C} at 18.04 (s, 2',6'- CH_3), 21.04 (s, 4'- CH_3), 121.28 (s, C-4 and C-5), 129.69 (s, Mes C-3',5'), 135.73 (s, Mes C-2',6'), 137.55 (s, Mes C-4), 139.73 (s, Mes C-1), 219.69 (s, NCN carbene-C); and the ^{15}N NMR (30MHz, THF- d_8 , $\text{NH}_4^{15}\text{NO}_3$) has δ at -178.85 . [Arduengo et al. *J Am Chem Soc* **114** 5530 1992, DOI: 10.1021/ja00040a007.] It is useful for making a variety metal complexes of ruthenium (among other metals) for catalysing olefin metathesis [Grubbs *Tetrahedron* **60** 7117 2004, DOI: 10.1016/j.tet.2004.05.124], and other metathesis reactions [Love et al. *Angew Chem Int Ed* **41** 4035 2002, DOI: 10.1002/1521-3773(20021104)41:21<4035; Schrodi & Pederson *Aldrichimica Acta* **40** 45 2007, Kanemitsu & Seeberger *Org Lett* **5** 4541 2003, DOI: 10.1021/ol035463z].

4-Bromobiphenylboronic acid MIDA ester {[2-(4-bromophenyl)-6-methyl-1,3,6,2-dioxazaborocane-4,8-di-

one], (4-bromophenyl)[*N*-[(carboxy- κO)methyl]-*N*-methylglycinato(2-)- $\kappa N, \kappa O$] boron [943552-04-3] **M** **C₁₁H₁₁BBrNO₄**, **311.9**, **m 238-240°**, **248-253°**. This ester is obtained from 4-bromophenylboronic acid (24.99g, 124.4mmol) and MIDA (18.31g, 124.4mmol, see [4408-64-4]) to which is added a freshly prepared mixture of DMSO (6.3ml) and toluene (118.7ml) in a flask to give a white solid suspension in a clear colourless solution. The flask is fitted with a Dean-Stark trap and a reflux condenser, and the mixture is boiled under reflux with stirring (magnetic Teflon coated bar) for 6 hours during which time the mixture became clear and about 2.1ml of H₂O are collected in the trap. Heating is stopped and the reaction is allowed to cool, with stirring, to ~25° during 1 hour; the apparatus is washed down with toluene and the combined liquids are evaporated (40°/15mm) to remove most of the toluene and give a crude tan, chunky, solid. Acetone (15ml) is added to the residue and swirled vigorously to give a white solid in a tan solution. To this mixture is added Et₂O (150ml, dried through neutral Al₂O₃ columns) in small portions (25ml) with stirring each time to allow further separation of white solid, and is finally filtered onto a medium porosity glass frit. The solid is washed with Et₂O (3 x 50ml), dried by suction, and is dried further in a vacuum (~25°/1mm) for 4 hours to give **4-bromophenylboronic acid MIDA ester** as an air stable, free-flowing white powder (36.3g, 94%) with **m 238-240°**. Alternatively, the crude tan solid can be purified by silica gel chromatography to provide the white MIDA ester in almost quantitative yield. An analytical sample has ¹H NMR (400MHz, CD₃CN) with δ_H at 2.50 (s, 3H), 3.89 (d, *J* = 16.0Hz, 2H), 4.07 (d, *J* = 16.0Hz, 2H), 7.41 (d, *J* = 8.0Hz, 2H), 7.55 (d, *J* = 8.0Hz, 2H); the ¹¹B NMR (100MHz, CD₃CN) has δ_B at 12.0; and for IR, ¹³C NMR and EI-MS see reference. [Ballmer, Gillis and Burke *Org Synth* **86** 344 2009, DOI: 10.15227/orgsyn.086.0344.] The **2-bromophenyl-** and **3-bromophenyl-** MIDA esters have **m 205-315°** and **m 237-242°** respectively.

trans-(2-Bromovinyl MIDA boronate {trans-2-bromovinylboronic acid MIDA ester 6-methyl-2-trans-bromovinyl-1,3,6,2-dioxazabronane-4,8-dione, [(1Z)-2-bromoethenyl][*N*-[(carboxy- κO)methyl]-*N*-methylglycinato(2-)- $\kappa N, \kappa O$]-boron} [*trans E* 1104636-68-1; *cis Z* has 1105069-27-9] **C₇H₉BBrNO₄**, **M 261.9**. The ester is prepared using Schlenk equipment in subdued daylight under dry N₂ by adding dropwise *via* a syringe freshly distilled (*E*)-(2-bromoethenyl)dibromoborane (21.0g, 75.9mmol, prepared from BBr₂ and acetylene, Hyuga et al. *Chem Lett* **16** 1757 1987, DOI: 10.1246/cl.1987.1757) to a stirred mixture of MIDA (16.9g, 114mmol, 1.5 equivalents, see [4408-64-4]) and 2,6-lutidine (17.7g, 151.9mmol, 2 equivalents) in DMSO (250ml) at 0° over 15 minutes, and allowed to warm to ~25° then kept at that temperature for 48 hours. The yellow mixture is treated with H₂O (300ml) and extracted with 1:1 THF:Et₂O (3 x 500ml), the combined extracts are washed with brine (3 x 350ml), dried (MgSO₄), filtered and evaporated *in vacuo*. The light yellow solid residue is purified by flash chromatography on silica gel and eluting with Et₂O/petroleum ether (1:1), then EtOAc, then EtOAc/MeCN (9:1) to give the desired boronate as colourless crystals (12g, 60%). On TLC it has R_F 0.46 (Merck silica gel plate grade 9385, 60Å, 230-400 mesh, eluting with EtOAc and visualised with KMnO₄). [A preparation similar to the one for vinyl MIDA boronate (this chapter) using the MIDA-di Na salt gave similar yields [cf. Uno, Gillis and Burke *Tetrahedron* **65** 3130 2009, DOI: 10.1016/j.tet.2008.11.010]. The material stored under air at room temperature for 1.5 years showed no signs of deterioration. Crystals grown by slow evaporation of an EtOAc solution at ~25° were suitable for X-ray analysis which revealed a **trans configuration** while confirming the expected structure with a pyramidalised boron centre. Its ¹H NMR (500MHz, CD₃CN, δ = 1.93 centre line, TMS) has δ_H at 6.69 (d, *J* = 15.0Hz, 1H), 6.63 (d, *J* = 14.5Hz, 1H), 3.97 (d, *J* = 17.0Hz, 2H), 3.82 (d, *J* = 17.0Hz, 2H), 2.80 (s, 3H); the ¹¹B NMR (100MHz, CD₃CN, BF₃·Et₂O internal standard) has δ at 10.5; and for IR, ¹³C NMR and HRMS (EI) see references. It is very useful in the reactions stated for vinyl MIDA boronate and for **iterative cross-coupling**. [Lee, Gray, Peak and Burke *J Am Chem Soc* **130** 466 2008, DOI: 10.1021/ja078129x; Uno, Gillis and Burke *Tetrahedron* **65** 3230 2009, DOI: 10.1016/j.tet.2008.11.010; Gillis & Burke *Aldrichimica Acta* **42** 17 2009].

Chloro(dimethylsulfide)gold(I) [(dimethylsulfide)gold(I) chloride, aurochloro dimethylsulfide, AuCl.Me₂S] [29892-37-3] (CH₃)₂SAuCl, **M 295.6**, **m >100°(dec)**, **100-102°(dec)**, **120°(dec)**. All procedures for preparing this complex should be carried out in an efficient fume cupboard because Me₂S is **toxic** and has a **foul** odour. The gold complex has been prepared by adding Me₂S (1 mole, see [75-18-3]) slowly to a stirred solution of NH₄AuCl₄ (1mole) in dilute *Aqua Regia* when the complex separates and is collected [Allen & Wilkinson *Spectrochimica Acta* **28B** 2257 1972, DOI: 10.1016/0584-8539(72)80200-9]. Alternatively, Me₂S is added to a solution of auric chloride (AuCl₃) in HCl [which forms **aurichloric acid (chloroauric acid)** HAuCl₄ see [16903-35-8]) when much heat is evolved and a flocculent white precipitate of **AuCl.Me₂S** separates. It is

filtered off, washed with H₂O (in which it is insoluble) and a little EtOH (in which it is slightly soluble), and dried at room temperature in air in the dark. It is air stable and can be recrystallised from *C₆H₆/Me₂CO. Clearly, the Me₂S reduces Au(III) to Au(I) before the complex separates. It is rapidly decomposed by sunlight to give metallic gold, Me₂S and HCl. In the absence of sunlight it can be kept in the solution in which it has been prepared for long periods, or in the presence of a slight excess of Me₂S in which colourless crystalline needles are formed. It is soluble in many organic solvents, e.g. CH₂Cl₂, *C₆H₆, but solutions deposit metallic gold in time. The gold content can be determined by exposing solutions to sunlight, and the gold that deposits can be collected and weighed. On heating in N₂, Me₂S begins to evolve above 100° until 200°, after which a residue of gold is obtained. [Phillips *J Am Chem Soc* **23** 250 1901, DOI: 10.1021/ja02030a008.] In a third synthesis, a suspension of freshly prepared Au(I)Cl (0.23g, 1mmol, [10294-29-8]) in dry *C₆H₆ (25ml) is treated with an excess of Me₂S which causes the solid to dissolve on stirring at 20°. After half an hour, the solution is evaporated *in vacuo*, and the crystalline residue is recrystallised from *C₆H₆/Me₂CO (1:1) to give a 70% yield of the pure complex, **m 100-102°(dec)**. [Dash & Schmidbaur *Chem Ber* **106** 1221 1973, DOI: 10.1002/cber.19731060418.]

For the **Au(III) complex**, aurochloric acid (10g) in H₂O (5ml) and a 1:1 v/v mixture of HNO₃ (d 1.4) and HCl (d 1.2) (1ml) are cooled in an ice bath for 10 minutes, and then Me₂S (0.5g) in Me₂CO (5ml) is added gradually (1ml at a time, at 3-5 minutes interval), heat is evolved and a yellow oil separates which soon solidifies to an orange yellow solid (some white *aurochloro dimethylsulfide* separates simultaneously). After allowing to stand in contact with a further quantity of acid mixture, all the solid is converted to *aurichloro dimethylsulfide* (**AuCl₃.Me₂S**). This is filtered off, washed with a 1:1v/v mixture of N/10 HNO₃ and N/10 HCl then H₂O and finally with EtOH, and dried in air. It is recrystallised from CHCl₃/Et₂O to give yellow prismatic crystals of analytically pure *auri complex AuCl₃.Me₂S* with **M 365.5** [29826-91-3] and **m 160°**, which is soluble in CHCl₃, Et₂O, Me₂CO, and warm *C₆H₆. When this auri complex is warmed with EtOH on a water bath, it is converted into the *auro complex AuCl.Me₂S* and the colour changes to white. The latter can be converted back to the former complex by treatment with aqua regia or chlorine water. [Ray & Sen *J Indian Chem Soc* **7** 67 1930.]

AuCl.Me₂S is *monomeric* with M⁺ at *m/e* at 294; its IR (Nujol mulls between polyethylene discs or CsI plates) has bands at 730vw (SC₂ *asym*), 675vw (SC₂ *sym*), 345s (Au-S, str), 326s and 319sh (Au-Cl str), 279m (SC₂ *def*), 198s (CSAu *def*), 109m, 93m and 83m (SAuCl *bend and lattice modes*) cm⁻¹ [Goggin et al. *JCS Dalton* 1904 1972, DOI: 10.1039/DT9720001904], the ¹H NMR (100MHz, CH₂Cl₂, TMS) has one peak at τ = 8.1 ppm [Allen & Wilkinson *Spectrochim Acta* **28A** 2257 1972, DOI:10.1016/0584-8539(72)80200-9], and the ¹H NMR (60MHz, CHCl₃, TMS) also has one peak at δ = -2.81 [Dash & Schmidbaur *Chem Ber* **106** 1221 1973, DOI: 10.1002/cber.19731060418].

It is a very useful compound for preparing a variety of gold complexes and gold catalysts [e.g. IMesAuCl in Part 1 of this Chapter].

Chlorotris(triphenylphosphine)cobalt [CoCl(PPh₃)₃] [26305-75-9] [(C₆H₅)₃P]₃CoCl, **M 881.2**, **m 135-139°(dec)**, **176-179°(dec.)**, **177°(dec.)**, **188°(dec)**. The complex is prepared under N₂, and preferably in Schlenk-type equipment. CoCl₂.6H₂O (9.6, 40.3mmol), PPh₃ (32g, 122mmol) and EtOH (600ml) are stirred while being purged with N₂ for several minutes, then vigorously at 60-70° for 30 minutes to ensure complete formation of the blue-coloured fine powder of CoCl₂(PPh₃)₂. Then under vigorous stirring the mixture is cooled to ~30° and NaBH₄ (1.28g, 33.9mmol) is added during ~10 minutes (in ~10 portions) whereby the colour of the solution changes from blue to green, then to brown. The green-brown precipitate is collected by filtration (in air), washed with portions of EtOH until the filtrate is no longer blue in colour, then with H₂O, EtOH again and finally with hexane, and dried *in vacuo* to give CoCl(PPh₃)₃ (24g, 67%). In the solid state it decomposes quite slowly in air, but is stable indefinitely under argon or N₂ in a refrigerator. It is soluble in *C₆H₆ and CH₂Cl₂ and the spectra in *C₆H₆ have λ_{max} at 745, ~940sh, 1100, 2200nm. [Wakatsuki et al. *Inorg Synth* **26** 189 1989, DOI: 10.1002/9780470132579.ch34; Aresta et al. *Inorg Chim Acta* **3** 227 1969, DOI: 10.1016/S0020-1693(00)92484-8.]

CoCl(PPh₃)₃ is a useful stoichiometric reagent for radical dimerisation of halogenated organic compounds as in the biomimetic synthesis of the bis-sesquiterpene lactones (±)-biatractylolide and (±)-biepiasterolide [Bagal et al. *J Org Chem* **69** 9100 2004, DOI: 10.1021/jo0488053], and alkaloid dimerisation in the total synthesis of the alkaloids (-)-calycanthine and (+)-chimonanthine [Movassaghi & Schmidt *Angew Chem Int Ed* **46** 3725 2007, DOI: 10.1002/anie.200700705]. It is used for the preparation of COPs (cobalt oxazoline palladacycles) catalysts

[see above, Stevens & Richards *Organometallics* **18** 1346 1999, DOI: 10.1021/om980812s].

(1,3,5,7-Cyclooctatetraene)dilithium (2,4,6-cyclooctatriene-1,2-diyl dilithium) [40698-91-7] $\text{C}_8\text{H}_8\text{Li}_2$, **M 118.1**. It is prepared by suspending lithium foil (1.0g, 144mmol) under N_2 in Et_2O (20ml) and stirring at 0° with 1,3,5,7-cyclooctatetraene (5.0g, 48mmol) for 16 hours. The small amount of white precipitate is allowed to settle, and an aliquot sample of the orange solution is sucked into a syringe, and the molarity of the solution is determined by hydrolysis with H_2O and titrating with standard acid. A saturated solution of $\text{Li}_2\text{C}_8\text{H}_8$ in Et_2O is $\sim 0.24\text{M}$. Store in the cold and away from air and moisture. The solution is not more flammable than an Et_2O solution, but the dry solid $\text{Li}_2\text{C}_8\text{H}_8$ is **pyrophoric** in air. [Spencer et al. *Inorg Synth* **19** 213 1979, DOI: 10.1002/9780470132500.ch49; Crascall et al. *Inorg Synth* **28** 126 1990, DOI: 10.1002/9780470132593.ch34.]

(1*R*,2*R*)-(+)-1,2-Diaminocyclohexane-*N,N'*-bis(2-diphenylphosphinobenzoyl) [(*R,R*)-DACH-phenyl Trost ligand] [138517-61-0] and **(1*S*,2*S*)-(-)-1,2-diaminocyclohexane-*N,N'*-bis(2-diphenylphosphinobenzoyl)** [(*S,S*)-DACH-phenyl Trost ligand] [169689-05-8] $\text{C}_{44}\text{H}_{40}\text{N}_2\text{O}_2\text{P}_2$, **M 690.8**, **m 137-142°**, $[\alpha]_{\text{D}}^{20}$ *R,R*- +134, *S,S*- -134 (c 1.0, MeOH), $[\alpha]_{\text{D}}^{20}$ *R,R*- +55.1, *S,S*- -55.1 (c 2.85, CH_2Cl_2). The (1*R*,2*R*)-enantiomer is prepared from (1*R*,2*R*)-1,2-diaminocyclohexane (0.535g, 4.68mmol), 2-(diphenylphosphino)benzoic acid (3.02g, 9.83mmol, [17261-28-8]), DMAP (61.0g, 0.5mmol) and DCC (2.13g, 10.3mmol) in CH_2Cl_2 (30ml) for 6 hours. The residue is chromatographed on silica gel and eluted with a 15–30% gradient of EtOAc/hexanes followed by recrystallisation from $\text{CH}_2\text{Cl}_2/\text{Et}_2\text{O}$ to give the **(+)-DACH-phenyl Trost ligand** (2.96g, 90%). It has an R_F of 0.43 on silica TLC (1:1 EtOAc-pentanes) and its ^1H NMR (200MHz, CDCl_3) has δ at 7.57 (m, 2H), 7.15-7.26 (m, 24H), 6.91 (m, 2H), 6.31 (br d, $J = 7.7\text{Hz}$, 2H, N-H), 3.77 (m, 2H), 1.87 (m, 2H), 1.62 (m, 2H), 0.9-1.3 (m 6H); has correct elemental analyses for C, H, N and P; and for IR and ^{13}C NMR see ref. [Trost et al. *J Am Chem Soc* **114** 9327 1992, DOI: 10.1021/ja00050a013.]

The Pd complexes have found extensive use in catalysing Asymmetric Allylic Alkylation (AAA) and palladium-catalysed Dynamic Kinetic Asymmetric Transformations (DKYATs), i.e. kinetic resolutions [Trost & Fandrick *Aldrichimica Acta* **40** 59 2007].

(-)-*N,N'*-(1*R*,2*R*)-1,2-Diaminocyclohexanediylbis(2-pyridinecarboxamide) [(*R,R*)-DACH-pyridyl Trost ligand] [218290-24-5] and **(+)-*N,N'*-(1*S*,2*S*)-1,2-diaminocyclohexanediylbis(2-pyridinecarboxamide)** [(*R,R*)-DACH-pyridyl Trost ligand] [172138-95-3] $\text{C}_{18}\text{H}_{20}\text{N}_4\text{O}_2$, **M 324.4**, **m 171-176°, 174-176°**, $[\alpha]_{\text{D}}^{20}$ *R,R*- -97.0, *S,S*- +98.0 (c 1.0, MeOH). The (*RS,RS*)-1,2-diaminocyclohexanediylbis(2-pyridinecarboxamide) is readily prepared by treating a mixture of 1,2-diaminocyclohexane (5.7g, 50mmol) in pyridine (20ml) and pyridine-2-carboxylic acid (12.3g, 100mmol) in pyridine (40ml) with triphenylphosphite (31.0g, 100mmol), and heating on a steam bath for 12 hours. The resulting brown oil is extracted into CHCl_3 , washed twice with aqueous NaHCO_3 , and H_2O , dried (MgSO_4), filtered, evaporated to a small volume and set aside overnight whereby brown crystals of the product are obtained (7.6g, 47%). Recrystallisation from CHCl_3 gives **analytically pure (\pm)-ligand** as white crystals, **m 201-202°**, with IR (KBr) ν_{max} at 3300, 3050, 2940, 2850, 1655, 1535 cm^{-1} ; and the ^1H NMR (60MHz, CDCl_3) has δ at 8.6 (m), 8.4 (br), 8.1 (m, combined 8H), 7.8 (m, 2H), 7.4 (m, 2H), 4.1(br, 2H), 2.3 (br), 1.6 (br, combined 6H) downfield from TMS (peaks are not well separated at 60MHz). [Barnes et al. *J Chem Eng Data* **23** 349 1978, DOI: 10.1021/jc60079a006.] **(1*R*,2*R*)-DACH-pyridyl Trost ligand** and **(1*S*,2*S*)-DACH-pyridyl Trost ligand** are prepared similarly, but starting from (1*R*,2*R*)-(-)-[20439-47-8] and (1*S*,2*S*)-(+)-[21436-03-3] 1,2-diaminocyclohexanes respectively [cf. their resolution in Chapter 3, ‘Alicyclic Compounds’].

These ligands readily form complexes with Cu, Zn, Co, Pt and Pd [Mulqi et al. *Inorg Chim Acta* **53** L91c 1981, DOI: 10.1016/S0020-1693(00)84752-0; Adolfsson & Moberg *Tetrahedron: Asymmetry* **6** 2023 1995, DOI: 10.1016/0957-4166(95)00263-O] and the Pd complexes have found extensive use in catalysing Asymmetric Allylic Alkylation (AAA) and palladium-catalysed Dynamic Kinetic Asymmetric Transformations (DKYATs), i.e. kinetic resolutions [Trost & Fandrick *Aldrichimica Acta* **40** 59 2007].

Di-*tert*-butylchlorophosphine [13716-10-4] $[(\text{CH}_3)_3\text{C}]_2\text{PCl}$, **M 180.7**, **b 48°/3mm, 72-73°/13mm, d_4^{25} 0.951, n_{D}^{20} 1.482**. This chlorophosphine is prepared from two equivalents of *tert*-butylmagnesium bromide and PCl_3 followed by the usual workup and fractional distillation in a vacuum, and redistillation of the desired fraction. With MeLi it provides **di-*tert*-butylmethylphosphine** **b 95-105°/3mm (170-172°/atm)**, which with MeI provides **di-*tert*-butyl-dimethylphosphonium iodide** **m 95-105°** (from EtOH). Hydrolysis of the chlorophos-

phine with H₂O provides **di-tert-butylphosphine oxide** [684-19-5] **M 162.2, m 55-59°** (hygroscopic crystals), **b 112°/9mm**, with IR which has ν_{\max} (CHCl₃) 2950s, 2290, 1470s, 1394w, 1370, 1142s, 918, 815 (P-Bu^t) and 656 cm⁻¹; the ¹H NMR (CD₂Cl₂) has τ at 3.53 (d, $J_{\text{HP}} = 453\text{Hz}$), 8.65 (d, $J_{\text{HCCP}} = 15\text{Hz}$, Bu^t). **Note** that this phosphine oxide is tautomeric with **di-tert-butylphosphinous acid**. [Hoffmann & Schellenbeck *Chem Ber* **99** 1134 1966, DOI: 10.1002/cber.19660990408; Hoffmann & Schellenbeck *Chem Ber* **100** 692 1967, DOI: 10.1002/cber.19671000241; Issleib & Krech *J Organometal Chem* **13** 283 1968, DOI: 10.1016/S0022-328X(00)82755-2; Crofts et al. *J Chem Soc C* 332 1970, DOI: 10.1039/J39700000332.]

The complex from Pd(OAc)₂ and this bulky **di-tert-butylchlorophosphine** ligand, after condensation with *P*-imino-azaphosphatane, catalyses efficiently the cross-coupling of arylboronic acids with aryl halides in the Suzuki-Miyaura reaction to provide specific unsymmetrical biphenyls in well over 90% yields [Kingston & Verkade *J Org Chem* **72** 2816 2007, DOI: 10.1021/jo0624521.]

4,4'-Di-tert-butyl-2,2'-bipyridyl (dtbpy) [72914-19-3] C₁₈H₂₄N₂, **M 268.4, m 159-160°, 159-161°, b 235°/32mm, pK_{Est} ~4.2**. The bipyridyl has been prepared by reaction of *tert*-butylpyridine with NaNH₂ (ratio 2:1) in xylene at 144-218° during 26 hours, then cooled, hydrolysed with H₂O, the organic layer is separated, dried, and distilled *in vacuo*. The bipyridyl solidified and can be sublimed. [McGill USP 4177349, *Chem Abstr* **92** 110871 1980, *Beilstein* **23/8** IV 181.] By coordinating with Ir(COD) and Ir(COE), complexes formed with **dtbpy** assist in C-H borylations [see Ir complexes above and Ishiyama et al. *Angew Chem Int Ed* **41** 3056 2002, DOI: 10.1002/1521-3773(20020816)41:16<3056; Ishiyama et al. *J Am Chem Soc* **124** 390 2002, DOI: 10.1021/ja0173019.]

Di-tert-butylneopentylphosphine [DTBNpP, (Pbu^t₂(-CH₂Cme₃)] [60633-21-8] C₁₃H₂₉P, **M 216.3, b 40°/0.1 mm, d²⁵ 0.839 g/ml, n_D²⁰ 1.478**. This ligand, with increased steric bulk, can be obtained by adding a solution of *di-tert-butylchlorophosphine* (108.4g, 114.0ml, 0.6mol, see [13716-10-4]) in THF (250ml) to neopentylmagnesium chloride (from 16g, 0.67g-atom of Mg turnings, 74ml, 64g, 0.6mol of neopentyl chloride in 250ml of dry THF containing a small amount of 1,2-dibromoethane as initiator, and stirred until all the Mg has dissolved, then cooled to 0°) at 0° with stirring, allowing to warm to ~25°, boiling under reflux for *ca* 15 minutes, then the solvent is removed at ~40°/40mm). The residue is diluted with Et₂O, the mixture is hydrolysed by stirring with saturated aqueous NH₄Cl, the layers are separated and the organic phase is dried (MgSO₄), filtered, the Et₂O is removed at ~25°/50mm, and the residual phosphine is distilled at high vacuum. There are also alternative syntheses for DTBNpP. **Work with protective gloves, with eyes and face protection, and in an efficient fume cupboard.** [cf. King et al. *J Org Chem* **41** 972 1976, DOI: 10.1021/jo00868a016.] It is, however, better to store and use it in the form of its salts, e.g. DTBNpP.HBF₄ (see below). In that case an equivalent amount of base, e.g. *t*-BuONa, must be added to the reaction mixture to liberate the free ligand. With a Pd source, e.g. Pd(OAc)₂ or Pd₂(dba)₃, catalysed H—B coupling (Hartwig—Buchwald) of various aryl halides has been achieved with a variety of aromatic primary or secondary amines, or saturated heterocyclic amines, e.g. morpholine, to form di- or tri-substituted amines in toluene solutions at 23° to 140° in high yields. [Hill et al. *J Org Chem* **71** 5117 2006, DOI: 10.1021/jo060303x.]

Unlike with Pd complexes, when DTBNpP is mixed with Pt(PhCN)₂Cl₂ (1:1 mol per Pt atom) in CH₂Cl₂ at 20° for a few minutes, a white binuclear complex is formed. It was characterised by spectroscopic and X-ray diffraction methods to have the structure [PtCl(Pbu^t₂CH₂Cme₂CH₂)]₂, i.e. one of the methyl groups of the neopentyl moiety is folded back and is metallated by the Pt to form a five membered Pt heterocyclic ring — and the molecule is **dimeric**. Such metallation does not occur with Pbu^t₂Prⁿ under similar conditions, and in this case the **monomeric cis** and **trans** [PtCl₂(PhCN)(Pbu^t₂Prⁿ)] square planar complexes are formed. [Mason et al. *JCS Chem Commun* 292 1976, DOI: 10.1039/C39760000292.]

Di-tert-butylneopentylphosphonium tetrafluoroborate (DTBNpP⁺.HBF₄⁻) [886059-84-3] C₁₃H₃₀P⁺ BF₄⁻, **M 304.2, m 258-262°(dec)**. This air stable salt is prepared in the same manner as [(*tert*-Bu)₃PH⁺ BF₄⁻, [131274-22-1] by mixing DTBNpP and aqueous HBF₄. It is easier to handle [cf. Netherton & Fu *Org Lett* **3** 4295 2001, DOI: 10.1021/ol016971g], and is a useful ligand in Pd-catalysed cross-coupling amination reactions of aryl bromides and chlorides [Hill et al. *J Org Chem* **71** 5117 2006, DOI: 10.1021/jo060303x].

3,3'-Di-tert-butyl-5,5',6,6'-tetramethyl-1,1'-biphenyl-2,2'-diol {H₂[BIPHEN]}, **1,1'-biphenyl-2,2'-diol, 3,3'(1,1'-dimethylethyl)-5,5',6,6'-tetramethyl-}** [R- 329735-68-4, S- 205927-03-3, RS- 101203-31-0]

C₂₄H₃₄O₂, M 354.5, m 160.5-161.5°. This biphenyl is prepared by the oxidation of 2-*tert*-butyl-4,5-dimethylphenol. The oxidation is carried out by slowly adding a solution of K₂Cr₂O₇ (130g, 0.442 mole, i.e. 1.02 equivalents being twice the theoretical amount to form the biphenyl) in a mixture of concentrated H₂SO₄ (260ml) and H₂O (780ml, **CARE** since much heat is evolved on mixing) at ~25°, to a solution of 2-*tert*-butyl-4,5-dimethylphenol (231.7g, 1.3 moles, see [1445-23-4]) in AcOH (1300ml) over 10 minutes at 55-60° and keeping at this temperature range. The colour of the solution changes from orange to green and a tan coloured solid separates during the reaction which is then cooled to room temperature. The solid is collected, washed sequentially with H₂O (2 x 250ml) and MeOH (3 x 200ml) until the solid is almost colourless, dried *in vacuo* and recrystallised from hot MeOH or dioxane/MeOH to give racemic **3,3'-di-*tert*-butyl-5,5',6,6'-tetramethyl-1,1'-biphenyl-2,2'-diol** [101203-31-0] (~115g, ~50%) **m 160.5-161.5°** (m 140-142° before recrystallisation). Its ¹H NMR (400MHz, CDCl₃, TMS) has δ at 7.12 (s, 2H, aromatic CH), 4.79 (s, 2H, OH), 2.24 (s, 6H, CH₃), 1.80 (s, 6H, CH₃), 1.38 (s, 18H, C(CH₃)₃); and for IR and ¹³C NMR see references. [Albert *J Am Chem Soc* **76** 4983 1954, DOI: 10.1021/ja01648a068; Alexander et al. *J Am Chem Soc* **120** 4041 1998, DOI: 10.1021/ja974353i; Alexander et al. *Organometallics* **19** 3700 2000, DOI: 10.1021/om000336h.]

Optical Resolution of H₂[BIPHEN]: Two procedures have been used: (i) preparation of the respective 2,2'-cyclic phosphoric acid (via reaction with POCl₃ with the di-Na salt in THF) followed by hydrolysis and recrystallisation of the (±)-acid (-)-**cinchonidine salt** in EtOH, followed by hydrolysis of the optically salts, conversion of the optically active cyclic acids to their respective optically active cyclic methyl esters, and finally cleavage with 'Red-Al' [Na AlH₂(OCH₂CH₂OMe₂)] in toluene to give optically pure H₂[BIPHEN]s. [Alexander et al. *J Am Chem Soc* **120** 4041 1998, DOI: 10.1021/ja974353i]; and (ii) which is described here in some detail as it can be used generally for the resolution of racemic 2,2'-dihydroxybiphenyls. **Firstly:** A solution of (1*R*,2*S*,5*R*)(-)-menthol (44g, 282mmol) in CH₂Cl₂ (100ml) is added to a solution of PCl₃ (58g, 423mmol, 1.5 equivalents) in CH₂Cl₂ (200ml) at 0° during 30 minutes. The cold bath is removed, the mixture is kept at ~25° for 1 hour, and the volatiles are removed at <25°/100mm. The residual oil, which consists of **menthyl phosphorochloridite** (see *l*-MenOPCl₂ [95456-31-8] in 'Miscellaneous As, B, P,...', in Chapter 3), is dissolved in CH₂Cl₂ (250ml), and a mixture of Et₃N (118ml, 847mmol, 3 equivalents) in CH₂Cl₂ (400ml) and H₂[Biphen] (100g, 282mmol) is added during 30 minutes. After 2 hours the mixture is filtered; H₂O₂ (30%, 200ml, **extremely violent reaction** take great **CARE**, wear body protection) is added **very** slowly with stirring, the biphasic mixture is then stirred vigorously for 2 hours, the layers are allowed to settle, the organic phase is washed with H₂O and brine (200ml), dried (MgSO₄), filtered and evaporated *in vacuo* to give a mixture of (*R*)- and (*S*)- BiphenP(O)(OMen) (124g, 85%) with ³¹P NMR (121MHz, THF, external ref 85% H₃PO₄) which has δ at -4.89 and -3.37 respectively due to the diastereomeric menthyl phosphates. **Secondly:** The diastereomeric mixture of phosphates is dissolved in the minimum volume of refluxing AcOH (~450ml), allowed to cool, and after 16 hours white crystals are filtered off, washed with cold AcOH (2 x 50ml) and dried *in vacuo* to give (*S*)-BiphenP(O)(OMen) (42g, 97-99% de). Recrystallisation from refluxing AcOH gives **pure S-diastereomer** (37.8g, >99%de, in 61% yield). The mother liquors from the first crystallisation are evaporated to give **pure (R)-BiphenP(O)(OMen)** (26.8g, >99%de, 43% *R*-diastereomer) after two recrystallisations from refluxing MeOH and cooling to 0°. [Partly racemic ester from the mother liquors are re-used in subsequent resolution processes.] **Thirdly:** To the preceding (*S*)-biphenP(O)(OMen) (37.83g, 70.3mmol) in toluene (500ml) in a 2L Schlenk flask at 0° is treated dropwise with 'Red-Al' (53ml, 65%wt in toluene, introduced into the separating funnel *via* a cannula) at 0°, and stirred for 16 hours. The mixture is carefully quenched with H₂O (75ml) then bleach (75ml), the slurry is filtered through a Celite pad, the pad is washed with toluene (250ml), the combined filtrates are allowed to settle, the toluene layer is separated, washed with bleach (200ml), brine (200ml), dried (MgSO₄), filtered, then the toluene is removed *in vacuo* at 0°, any menthol (minty odour) is removed by trituration with MeOH (50ml), to provide after drying, the resolved (*S*)-H₂[BIPHEN] (17.5g, 70%, >99% ee) which has [α]_D²⁰ -53.0 (c 0.352, THF, also reported [α]_D²⁰ -83.0 with c 0.4). The absolute configuration of this diastereoisomer was deduced from the X-ray crystallographic structure of its *syn*-Mo(*N*-2,6-di-*iso*-PrC₆H₃)(CHCMe₂Ph)[(*S*)-BIPHEN] complex (see Schrock-Hoveyda catalyst [205815-80-1] in Part 1). (*R*)-H₂[BIPHEN] is prepared from (*R*)-BiphenP(O)(OMen) using an identical procedure and the optical purity is deduced from that of its precursor as **no racemisation** occurs in these reactions. **Note** that the NMR spectra of the *RS*- (see above), *R*- and *S*-H₂[BIPHENS] are the same. About 20g each of the *R*- and *S*-H₂[BIPHENS] are usually obtained from 100g of *RS*-H₂[BIPHEN]Me; and ~60g of the latter are pooled from the mother liquors which can be recycled. [Alexander et al. *Organometallics* **19** 3700 2000, DOI: 10.1021/om000336h; Alexander et al. *J Am Chem Soc* **120** 4041 1998, DOI: 10.1021/ja974353i.]

By using identical compounds and reactions, but in which the *tert*-butyl group is replaced by a 1-adamantyl group the corresponding **1-adamantyl (H_2BIAD) derivatives** are obtained in almost similar yields and used to prepare related molybdenum catalysts. For example (*S*)-**3,3'-bis(1-adamantyl)-5,5',6,6'-tetramethyl-1,1'-biphenyl-2,2'-diol** {(*S*)- $H_2(BIAD)$ {*S*-(*-*)-**3,3'-biphenyl-2,2'-diol, 5,5',6,6'-tetramethyl-3,3'-bis(tricyclo-[3.3.1.1^{3,7}]dec-1-yl, [255728-57-5]**} with $[\alpha]_D^{20}$ -32.1 (c 3.3, THF), whose absolute configuration was also deduced by X-Ray crystallography of its related respective Mo complex catalyst, was prepared starting from 2-(1-adamantyl)-4,5-dimethylphenol **m 135-138°**. [Alexander et al. *Organometallics* **19** 3700 2000, DOI: 10.1021/om000336h.]

Dicyclohexyl(2-methylphenyl)phosphine (Cy_2P -*o*-Tol) [173593-25-4] $C_{19}H_{29}P$, **M 288.4, m 90-93°, pK_{Est} ~4.0**. This phosphine is prepared and purified in the same way as Cy_2PPh (following entry), and is used in the same manner as a phosphorus ligand for catalyst systems in coupling reactions. It has the advantage that it is less susceptible to oxidation, has a higher melting point and is easier to handle.

Dicyclohexylphenylphosphine (Cy_2PPh) [6476-37-5] $(C_6H_{11})_2PC_6H_5$, **M 274.4, m 56-57°, 57-58°, pK^{25} 3.40**. Cy_2PPh is a 'foul smelling' solid which is a good ligand for Cr, Hg and Group VIII metals. It can be prepared by reaction of $PhPCl_2$ with $CyMgCl$ in 50% yield [Issleib & Völker *Chem Ber* **94** 392 1961, DOI: 10.1002/cber.19610940215], or in 90% yield from $PhPCl_2$ and $CyLi$ [Screttas & Isbell *J Org Chem* **27** 2573 1973, DOI: 10.1021/jo01054a064], and is recrystallised from oxygen-free Me_2CO or $EtOH$. **In detail:** to the Grignard solution, at ice-salt bath temperature, made from Mg turnings (24.1g, 0.99mol) in dry Et_2O (350ml) and $CyBr$ (162g, 122ml, 0.99mol) in dry Et_2O (130ml), is added dropwise (2 hours) a solution of $PhPCl_2$ (81g, 61.4ml, 0.45mol) in dry Et_2O (100ml). The mixture is then boiled under reflux for 30 minutes and hydrolysed by stirring with saturated, deoxygenated, aqueous NH_4Cl . The Et_2O layer is separated and evaporated at ~20°/30mm leaving an oil that crystallises after all the solvent is removed. It may be recrystallised by slowly cooling a hot, oxygen-free $EtOH$ solution (1ml/g), with gentle stirring to prevent formation of an oil, to give analytically pure (Cy_2PPh) as white needles (98g, 79.5%). [Bianco et al. *Inorg Synth* **18** 169 1978, DOI: 10.1002/9780470132494.ch30.] It is a phosphorus ligand used in catalytic systems for coupling reactions. **Dicyclohexylphenylphosphine oxide (Cy_2PPhO)**, crystallises from petroleum ether, $EtOH$ or Me_2CO and melts at **157.5°** (m 165° was also reported).

Dicyclohexylphosphine (Cy_2PH) [829-84-5] $(C_6H_{11})_2PH$, **M 198.3, b 105-108°/3mm, 128°/8mm, 281-283°/atm, n_D^{25} 1.5142, pK_a 4.55**. This phosphine is a useful *air sensitive ligand* and should be used preferably in an inert atmosphere. It has been prepared in 80% yield by heating $Cy_2PSSPCy_2$ with excess Cu and distilling [Niebergall & Langenfeld *Chem Ber* **95** 64 1962, DOI: 10.1002/cber.19620950114], or in 55% yield by the addition of PH_3 to cyclohexene under pressure in the presence of α, α -azobis(isobutyronitrile) followed by fractional distillation [Rauhut et al. *J Org Chem* **26** 5138 1961, DOI: 10.1021/jo01070a087]. It is a phosphorus ligand in catalytic systems used in coupling reactions. It forms a pale yellow **Li salt** [19966-81-5] which is slightly soluble in dioxane but insoluble in Et_2O , $*C_6H_6$ and petroleum ether. [cf. Edmundson *Dictionary of Organophosphorus Compounds*, Chapman & Hall, London, 1988, p 221, ISBN 0-412-25790-4.]

Dicyclohexylphosphine oxide (Cy_2PHO) [14717-29-4] $(C_6H_{11})_2PHO$, **M 214.3, m 72.5-74.5°, 73-75°, b 324°/atm**. The oxide crystallises from hexane or $*C_6H_6$. It is *tautomeric* with dicyclohexylphosphinous acid (Cy_2P-OH), whose *ethyl ester* ([80413-46-3] **M 242.3**, is obtained from CyP_2Cl [16523-54-9] and $EtOH$) distils at **111-113°/1mm, n_D^{20} 1.4950** [Kabachnik et al. *Izv Akad Nauk SSSR, Ser Khim*, 949, 923 (*Engl Trans*) 1967]; and the *trimethylsilyl ester*, **M 286.5**, distils at **108-110°/1mm, n_D^{20} 1.4919** [Foss et al. *Zh Obhsch Khim* **49** 2418, 2134 (*Engl Trans*) 1979, cf. Edmundson *Dictionary of Organophosphorus Compounds*, Chapman & Hall, London, 1988, p 222, ISBN 0-412-25790-4]. It is used for preparing six-coordinate octahedral adducts with lanthanide(II) chlorides, see Kapoor & Saraswati [*Inorganica Chim Acta* **110** 63 1985, DOI: 10.1016/S0020-1693(00)81356-0].

2-Dicyclohexylphosphino-2',6'-dimethoxybiphenyl (S-Phos) [657408-07-6] $C_{26}H_{35}O_2P$, **M 410.5, m 162-162.5°, 164-166°**. This Buchwald phosphine can be synthesised in one pot under argon by adding *n*-BuLi (6.20ml, 2.5M solution in hexanes, 15.5mmol, 1.1 equivalents) *via* a syringe over 5 minutes to a cold (0°) solution of 1,3-dimethoxybenzene (2.00ml, 15.3mmol, 1.1 equivalents, [151-10-0]) in dry THF (35ml), allowing

the temperature to rise to $\sim 25^\circ$, then it is stirred for 3.5 hours. The mixture is re-cooled (0°) and 2-bromochlorobenzene (1.60ml, 13.7mmol, 1.0 equivalents, [694-80-4]) is added dropwise *via* a syringe over 15 minutes while stirring vigorously; and the burgundy coloured solution is stirred for a further 15 minutes at 0° . At this stage GC analysis of an aliquot quenched with $\text{Et}_2\text{O}/\text{H}_2\text{O}$ should show that all the bromochlorobenzene has been used up and a clean conversion to 2-bromo-2',6'-dimethoxybiphenyl has occurred. The mixture is cooled to -78° and *n*-BuLi (6.20ml, 2.5M solution in hexanes, 15.5mmol, 1.1 equivalents) is added dropwise *via* a syringe during 5 minutes and stirred further (with swirling if necessary) at -78° for further 30 minutes. Neat chlorodicyclohexylphosphine (3.03ml, 13.7mmol, 1.0 equivalents [16523-54-9]) is then added dropwise *via* syringe, stirred at -78° for 1 hour and allowed to warm to $\sim 25^\circ$. The mixture is filtered through a pad of Flash silica gel topped with a layer of Celite, eluted with EtOAc (400ml) and the combined filtrates are evaporated *in vacuo* to give a yellow solid residue which is recrystallised from Me_2CO to give **S-Phos** (2.90g) as a white solid. A further crop can be obtained by concentrating the mother liquors and triturating with MeOH to provide a total yield of 3.32g (59%) as a white solid **m 162-162.5°**. Its ^1H NMR (300MHz, C_6D_6) has δ at 7.59 (dm, *J* for the doublet 7.2Hz, 1H), 7.39-7.42 (m, 1H), 7.15-7.25 (m, 3H), 6.43 (d, *J* = 8.5 Hz, 2H), 3.33 (s, 6H), 1.60-1.94 (m, 12H), 1.06-1.36 (m, 10H) from TMS; the ^{31}P NMR (121MHz, C_6D_6) has δ at -8.6; and for IR and ^{13}C NMR see reference. [Barder et al. *J Am Chem Soc* **127** 4685 2005, DOI: 10.1021/ja042491j.]

Complexation of S-Phos to form a Pd-catalyst [e.g. with $\text{Pd}(\text{OAc})_2$] provides a system with *unprecedented scope*, reactivity and stability for Suzuki-Miyaura coupling processes, e.g. *it generates truly hindered highly chiral biaryls and heterobiaryls* [Walker et al. *Angew Chem Int Ed* **43** 1871 2004, DOI: 10.1002/anie.200353615; and for quinine-quinidine syntheses see Raheem et al. *J Am Chem Soc* **126** 706 2004, DOI: 10.1021/ja039550y; Huang et al. *J Am Chem Soc* **125** 6653 2003, DOI: 10.1021/ja035483w; Nguyen et al. *J Am Chem Soc* **125** 11818 2003, DOI: 10.1021/ja036947t; Gelman & Buchwald *Angew Chem Int Ed* **42** 5993 2003, DOI: 10.1002/anie.200353015]; and is useful for C-N bond formation [Strieter & Buchwald *Angew Chem Int Ed* **45** 925 2006, DOI: 10.1002/anie.200502927].

2-Dicyclohexylphosphino-2',4',6'-triisopropylbiphenyl (X-Phos, 2,4,6-triisopropyl-2'-dicyclohexylphosphinobiphenyl) [564483-18-7] $\text{C}_{33}\text{H}_{49}\text{P}$, **M 476.7**, **m 182-184°**, **187-190°**. This *Buchwald ligand* is prepared by reacting 2,4,6-triisopropylphenylmagnesium bromide with *o*-bromochlorobenzene in the presence of CuCl, and the resulting biphenyl is condensed with dicyclohexylphosphorus chloride. X-Phos is purified by solubilising it in $\text{Et}_2\text{O}/\text{CH}_2\text{Cl}_2$ (2:1, *ca* 30ml/g) by sonication, seeding it, and setting aside at -40° for 24 hours to give a first crop which is collected and washed with cold Et_2O . A second crop can be obtained by concentrating the mother liquors then mixing with hot MeOH ($\sim 10\text{ml}$, at 65°), cooling and while sonicating $\text{Et}_2\text{O}/\text{CH}_2\text{Cl}_2$ is added until homogeneous and set aside as before. The combined crops are dried *in vacuo* for 24 hours to give white crystals **m 182-184°**. The ^1H NMR (300MHz, C_6D_6) has δ_{H} at 7.48-7.42 (m, 1H), 7.24-7.06 (m, 5H), 2.85 (septet, *J* = 6.9Hz, 1H), 2.71 (septet, *J* = 6.9Hz, 2H), 1.94-1.52 (m, 12H), 1.42 (d, *J* = 6.9Hz, 6H), 1.24 (d, *J* = 6.9Hz, 6H), 1.30-1.02 (m, 10H), 1.12 (d, *J* = 6.9Hz, 6H); the ^{31}P NMR (121MHz, C_6D_6) has δ_{P} at -11.5; and for IR and ^{13}C NMR see references. [Huang et al. *J Am Chem Soc* **125** 6653 2003, DOI: 10.1021/ja035483w; Tomori et al. *J Org Chem* **65** 5334 2000, DOI: 10.1021/jo000691h.] X-Phos has been used for the amidation of arylsulfonates, in Pd catalysed Suzuki coupling [Johnson et al. *J Org Chem* **71** 7899 2006, DOI: 10.1021/jo061243y], in annulation of *o*-haloanilines [Jia & Zhu *J Org Chem* **71** 7826 2006, DOI: 10.1021/jo061471s], and in the synthesis of regioregular polythiophenes [Liversedge et al. *Tetrahedron Lett* **47** 5143 2006, DOI: 10.1016/j.tetlet.2006.05.063.] It is also available commercially in tablets containing 2 μmol of reagent per tablet.

R(+)- and S(-)- *N,N*-Dimethyl-1-ferrocenylethylamine (α -ferrocenylethyldimethylamine) [(*R*) 31886-58-5; (*S*) 31886-57-4; (*RS*) 31904-34-4] $\text{C}_{14}\text{H}_{19}\text{FeN}$, **M 257.2**, ***R* or *S* b 120-121°/0.7mm**, ***d*²⁵ 1.222g/ml**, ***n*_D²⁰ 1.589**, ***RS* b 111°/0.65mm (part dec)**, ***n*_D²⁵ 1.5883**, ***R* [α]_D²⁵ + 14.1 (c 1, EtOH)**, ***S* [α]_D²⁵ -14.1 (c 1.6, EtOH)**. The *RS*-amine has been prepared from ferrocenyldimethylaminoacetonitrile and MeMgI [Hauser & Lindsay *J Org Chem* **22** 906 1957, DOI: 10.1021/jo01359a013], but it is easily obtained by adding a solution of 1-ferrocenylethanol (23.0g, see [1277-49-2]) in toluene (150ml) at -20° dropwise to a stirred solution of phosgene (12.5g) in toluene (100ml) at -20° during ~ 30 minutes, then allowing to warm to $\sim 20^\circ$; and without isolating the chloro-derivative, the mixture is added to a solution of Me_2NH (22.5g) in propan-2-ol (200ml) at $\sim 20^\circ$. The temperature is then allowed to rise to $\sim 25^\circ$, the solution is filtered, evaporated to dryness, treated with $^*\text{C}_6\text{H}_6$ which is washed with 8.5% aqueous H_3PO_4 , more $^*\text{C}_6\text{H}_6$ is added, neutralised with Na_2CO_3 , the $^*\text{C}_6\text{H}_6$ layer is

dried (K_2CO_3), filtered and evaporated to give the **RS-amine** (24.2g, 95% crude), which after distillation (b 110°/0.45mm with some decomposition) gives the base as a **clear amber oil** (17.5g, 68%). Its **picrate** [39699-90-6] forms red plates **m 136-137°** from EtOH. The crude (\pm)-base is satisfactory for optical resolution. The **resolution** is carried out by stirring a solution of the (\pm)-base (51.4g) and *R*(+)-tartaric acid (30.0g) in MeOH (100ml) at 55°, seeding, lowering the temperature slowly (2°/hour), and after 24 hours the **(-)-base-(+)-acid** (30.0g) is obtained which gives the (-)-base (19.0g, $[\alpha]_D^{25}$ -11.0, c 1.5, EtOH). This is purified further by mixing it again with the (+)-acid (11.1g), each in MeOH (50ml) at 55°, seeded, and cooled as before to give pure (-)-base-(+)-acid from which **pure (-)-base** (17.0g, b 120-121°/0.7mm, $[\alpha]_D^{25}$ -14.1, c 1.6, EtOH) is isolated. From the mother liquors, the crude (+)-base is converted to the (+)base-(+)tartrate, recrystallised twice from Me_2CO/H_2O (10:1), from which **pure (+)-base**, $[\alpha]_D^{25}$ -14.1 is obtained as a **clear amber oil**. [Marquarding et al. *J Am Chem Soc* **92** 5389 1970, DOI: 10.1021/ja00721a017.] The absolute configuration was determined by X-ray analysis of the **(S,R,S)-2-(p-methoxyphenyl)hydroxymethyl-N,N-dimethyl-1-ferrocenylethylamine** [yellow powder from *n*-heptane has **m 110-111°**, $[\alpha]_D^{22}$ -120.8 (c 1.6, EtOH)] which is obtained by lithiation of the (+)-base (which is highly stereoselective), followed by reaction with *p*-anisaldehyde [Battelle et al. *J Am Chem Soc* **95** 482 1973, DOI: 10.1021/ja00783a030.]

This **ferrocene** is a very useful compound for preparing a variety of chiral ferrocene derivatives [see Marquarding et al. *J Chem Res (S)* **82** 1977 and (*M*) **0915** 1977], and is a remarkable phosphine ligand for catalysts by virtue of its highly stereoselective lithiation (with BuLi) in the position *ortho* to the dimethylaminoethyl group that allows substitution to give chiral 2-phosphines [Hayashi et al. *Tetrahedron Lett* **4405** 1974, DOI: 10.1016/S0040-4039(01)92175-6], e.g. 2—SiMe₃, 2—CH₂OH and 2—CPh₂OH derivatives [Marquarding et al. *J Am Chem Soc* **92** 5389 1970, DOI: 10.1021/ja00721a017]. The chiral 2-lithio derivative (prepared with *sec*-BuLi/Et₂O) reacts with sulfur, selenium or tellurium to give the yellow, red or red-brown **dichalcogenides** (*R,S*)-{[E-C₅H₃-CHMe(NMe₂)]Fe(C₅H₅)₂}₂ (from the *R*(+)-base) and (*S,R*)-{[E-C₅H₃-CHMe(NMe₂)]Fe(C₅H₅)₂}₂ (from the *R*(+)-base), where E is S, Se or Te respectively. These dichalcogenides are effective chiral ligands for the rhodium(I) {using [Rh(COD)Cl]₂} -catalysed asymmetric hydrosilylation of alkyl and aryl ketones with high enantiomeric excesses. The iridium(I) catalyst exhibits lower selectivity [Nishibayashi et al. *JCS Chem Commun* **1375** 1994, DOI: 10.1039/C39940001375]. The tellurium dichalcogenide (*R,S*)-{[Te-C₅H₃-CHMe(NMe₂)]Fe(C₅H₅)₂}₂, [*R,S*-(+)-(Fc*Te)₂] is prepared from lithiated *R*(+)-*N,N*-dimethyl-1-ferrocenylethylamine (with *sec*-BuLi/Et₂O) in Et₂O under N₂ at 0° which is then treated with Te powder in portions, and the mixture is subjected to ultrasonic irradiation for 1 hour. The mixture is poured into H₂O, and air is bubbled through it for 3 hours at ~25°. The solid is collected and purified by column chromatography on activated Al₂O₃, and eluting with EtOAc/hexane to give analytically pure [*R,S*-(+)-(Fc*Te)₂] as a red-brown solid (42%) with **m 57-58°**, $[\alpha]_D^{25}$ -622 (c 1, CHCl₃), the ¹H NMR (270MHz, CDCl₃) has δ_H at 1.25 (d, *J* = 6.9Hz, 6H), 2.18 (s, 12H), 4.01 (q, *J* = 6.9Hz, 2H), 4.06 (s, 10H), 4.18 (q, *J* = 1.3Hz, 2H), 4.23 (q, *J* = 1.1Hz, 2H), 4.48 (q, *J* = 1.3Hz, 2H); and for ¹³C NMR see reference. [*S,R*-(+)-(Fc*Te)₂] is prepared similarly, but from *S*(-)-*N,N*-dimethyl-1-ferrocenylethylamine in 47% yield and has identical physical properties except for $[\alpha]_D^{25}$ +613 (c 1, CHCl₃). After conversion into their anions (e.g. Te⁻Fc*, with NaBH₄/EtOH), they react with allylic bromides, e.g. geranyl bromide, to form **allylic-TeFc*** which produce the corresponding allylic alcohols (on treatment with Bu^oOOH/toluene) in 14-22% enantiomeric excess by chirality transfer, possibly *via* a [2.3]-sigmatropic rearrangement [Chiba et al. *Tetrahedron Lett* **36** 1519 1995, DOI: 10.1016/0040-4039(95)00074-M].

Similarly, but using Se instead of Te, *R*(+)- and *S*(-)-*N,N*-dimethyl-1-ferrocenylethylamine provides [*R,S*-(Fc*Se)₂] (**m 98-100°** from hexane), and [*S,R*-(Fc*Se)₂] (**m 103°** from hexane) respectively are produced as red solids, and these catalysts induce highly enantioselective selenoxide elimination to form axially chiral allene carboxylic esters with high enantiomeric excesses [Nishibayashi et al. *Tetrahedron Lett* **35** 3115 1994, DOI: 10.1016/S0040-4039(00)76844-4].

[*R,S*-(Fc*S)₂] (**m 169-170°** from hexane) is obtained as a yellow solid in the same way, but using S instead of Se or Te. [See above for the Rh(I)-catalysed asymmetric hydrosilylation reaction of ketones with these dichalcogenide ligands.]

(1*R*,2*R*)-(+)- and (1*S*,2*S*)-(-)- 1,2-Diphenylethylenediamine [(1*R*,2*R*)-(+)- and (1*S*,2*S*)-(-)- 1,2-diamino-1,2-diphenylenediamine, 1,2-diamino-1,2-diphenylethane, stilbenediamine, DPEN, STEIN) [(*R*) 35132-20-8]; (*S*) 29841-69-8; (*RS* \pm) 16635-95-3] C₁₄H₁₆N₂, M 212.3, *R* or *S* m 79-83°, *RS* m 82°, $[\alpha]_D^{20}$ *R,R* +102, *S,S* -102° (c 1, EtOH), $[\alpha]_D^{23}$ *R,R* +106, *S,S* -106 (c 1.1, MeOH), pK_(Est1)²⁵ ~5.86, pK_(Est2)²⁵ ~8.92. The racemate is

prepared in two steps. In the *first step*, a mixture of benzil (158g, 0.75mol, [134-81-6]), anhydrous NH_4OAc (400g) and cyclohexanone (80ml, 0.77mol) in glacial AcOH (1.0L) are stirred and heated under reflux (colour changing from light yellow to dark green) for 1.5 hours, and while hot it is poured into H_2O (3.0L) with vigorous stirring. After cooling to $\sim 25^\circ$ overnight, the crystals are collected, washed with H_2O (4 x 300ml), and crushed in a mortar (*in vacuo*) to give 2,2-*spirocyclohexane-4,5-diphenyl-2H-imidazole (2,3-diphenyl-1,4-diazaspiro-[4,5]deca-1,3-diene)* [5396-98-5] ($\sim 208\text{g}$, 96%) as yellow-green crystals, **m 107-108°**, on recrystallisation from hexane or aqueous MeOH . Its ^1H NMR (400MHz, CDCl_3 , TMS) has δ_{H} at 1.65-1.92 (m, 6H), 1.95-2.00 (m, 4H), 7.33-7.53 (m, 10H); and for ^{13}C NMR see references. In the *second step*, under argon and a Dry-ice condenser, the preceding crude spiro-imidazole (72.0g, 0.25mol) in dry THF (400ml, distilled from Na/benzophenone) is stirred until clear, cooled to -78° (Dry-ice/ Me_2CO bath), a stream of gaseous NH_3 is passed through until the volume of the homogeneous solution increases to 400ml. Lithium metal (6.94g, 1.0mol, *via* a powder funnel, from wire cut with scissors under a gentle stream of argon) is added at such a rate that the temperature is kept below -65° . The mixture is stirred for 30 minutes, absolute EtOH (30ml, 1.0mol) is added carefully, the mixture is stirred further for 20 minutes and NH_4Cl (70g) is added. The cooling bath is removed, the reaction temperature is made to rise to 0° , H_2O (400ml) is carefully added and the liquid phases are allowed to separate. The aqueous phase is extracted with Et_2O (3 x 300ml), the combined organic phases are washed with saturated aqueous NaCl (brine), dried (Na_2SO_4), filtered, evaporated down (to 200ml), and while being kept at 0° , 2 N aqueous HCl (300ml) is added slowly, then the two phases are stirred vigorously for 1 hour at $\sim 25^\circ$, H_2O (500ml) is added and the phases are separated. The organic layer is treated with H_2O (150ml) and extracted with CH_2Cl_2 (to remove any cyclohexanone). All the aqueous acidic phases are combined, and carefully (cool if necessary) basified with 2N aqueous NaOH (300ml) and extracted with CH_2Cl_2 (4 x 300ml); the combined extracts are washed with brine, dried (Na_2SO_4), filtered and evaporated *in vacuo* to give **crude $\pm\text{STEIN}$** ($\sim 40\text{g}$, $\sim 90\%$) as a light yellow solid **m 81-82°**. Its ^1H NMR (400MHz, CDCl_3 , TMS) has δ_{H} at 1.59 (br s, 4H), 4.10 (s, 2H), 7.2-7.3 (m, 10H); and for ^{13}C NMR see references.

The *optical resolution* is carried out by adding carefully a hot (70°) homogeneous solution of (L)-(+)-tartaric acid (30.0g, 0.20mol) in EtOH (230ml) to a hot (70°) solution of $\pm\text{STEIN}$ (42.5g, 0.20mol) in EtOH (230ml) (care exothermic, EtOH may boil) with stirring. The tartrate salts separate immediately and the mixture is cooled to $\sim 25^\circ$, the salts are filtered off, washed with EtOH (2 x 60ml), dried *in vacuo*, dissolved in boiling H_2O (230ml), EtOH (230ml) is added and the clear solution is cooled slowly to $\sim 25^\circ$. The crystals are collected by filtration, washed with EtOH (40ml), dried *in vacuo*, and recrystallised twice as before [from boiling H_2O (230ml), and EtOH (230ml)] to give the colourless (-)-*diamine-(+)-tartrate salt* ($\sim 24\text{g}$, $\sim 66\%$) with $[\alpha]_{\text{D}}^{23} -10.8$ (c 1.2, H_2O). To this (-)-(+)-salt suspended in H_2O (300ml) and vigorously stirred at $0-5^\circ$, is added dropwise an aqueous solution of 50% NaOH (23ml) followed by CH_2Cl_2 (150ml), and is stirred for 30 minutes; the phases are separated, the aqueous phase is extracted with CH_2Cl_2 (2 x 50ml), the organic layers are combined, washed with brine, dried (Na_2SO_4), filtered and evaporated *in vacuo* to give a colourless solid that is recrystallised from hexane to provide $>98\%$ optically pure *S,S*-(-)-*diphenylethylenediamine* as colourless crystals with the properties stated in the title and with NMR spectrum same as the racemate.

To obtain the enantiomeric diamine, all the filtrates from above are combined, evaporated to dryness *in vacuo*, the residue is stirred vigorously in H_2O (250ml), treated with aqueous 50% NaOH (25ml) then CH_2Cl_2 (200ml), stirred further for 30 minutes, the phases are separated, the aqueous phase is extracted with CH_2Cl_2 (2 x 50ml), the combined extracts are washed with brine, dried (Na_2SO_4), filtered and evaporated *in vacuo* to give enriched *R,R*-(+)-diamine ($\sim 26\text{g}$) as pale yellow crystals. This is treated with (D)-(-)-tartaric acid as in the above and gives relatively impure (+)-*diamine-(-)-tartrate salt* ($\sim 31\text{g}$, $\sim 85\%$, $[\alpha]_{\text{D}}^{23} +4$ (c 1.2, H_2O). However, recrystallisation of this salt did not improve its optical purity, but treatment with 50% NaOH and extraction as above etc, and crystallisation from hexane gave optically pure *R,R*-(+)-*diphenylethylenediamine* as colourless crystals with $[\alpha]_{\text{D}}^{23} +106^\circ$ (c 1.1, MeOH), and NMR spectrum as obtained with the racemate. The optical purity can be confirmed by the ^1H NMR spectra of their (L)-mandelate salts (cf. Benson et al. *J Org Chem* **53** 5335 1988, DOI: 10.1021/jo00257a024). These diamine ligands were used successfully in highly stereoselective epoxidation [Zhang et al. *J Am Chem Soc* **112** 2801 1990, DOI: 10.1021/ja00163a052], in aldol and Diels-Alder reactions [Corey et al. *J Am Chem Soc* **111** 5493 1989, DOI: 10.1021/ja00196a081], in allylation reactions [Corey et al. *J Am Chem Soc* **111** 5495 1989, DOI: 10.1021/ja00196a082], in osmylation reactions [Corey et al. *J Am Chem Soc* **111** 9243 1989, DOI: 10.1021/ja00208a025], as well as in enantioselective Michael additions [Bruner & Hammer *Angew Chem Int Ed* **23** 312 1984, DOI: 10.1002/anie.198403121] and in asymmetric hydrogenations [Fiorini & Giongo *J Mol Cat* **5** 303 1979, DOI: 10.1016/0304-5102(79)80027-9]. [Corey et al.

J Am Chem Soc **111** 5493 1989, DOI: 10.1021/ja00196a081; Pikul & Corey *Org Synth* **71** 22 1993, DOI: 10.15227/orgsyn.071.0022; and references herein.]

This diamine has been used successfully as a **Trost ligand** with Pd(dba)₂ to catalyse a Heck reaction, but it is not, albeit, as commonly used as DACH (*trans*-1,2-diaminocyclohexane) ligands.

(1*S*,2*S*)-1,2-diphenylethylenediamine bis-triflamide [121788-77-0] C₁₈H₁₄F₆N₂O₂, **M 404.3, m 213-214°**, [α]_D²³ **-6.6 (c 1.4, CHCl₃)**. The bis-amide is obtained from (1*S*,2*S*)-(-)-1,2-diphenylethylenediamine (see DPEN, [29841-69-8]) by treatment with triflic anhydride [358-23-6], Et₃N and 4-dimethylaminopyridine (DMAP, see [1122-58-3]) in CH₂Cl₂; and purified by flash chromatography on silica gel (15% EtOAc-hexane, v/v). Its ¹H NMR (400MHz, CDCl₃, TMS) has δ_H at 4.81(s, 2H), 6.80 (br s, 2H), 7.25 (s, 6H), 7.0 (s, 4H); and for ¹³C NMR see references. It is a useful reagent for enantioselective Diels-Alder and aldol reactions using a chiral controller system. [Corey et al. *J Am Chem Soc* **111** 5493 1989, DOI: 10.1021/ja00196a081; Pikul & Corey *Org Synth* **71** 30 1993, DOI: 10.15227/orgsyn.071.0030 and references therein.]

meso-1,2-Diphenylethylenediamine (meso-DPEN, stilbene-1,2-diamine) [951-87-1] C₁₄H₁₆N₂, **M 212.3, m 118°, 119°, 121°, 122°**. A 1:1 mixture of *meso*- and *rac*-DPEN is obtained by reaction of benzaldehyde, hexamethyldisilazane and BuLi, in THF, in the presence of TiCl₄ and TiO at ~25°/15hrs which gave, after workup and Kügelrohr distillation (bath temperature 150°/0.1mm), an oil (25% yield) that on repeated crystallisation from Et₂O/hexane gave crystalline *meso*-DPEN, **m 115-117°**. *Rac*-DPEN is recovered from the mother liquors [Betschart & Seebach *Helv Chim Acta* **70** 2215 1987, DOI: 10.1002/hlca.19870700826]. Purify it further by recrystallisation from Et₂O (leaflets). The *meso*-**dihydrochloride** has **m 156°(dec)** (from HCl), the **picrate** has **m 225°**, and the *meso*-**N,N,N',N'-tetramethyl derivative** [94533-51-4], **m 195-197°**, crystallises from hexane. Store it at <10°. A simpler preparation of *meso*-DPEN has been described which involves boiling ammonium acetate (0.52 mol) and benzaldehyde (0.99 mol) for 3 hours, and the resulting thick white precipitate is washed with EtOH and recrystallised from *n*-BuOH to give **N-benzoyl-N'-benzylidene-meso-1,2-diphenylethylene diamine (m 259°, 44% yield)**. This *derivative* is then heated with H₂O (100ml) and H₂SO₄ (54ml) while steam is passed through to remove PhCHO and PhCO₂H, concentrated until the distillate is no longer acidic, ice-cooled, neutralised with 880 NH₃/H₂O (1:1), and the flaky white crystals are extracted into Et₂O, dried (KOH), filtered and evaporated to give pure *meso*-DPEN (**m 120°, 75% yield**) after recrystallisation from petroleum ether (b 60-80°). [Irvine & Parkins *J Inorg Nucl Chem* **27** 270 1965, DOI:10.1016/0022-1902(65)80230-5]. [Beilstein **13** H 543, **13** IV 403.] *meso*-DPEN, and related 1,2-diphenylethylenediamines are highly sensitive **fluorogenic reagents for catecholamines** (norepinephrine, epinephrine and dopamine) [Umegae et al *Analyt Chim Acta* **208** 59 1988, DOI: 10.1016/S0003-2670(00)80736-9].

R-(+)- and S-(-)-2-[2-(Diphenylphosphino)phenyl]-4-isopropyl-2-oxazoline [**R-(+)- and S-(-)-2-[2-(diphenylphosphino)phenyl]-4,5-dihydro-4-isopropylloxazole** [*R*- 164858-78-0; *S*- 148461-14-7] C₂₄H₂₄NOP, **M 373.4, m 85-89°, [α]_D²⁰ *R* +48 and *S* -48 (c 1.4, CHCl₃)**. The 4*S*-(-)- enantiomer can be obtained by decomposing the 4*S*-(+)-Zinc II dichloride complex [crystallises from *tert*-butyl-methyl-ether/CHCl₃ as colourless crystals **m 221-223°, [α]_D²⁰ +53 (c 1.52, CHCl₃)**] (9.5g, 17.5mmol) in CHCl₃ (130ml) under argon with 2,2'-dipyridyl (2.72g, 1.74mmol) at ~25° for 1 hour (stirring), then the mixture is applied directly onto a silica-gel column (6 x 7 cm), eluted with CHCl₃ or EtOAc (800ml), and the eluate is evaporated to yield the desired **4*S*-(-)-oxazoline** (6.78g 95%) as a colourless solid which can be recrystallised from petroleum ether/Et₂O. On TLC (silica gel 60 Merck, 0.25mm F₂₄₅) it has R_F 0.40 (hexane/EtOAc 6/1); the ¹H NMR (300MHz, CDCl₃) has δ_H at 0.70 (d, *J* = 6.8Hz, 3H, CH(CH₃)₂), 0.81 (d, *J* = 6.8Hz, 3H, CH(CH₃)₂), 1.43-1.54 (m, 1H, CH(CH₃)₂), 3.80-3.90 (m, 2H, H₂C(5)), 4.09-4.19 (m, 1H, HC(4)), 6.85-6.89 (m, 1H, HC(4')), 7.24-7.37 (m, 12 H, arom. H), 7.88-7.92 (m, 1H, HC(6')); the ³¹P NMR (121MHz, CDCl₃, triphenyl phosphate external standard at -18ppm) has δ at -5.8; and for IR and ¹³C NMR see references below.

S-(+)-2-[2-(diphenylphosphino)phenyl]-4,5-dihydro-4-phenyloxazole [148461-15-6] C₂₇H₂₂NOP, **M 407.4, m 89-93°, [α]_D²⁰ +29.5 (c 1, CHCl₃)** and **R-(-)-2-[2-(diphenylphosphino)phenyl]-4,5-dihydro-4-phenyloxazole** [167171-03-1] **M 407.4, m 89-93°, [α]_D²⁰ -29.5 (c 1, CHCl₃)** were prepared as for the 4-isopropylloxazole above and have R_F 0.27 (hexane/EtOAc 5/1). [Koch et al. *Recl Trav Chim Pays-Bas* **114** 206 1995, DOI: 10.1002/recl.19951140413.]

With an iridium catalyst they promote asymmetric alkylation [Janssen & Helmchen *Tetrahedron Lett* **38** 8025 1997, DOI: 10.1016/S0040-4039(97)10220-9], and are chiral ligands for asymmetric reduction reactions

[Lightfoot et al. *Angew Chem* **110** 3047 1998, DOI: 10.1002/(SICI)1521-3757(19981016)110:20<3047>. These ligands, when complexed with Ni(COD)₂, promote the regio and stereoselective addition of allylic sulfides to alkynes; a reaction which tolerates a variety of functional groups [Hua et al. *Org Lett* **9** 263 2007, DOI: 10.1021/ol062686r]. These types of ligands also promote Pd-catalysed allylic substitution reactions [von Matt & Pflatz *Angew Chem* **105** 614 1993, DOI: 10.1002/ange.19931050430; Dawson et al. *Tetrahedron Lett* **34** 3149 1993, DOI: 10.1016/S0040-4039(00)93403-8].

Diphenylphosphine [829-85-6] (C₆H₅)₂PH, **M 186.2, b 100-102°/1.5mm, 150-151°/12mm, 156-157°/16mm, 165°/25mm, 280°/atm, d²⁵ 1.07g/ml, n_D²⁰ 1.6269, pK 0.03**. This toxic and flammable phosphine is prepared under N₂ by adding dropwise a solution of chlorodiphenylphosphine (50g, [1079-66-9]) in dry Et₂O (75ml) to a solution/suspension of LiAlH (2.6g) in absolute Et₂O (75ml), then boiling the mixture under reflux for 1 hour, adding H₂O (6ml, carefully and with cooling) and stirring for 2-3 hours. The Et₂O layer is collected, dried (CaCl₂), decanted, evaporated and the residue is distilled *in vacuo* to give the phosphine (70%) as a colourless oil [Kuchen & Buchwald *Chem Ber* **91** 2871 1958, DOI: 10.1002/cber.19580911246]. Alternatively, Ph₂PCl is reacted with Na in dry *n*-Bu₂O under N₂ to give Ph₂PNa which is treated with EtOH, boiled under reflux for 30 minutes, poured into H₂O, extracted with Et₂O, the extract is dried (CaCl₂), decanted, evaporated and the residue is distilled *in vacuo* to give the phosphine (72%) [Kuchen & Buchwald *Chem Ber* **92** 227 1959, DOI: 10.1002/cber.19590920126]. [Beilstein **16** H 758, **16** II 371, **16** III 833, **16** IV 950.]

2-(Diphenylphosphino)benzoic acid [(2-carboxyphenyl)diphenylphosphine] [17261-28-8] (C₆H₅)₂P C₆H₄CO₂H, **M 306.3, m 174-177°, 175-176° (187-188°), pK_{Est} ~2.7**. The benzoic acid is prepared by first adding Na (15.3g, 667mmol, cut in ~1g pieces) carefully to liquid NH₃ (1.5L) with stirring until a blue colour is persistent (**use an efficient fume cupboard**), then adding PPh₃ (87.4g, 330mmol) in small portions during 30-40 minutes with swirling to dissolve any Na on the sides of the flask. After 2.5 hours the red-orange colour of the solution of NaPPh₂ is treated with 2-chlorobenzoic acid (52.2g, 330mmol) in small portions over 30-40 minutes followed by anhydrous THF (500ml, using a syringe), and the resulting solution is allowed to warm to room temperature overnight under a blanket of argon. As the solution warms, the colour of the red solid turns to gold as mild evolution of heat results. The residue is dissolved in H₂O (1.4L) and extracted with Et₂O (400ml) which is discarded. The aqueous layer is acidified to pH 2 with concentrated HCl (~60ml), and extracted with CH₂Cl₂ (3 x 200ml), the extract is washed with H₂O (500ml), evaporated to *ca* 125ml, and MeOH (~50ml) is added to precipitate the pale-yellow crystalline acid (requiring inducement sometimes) which is collected and dried to give the phosphino-acid (50g, 49%) which provides analytically pure acid (C, H and P) on recrystallisation from a minimum amount of boiling MeOH. The **air stable** compound forms yellow needles from aqueous EtOH, is soluble in halogenated solvents and Et₂O, but is less soluble in other solvents. Its IR (Nujol) has ν_{max} at 3200 (OH), 1690 (CO), 1275 cm⁻¹. The **methiodide** crystallises from Me₂CO/*C₆H₆ with **m 169-173°(dec)**, and the **phosphine oxide** [2572-40-9] **M 322.3 has m 262-264° (252-254° and 274-275° are also reported), pK_a 5.06 and 5.37** (in 50% aqueous EtOH) and crystallises from EtOH, aqueous EtOH or aqueous AcOH. With diazomethane, **methyl 2-(diphenylphosphino)benzoate** is obtained in 85% yield; it forms colourless crystals from MeOH with **m 96-97°**, its IR (Nujol) has ν_{max} at 1712 (C=O) cm⁻¹, and the ¹H NMR (90MHz, CDCl₃) has δ_H at 7.6-6.9 (m, 14H), 3.70 (s, 3H), and the ³¹P NMR (40.5MHz, CDCl₃) has δ at 5.10 upfield of 85% H₃PO₄. [Rauchfuss et al. *J Am Chem Soc* **103** 6769 1981, DOI: 10.1021/ja00412a050; Hoots et al. *Inorg Synth* **21** 175 1982, DOI: 10.1002/9780470132524.ch39.] It is used for preparing the DACH-Trost phenyl ligand above.

S-(-)-2-Ferrocenyl-4-(1-methylethyl)oxazoline [S-(-)-4-isopropyl oxazo-2-yl ferrocene] [162157-03-1] C₁₆H₁₉NOFe, **M 297.0, m 71.5-72.5° (m 61-62°), [α]_D²² -129 (c 1.5, EtOH), CD (CHCl₃) λ_{max} (Δε) 464 (-080), 358 (+1.0)nm**. This complex is prepared in three steps from ferrocenecarboxylic acid (1.33g, 4.49mmol, 1271-42-7), **firstly** by converting it into the acid chloride with oxalyl chloride (0.79g, 9mmol) in CH₂Cl₂ (15ml) under N₂ at ~25° until gas evolution ceases (~10 minutes), stirring for 10 minutes further, and the solution is evaporated to dryness *in vacuo*. **Secondly** the crude dark red oily acid chloride (that solidified on standing) is dissolved in CH₂Cl₂ (10ml) and added (syringe) to a solution of S-(+)-valinol (0.554g, 5.37mmol) and Et₃N (1.25ml, 9mmol) in CH₂Cl₂ (7ml) under N₂ at ~25°. After standing overnight, the mixture is washed with H₂O (2 x 20ml), dried (Na₂SO₄), filtered, and evaporated to dryness *in vacuo*. The residue is purified by column chromatography on SiO₂ (40-83mm), eluting with 3% MeOH/CH₂Cl₂, to give analytically pure **2S-N-(1-hydroxy-3-methylbutyl)-ferrocenamide** (1.184g, 84%) with **m 109-110°, [α]_D²¹ -8 (c 1.34, EtOH)**. It has IR

(*nujol*) with ν_{\max} at 3284 (N-H), 3192 (O-H), 1611 (C=O amide I), 1551 (amide II) cm^{-1} ; the ^1H NMR (360MHz, CDCl_3) has δ_{H} at 0.97 (d, 3H, $J = 6.8\text{Hz}$, $-\text{CH}_3$), 0.98 (d, 3H, $J = 6.8\text{Hz}$, $-\text{CH}_3$), 1.93 (octet, 1H, $J = 6.8\text{Hz}$, $-\text{CH}(\text{CH}_3)_2$), 2.71 (brt, 1H, $-\text{OH}$), 3.56-3.83 (m, 3H, $-\text{CH}_2\text{OH}$ and $-\text{NHCH}-$), 4.16 (s, 5H, C_5H_5), 4.30 (brs, 2H, Fc), 4.60 (brs, 1H, Fc), 4.62 (brs, 1H, Fc), 5.80 (brd, 1H, $-\text{NH}-$); and for ^{13}C NMR and EI-MS see reference. **Thirdly**, to a light orange solution of the preceeding *S*-(-)-amide (0.817g, 2.59mmol) and Ph_3P (2.49g, 9.5mmol) in dry MeCN (60ml) is added Et_3N (1.6ml, 11.5mmol) followed by CCl_4 (2.2ml, 22.8mmol), and the mixture is stirred at $\sim 25^\circ$ overnight. Water (80ml) is added to the mixture, extracted with petroleum ether (5 x 50ml), the combined organic layers are dried (MgSO_4), filtered, and evaporated to dryness *in vacuo*. The residue is contaminated with Ph_3PO , and is purified by column chromatography [SiO_2 , 40-83mm, and eluting with 30% EtOAc/petroleum ether (b 40-60 $^\circ$)] to give analytically pure **ferrocenyl oxazoline** as dark yellow crystals (0.685g, 89%) with **m 71.5-72.5 $^\circ$** . *Note that much lower yields of product are obtained if the solvent is removed from the reaction mixture before work-up.* It has IR (*nujol*) with ν_{\max} at 1657 (C=N) cm^{-1} ; the ^1H NMR (360MHz, CDCl_3) has δ_{H} at 0.87 (d, 3H, $J = 6.8\text{Hz}$, $-\text{CH}_3$), 0.94 (d, 3H, $J = 6.8\text{Hz}$, $-\text{CH}_3$), 1.78 (hextet, 1H, $J = 6.6\text{Hz}$, $-\text{CH}(\text{CH}_3)_2$), 3.41 (q, 1H, $J = 7.0\text{Hz}$, $-\text{NHCH}-$), 3.89-3.95 (m, 1H, $-\text{OCHH}-$), 4.00 (t, 1H, $J = 7.7\text{Hz}$, $-\text{OCHH}-$), 4.06 (s, 5H, C_5H_5), 4.26 (brs, 2H, Fc), 4.66 (brs, 1H, Fc), 4.70 (brs, 1H, Fc); and for ^{13}C NMR and EI-MS see references. [Richards & Mulvaney *Tetrahedron Asymm* **7** 1419 1996, DOI: 10.1016/0957-4166(96)00159-0.] The *R*-enantiomer is prepared in the same way by using *R*-(+)-valinol. Similarly prepared while using the respective chiral *S*-2-aminoethanols were *S*-(-)-4-methyloxazol-2-yl ferrocene [red solid, 96% yield, **m 84-85 $^\circ$** , $[\alpha]_{\text{D}}^{27}$ -60 (c 0.2, EtOH), CD (CHCl_3) λ_{\max} ($\Delta\epsilon$) 461 (+0.34), 336 (-0.25)nm], *S*-(-)-4-ethyloxazol-2-yl ferrocene [orange solid, 34% yield, **m 47-48 $^\circ$**], *S*-(-)-4-tert-butyloxazol-2-yl ferrocene [orange-red solid, 34% yield, **m 127-128 $^\circ$** , $[\alpha]_{\text{D}}^{24}$ -150 (c 2.1, CH_2Cl_2)], and among others, where the 4-alkyl group was replaced by isobutyl, *sec*-butyl, benzyl, phenyl, 4,5-diphenyl, 1-hydroxy-1-methylethyl, 1-methoxy-1-methylethyl, 1-hydroxy-1-ethylpropyl, 1-methoxy-1-ethylpropyl, hydroxymethyl, methoxymethyl, and several more, were synthesised in the same way [Sammakia et al. *J Org Chem* **60** 10 1995, DOI: 10.1021/jo00106a005; Nishibayashi et al. *J Organomet Chem* **545-546** 381 1997, DOI: 10.1016/S0022-328X(97)00368-9; Richards & Mulvaney *Tetrahedron Asymm* **7** 1419 1996, DOI: 10.1016/0957-4166(96)00159-0.] An independent synthesis of the title compound was achieved from *S*-(-)-valinol, ZnCl_2 , cyanoferrocene in refluxing chlorobenzene under N_2 for 72 hours followed by column chromatography (SiO_2 , eluting with hexane/EtOAc) which gave a red solid (40% yield based on cyanoferrocene) **m 61-62 $^\circ$** [note difference in m.p.] but with similar ^1H and ^{13}C NMR. Other chiral derivatives, some mentioned above, were also obtained in this way. [Nishibayashi et al. *J Organomet Chem* **545-546** 381 1997, DOI: 10.1016/S0022-328X(97)00368-9.]

Iridium(III) acetylacetonate [tris(pentane-2,4-dionato)iridium, $\text{Ir}(\text{acac})_3$] [15635-87-7] ($\text{C}_5\text{H}_7\text{O}_2$) $_3\text{Ir}$, **M 489.5, **m 269-271 $^\circ$** . Iridium sulfate solution is made from freshly precipitated iridium dioxide (from 1.0g of potassium hexachloridate) by heating it with 1N H_2SO_4 (25ml) until the acid begins to fume. The solution is diluted to its original volume, and undissolved iridium oxide is removed by centrifugation. The light green solution is treated with 10% aqueous NaOH until the green iridium III hydroxide precipitate just begins to redissolve. This precipitation should be rapid in order to avoid oxidation; the pH of the solution is adjusted to 6, and the mixture is heated at 60 $^\circ$ with acetylacetone (2ml) for one hour. The solution then becomes red and deposits a yellow crystalline precipitate. This iridium complex is collected and crystallises from aqueous MeOH in rhombic plates (0.1g, 10%, **m 269 $^\circ$**). It is insoluble in H_2O , slightly soluble in EtOH and petroleum ether, but freely soluble in $^*\text{C}_6\text{H}_6$ and CHCl_3 . It is stable in boiling dilute acids and 10% aqueous NaOH. It sublimes at 260 $^\circ$ /1mm, but decomposes above 290 $^\circ$ depositing an iridium mirror. Molecular weight determination (~ 400 , by Rast in camphor) indicates that it is *monomeric*. [Dwyer & Sargeson *J Am Chem Soc* **75** 984 1953, DOI: 10.1021/ja01100a503; *Beilstein* **1** IV 3678.] It is a useful precursor for making Iridium complexes.**

3,4-O-isopropylidene-(3*S*,4*S*)-dihydroxy-(2*R*,5*R*)-bis(diphenylphosphino)hexane (RSSR-dimeDIOP, or R,S,S,R-DIOP*, the DIOP starred in dedication to Professor Kagan) [258873-45-9] $\text{C}_{33}\text{H}_{36}\text{O}_2\text{P}_2$, **M 526.2, **low melting oil**, $[\alpha]_{\text{D}}^{24}$ (+) 41.8 (c 0.88, PhCH_3), $\text{pK}_{\text{Est}} \sim 0.0$. The starting material for this ligand is the readily available D-mannitol that is converted to the isopropylidene di-epoxide, which in turn is converted to 3,4-O-isopropylidene-(3*S*,4*S*)-dihydroxy-(2*S*,5*S*)-hexane. The latter, (2.2g, 11.6mmol) and Et_2NH (4.9ml, 34.8mmol) in CH_2Cl_2 (30ml) is added dropwise to a solution of MeSO_2Cl (2.0ml, 25.8mmol) in CH_2Cl_2 (10ml) at 0 $^\circ$, kept at 0 $^\circ$ for 30 minutes, stirred at $\sim 25^\circ$ for 30 minutes, and treated with saturated aqueous NH_4Cl . The layers are separated, the aqueous phase is extracted with CH_2Cl_2 , the combined organic phases are dried (Na_2SO_4), evapor-**

ated and the residue is purified by flash chromatography on silica gel, eluting with $\text{CH}_2\text{Cl}_2/\text{Et}_2\text{O}$ (9/1) to give **3,4-O-isopropylidene-(3*S*,4*S*)-bis(mesyloxy)-(2*S*,5*S*)-hexane** (3.85g, 96%) as a colourless oil. This is a useful intermediate for making a variety of phosphine ligands and has $[\alpha]_D^{24}$ (-) **1.3** (c **1.04**, CHCl_3); its ^1H NMR (360MHz, CDCl_3) has δ at 4.82-4.76 (m, 2H), 3.99-3.96 (m, 2H), 3.03 (s, 6H), 1.45 (d, $J = 6.6$, 6.0Hz), 1.37 (s, 6H); and for ^{13}C NMR and EI-MS see references. The preceding dimesylate (1.04g, 3.0mmol) in THF (20ml) is added to the orange solution at -78° of Ph_2PLi [prepared under N_2 , from Ph_2PH (1.15ml, 6.6mmol) in THF (50ml) to which is added *via* a syringe *n*-BuLi in hexane (4.0ml, 6.4mmol) at -78° during 5 minutes, warmed to $\sim 25^\circ$ while stirring for 1 hour, and cooling to -78° again] during 20 minutes. The orange solution is warmed to $\sim 25^\circ$, stirred overnight, the resulting white suspension is hydrolysed with saturated aqueous NH_4Cl , the aqueous layer is extracted with CH_2Cl_2 , the combined organic layers are dried (Na_2SO_4), filtered, evaporated and the residue is purified by flash chromatography on silica gel, eluting with hexane/ Et_2OAc (95/5), or hexane/ Et_2O (95/5) to give **(+)-RSSR-DIOP*** (1.06g, 67%) as a colourless oil or low melting white solid respectively. Its ^1H NMR (360MHz, CDCl_3) has δ at 7.56-7.52 (m, 8H), 7.38-7.33 (m, 12H), 3.78-3.76 (m, 2H), 2.50-2.46 (m, 2H), 1.14 (s, 6H), 0.91 (d, $J = 7.0\text{Hz}$, 3H), 0.87 (d, $J = 6.9\text{Hz}$, 3H); the ^{31}P NMR (CDCl_3) has δ at -6.3 (-5.42); and for ^{13}C NMR and HRMS see references. [Li & Zhang *J Org Chem* **65** 5871 2000, DOI: 10.1021/jo0004613; Yan & RajanBabu *Org Lett* **26** 4137 2000, DOI: 10.1021/ol006591f.]

Lithium tetrachloropalladate(II) hydrate ($\text{Li}_2\text{PdCl}_4 \cdot x\text{H}_2\text{O}$) [123334-21-4] **$\text{Li}_2\text{PdCl}_4 \cdot x\text{H}_2\text{O}$, M 262.1 (anhydrous)**. It is commercially available as a powder or in chunks. A 0.5M solution of water-free Li_2PdCl_4 in dry MeOH for the preparation of Pd catalysts (e.g. see *Oxime Palladacycles* above) is made by stirring anhydrous LiCl (42.4g, 1 mole, dried at 130° to constant weight, several hours, see [7447-41-8]) and anhydrous PdCl_2 (88.6g, 0.5 mole, see [7647-10-1]) in dry MeOH (1L) until the solution is homogeneous. Store the solution under N_2 or argon. [Onue, Minami and Nakagawa *Bull Chem Soc Jpn* **43** 3480 1970, DOI: org/10.1246/bcsj.43.3480]

η^5 -(*S*)-(p*R*)-2-(2'-(4'-Methylethyl)-oxazolinyl)cyclopentadienyl,1-*C*-3'-*N*)(η^4 -tetraphenylcyclobutadiene)cobalt [222400-02-4] **$\text{C}_{39}\text{H}_{34}\text{NOCo}$, M 591.6, m 160-162°, $[\alpha]_D^{24}$ -55.2 (c 0.09, CHCl_3)**. This cobalt complex is the key intermediate for the preparation of all the cobalt oxazoline palladacycle (COP) catalysts described in Part 1. Although originally prepared in four distinct steps [Stevens & Richards *Organometallics* **18** 1346 1999, DOI: 10.1021/om980812s], the synthesis has been slightly modified and can be completed in two steps without purifying intermediates. **Firstly**, **η^5 -(carbomethoxycyclopentadienyl)-(η^4 -tetraphenylcyclobutadiene)cobalt** [182627-81-2] is prepared (using Schlenk equipment flushed with argon) in a triple-necked flask with two rubber septa containing sodium cyclopentadienide (20ml, 2.0M in THF, 40mmol, [4984-82-1], should be light red/orange in colour; discard if black or contains solid residues), anhydrous THF (20ml *via* a syringe, pre-filtered through activated Al_2O_3), then dimethyl carbonate (10.2ml, 120mmol, *via* a syringe), and the mixture is stirred and boiled under reflux for 4 hours before cooling to $\sim 25^\circ$. [The intermediate can be isolated but this is not necessary.] To the cooled mixture is added dry toluene (160ml, *via* a syringe), followed by chlorotris(triphenylphosphine)cobalt(I) (30.8g, 34.8mmol, see above [26305-75-9], preferably freshly prepared for higher yields) and diphenylacetylene (14.2g, 80mmol, recrystallised from EtOH), and heated at reflux for 5 hours, cooled to $\sim 25^\circ$ before transferring to a single-necked flask and evaporating to dryness *in vacuo*. The residue is suspended in hexanes (150ml), the solid is filtered off, washed with hexanes ($\sim 3.5\text{L}$) until the filtrate is colourless. The dark mustard-coloured solid is dissolved from the filter by washing with CH_2Cl_2 until the filtrate is colourless (discard any insoluble black solid) and evaporate the combined filtrates *in vacuo* to give the **Cobalt-ester** (12.7g, 67%) as a mustard-coloured solid. It can be purified by column chromatography on Matrix silica 60 (35-70 μm , eluting with 20% EtOAc/petroleum ether b 40-60°) to give the **Cobalt-ester** as an orange crystalline solid (m **240-244°**, lit m **246-248°**, Mabrouk & Rausch *J Organomet Chem* **523** 111 1996, DOI: 10.1016/S0022-328X(97)00368-9). Its ^1H NMR (500MHz, CDCl_3) has δ_{H} at 3.23 (s, 3H), 4.79 (t, $J = 2.0\text{Hz}$, 2H), 5.21 (t, $J = 2.0\text{Hz}$, 2H), 7.22-7.30 (m, 12H), 7.43-7.45 (m, 8H); and for IR and ^{13}C NMR see references. [Anderson, Overman, Richards, Watson and White *Org Synth* **84** 139 2007, DOI: 10.15227/orgsyn.084.0139.]

Secondly, **η^5 -(*S*)-(p*R*)-2-(2'-(4'-methylethyl)oxazolinyl)cyclopentadienyl,1-*C*-3'-*N*)(η^4 -tetraphenylcyclobutadiene)cobalt** is prepared by stirring under argon the above crude **Cobalt-ester** (10g, 18.6mmol), lithium iodide (4.96g, 37.2mmol, see [10377-51-2]) and 2,4,6-collidine (100ml, b **175-178°/atm**, see [108-75-8]), then boiling under reflux (air condenser) for ~ 16 hours until the **Cobalt-ester** (R_F 0.8, silica plate eluted with 20% EtOAc:hexanes) has completely reacted. The solution is cooled to $\sim 25^\circ$, diluted with CH_2Cl_2 , washed with 2N

aqueous HCl (4 x 150ml), dried (Na₂SO₄), filtered and evaporated *in vacuo* to give the solid orange coloured **Cobalt-acid** [¹H NMR (500MHz, CDCl₃) at δ_H: 4.84 (2d, *J* = 2.0Hz, 2H), 5.23 (2d, *J* = 2.0Hz, 2H), 7.20-7.24 (m, 12H), 7.42-7.44 (m, 8H)] which is used further. Under argon, with stirring, the preceding **cobalt-acid** in CH₂Cl₂ (124ml, pre-filtered through activated Al₂O₃), oxalyl chloride (3.25ml, 37.2mmol) followed by anhydrous DMF (3 drops sequentially, as evolution of gas occurs) are kept at ~25° for 30 minutes, evaporated *in vacuo*, CH₂Cl₂ (100ml) is added to dissolve the residue, evaporated again *in vacuo* and the process is repeated three times to give dry **Cobalt-acid chloride** as a red-brown residue. A Schlenk flask with a rubber septum, flushed with argon *via* inlet and outlet needles, is loaded with *S*-valinol hydrochloride (3.6g, 26mmol, prepared from the free base, see [2026-48-4], by treating with 2N HCl in Et₂O), then dry Et₃N (15.5ml, 112mmol) and CH₂Cl₂ (86ml) are added *via* the septum, followed, *via* a cannula, a solution of the preceding crude **Cobalt-acid chloride** in CH₂Cl₂ (100ml). The mixture is stirred at ~25° for 2 hours and cooled to 0° in an ice-bath. MeSO₂Cl (3.6ml, 47mmol, [124-63-0]) is then added to the mixture in one portion *via* a syringe and allowed to warm to ~25°. After 16 hours the mixture is washed with saturated aqueous NaHCO₃ (150ml), brine (150ml), the organic layer is dried (Na₂SO₄), filtered, and concentrated *in vacuo*. The residue is purified by dissolving it in the minimum volume of CH₂Cl₂ (~50ml), adding it to a silica gel column and eluting with hexanes/EtOAc (9:1) to give **η⁵-(S)-(pR)-2-(2'-(4'-methylethyl)-oxazolinyl)cyclopentadienyl,1-C-3'-N)(η⁴-tetraphenylcyclobutadiene)cobalt** (8.2g, 75%) as a yellow solid, which can be recrystallised to analytical purity from 10% EtOAc/petroleum ether to **m 160-162°**. Its ¹H NMR (400MHz, CDCl₃) has δ_H at 0.77 (d, *J* = 6.7Hz, 6H, CH₃), 0.97 (d, *J* = 6.7Hz, 3H, CH₃), 1.40 (oct, *J* = 6.7Hz, 1H, CH(CH₃)₂), 3.41-3.56 (m, 3H, -CH-CH₂-), 4.62 (brs, 1H, CpH), 4.80 (brs, 1H, CpH), 5.09 (brs, 1H, CpH), 5.20 (brs, 1H, CpH), 7.17-7.27 (m, 12H, *m*+*p*-PhH), 7.44-7.47 (m, 8H, *o*-PhH); and for IR and ¹³C NMR see references. The **R-enantiomer** is prepared in precisely the same way except that *R*-valinol hydrochloride is used instead, and it differs only in having optical rotation of opposite sign. [Stevens & Richards *Organometallics* **18** 1346 1999, DOI: 10.1021/om980812s; Anderson, Overman, Richards, Watson and White *Org Synth* **84** 139 2007, DOI: 10.15227/orgsyn.084.0139.]

N-Methyliminodiacetic acid (MIDA) [4408-64-4] CH₃N(CH₂CO₂H)₂, **M 147.1**, **m 220° (dec)**, **223-225° (dec)**, **226° (dec)**, **pKa** (thermodynamic) **acidic: 2.138 (0°), 2.142 (10°), 2.146 (20°), 2.150 (30°), 2.154 (50°); basic: 10.474 (0°), 10.287 (10°), 10.088 (20°), 9.920 (30°), 9.730 (40°)**. It has been prepared from MeNH₂ and formaldehyde cyanohydrin followed by hydrolysis of the dinitrile formed [Eschweiler *Justus Liebigs Ann Chem* **279** 39 1894, DOI: 10.1002/jlac.18942790105], and in 63-71% yield from chloroacetic acid and MeNH₂ [Berchet *Org Synth Coll Vol* **2** 397 1943, DOI: 10.15227/orgsyn.018.0056]. However, a more convenient preparation is from iminodiacetic acid [142-73-4] (13.3g), formic acid (4.6g), aqueous formaldehyde (10ml, 36%) and H₂O (5ml) at 100° until evolution of CO₂ ceases, and after 2 hours the mixture is refluxed for 10 minutes, diluted with excess EtOH, cooled and the precipitated acid is filtered off. Recrystallisation from aqueous EtOH (charcoal) provides **pure colourless MIDA** (14.7g, 90%). [Childs et al. *J Chem Soc* 2174 1948, DOI: 10.1039/JR9480002174]. It has been purified *via* the Ba salt [Berchet *Org Synth Coll Vol* **2** 397 1943, DOI: 10.15227/orgsyn.018.0056] and the stability constants of the complexes with Cu²⁺, Mg²⁺, Ca²⁺, Sr²⁺, Ba²⁺, Zn²⁺, Cd²⁺, Hg²⁺, and Ni²⁺ have been determined [Swarzenbach et al. *Helv Chim Acta* **38** 1147 1955, DOI: 10.1002/hlca.19550380509]. The **diethyl ester** has **b 92-94°/0.04mm** [Viscontini et al. *Helv Chim Acta* **35** 451 1952, DOI: 10.1002/hlca.19520350205], and the **monoamide** has **m 168°** after crystallisation (needles) from aqueous EtOH [Eschweiler *Justus Liebigs Ann Chem* **279** 39 1894, DOI: 10.1002/jlac.18942790105]. **MIDA (m 215-216° dec)** has been prepared on a ~100g scale by the method described by Childs et al. (above) with slight modification in 88% yield, and had IR (thin film) with ν_{max} at 2998, 2955, 1682 (C=N), 1477, 1380, 1328, 1223, 1172, 1126, 1065, 1018, 982, 958, 903, 886, 723 cm⁻¹; the ¹H NMR (400MHz, D₂O) has δ_H at 2.98 (s, 3H), 3.96 (s, 4H); the ¹³C NMR (100MHz, 95:5 DMSO-*d*₆: D₂O, TMS) has δ_C at 170.0, 56.7, 41.7; the LRMS (ESI+) has *m/z* (relative intensity) 219.1 (24%), 148.1 (M⁺, 100%), 102.1 (8%); and the HRMS (ESI+) for C₅H₁₀NO₄ [M+H⁺] has calculated 148.0604, and found 148.0603. [Ballmer, Gillis and Burke *Org Synth* **86** 344 2008, DOI: 10.15227/orgsyn.086.0344]. [Beilstein **4** H 367, **4** II 800, **4** III 1177, **4** IV 2428].

MIDA boronic esters: MIDA is a very useful protecting group for boronic acids and forms stable (more stable than the free acid) **boronates** which are readily prepared from the boronic acid, MIDA in toluene-DMSO (2.5:1) under reflux using Dean-Stark conditions to eliminate H₂O requiring ~3 hours. When these boronic acid surrogates are suitably substituted on boron they can be made to undergo Suzuki-Miyaura, Stille, Heck, Sonogashira and Negishi coupling, as well as Miyaura borylations. The boronates are easy to handle, stable in

air, stable under cross-coupling conditions, surviving temperatures of up to 80°, and acidic reagents like trifluoromethanesulfonic, acetic acid and oxidising agents such as Jones reagent. The MIDA group can be *easily hydrolysed* by mild aqueous base (see below) thus also allowing the use of controlled *iterative cross-coupling* (ICC) of protected haloboronic acids. [Gillis & Burke *Aldrichimica Acta* **42** 17 2009]. A variety of MIDA boronate synthons are available commercially.

Hydrolysis of 4-(*p*-tolyl)phenylboronic acid MIDA ester (m 214-216°) to the boronic acid is given here as a typical example for the hydrolysis procedure. The MIDA ester (10.1g, 31.3mmol, 1 equivalent) in THF (220ml), and aqueous 1M NaOH (93.5ml, 93.5mmol, 2.99 equivalents) gives a biphasic mixture with a clear colourless lower layer and a milky white upper layer. The container is capped, and the mixture is stirred vigorously at ~25° for 10 minutes whereby the lower colourless layer becomes clear and the upper layer becomes clear yellow. A saturated aqueous solution of NH₄Cl (250ml) is added (to act as acidifier) and the mixture is stirred vigorously for 5 minutes, shaken with Et₂O (4 x 50ml), and the aqueous layer is extracted with freshly prepared THF/Et₂O (1:1, 400ml). The combined organic layers are dried (MgSO₄), filtered through Celite, and concentrated by rotary evaporation in a vacuum (40°/20mm), and residual solvent is removed by three azeotropic cycles with MeCN (3 x 50ml) in a rotary evaporator (40°/20mm), and then at higher vacuum (25°/1mm) for 12 hours to give the boronic acid as a fine off-white powder (6.24g, 94% yield, **m 136-138° dec**) with the expected IR, NMR and Mass spectra which indicate ~92% purity. [Ballmer, Gillis and Burke *Org Synth* **86** 344 2008, DOI: 10.15227/orgsyn.086.0344].

MIDA-disodium salt is prepared from MIDA (147g, 1 mol), which is dissolved (caution: exothermic) in NaOH (120g, 3 mol) in H₂O (300ml) to give a clear light yellow solution that is evaporated *in vacuo*. The residue is treated with MeOH (300ml), heated to reflux, filtered through a glass frit, heated to reflux and filtered again, and the process repeated once more. The filtrate is co-evaporated with MeCN (3 x) and lyophilised to give analytically pure **diNa salt** (173g, 91%) as a white powder with ¹H NMR (300MHz, CD₃OD, TMS) δ_H at 2.95 (s, 4H), 2.24 (s, 3H); and ¹³C NMR (125MHz, CD₃OD, TMS) δ_C at 179.2, 63.7, 43.9. It is used for preparing MIDA-boronate esters [Uno et al. *Tetrahedron* **65** 3130 2009, DOI: 10.1016/j.tet.2008.11.010; cf. also Coombs & Margerum *Inorg Chem* **9** 1711 1970, DOI: 10.1021/ic50089a020; Davis & Richardson *Inorg Chem* **23** 184 1984, DOI: 10.1021/ic00170a014; Stringfield & Shepherd *Inorg Chim Acta* **309** 28 2000, DOI: 10.1016/S0020-1693(00)00238-3].

Norbornadiene (NBD, bicycle[2.2.1]hepta-2,5-diene) [121-46-0] C₇H₈, M 92.1, m -19.1°, b 89.5-90°/atm, 90.3°/atm, d₄²⁰ 0.9064, n_D²⁵ 1.4684. NBD is a versatile diene that coordinates with metals, helps in solubilising them in organic solvents, and is used for making a variety of *pre-catalysts* (see in this chapter under 'Homogeneous Catalysts' Part-1). It is prepared by a general method whereby *cis*- or *trans*- **1,2-bis(phenylsulfonyl)ethylene** {0.22g, 0.714mmol; *trans* [963-16-6] M 308.4, **m 221-223°, 226.5-227°, 229.5°** from EtOH or AcOH; *cis*-isomer has **m 89.5-90°, 101-101.5°**, from EtOH, *Beilstein* **6** IV 1500, Truce & McManimie *J Am Chem Soc* **75** 1672 1953, DOI: 10.1021/ja01103a045} and freshly distilled cyclopentadiene (1.6g, 2.42mmol) in toluene (5ml) are heated at 130° for 24 hours, evaporated to dryness *in vacuo*, and the residue is recrystallised from *C₆H₆-EtOH (1:1) to give the Diels-Alder adduct **3,6-methano-4,5-bis(phenylsulfonyl)cyclohex-1,2-ene, m 257-258°**, in good yield. Lower yields are obtained in *C₆H₆ (~25°/40 hours). Elimination of the β-disulfone groups is achieved by stirring a mixture of the *bis*(diphenylsulfonyl) compound (2.5mmol) and NaH₂PO₄·2 H₂O (5g) in MeOH (~40ml) at room temperature overnight with freshly prepared sodium amalgam (from *ca* 0.4g, 17mmol of Na, [11110-52-4]). The mixture is extracted with pentane (b 36°/atm) and the extract is fractionated at atmospheric pressure to give NBD in 65% yield. Store and stabilise with 0.50.25% of BHT under N₂. [De Lucchi & Modena *JCS Chem Commun* 914 1982, DOI: 10.1039/C39820000914; cf. Parham & Heberling *J Am Chem Soc* **77** 1175 1955, DOI: 10.1021/ja01610a027.] Industrially, it is prepared from cyclopentadiene and acetylene at 340°/6atm [Shell Devel. Co., USP 2 875256 1953].

NBD has UV with λ_{max} (logε) at 198.5 (3.78), 202infl (3.41), 213infl (3.24), 219infl (3.01), 227infl (2.41)nm (in EtOH), and 194.5 (3.53)nm (in cyclohexane) [Stich, Rotzler & Reichstein *Helv Chim Acta* **42** 1480 1959, DOI: 10.1002/hlca.19590420511]; its IR (thin film) has ν_{max} of the more intense bands at 2987, 1543, 1311, 1228, 875, 799, 727, 502 cm⁻¹, and in CS₂ and CCl₄ it has ν_{max} at 3080(s), 2960 (vs), 2880 (s), 1646 (m), 1452 (ms), 1335 (ms), 1310 (vs), 1271 (m), 1229 (s), 1206 (s), 1150 (s), 1105 (ms), 1063 (m), 1016 (m), 935 (s), 911 (s), 885 (sh), 870 (vs), 795 (sh), 715 (vs), 650 (s) cm⁻¹ [Abel, Bennett and Wilkinson *J Chem Soc* 3178 1959,

DOI: 10.1039/JR9590003178.]; the ^1H NMR (60MHz, CDCl_3) has δ_{H} at 1.99 (s, 2H, CH_2). 3.56 (s, 2H, bridgehead H), 6.74 (s, 4H, olefin H) from TMS, and for ^{13}C NMR see references. The photoelectron spectrum shows intense bands at 8.69, 9.55, 11.26, 12.51–12.75, 14.24 and weak bands at 15.66–17.16 eV; and the interaction between the two non-conjugated π -bonds has been shown to be 0.85eV [Bischof et al. *Helv Chim. Acta* **52** 1745 1969, DOI: 10.1002/hlca.19690520631.]

NBD-silver nitrate complex is formed by shaking for 5 minutes a mixture of AgNO_3 (1.36g) in H_2O (10ml) and NBD (3ml). The white complex is collected and recrystallised from EtOH to give pure **NBD-(AgNO_3) $_2$** (0.95g) which decomposes in air, in H_2O , and on heating to give a strong odour of NBD. Store it cold and under N_2 . It is soluble in warm MeOH, EtOH, CCl_4 , CHCl_3 , and $^*\text{C}_6\text{H}_6$, but insoluble in Me_2CO , Et_2O and light petroleum. Its IR (Nujol and hexachlorobutadiene mulls) has ν_{max} at 3020 (w), 2960 (w), 2885 (w), 1470 (m), 1385 (v, brs), 1325 (s), 1310 (s), 1243 (m), 1189 (m), 1046 (m), 1014 (w), 969 (m), 950 (vw), 929 (m), 890 (w), 877 (w), 808 (m), 795 (w), 790 (w), 725 (ms) cm^{-1} [Abel, Bennett and Wilkinson *J Chem Soc* 3178 1959, DOI: 10.1039/JR9590003178.]

Similarly the **NBD-cuprous bromide** complex is formed by shaking for 5 minutes a mixture of anhydrous CuBr (2.6g) in EtOH (30ml) and NBD (5ml). The white crystalline complex is filtered off, and washed with EtOH (5ml) and Et_2O (5ml) to give **NBD-(CuBr) $_2$** as fine white crystals which slowly become green in air. The complex, which is insoluble in most organic solvents, releases CuBr under reduced pressure, and is decomposed by H_2O to give Cu_2O . Its IR (Nujol and hexachlorobutadiene mulls) has ν_{max} at 3027 (vw), 3000 (vw), 2930 (m), 2860 (w), 1471 (w), 1453 (m), 1310 (ms), 1265 (w), 1245 (vw), 1234 (w), 1180–1080 (br-w), 1050 (vw), 993 (m), 976 (m), 950 (w), 938 (w), 918 (ms), 889 (w), 867 (w), 777 (w), 765 (w), 739 (m), 719 (m) cm^{-1} [Abel, Bennett and Wilkinson *J Chem Soc* 3178 1959, DOI: 10.1039/JR9590003178.] [Beilstein **5** IV 879.]

For the **bis(phenylsulfonyl)ethylenes** see also Truce & McManimie *J Am Chem Soc* **76** 5745 1954, DOI: 10.1021/ja01651a036; Montaniari *Gazetta* **86** 429 1956, Parham & Heberling *J Am Chem Soc* **77** 1175 1955, DOI: 10.1021/ja01610a027; Modena & Montaniari *Gazetta* **86** 436 1956, Adams & Ferretti *J Am Chem Soc* **81** 4927 1959, DOI: 10.1021/ja01527a042; and for UV see Angeletti & Montaniari *Boll Scient Fac Chim Ind Univ Bologna* **15** 44, 45 1957; in EtOH the *cis*-isomer has λ_{max} at 241–243nm, and the *trans*-isomer λ_{max} at 246nm.

Pentaphenylferrocenyl(di-tert-butyl)phosphine [**P(C $_5\text{H}_4\text{FeC}_5\text{Ph}_5$)(*t*-butyl) $_2$** , **Q-Phos**] [312959-24-3] [(C_6H_5) $_5\text{C}_5\text{Fe}[\text{C}_6\text{H}_4(\text{P}(\text{C}_6\text{H}_9)_2)_2]$, **M 710.7**, **m 211–219°**. This catalyst ligand is prepared in two steps in high yield, and all reactions are performed in a drybox. **Firstly** (*di-tert-butylphosphino*)ferrocene is prepared by adding *t*-BuLi (31.6ml, 53.8mmol) in THF to a solution of Cp_2Fe (10.0g, 53.8mmol) in THF (25ml) during 5 minutes at 0°, stirring for 20 minutes and the solvent is removed *in vacuo*. The residue is dissolved in pentane (100ml) and THF (5ml), (*t*-Bu) $_2\text{PCl}$ (5.33g, 29.5mmol) is added and the mixture is stirred for 3 hours after which time degassed MeOH (1ml) is added, and most of the solvents are removed *in vacuo*. The residue is filtered through a plug of silica gel under N_2 and unreacted Cp_2Fe is eluted with pentane first and then the phosphine is eluted all at once with Et_2O , the solvent is removed *in vacuo*, and the required *phosphine* is crystallised from pentane (yield 7.58g, 78%). In the **second step** this *phosphine* (1.00g, 3.03mmol), $\text{Pd}(\text{OAc})_2$ (0.35g, 0.156mmol) and *t*-BuONa (2.93g, 30.5mmol) are dissolved in PhCl (34.10g, 303.0mmol) and heated at 110° for 18 hours. The mixture is filtered through Celite, the PhCl is removed *in vacuo*, and the residue is subjected to chromatography on Silica gel, and eluting with pentane/ Et_2O (80:1), to **give** **P(C $_5\text{H}_4\text{FeC}_5\text{Ph}_5$)(*t*-butyl) $_2$** (1.47g, 68%) as a pink-red solid. Its ^1H NMR (C_6D_6^*) has resonances at δ : 1.07 (d, 11.0Hz, 18H, Me_3), 4.42 (t, 1.7Hz, 2H, C_5H_4), 4.67 (d, 1.0Hz, 2H, C_5H_4), 6.95–6.97 (m, 15H, *m,p*- C_6H_5), 7.44–7.48 (m, 10H, *o*- C_6H_5); the $^{31}\text{P}\{^1\text{H}\}$ NMR (C_6D_6^*) has a resonance at δ 25.49 (s); and for ^{13}C NMR see references. Similarly prepared were **P[C $_5\text{H}_4\text{FeC}_5(\text{C}_6\text{H}_4\text{-}p\text{-CF}_3)_5$](*t*-butyl) $_2$** (37.3% yield, from pentane), **P[C $_5\text{H}_4\text{FeC}_5(\text{C}_6\text{H}_4\text{-}p\text{-OMe})_5$](*t*-butyl) $_2$** (36% yield, from Et_2O at -30°), **P(C $_5\text{H}_4\text{FeC}_5\text{HPh}_4$)(*t*-butyl) $_2$** (47.4% yield, orange crystals from pentane at -35°), and **P[C $_5\text{H}_4\text{FeC}_5(3,5\text{-Me}_2\text{C}_6\text{H}_3)_5$](*t*-butyl) $_2$** (30% yield, red solid from pentane/ Et_2O) by using *p*-trifluoromethylchlorobenzene, *p*-methoxychlorobenzene, a limited amount of PhCl and 5-bromo-*m*-xylene respectively. These are **air-stable ferrocenyl dialkylphosphines** for palladium-catalysed C-C, C-N and C-O bonding-forming cross-coupling reactions. The ligand **P(C $_5\text{H}_4\text{FeC}_5\text{Ph}_5$)(*t*-butyl) $_2$** exhibits turnovers of ~1000 for aminations with unactivated aryl bromides or chloride, it catalysed the formation of selected aryl ethers under mild conditions, it catalysed the aryl halides coupling with acyclic or cyclic secondary alkyl- and arylamines, and it catalysed coupling of the latter amines with aryl- and primary alkylboronic acids. The ligand is **air stable** in the solid state and in solution, and produces highly active Pd catalysts which are **stable** in the solid state and react only slowly with oxygen in solution. [Kataoka et al. *J Org Chem* **67** 5553 2002, DOI: 10.1021/jo025732j;

cf. also Shelby et al. *J Am Chem Soc* **122** 10718 2000, DOI: 10.1021/ja002543e.]

4-(3-Phenylpropyl)pyridine 1-oxide (P₃NO) [34122-28-6] C₁₄H₁₅NO, M 213.3, m 58-65°, 60°, pK_{Est}~1.3. The *N*-oxide is prepared by adding 4-(3-phenylpropyl)pyridine (12.5g, [2057-49-0], in ‘Heterocyclic Compounds’ Chapter 3) during 5 minutes to a slurry of oxone (37.5g, [70693-62-8]), H₂O (63.5ml) and MeOH (125ml) (pH ~1.4) under N₂, and the stirred mixture is kept at pH 5.5 by addition of 5N aqueous NaOH while the temperature is kept at ≤ 35° (the temperature being controlled by external water cooling and by the rate of NaOH addition). The reaction is monitored by HPLC, and at completion, the salts are removed by filtration and the cake is washed with MeOH (50ml). The combined filtrate and washes are treated with 1M aqueous sodium metabisulfite solution (12ml), stirred for 0.5 hours and the pH is adjusted to 10.0 by addition of 5N aqueous NaOH, and set aside for 1 hour. The mixture is concentrated *in vacuo* (at 50°) to a final volume of ~75ml, cooled to 20°, the solid is filtered off, washed with H₂O (50ml), and dried under N₂ to give the desired **1-oxide** in 93% yield. *Alternatively*, it can be prepared by the oxidation of the *phenylpropylpyridine* with 30% H₂O₂/AcOH using the standard method of Ochiai [J Org Chem **18** 534 1953, DOI: 10.1021/jo01133a010; see also Boekelheide & Linn *J Am Chem Soc* **76** 1286 1954, DOI: 10.1021/ja01634a026]. Its ¹H NMR (300MHz, CDCl₃) has δ at 1.8-1.98 (m, 2H, 2'-propyl-CH₂), 2.3-2.6 (m, 4H, ph-CH₂, py-CH₂), 7.05 (m, 2H, 3,5-pyridine), 7.12-7.30 (m, 5H, benzene H), 8.11 (m, 2H, 2,6-pyridine H). [Senanayake et al. *Tetrahedron Lett* **37** 3271 1996, DOI: 10.1016/0040-4039(96)00565-5.] This reagent stabilises and enhances Jacobsen Mn-salen catalysts and is more readily obtained in quantity than the commonly used 4-phenylpyridine 1-oxide [see Jacobsen et al. *J Am Chem Soc* **113** 7063 1991, DOI: 10.1021/ja00018a068].

Pinacolborane (4,4,5,5-tetramethyl-1,3,2-dioxaborolane) [25015-63-8] C₆H₁₃BO₂, M 128.0, b 42-43°/50mm, d₄²⁵ 0.882, n_D²⁰ 1.396. This borolane is prepared by stirring a cooled solution (at 0°) of pinacol (2.36g, 20mmol [76-09-5]) in CH₂Cl₂ (2ml) to which is added, dropwise, BH₃-SMe₂ (20mmol, 10.0M in SMe₂) whereby effervescence occurs (use efficient fume hood, and gases have a foul odour, are flammable, and should be absorbed through alkaline solution or oxidising solutions containing H₂O₂ or KMnO₄). After stirring for ~1 hour at 0°, the mixture is warmed to ~25°, when gas evolution ceases (*ca* another 1 hour) and is distilled to give the borolane as a clear oil (1.61g, 63% yield). Its IR (CH₂Cl₂) has ν_{max} at 3272s, 2980s, 1530m, 1482s, 1333s, 1143s, 983s, 851s cm⁻¹; and the ¹H NMR (300MHz, CDCl₃) has δ at 1.27 (s, 12H); the ¹³C NMR (75.5MHz, CDCl₃) has δ at 83.1, 24.5; and the ¹¹B NMR (115.5MHz, CDCl₃) has δ at 28.07. It is a very **efficient hydroboration reagent**, requires mild reaction conditions, has higher functional group tolerance, higher regio- and stereoselectivity, with excellent stability of the resulting pinacol boronic esters produced. [Tucker et al. *J Org Chem* **57** 3482 1992, DOI: 10.1021/jo00038a044.]

Rhodium(I) chloride 1,5-cyclooctadiene complex dimer {chloro(1,5-cyclooctadiene)rhodium dimer, di-μ,μ'-chlorobis[(1,2,5,6-η)-1,5-cyclooctadiene]dirhodium (I), [Rh(COD)Cl]₂} [12092-47-6] C₁₆H₂₄Cl₂Rh₂, M 493.1, m ~243° (dec, see below). The complex is prepared by refluxing a solution of RhCl₃·3H₂O (1.93g, 7.3mmol), H₂O (3ml), EtOH (35ml) and COD (6ml, 48mmol) overnight. After cooling, the crude solid is collected, washed with a little H₂O, dried and recrystallised from CH₂Cl₂/hexane or acetic acid, then dried *in vacuo* to give 82% yield (1.5g) of dimer. On heating it darkens from about 220°, melts at ~256° and decomposes with effervescence at 258°. M by ebullioscopy in 0.9% CHCl₃ is ~513 confirming the **dimeric** structure. It is soluble in CH₂Cl₂ and THF, moderately in CHCl₃, AcOH and Me₂CO, slightly soluble in hexane, Et₂O, MeOH, EtOH, and *C₆H₆ but insoluble in H₂O. Its magnetic susceptibility χ, is -0.52 x 10⁻⁶ ±4% per g. *Alternatively*, NaBH₄ (0.5g) in EtOH (100ml) is added, over 3 hours, to a stirred solution of RhCl₃·3H₂O (5g) and COD (3ml) in EtOH (125ml) at 20°. The **exothermic reaction** deposits some Rh, and after 2 days the mixture is filtered, and the residue is extracted with CH₂Cl₂. The extract is evaporated to dryness and the orange residue is recrystallised from AcOH (yield 0.67g). The reaction mixture still contained much unreacted RhCl₃. [Chatt & Venanzi *J Chem Soc* 4735 1957, DOI: 10.1039/JR9570004735; Schenck et al. *Inorg Chem* **24** 2334 1985, DOI: 10.1021/ic00209a003; cf. Giordano et al. *Inorg Synth* **19** 218 1979, DOI: 10.1002/9780470132500.ch50.] The related **bis-(Rhodium COD-μμ'-diacetato)** complex [Rh(COD)OAc]₂, M 540.1, is obtained from an acetone solution of the preceding chloro complex (1g) and KOAc (1g) which is boiled under reflux for 2 hours. After filtration and evaporation of the filtrate to dryness, the residue is recrystallised from EtOAc to give **pure diacetato** complex as orange crystals, m 197-198° (yield 0.54g), M 540 (ebullioscopically in 1.4% *C₆H₆; M 587 in 1.5% *C₆H₆; M 554 in 0.9% *C₆H₆). A 10⁻³M solution in PhNO₂ is non-conducting.

Rhodium(I) chloride norbornadiene complex dimer {bicyclo[2.2.1]hepta-2,5-diene-rhodium(I) chloride dimer, di- μ '-chlorobis[2,5-norbornadiene]dirhodium(I), $[\text{Rh}(\text{NBD})\text{Cl}]_2$ } [dimer 12257-42-0; polymer 42740-82-9] $\text{C}_{14}\text{H}_{16}\text{Cl}_2\text{Rh}_2$, **M 460.99, m $\sim 240^\circ$ (dec).** The yellow dimer is obtained by shaking rhodium 'trichloride' (0.7g, $\text{RhCl}_3 \cdot 3\text{H}_2\text{O}$) and norbornadiene (2ml) in aqueous EtOH (10ml) for 2 days. It is collected and recrystallised from hot CHCl_3 /light petroleum to give analytically complex (0.62g). **M** by ebullioscopy in $^*\text{C}_6\text{H}_6$ is 481 consistent with a **dimeric** structure. The fine yellow crystals are soluble in CHCl_3 and $^*\text{C}_6\text{H}_6$, but almost insoluble in light petroleum and Et_2O . Its IR (CS_2) has ν_{max} at 3060(m), 3000(m), 2960(m), 2920(m), 2855(m), 1307(s), 1171(m), 1157(w), 1068(w), 1029(w), 995(w), 932(m), 882(m), 721(w), 680(s), 630(s) cm^{-1} . [Abel, Bennett & Wilkinson *J Chem Soc* 3178 1959, DOI: 10.1039/JR9590003178.]

Ruthenium(II)benzenedichloride dimer {di- μ -chloro-bis[(η -benzene)chlororuthenium(II)], $\text{Ru}_2(\text{C}_6\text{H}_6)_2\text{Cl}_4$, $[\text{Ru}(\text{C}_6\text{H}_6)\text{Cl}_2]_2$ } [37366-09-9] $\text{Rh}_2(\text{C}_6\text{H}_6)_2 2\text{Cl}_2$, **M 500.2, m 242° .** This **versatile reagent** is prepared by refluxing $\text{RuCl}_3 \cdot 3\text{H}_2\text{O}$ (2.0g) in EtOH (100ml) with cyclohexadiene (10ml, of 1,3- or 1-4 prepared by reduction of $^*\text{benzene}$ with Na/Liquid NH_3) [Kaiser *Synthesis* 391 1972, DOI: 10.1055/s-1972-21889; Birch & Smith *Q Rev Chem Soc* 12 17 1958, DOI: 10.1039/QR9581200017] for 4 hours. The brown precipitate is filtered off, washed with MeOH and dried *in vacuo* to give the **complex** (1.83g, 95%). The complex is diamagnetic, it is **dimeric** in freezing CHBr_3 , and is sparingly soluble in organic solvents except Me_2SO with which it forms a red complex with the **monomeric** structure $\text{RuCl}_2(\text{C}_6\text{H}_6)(\text{dmsO})$. It dissolves in pyridine, tertiary phosphines, phosphites, tertiary arsines (**L**) which cleave the chlorine bridges of the dimer to form orange or red air stable complexes $\text{RuCl}_2(\text{C}_6\text{H}_6)(\text{L})$ which are **monomeric** in CHCl_3 . Large excess of the ligand **L** should be avoided in the reaction, otherwise the benzene rings may be displaced. Its IR has Ru—Cl stretching bands with ν_{max} at 295, 259 and 248sh cm^{-1} ; and the ^1H NMR (CDCl_3 , TMS) has a singlet at τ_{H} 4.02 which is *ca* 2.0 ppm upfield of the free benzene H resonance. It is a useful **precursor** for the preparation of Ru(II) complexes, some of which are good catalysts. [Bennett & Smith *JCS Dalton Trans* 233 1974, DOI: 10.1039/DT9740000233; Bennett et al. in *Comprehensive Organometallic Chemistry* Wilkinson ed, Vol 4 pp 748-750 Pergamon Press Oxford 1982.]

The complex $\text{Ru}_2(\text{C}_6\text{H}_6)_2$ has also been prepared by evaporating a coat of Ru powder (10-15%) in slow curing 'Araldite' cement on a 1mm tungsten wire heated to $\sim 2400^\circ$. The Ru vapour thus produced is co-condensed with $^*\text{benzene}$ to form thermally unstable orange-yellow crystals of $\text{Ru}_2(\text{C}_6\text{H}_6)_2$ which decompose below 0° . The ^1H NMR (CFCl_3) of the orange crystals show H resonances at τ 4.22 (2H), 4.84 (6H), 5.10 (2H) and 7.11(2H) consistent with the 1—4- η , 1—6- η formulation of the $\text{Ru}_2(\text{C}_6\text{H}_6)_2$ structure. [Timms & King *JCS Chem Commun* 898 1978, DOI: 10.1039/C39780000898.]

Ruthenium dichloride 1,5-cyclooctadiene complex [dichloro- μ -(η^4 -cycloocta-1,5-diene)ruthenium(II), $\text{RuCl}_2(\text{COD})_n$] [50982-12-2] $[\text{Rh}(\text{C}_8\text{H}_{12})\text{Cl}_2]_n$, **M 280.2 (monomer).** Analytically pure complex is obtained by heating a mixture of pure $\text{RuCl}_3 \cdot 3\text{H}_2\text{O}$ (50.0g, 0.2mol) and cycloocta-1,5-diene (50ml, 0.4mol, use a fume hood) in absolute EtOH (400ml) under reflux for 24 hours whereby the brown complex precipitates. After cooling, the solid is filtered off, washed with Et_2O (50ml) and dried *in vacuo* to give pure $[\text{RuCl}_2(\text{COD})]$ ($\sim 52\text{g}$, $\sim 90\%$). It is stable in air, only slightly soluble in organic solvents, reacts by halogen-bridge cleavage and is a useful synthetic **precursor** of ruthenium(II) complexes. [Ohta, Takaya and Noyori *Inorg Chem* 27 566 1988, DOI: 10.1021/ic00276a025; Albers et al. *Inorg Synth* 26 249 1989, DOI: 10.1002/9780470132579.ch44; Bennett et al. in *Comprehensive Organometallic Chemistry* Wilkinson ed., Vol 4 pp 748-750 Pergamon Press Oxford 1982.]

Tetrakis(acetonitrile)copper(I) hexafluorophosphate $[\text{Cu}(\text{MeCN})_4]^+ \text{PF}_6^-$ [64443-05-6] $[(\text{CH}_3\text{CN})_4\text{Cu}]\text{PF}_6$, **M 372.7, m 160° (dec).** In an efficient fume cupboard, 60-65% aqueous hexafluorophosphoric acid (10ml, $\sim 113\text{mmol}$, [16940-81-1], M 146.0) is added in 2ml portions to a stirred suspension of Cu(I) oxide (4.0g, 28mmol) in MeCN (80ml) when a **very exothermic** reaction occurs (use a reflux condenser), causing the solution to boil. After stirring for a further 3 minutes the solution is filtered (use a medium-porosity frit) from some black solid, and $\text{Cu}(\text{MeCN})_4^+ \text{PF}_6^-$ may begin to crystallise out, and is complete on standing at -20° , or by adding an equal volume of dry Et_2O and cooling to 0° . (If some white solid separates with the black solid it should be washed off with MeCN and the MeCN solutions combined and cooled at -20° , or diluted with Et_2O .) The salt is collected by filtration, washed with Et_2O and immediately dissolved in MeCN (100ml), filter off any undissolved blue Cu^{2+} species, add Et_2O (100ml) and set aside at -20° for several hours to crystallise out. The

crystalline complex would need a second recrystallisation if it has a blue tinge. It may be recrystallised from MeCN (80ml) and Et₂O (80ml). The white complex salt is dried *in vacuo* for ~30 minutes immediately after washing with Et₂O to give the **analytically pure salt** (~12.5g, ~60% depending on crystallisation losses). It is a free-flowing white, microcrystalline powder that does not darken on long storage under N₂ or argon. On exposure to air for >1 hour minor surface oxidation occurs and it is **slightly hygroscopic**. The MeCN ligands are not removed easily *in vacuo* at ~25° (but dissociation occurs at ~5 torr/80° and ~25 torr/110°). The IR (Nujol) has ν_{\max} at 2277m and 2305m cm⁻¹ (for MeCN), and 850vs and 557s cm⁻¹ for PF₆⁻. The related salt, **Cu(MeCN)₄⁺ BF₄⁻**, can be prepared similarly by using an equivalent amount of HBF₄ (as 48% aqueous acid [16872-11-0], M 87.9) instead of HPF₆. These complexes are suited for the non-aqueous-media synthesis of Cu(I) complexes [see (IPr)₂Cu⁺ PF₆⁻ and (IPr)₂Cu⁺ BF₄⁻ above]; as well as for preparing other complexes such as [Rh(C₂H₄)₃(MeCN)₂]⁺ + CuCl by reaction of Cu(MeCN)₄⁺ with [Rh(C₂H₄)₂Cl]₂ and C₂H₄ in CH₂Cl₂ [Maspero et al. *J Organomet Chem* **38** C43 1972, DOI: 10.1016/S0022-328X(00)83316-1]. The X-ray structure of the ClO₄⁻ salt showed that the copper is almost ideally **tetrahedral** coordinated with nearly linear MeCN molecules [Csöregi et al. *Acta Cryst* **B31** 314 1975, DOI: 10.1107/S0567740875002634]. [Kubas et al. *Inorg Synth* **19** 90 1979, DOI: 10.1002/9780470132500.ch18; Kubas et al. *Inorg Synth* **28** 68 1990, DOI: 10.1002/9780470132593.ch15].

cis-cis-cis-1,2,3,4-Tetrakis(diphenylphosphinomethyl)cyclopentane [(1R*,2R*,3S*,4S*)-tetrakis(diphenylphosphinomethyl)cyclopentane, Tedicyp] [333380-86-2] **C₅₇H₅₄P₄**, **M 862.9**, **m 79°**. The intermediate **cis-cis-cis-1,2,3,4-tetrakis(boranatodiphenylphosphanylmethyl)cyclopentane** (white crystals **m 149°**, after chromatography on silica gel eluting with AcOEt/petroleum ether, 1:4) is obtained from Ph₂PLi (from Ph₂PCl see [1079-66-9] and Li) and **cis-cis-cis-1,2,3,4-tetrakis(tosyloxymethyl)cyclopentane** in THF under argon (~20°/5 hours), then borane in THF is added. This boranato intermediate formed is added to anhydrous Et₂NH, heated at 55-60° for 10 hours, evaporated to dryness, this process is repeated twice, and the residue is chromatographed on silica gel (eluted with Et₂O/petroleum ether, 1:4) to give white crystals of Tedicyp (98%) with **m 79°**. It is not very stable in air and must be stored under argon. It has ¹H NMR (400MHz, THF-*d*₈) with δ at 7.71-7.23 (40H, m), 2.91-2.52 (2H, m), 2.38-1.72 (10H, m), 1.25 (2H, m); the ³¹P NMR (162MHz, THF-*d*₈) has δ at -16.3 and -17.7; and for ¹³C NMR see references. It is used for preparing the Pd-Tedicyp catalyst with [Pd(η^3 -C₃H₅)Cl]₂ (see above) for promoting allylic alkylations [Laurenti et al. *J Org Chem* **66** 1633 2001, DOI: 10.1021/jo001146j], cross coupling reactions, and catalysing Heck reactions of α - or β -substituted enol ethers with aryl bromides [Battace et al. *Eur J Org Chem* 3122 2007, DOI: 10.1002/ejoc.200700152] with high efficiency. [cf. for efficiency only see Wallow & Novak *J Org Chem* **59** 5034 1994, DOI: 10.1021/jo00096a056]

The tetra-phosphine oxide **cis-cis-cis-1,2,3,4-tetrakis(diphenylphosphinoylmethyl)cyclopentane**, **C₅₇H₅₄O₄P₄**, **M 926.29**, is obtained by oxidising Tedicyp in THF with excess H₂O₂ (~20°/4 days) then purified by chromatography on silica gel (eluting with MeOH/CH₂Cl₂, 5:95) to yield (66%) white crystals of the **tetroxide** with **m 165°**. It has ¹H NMR (400MHz, CDCl₃) with δ at 7.79-7.62 (16H, m), 7.32-7.28 (8H, m), 2.85 (2H, br t, *J* = 12.9Hz), 2.28 (4H, d, *J* = 11.9Hz), 2.26-2.15 (2H, m), 2.10-2.05 (2H, m), 1.88 (2H, br q, *J* = 12.9Hz), 1.60 (1H, dt, *J* = 13.8, 7.5Hz), 1.36 (1H, dt, *J* = 13.8, 10.2Hz); and for ¹³C NMR see reference. [Laurenti et al. *J Org Chem* **66** 1633 2001, DOI: 10.1021/jo001146j.]

R-(+)- and S-(-)- 2,2',6,6'-Tetramethoxy-4,4'-bis(diphenylphosphino)-3,3'-bipyridine (R-(+)- P-Phos and S-(-)- P-Phos) [*R*- 221012-82-4 and *S*- 362524-23-0] **C₃₈H₃₄N₂O₄P₂**, **M 644.6**, **m 261-265°**, [α]_D²⁰ **+98 and -98 (c 1, CHCl₃)**. This group of ligands have been prepared from (3-bromo-2,6-dimethoxy-pyrid-4-yl)(diphenyl)-phosphine oxide by an Ulmann reaction (Cu/DMF/140°) to give **RS- 2,2',6,6'-tetramethoxy-3,3'-bipyridine-4,4'-diphenylphosphine dioxide** (85%) which is resolved into the (+)- and (-)- enantiomers by HPLC on a preparative Diacel AD column (25m x 1.2m), eluting with 2-propanol/hexane (20:80) at a flow rate of 3.0ml/minute: *t_R* = 12.24 minutes, *t_S* = 25.06 minutes. The phosphine oxide groups are then each reduced by Cl₃SiH/Et₃N/toluene at 140° to give *R*-(+)- and *S*-(-)- *R*-Phos in 99% yields. Alternatively, the above *RS*-dioxide, which cannot be resolved easily, is converted into the **5,5'-dibromo derivative** (Br₂/NaOAc/AcOH; 0°-60°, 97%) which can now be resolved *via* the (+) and (-)-di-*O*-benzoyl tartrate salts (in EtOAc/CHCl₃, 1/1) to provide the enantiomeric **5,5'-dibromo-dioxides** that are de-brominated, and reduced (Cl₃SiH/Et₃N/toluene during an extended period from 25° to 140°) to give *R*-(+)- and *S*-(-)- *P-Phos* in 91% yields. [Pai et al. *J Am Chem Soc* **122** 11513 2000, DOI: 10.1021/ja000163n.] This ligand is employed in transition-metal-catalysed

asymmetric reactions including hydrogenation, hydrosilylation and C—C bond formation [Wu & Chan *Acc Chem Res* **39** 711 2006, DOI: 10.1021/ar0680015.]

***R*-(+)- and *S*-(-)- 2,2',6,6'-Tetramethoxy-4,4'-bis[di(3,5-xylyl)phosphino]-3,3'-bipyridine** [*R*-(+)- Xylyl-P-Phos and *S*-(-)- Xylyl-P-Phos) [*R*- 442905-33-1 and *S*- 443347-10-2] $C_{46}H_{50}N_2O_4P_2$, *M* 756.9, *m* (158-162°), 190-194°, $[\alpha]_D^{20}$ +125 and -125 (c 1, $CHCl_3$). The Xylyl-P-Phos ligands are prepared in a slightly different way from the P-Phos ligands above. Thus 3-bromo-2,6-dimethoxy-pyridine is converted into (3-bromo-2,6-dimethoxy-pyrid-4-yl)di(3,5-dimethylphenyl)phosphine [**a**: LDA/THF/-78°; **b**: (3,5-dimethylphenyl)₂PCl/-78°] in 56% yield, then oxidised ($H_2O_2/Me_2CO/0^\circ$) to the phosphine oxide (99%), followed by an Ullmann reaction as above (Cu/DMF/140°) to give *RS*- 2,2',6,6'-tetramethoxy-3,3'-bipyridine-4,4'-di(3,5-dimethylphenyl)phosphine dioxide (85% yield) that is resolved via (-)- and (+)- di-*O*-benzoyl tartrate salts (**a**: see above fractional crystallisation; **b**: 10% aqueous NaOH) to give the *R*- (75%) and *S*- (88%) Xylyl-P-Phos dioxides which are reduced (Cl_3SiH/Et_3N /toluene) to *R*(+)- and *S*- Xylyl-P-Phos in >90% yields. The absolute configuration *S*- was determined from the X-ray crystal structure of the diastereoisomerically pure 1:1 salt of (-)-Xylyl-P-Phos and (+)-(2*S*,3*S*)-dibenzoyl-*O*-tartrate. The latter have ¹H NMR (500MHz, $CDCl_3$) with δ_H at 2.20 (s, 12H, $PhCH_3$), 2.25 (s, 12H, $PhCH_3$), 3.37 (s, 6H, O CH_3), 3.83 (s, 6H, O CH_3), 6.06 (d, *J* = 1.5Hz, 2H, PyH), 6.79-6.92 (m, 12H, PhH); the ³¹P NMR (200MHz, $CDCl_3$) with δ_P at -11.99; the C, H and N elemental analyses fit for the latter empirical formula; and for ¹³C NMR and LSMS see reference. [Wu et al. *Tetrahedron Lett* **43** 1539 2002, DOI: 10.1016/S0040-4039(02)00062-X.] This ligand is employed in transition-metal-catalysed asymmetric reactions including hydrogenation, hydrosilylation and C—C bond formation [Wu & Chan *Acc Chem Res* **39** 711 2006, DOI: 10.1021/ar0680015.]

Tricyclohexylphosphine (PCy₃) [2622-14-2] $C_{18}H_{33}P$, *M* 280.4, *m* 76-78°, 77°, 82°, 82-83°, pK_{Est} ~9.5. It recrystallises from EtOH [Boere et al. *J Am Chem Soc* **109** 7781 1987, DOI: 10.1021/ja00259a029]. [Beilstein **16** IV 947.] Used in organometallic chemistry and characterised by its high basicity and large ligand cone angle. When applied with Ni it is a critical intermediate in forming cyclopentane compounds [Ogoshi et al. *J Am Chem Soc* **128** 5350 2006, DOI: 10.1021/ja060220y], when used with Pd it mediates coupling of malononitrile with aryl halides [Schnyder et al. *Synlett* 3167 2006, DOI: 10.1055/s-2006-944215], and when used with a Pd(0)-triolefinic macrocycle catalyst it allows Suzuki coupling of aryl bromides and chlorides [Moreno-Manas et al. *Synlett* 3001 2006, DOI: 10.1055/s-2006-948173].

6*RS*(±)-4,4,6-Trimethyl-2-vinyl-1,3,2-dioxaborinane (vinylboronic acid 2-methyl-2,4-pentanediol ester) [4627-10-5] $C_8H_{15}BO_2$, *M* 154.0, *b* 50-55°/0.46mm, *d*²⁵ 0.893g/ml, *n*_D²⁰ 1.429. It is prepared by a modification of a published method [Hoffmann & Landmann *Chem Ber* **119** 2013 1986, DOI: 10.1002/cber.19861190622] where *n*-octanol is replaced by 2-methylpentane-2,4-diol (MPD). To a solution of (MeO)₃B (30ml, 270mmol) in dry Et₂O (125ml) at -78° is added over 45 minutes a 2.1M solution of vinylmagnesium chloride (118ml, 248mmol) in THF with stirring, and allowing the temperature to rise to ~25°. The mixture is then acidified, under cooling, with aqueous HCl (concentrated, 21ml, 0.25mol in 62.5ml of H₂O), phenothiazine (0.1g, as stabiliser) is added, the phases are separated, the aqueous phase is extracted with MPD (3 x 50ml). The combined organic phases are evaporated *in vacuo* from a bath at 80° and the residual oil is distilled at high vacuum to give pure *borinane* in ~60% yield. [Note that all the liquids in the reaction mixture distil at lower temperatures than the desired racemic vinyl-dioxaborinane.] It is superior to vinylboronic pinacol ester in preparation, storage, stability, and reactivity, providing improved selectivity for Heck versus Suzuki coupling with aryl and heteroaryl bromides and iodides [Lightfoot et al. *Tetrahedron Lett* **44** 7645 2003, DOI: 10.1016/j.tetlet.2003.08.032]. By adopting different reaction conditions with Pd(PPh₃)₄/*t*-BuOK, selective Suzuki-Miyaura coupling with a range of aryl and heteroaryl halides (Cl, Br and I) was achieved [Lightfoot et al. *Synlett* 529 2005, DOI: 10.1055/s-2005-862354; cf. Lightfoot et al. *Org Biomol Chem* **3** 3167 2005, DOI: 10.1039/B507900D].

Tri-*tert*-butylphosphonium tetrafluoroborate [(*tert*-Bu)₃PH⁺ BF₄⁻] [131274-22-1] [(CH₃)₃C]₃P⁺ HBF₄⁻, *M* 290.1, *m* 261°(dec), pK^{25} 11.4 (phosphine). The salt is prepared by adding HBF₄ (48% aqueous solution, 1.0ml, 7.6mmol) to a solution of pure (*t*-Bu)₃P (225mg, 1.11mmol, see above) in CH_2Cl_2 (15ml), stirring vigorously for 5 minutes, the organic layer is separated, dried (MgSO₄), filtered and evaporated to dryness to give analytically pure salt (302mg, 94%) as a white powder. Its ¹H NMR (400MHz, $CDCl_3$) has δ_H at 6.07 (d,

$^1J_{\text{PH}} = 465\text{Hz}$, 1H), 1.65 (d, $^3J_{\text{PH}} = 15.3\text{Hz}$, 27H); the $^{31}\text{P}\{^1\text{H}\}$ NMR (121MHz, CDCl_3) has δ_{P} at 51.7; and for IR, and ^{13}C NMR see references. The salt is more *air-stable* than the free phosphine, and can be heated at 120° for 24 hours without decomposition (no change in NMR spectra), or loss of catalytic activity with transition metal complexes. It is *not hygroscopic*. In the presence of $\text{Pd}_2(\text{dba})_3$ in THF it catalyses Suzuki cross-coupling reactions between aryl halides and arylboronic acids, Stille cross-coupling reactions between aryl halides and tributylSn compounds, and Heck reactions between aryl halides and olefins [Netherton & Fu *Org Lett* **3** 4295 2001, DOI: 10.1021/ol016971g]. The salt has also been used successfully in the Heck coupling [with $\text{PdCl}_2(\text{COD})$, LiCl, Cy_2NMe] of non-activated alkenyl tosylates and phosphates as substrates with electron-poor alkenes and styrene derivatives [Hansen et al. *Angew Chem Int Ed* **45** 3349 2006, DOI: 10.1002/anie.200600442]. This ligand is useful in the Pd-catalysed enantioselective α -arylation of *N*-BOC-pyrrolidine [Campos et al *J Am Chem Soc* **128** 3538 2006, DOI: 10.1021/ja0605265].

Vinyl MIDA boronate {vinylboronic acid MIDA ester, 6-methyl-2-vinyl-1,3,6,2-dioxazaborocane-4,8-dione, [N-[(carboxy- κO)methyl]-N-methylglycinato(2-)- $\kappa\text{N},\kappa\text{O}$]ethenyl boron } [1104636-73-8] $\text{C}_7\text{H}_{10}\text{BNO}_4$, M 183.0, m 152-156°. The boronate is prepared in Schlenk equipment by adding dropwise vinyltrimethylsilane (4.49ml, 31.5mmol, freshly distilled see [754-05-2]) to a stirred solution of BBr_3 (1.0M in CH_2Cl_2 , 30mmol) at 0°, and maintained at this temperature for 20 minutes, then allowed to warm to ~25° with stirring for a further 2 hours. This mixture is added *via* a cannula to a stirred suspension of MIDA sodium salt (5.73g, 30.0mmol, see above) in MeCN (50ml) at 0°, at such a rate as to keep the temperature below 5°; then the temperature is allowed to warm to ~25° while stirring for 1 hour. The resulting white suspension is filtered through a pad of Celite and the filtrate cake is extracted 3 times with Me_2CO . Et_2O is added to the combined orange filtrates which allowed the colourless free flowing **vinyl MIDA boronate** (4.74g, 86%) to crystallise out. On TLC it has R_{F} 0.26 (Merck silica gel plate grade 9385, 60Å, 230-400 mesh, with EtOAc). Its ^1H NMR (500MHz, $\text{Me}_2\text{CO}-d_6$, $\delta = 2.04$ centre line) has δ_{H} at 5.96 (dd, $J = 19.0, 13.5\text{Hz}$, 1H), 5.72-5.63 (m, 2H), 4.21 (d, $J = 17.0\text{Hz}$, 2H), 4.01 (d, $J = 17.0\text{Hz}$, 2H), 3.0 (s, 3H); and for IR, ^{13}C NMR and EI-MS see reference. Single crystal X-ray analysis confirmed the predicted structure as having a pyramidalised boron centre. The vinylboronate shows no signs of deterioration when kept on the benchtop in air for more than 3 months. It is a versatile reagent that can be prepared on a multigram scale. It readily reacts with $\text{CH}_2\text{N}_2/\text{Pd}(\text{OAc})_2$ to provide **cylopropyl MIDA boronate** [Et_2O , 0-23°, 1 hour, 93%], and with *m*-CPBA it yields **oxiranyl MIDA boronate** [CH_2Cl_2 , 0-23°, 18 hours, 74%] [Uno, Gillis and Burke *Tetrahedron* **65** 3130 2009, DOI:10.1016/j.tet.2008.11.010]; and can be used in Suzuki, Heck, Stille, Negishi and Sonogashira couplings and Miyaura borylations [Gillis & Burke *Aldrichimica Acta* **42** 17 2009].

Xantphos [4,5-Bis(diphenylphosphino)-9,9-dimethylxanthene] [161265-03-8] $\text{C}_{39}\text{H}_{32}\text{OP}_2$, M 578.2, m 221-222°, 224-228°. Xantphos is prepared, using Schlenck techniques under N_2 , by adding *sec*-BuLi (13.6ml of 1.3M in 98/2 cyclohexane/hexane, 14.3mmol) dropwise to a stirred solution of 9,9-dimethylxanthene (1.0g, 4.76mmol, see [19814-75-6]) and TMEDA (2.3ml, 13.3mmol) in dry Et_2O (54ml), and stirring for 16 hours at ~25° to form the 4,5-dilithium derivative. To this mixture is added a solution of chlorodiphenylphosphine (2.8ml, 14.3mmol, see [1079-66-9]) in hexane (16ml) dropwise, and stirring is continued for 16 hours, then the solvent is removed *in vacuo*. The residual thick oil is dissolved in CH_2Cl_2 , washed with H_2O , dried (MgSO_4), filtered, the solvent is removed *in vacuo*, the residue is washed with hexanes, and recrystallised from propan-1-ol to give **xantphos** (2.05g, 74.6%) as an air stable yellow-white powder. Its ^1H NMR (300MHz, CDCl_3) has δ at 7.4 (dd, 2H, $J = 7.8, 1.0\text{Hz}$, CPCHCH), 7.15-7.26 (aryl, 20H, $[\text{P}(\text{C}_6\text{H}_5)_2]_2$), 6.96 (t, 2H, $J = 7.7\text{Hz}$, CHCHCH), 6.54 (dd, 2H, $J = 7.4, 1.4\text{Hz}$, CHCHCC), 1.65 (s, 6H, CH_3); the $^{31}\text{P}\{^1\text{H}\}$ NMR (121.5MHz, CDCl_3 , referenced to external 85% H_3PO_4) has δ at -17.5; and for IR, ^{13}C NMR and HRMS see references. Out of several diphenylphosphino ligands studied, xantphos induced the *highest selectivity* in rhodium-catalysed hydroformylation of 1-alkenes to form linear aldehydes. The hydroformylation is carried out in a stainless steel bomb under pressure of ~10 bars of CO/H_2 (1:1) in toluene with phosphorus ligand and $\text{Rh}(\text{acac})(\text{CO})_2$ as metal source. [Kranenburg et al. *Organometallics* **14** 3081 1995, DOI: 10.1021/om00006a057.] The synthesis of heterocycles by C—N cross coupling of bromothiophenes with amino pyridines is achieved with this ligand by Pd catalysis [Begouib et al. *Synthesis* 2794 2006, DOI: 10.1055/s-2006-942510]. This ligand is also used in a ruthenium-catalysed alkylation of active methylene compounds with alcohols [Slatford et al. *Tetrahedron Lett* **47** 6787 2006, DOI: 10.1016/j.tetlet.2006.07.069], and as a metal chelating ligand for catalysis [Manley & Bilodeau *Org Lett* **6** 2433 2004, DOI: 10.1021/ol049165t; and Gao et al. *Org Lett* **6** 1837 2004, DOI: 10.1021/ol049440b].

CHIRAL AUXILIARIES

—These compounds are involved in reactions, which direct stereospecificity and are decomposed, e.g. by hydrolysis, reduction, to generate the desired chiral products and the remains of the original auxiliary which can be recycled. Unlike the case of catalysis, the auxiliary is involved stoichiometrically and is not recycled during the reaction. It is one of the products of the reaction and can, *via* other reactions if necessary, be used to regenerate the original auxiliary. These have been used for decades and are of necessity chiral molecules themselves. The auxiliaries have to be linked to the molecule that has a reactive group which can be operated on, e.g. carbonyl, to generate an asymmetric centre, i.e. a chiral alcohol. This section includes more recently available auxiliaries, although many useful ones such as cinchonine, cinchonidine, borneols, camphors, menthol derivatives, substituted sugars, substituted chiral heterocycles (e.g. pyrrolidines), sterols etc. will be found scattered in Chapters 3, 4 and 6. [see Evans *Aldrichimica Acta* **15** 23 1982, Ager et al. *Chem Rev* **96** 835 1996, DOI: 10.1021/cr9500038; Ager et al. *Aldrichimica Acta* **30** 3 1997, Mukaiyama *Aldrichimica Acta* **29** 59 1996, K. Rück-Braun and H. Kunz *Chiral Auxiliaries in Cycloaddition Reactions* Wiley-VCH, Weinheim 1999, ISBN 3-527-29386-8.]

***R*-(\pm)- and *S*-(\pm)- Acetyl-cyclopentadienyl-ironcarbonyl triphenylphosphine complex** [*R* 36548-61-5 ; *S* 36548-60-4; *RS* 12101-02-9] (η^5 -C₅H₅)Fe(CO)(CH₃CO)(PPh₃) **M 454.3, m 142° (for +), m 140° (for -), m 145° (for \pm)**; *R*-[α]₄₃₆²⁷ -288, [α]₅₄₆²² -288 (c 0.045, C₆H₆), *S*-[α]₅₄₆²² +288 (c 0.045, C₆H₆). This racemic ‘chiral’ auxiliary can be made on a large scale from [η^5 -(C₅H₅)Fe(CO)₂]₂ [38117-54-3] which is cleaved with Na/Hg to give the anion [η^5 -(C₅H₅)Fe(CO)₂]⁻ (Na⁺) that is methylated to η^5 -(C₅H₅)Fe(CO)COMe (m **78-82°**, caramel coloured waxy crystals from sublimation *in vacuo* onto an ice-cooled glass finger) with MeI [Piper & Wilkinson *J Inorg Nucl Chem* **3** 104 1956, DOI:10.1016/0022-1902(56)80073-0; Aktogu et al. *J Organomet Chem* **262** 49 1984, DOI:10.1016/S0022-328X(00)99122-8; King *Organometallic Synthesis Vol 1*, Academic Press, NY, p 145 1965]. This methyl complex provides the racemic title compound when treated with PPh₃. Thus (C₅H₅)Fe(CO)₂Me (1.0g, 5mmol) and PPh₃ (1.3g, 5mmol) in redistilled THF (10ml) are refluxed (65°) under N₂ until the Fe-Me band (IR: strong C-H deformation band at 1170 cm⁻¹ in CS₂) disappears (~48 hours). The solution is filtered, the solvent is evaporated (to ~20mm), the residue is dissolved in pentane (10ml), passed through an Al₂O₃ column (5 x 20cm), and only one band (yellow orange to orange) is eluted with pentane which, on evaporation (to ~20mm), provides the analytically pure orange (\pm)- η^5 -C₅H₅Fe(CO)(CH₃CO)(PPh₃), **m 145°** in ~98% yield. This reaction is solvent dependent, i.e. no reaction occurs in boiling hexane (68°), and is only 50% complete in boiling Et₂O (34°) after 48 hours. The solid is **stable in air**, is soluble in organic solvents, e.g. pentane, hexane, Et₂O, THF, CH₂Cl₂, CHCl₃, and *C₆H₆, but is insoluble in MeOH or H₂O. Solutions in CHCl₃, and *C₆H₆ decompose rapidly in air to produce brown intractable solids. Hence these solutions should be prepared and used under N₂ or argon. Its IR (CHCl₃) has bands with ν_{\max} at 1598 (s, MeC=O) and 1920 (vs, br, Fe carbonyl) cm⁻¹; and the ¹H NMR (60MHz, CDCl₃, external TMS) has δ_{H} at 7.59 (m, Ph, 15H), 4.69 (s, C₅H₅, 5H) and 2.52 (s, COMe, 3H). [Bibler & Wojcicki *Inorg Chem* **5** 889 1966, DOI: 10.1021/ic50039a037; Butler et al. *Inorg Chem* **6** 2074 1967, DOI: 10.1021/ic50057a032.]

The versatility of this acetyl auxiliary has prompted its **optical resolution** into the pure enantiomers which proved to be very good chiral auxiliaries for preparing a variety of optically active molecules where high stereo control is achieved. Two independent resolutions were achieved, both involving 1*R*,2*S*,5*R*-(\pm)-2-isopropyl-5-methylcyclohexan-1-ol (*R*-*l*-menthol). The **first resolution** is from the reaction of sodium *R*-menthylate (NaOC₁₀H₁₉, *R*-menthyl refers to the radical produced by loss of the hydroxyl group) and (\pm)-(η^5 -C₅H₅)Fe(CO)₂(PPh₃)⁺PF₆⁻ [see below recovered from hydrolysis, decomplexation, of derivatives of the racemic title compound Aktogu et al. *J Organomet Chem* **262** 49 1984, DOI:10.1016/S0022-328X(00)99122-8] to give a diastereoisomeric mixture of (+)-(η^5 -C₅H₅)Fe(CO)(PPh₃)-(-)COOC₁₀H₁₉ {[α]₅₈₉²⁰ +30, [α]₅₇₉²⁰ +35, [α]₅₄₆²⁰ +70, [α]₄₃₆²⁰ -1450 (in 10⁻³M *C₆H₆)}, and (-)-C₅H₅Fe(CO)(PPh₃)-(-)COOC₁₀H₁₉ {[α]₅₈₉²⁰ -75, [α]₅₇₉²⁰ -80, [α]₅₄₆²⁰ -120, [α]₄₃₆²⁰ +1550 (in 10⁻³M *C₆H₆)}, (together with NaPF₆) which were separated by recrystallisation from pentane, with the latter being more soluble. [Brunner & Schmidt *J Organomet Chem* **21** P53 1970, DOI: 10.1016/S0022-328X(00)83614-1] Then the ester with [α]₅₄₆²⁰ -120 (560mg, 0.93mmol) in THF (20ml) at -30°, is treated dropwise with MeLi (1ml of 1.5M Et₂O solution), followed by stirring at -30°/1 hour, then at ~25°/1 hour. After quenching the reaction and evaporating, the brown residue is extracted into *C₆H₆, and purified through a column of Al₂O₃/3%H₂O and eluted with *C₆H₆. The greenish zone gives a menthol-free yellow solid, which on sublimation provides **analytically pure orange (+)- η^5 -(C₅H₅)Fe(CO)(CH₃CO)(PPh₃)** (47mg, 11% yield), **m 142°**, [α]₅₄₆²⁰ -228 (in 10⁻³M *C₆H₆), found M 465 (osmometry in *C₆H₆). Similarly the menthyl

ester with $[\alpha]_{546}^{20} +70$ gives **pure** $(-)\text{-}\eta^5\text{-(C}_5\text{H}_5\text{)Fe(CO)(CH}_3\text{CO)(PPh}_3\text{)}$ (80mg, 30% yield), **m 140°**, $[\alpha]_{546}^{20} +227$ (in $10^{-3}\text{M } ^*\text{C}_6\text{H}_6$), found M 464 (osmometry in $^*\text{C}_6\text{H}_6$). From the CD and ORD spectra it was concluded that these reactions occurred with **inversion of configuration** at the tetrahedral iron centre [Brunner & Schmidt *J Organomet Chem* **36** C18 1972, DOI: 10.1016/S0022-328X(00)85109-8].

In the **second resolution** *R*-($-$)-chloromethylmenthyl ether (**R**- is the configuration at 1-C-OH of menthol, see [26127-08-2]) was reacted with the lithium salt of $(\pm)\text{-}\eta^5\text{-(C}_5\text{H}_5\text{)Fe(CO)(CH}_3\text{CO)(PPh}_3\text{)}$ (generated with *n*-BuLi) to give the diastereomeric ethers $(-)\text{-}\eta^5\text{-(C}_5\text{H}_5\text{)Fe(CO)(PPh}_3\text{)(COCH}_2\text{CH}_2\text{O-(-)-C}_{10}\text{H}_{19}$ $\{[\alpha]_{\text{D}}^{20} +65$ (c 0.4, $^*\text{C}_6\text{H}_6\text{)}\}$, and $(+)\text{-}\eta^5\text{-(C}_5\text{H}_5\text{)Fe(CO)(PPh}_3\text{)(COCH}_2\text{CH}_2\text{O-(-)-C}_{10}\text{H}_{19}$ $\{[\alpha]_{\text{D}}^{20} -150$ (c 0.4, $^*\text{C}_6\text{H}_6\text{)}\}$ which are separable by chromatography and distinguishable by ^1H NMR (300MHz). Crystals of the latter diastereoisomer were subjected to x-ray crystallographic analysis which revealed that the absolute configuration at the tetrahedral **Fe centre** was **R**, by virtue that the absolute configuration of the *R*-menthyl moiety had been established. Since the original formation of these ethers occurs with alteration in the configuration at the Fe centre, then the desired *S*-($+$)- and *R*-($-$)- configurations of $(\pm)\text{-}\eta^5\text{-(C}_5\text{H}_5\text{)Fe(CO)(CH}_3\text{CO)(PPh}_3\text{)}$ are established. It was also shown that Brunner & Schmidt's reactions of the *R*-menthyl esters with *n*-BuLi to provide the title enantiomers occurred with **complete inversion of configuration** (as determined by 500MHz ^1H NMR spectroscopy). The enantiomers can be discriminated in solution (9mg in 700 μl of CDCl_3) containing the chiral shift reagent Eu(tfc) $\{\text{tris[3-(trifluoromethyl)hydroxymethylene-}(+)\text{-camphorato] europium (III)}$, see [34834-11-0] (0.48mg in 8 μl) where a clear separation of the methyl singlets in the 300MHz ^1H NMR spectra occurs: the *S*-($+$)- enantiomer has δ at 2.66 and the *R*-($-$)-enantiomer has δ at 2.58 (from TMS). [For absolute configuration and optical purity see Davies et al. *JCS Chem Commun* 607 1986, DOI: 10.1039/C39860000607.] The chiral auxiliary properties of the complexes $\eta^5\text{-C}_5\text{H}_5\text{Fe(CO)(CH}_3\text{CO)(PPh}_3\text{)}$ are displayed by deprotonation with *n*-BuLi (isoPr $_2\text{NLi}$ in THF has also been used) to give the blood-red enolate which undergoes a variety of reactions and sequences of reactions (homochiral synthesis) that are highly stereoselective. These include alkylation, homochiral succinylation, asymmetric aldol reactions, formation of α,β -unsaturated iron acyls and homochiral dienolates, asymmetric Michael addition reaction, asymmetric synthesis of β -amino acids and β -lactams, asymmetric synthesis of cyclopropanecarboxylic acids, asymmetric synthesis of chiral sulfoxides, and chirality recognition for homochiral synthesis. [Davies *Aldrichimica Acta* **23** 31 1990.] Decomplexation at the end of the reactions can be achieved by oxidative cleavage (Br/MeOH , H_2O or $n\text{-BuNH}_2$, NBS/EtOH) to provide the acid, amide or ester, also by alcoholysis (to provide different esters) from which the products can be isolated. From the aqueous solution $\eta^5\text{-C}_5\text{H}_5\text{Fe(CO)}_2\text{(PPh}_3\text{)}^+\text{PF}_6^-$ (evaporated to 20ml, from 1mmol of complex product) can be recovered by addition of NH_4PF_6 (0.6g, 3.7mmol), stirring for 1 hours, evaporating to dryness, and the residue is extracted with CH_2Cl_2 (3 x 10ml). The combined extracts are evaporated to 5ml and Et_2O is added to give pale yellow crystals of the complex salt (0.38g, 65%, **m 186°**). It can also be recrystallised from $\text{Me}_2\text{CO/Et}_2\text{O}$ or $\text{Me}_2\text{CO/hexane}$. Its ^1H NMR (60MHz, Me_2CO , TMS) has δ_{H} at 5.62 (d, $J_{\text{PH}} = 1.5\text{Hz}$, C_5H_5) and 7.68 (m, Ph, 15H); and for IR see references. [Aktogu et al. *J Organomet Chem* **262** 49 1984, DOI: 10.1016/S0022-328X(00)99122-8; Treichel et al. *Inorg Chem* **5** 1177 1966, DOI: 10.1021/ic50041a022.]

1*R*,2*S*,5*R*-($-$)-Chloromethylmenthylether [*R*-($-$)-chloromethylmenthyl ether] [26127-08-2] **M 204.7**, **b 62°/0.1mm**, **160-162°/13-16mm**, **d 0.9821**, **d** 25 **0.994**, **n** $_{\text{D}}^{20}$ **1.467**, **n** $_{\text{D}}^{27}$ **1.465**, **$[\alpha]_{\text{D}}^{24}$ -177.0** (c 1, CH_2Cl_2), and **1*S*,2*R*,5*S*-($+$)-chloromethylmenthyl-ether** [*S*-($+$)-chloromethylmenthyl ether] [103128-76-3] **C** $_{11}\text{H}_{21}\text{ClO}$, **M 204.7**, **b 72°/0.4mm**, **d** 21 **0.994**, **n** $_{\text{D}}^{20}$ **1.467**, **$[\alpha]_{\text{D}}^{24}$ +181.0** (c 2, CH_2Cl_2). The chloro methyl derivatives of *R*-(*l*)- and *S*-($+$)- menthol respectively are useful chiral auxiliary resolving agents (see previous entry) as they can be readily removed by mild hydrolysis and the recovered menthol can be recycled. The *R*-(*l*)-enantiomer is prepared by melting *l*-menthol (100g, 0.64moles) on a water bath and stirring vigorously with 40% w/w aqueous formalin (50g, 0.67moles) while HCl gas is bubbled through. The mixture warms up at first and has to be cooled at 0° until no more gas dissolves. The clear mixture separates into two layers, the upper layer is removed and the lower layer is dried over Na_2SO_4 . This is filtered and fractionated under reduced pressure from some unreacted menthol, formaldehyde trimer and HCl to give the **chloromethyl ether** (120-130g, 91-99%) as a slightly refracting oil which distils steadily at **160-162°/16mm** (and **62°/0.1mm**), but it decomposes at *ca* 230°/atm. The pure ether has **$[\alpha]_{\text{D}}^{21}$ -172.75** (c 6.78 CHCl_3). On heating in EtOH/charcoal it forms the **dimethylmethylyal** **C** $_{10}\text{H}_{19}\text{OCH}_2\text{OC}_{10}\text{H}_{19}$ derivative which crystallises from aqueous EtOH in greasy looking scales or from Et_2O in colourless needles **m 57°** (**b 337°/atm**), with **$[\alpha]_{\text{D}}^{24}$ -77.94** (c 0.8 EtOH). The **chloromethylmenthyl ethers** hydrolyse in H_2O to menthol, HCHO and HCl so they should be stored in an inert atmosphere in the cold [Wedekind *Chem Ber* **34** 813 1901, DOI: 10.1002/cber.190103401142; Deutsches Reichspatent No. 119008,

D.R.P. 189331 *Chem Zentralblatt* **1** 184 1908]. [*Beilstein* **6** H 32, **6** I 21.]

4*R*-(+)-4-Isopropyl-5,5-dimethyl-2-oxazolidinone (**4*R*-(+)-4-isopropyl-5,5-dimethyl-1,3-oxazolidin-2-one**) [223906-38-5] and **4*S*-(-)-4-isopropyl-5,5-dimethyl-2-oxazolidinone** (**4*S*-(-)-4-isopropyl-5,5-dimethyl-1,3-oxazolidin-2-one**) [168297-86-7] $\text{C}_8\text{H}_{15}\text{NO}_2$, **M 157.2**, **m 86-87°**, $[\alpha]_{\text{D}}^{20}$ (+) and (-) **47** (c **1**, H_2O). The starting chiral ethanolamine is prepared from the methyl esters of chiral α -aminoacids (*R*- or *S*- valine in this case) by reaction with $\text{MeMgI/Et}_2\text{O}$ to give chiral 2,2-dimethylvalinol which is converted into the desired chiral 2-oxazolidinones with $(\text{EtO})_2\text{CO/K}_2\text{CO}_3$ (as in the following entry), or by reaction with CCl_3COCl (in pyridine) or carbonyl-diimidazole (in CH_2Cl_2) as carbonyl equivalents. Similarly by using the esters of chiral alanine, norleucine or α -phenylglycine the respective optically active 5,5-dimethyl-2-oxazolidinones where the 4-isopropyl group is replaced by methyl, *n*-butyl and phenyl groups respectively can be prepared. These auxiliaries have been named '*second series Quat auxiliaries*', the first series being substituted 3,3-dimethyl-2-pyrrolidinones [Davies et al. *Tetrahedron Lett* **35** 2369 1994, DOI: 10.1016/0040-4039(94)85222-7; 2373 1994, DOI: 10.1016/0040-4039(94)85223-5]. The present 2-oxazolidinones can be *N*-acylated, (e.g. with BuLi , then RCH_2COCl or MeCH=CHCOCl), and the *N*-acyl moieties can be the targets for highly stereoselective enolate alkylation and conjugate addition reactions. The products can be hydrolysed (e.g. with LiOH , $\text{THF/H}_2\text{O}$ 3:1, 0° at $\sim 25^\circ$) to provide the respective chiral acids and regenerated oxazolidin-2-one. The *gem*-dimethyl groups enhance the face-stereoselective shielding of the attached *N*-acyl moiety leading to very high diastereomeric excess in the products. [Davies & Sanganee *Tetrahedron Asymm* **6** 671 1995, DOI: 10.1016/0957-4166(95)00057-V; cf. review by Mukaiyama *Aldrichimica Acta* **29** 59 1996.]

4*R*-(+)-4-Isopropyl-2-oxazolidinone (**4*R*-(+)-4-isopropyl-1,3-oxazolidin-2-one**) [95530-58-8] and **4*S*-(-)-4-isopropyl-2-oxazolidinone** (**4*S*-(-)-4-isopropyl-1,3-oxazolidin-2-one**) [17016-83-0] $\text{C}_6\text{H}_{11}\text{NO}_2$, **M 129.2**, **m 70-71.5°, 70-72°, 71-72°, *R*-** $[\alpha]_{\text{D}}^{20}$ **+18**, ***S*-** $[\alpha]_{\text{D}}^{20}$ **-18**, (c **6**, EtOH). These compounds are *Evans' type* of chiral auxiliaries. The *S*-enantiomer was prepared by stirring 1mol of *S*-valinol, 1.1mol of diethyl carbonate and 1mol of anhydrous K_2CO_3 at 125-126° (internal temperature) until 2.0mols of EtOH had distilled off (4-6 hours). The cooled mixture (to 20°) is dissolved in Et_2O , filtered through a pad of Celite to remove the K_2CO_3 , evaporated to a small volume and cooled slowly to 0° when the oxazolinone crystallises as white needles (**m 69-70°**, 85-95% yield). It is soluble in CH_2Cl_2 and recrystallises from hexanes/ EtOAc (4:1, v/v) by allowing it to stand at 6° overnight. On TLC (0.25mm silica gel 60-F₂₄₅ plates) it has R_F 0.19 (hexanes/ EtOAc 6:4, v/v). It has $[\alpha]_{589}^{20}$ -16.6, $[\alpha]_{577}^{20}$ -17.3, $[\alpha]_{546}^{20}$ -20.2, $[\alpha]_{435}^{20}$ -37.3, $[\alpha]_{365}^{20}$ -63.7 (c 5.81, EtOH); the IR (CH_2Cl_2) has ν_{max} at 1240, 1400, 1760, 2980, 3060, 3240-3340, 3480 cm^{-1} ; and the ^1H NMR (90MHz, CDCl_3) has δ at 6.7 (br s, 1H, NH), 4.42 (t, J = 8.6Hz, 1H, $\text{C}_5\text{-H}$), 4.07 (d of d, J = 8.5, 6.5Hz, 1H, $\text{C}_5\text{-H}$), 3.58 (d of t, J = 8.6, 6.5Hz, 1H, $\text{C}_4\text{-H}$), 1.9-1.6 (m, 1H, $\text{C}_4\text{-H}$), 0.95 (overlapping d's, J = 6.0Hz, 6H, $\text{CH}(\text{CH}_3)_2$). [Evans et al. *J Org Chem* **50** 1830 1985, DOI: 10.1021/jo00211a008; Evans et al. *J Am Chem Soc* **103** 2127 1981, DOI: 10.1021/ja00398a058.]

The **4*S*-(+)-4-isopropyl-3-propionyl-1:3-oxazolidine-2-one derivative** [77877-19-1] $\text{C}_9\text{H}_{15}\text{NO}_3$, **M 185.2**, has **b 102-106°/0.75mm**, d^{25} **1.094g/ml**, n_{D}^{20} **1.464**, $[\alpha]_{\text{D}}^{25}$ **+93** (c **8.7**, CH_2Cl_2). It is an auxiliary reagent also used as a chiral ligand in dirhodium (II) complexes [Doyle et al. *J Am Chem Soc* **115** 9968 1993, DOI: 10.1021/ja00075a013], and in aldol addition reactions [Pridgen et al. *J Org Chem* **58** 5107 1993, DOI: 10.1021/jo00071a020]. For a reviews on chiral auxiliaries for asymmetric synthesis see Ager et al. *Aldrichimica Acta* **30** 3 1997, and Mukaiyama *Aldrichimica Acta* **29** 59 1996.

4*R*-(+)-4-Isopropyl-2-oxazolidinethione [**4*R*-(+)-4-isopropyl-1,3-oxazolidin-2-thione**, (**4*R*)-4-(1-methylethyl)-2-oxazolidinethione**] [1217463-35-8] and **4*S*-(-)-4-isopropyl-2-oxazolidinethione** (**4*S*-(-)-4-isopropyl-1,3-oxazolidin-2-thione**) [104499-08-3] $\text{C}_6\text{H}_{11}\text{NOS}$, **M 122.16-51-545.2**, **m 48-52°, 51-53°, *R*-** $[\alpha]_{\text{D}}^{20}$ **+23.2**, ***S*-** $[\alpha]_{\text{D}}^{20}$ **-23.2**, (c **0.4**, CHCl_3). These compounds are *Evans' type* of chiral auxiliaries. The *S*-(-)-enantiomer is synthesised by adding CS_2 (0.9ml, 15mmol) to a solution of *S*-valinol (10mmol [cf. 2026-48-4]) in aqueous Na_2CO_3 (20ml) and stirring at 100° (bath at 110° under efficient reflux and fume-cupboard) for 15 minutes, cooling to 20° and extracting with CH_2Cl_2 (2 x 50ml). The extract is dried (Na_2SO_4), filtered, evaporated to dryness and the residue is recrystallised from EtOAc /cyclohexane or EtOAc /hexane. It has UV (EtOH) with λ_{max} at 244 nm (ϵ 18,800); the IR has ν_{max} (KBr) at 3160 and 1515 cm^{-1} ; and the ^1H NMR (300MHz, CDCl_3) has δ at 0.77 (d, 3H, J = 6.8Hz), 0.82 (d, 3H, J = 6.7Hz), 1.68 (m, 1H), 3.77 (d of t, 1H, J = 6.6 and 9.1Hz), 4.23 (d of d, 1H, J = 6.6 and 9.1Hz), 4.55 (t, 1H, J = 9.1Hz), 9.00 (br s, 1H); and the ^{13}C NMR (75.5MHz, CDCl_3) has δ at 189.54, 73.53, 62.52, 32.17, 18.04, 17.90. [Delaunay et al. *J Org Chem* **60** 6604 1995, DOI: 10.1021/

jo00125a059; Nagao et al. *JCS Perkin Trans I* 2361 1985, DOI: 10.1039/P19850002361.] They are selective and efficient chiral auxiliaries [Velázquez & Olivo *Current Org Chem* **6** 303 2002, DOI: 10.2174/1385272024605023] which can be directly reduced by reductive cleavage with diisobutylaluminium hydride to their corresponding aldehydes and the chiral auxiliary. [Crimmins & Chaudhary *Org Lett* **2** 775 2000, DOI: 10.1021/ol9913901].

4R-(+)-4-Isopropyl-2-thiazolidinethione (4R-(+)-4-isopropyl-1,3-thiazolidin-2-thione) [110199-16-1] and **4S-(-)-4-isopropyl-2-thiazolidinethione (4S-(-)-4-isopropyl-1,3-thiazolidin-2-thione)** [76186-04-4] $\text{C}_6\text{H}_{11}\text{NS}_2$, **M** 161.3, **m** 66-67°, 67-68°, 69-71°, **R**- $[\alpha]_{\text{D}}^{20} +37$, **S**- $[\alpha]_{\text{D}}^{20} -37$ (c 1, CDCl_3). These compounds are efficient *Evans' type* of chiral auxiliaries. The *S*-(-)-enantiomer was synthesised by adding CS_2 (3ml, 50mmol) to a solution of *S*-valinol (10mmol (see [2026-48-4]) in aqueous N KOH (50ml) and stirring at 100° (bath at 110° under reflux and efficient fume-cupboard) for 16 hours, cooling to 20° and extracting with CH_2Cl_2 (2 x 50ml). The extract is dried (Na_2SO_4), filtered, evaporated to dryness and the residue is recrystallised from CH_2Cl_2 (colourless needles) or Et_2O . **Note** that unlike the preparation of the 1,3-oxazolidine-2-thione above, the preparation of this 1,3-thiazolidine-2-thione required a larger excess of CS_2 , stronger base and much longer heating time to replace the alcoholic O by S. It has ^1H NMR (300MHz, CDCl_3) with δ at 1.00 (d, 3H, $J = 7.2\text{Hz}$), 1.03 (d, 3H, $J = 8.5\text{Hz}$), 2.01 (m, 1H), 3.32 (d of d, 1H, $J = 8.2$ and 11.0Hz), 3.53 (d of d, 1H, $J = 8.2$ and 11.0Hz), 4.11 (m, 1H), 9.05 (br s, 1H); and the ^{13}C NMR (75.5MHz, CDCl_3) has δ at 200.78, 70.20, 35.73, 31.98, 18.78, 18.18. [Delaunay et al. *J Org Chem* **60** 6604 1995, DOI: 10.1021/jo00125a059; Nagao et al. *JCS Chem Commun* 1418 1985, DOI: 10.1039/C39850001418; Nagao et al. *J Org Chem* **51** 2391 1986, DOI: 10.1021/jo00362a047; McKennon & Meyer *J Org Chem* **58** 3568 1993, DOI: 10.1021/jo00065a020.] These are selective and efficient chiral auxiliaries [Velázquez & Olivo *Current Org Chem* **6** 303 2002, DOI: 10.2174/1385272024605023], and the condensation products can be directly reduced to the corresponding aldehyde and the chiral auxiliary by reductive cleavage with diisobutylaluminium hydride [Crimmins & Chaudhary *Org Lett* **2** 775 2000, DOI: 10.1021/ol9913901].

1R(-)-Menthol [natural *l*-(-), 1R,2S,5R-(-)-1-hydroxy-2-isopropyl-5-methylcyclohexane] [2216-51-5] **M** $\text{C}_{10}\text{H}_{20}\text{O}$, 156.3, **m** 42-45°, 43°, 44-46.5°, 89°/2mm, 100-101°/7mm, 212°/atm, d^{25}_{D} 0.89, n^{25}_{D} 1.458, n^{60}_{D} 1.446, $[\alpha]_{\text{D}}^{20} - 50$ (c 10, EtOH), $[\alpha]_{546}^{18} - 58.7$ (c 2, EtOH), and **1S(+)-menthol [synthetic *d*-(+)-menthol, 1S,2R,5S-(+)-1-hydroxy-2-isopropyl-5-methylcyclohexane]** [15356-60-2] **m** 43-44°, 103-104°/9mm, d^{25}_{D} 0.89, n^{25}_{D} 1.458, n^{60}_{D} 1.446, $[\alpha]_{\text{D}}^{25} + 48$ (c 10, EtOH), $[\alpha]_{546}^{18} + 58.6$ (c 2, EtOH). The natural *l*-isomer is present in peppermint oil and has a strong odour of peppermint, and is sometimes called *peppermint camphor*. Crystallise menthol from CHCl_3 , petroleum ether or EtOH/water. It can be sublimed at 40° *in vacuo*, but distillation at 5-10mm is preferable with large quantities. It is best stored under N_2 in the dark. It is soluble in most organic solvents and is slightly soluble in H_2O . [Barrow & Atkinson *J Chem Soc* 638 1939, DOI: 10.1039/JR9390000638; *Beilstein* **6** III 133, **6** IV 150.] *l*-(-)-Menthol is a very useful resolving agent for acids [Brunel & Buono *J Org Chem* **58** 7313 1993, DOI: 10.1021/jo00077a072; see also resolution of the Fe-PPh₃ complex [12101-02-9] above], and has been used in crystallisation-induced asymmetric transformation of malonate esters [Ihara et al. *JCS Chem Commun* 9 1988, DOI: 10.1039/C39880000009]. It is a chiral auxiliary that can be recycled [Solladié et al. *Synthesis* 173 1987, DOI: 10.1055/s-1987-27877; Katagiri et al. *J Org Chem* **53** 226 1988, DOI: 10.1021/jo00236a057]. Similar purification and applications are applicable for *non-natural d*-(+)-menthol, with the advantage of producing the optical enantiomers of the products. The *racemic* form **1RS,2SR,5RS-(±)-1-hydroxy-2-isopropyl-5-methylcyclohexane (hexahydro-thymol)** [1490-04-6] $\text{C}_{10}\text{H}_{20}\text{O}$, **M** 156.3, **m** 28° and 38° (dimorphic), **b** 216.5°/atm, d^{30}_{D} 0.8911, n^{20}_{D} 1.4415, n^{60}_{D} 1.4461, is obtained by catalytic hydrogenation of thymol [89-83-8] followed by distillation. [Waters & Beal *J Am Pharm Assoc* **34** 52 1945, DOI: 10.1002/jps.3030340208; Huggett *J Soc Chem Ind* **60** 67 1941; *Beilstein* **6** III 137, **6** IV 152.]

LEWIS AND BRØNSTED/LOWRY ACIDS AND BASES

A few words are warranted here because these terms are frequently used in current literature. The definition of acids (which produce H^+ ions) and bases (which produce HO^- ions) was adequate to explain reactions (e.g. salt formation) in aqueous solutions, and led to the concepts of pH (S.P.L. Sørensen *Biochem Z* **21** 131, 201 1909) and pK (ionisation, cf: Chapter 1, see report by ad-hoc committee of New York Academy of Sciences conference on *Acid-Base Terminology* in *The Lancet* **286** 1010-1012 1965, DOI: 10.1016/S0140-6736(65)92864-3). The definition becomes unsatisfactory when applied to studies of reactions in non-aqueous media, particularly in the

catalytic context. Independently, J.N. Brønsted [*Recl Trav Chim Pays-Bas* **42** 718 1923, DOI: 10.1002/recl.19230420815; *Chem Rev* **5** 231 1928, DOI: 10.1021/cr60019a001] and T.M. Lowry [*J Soc Chem Ind* (London) **42** 42 1923, DOI: 10.1002/jctb.5000420302] developed the view that an acid is a substance that has a tendency to lose a proton, and a base is one that has a tendency to gain a proton. This led to the understanding of *conjugate* species, e.g. R_3NH^+ as potential acids, and defining the equilibrium: $Base + H^+ \rightleftharpoons H\text{-}Base^+$. They pointed out that the differences in nett charge is not as important as the chemical behaviour on which their definition is based. Basicity and acidity do not bear a simple relationship to the nett respective charges, and both their properties depend much more on the complex electronic constitutions of the reagents. Later G. N. Lewis (1928 and later work) began with the classical concept of acids and bases, and progressed to a broader definition of an acid as a substance that is able to accept a pair of electrons, and a base as a substance capable of supplying a pair of electrons. This broader definition has been used extensively for reactions in non-aqueous solutions as well as in aqueous solutions. [Note that these concepts originated from extensive studies of catalytic reactions, e.g. mutarotation, hydrolysis, etc]. Thus a **Lewis acid** is a substance that is electron deficient (e.g. BF_3), and a **Lewis base** is a substance that can donate electrons (e.g. amines, phosphines, boranes, ethers, sulfides etc) to form bonds or complexes with Lewis acids.

In the **Brønsted/Lowry definition**, a base donates an electron pair to a proton to form a covalent B—H bond (a positive charge, if involved, will reside on B). In the **Lewis definition**, a base donates a pair of electrons to an electron deficient atom (other than a proton). An acid does not donate a proton but accepts a base to form a new bond. An electron pair is required for forming a covalent or a dative bond.

In the broadest form, an acid is an *electrophile* whereas a base is a *nucleophile*. When these species are regenerated during reactions then they become catalytic. For general reading see Michael B. Smith *Organic Chemistry: An Acid-Base Approach* CRC Press October 2010, ISBN 10: 1420079204, 13: 978142007203.

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CHAPTER 6

PURIFICATION OF BIOCHEMICALS

INTRODUCTION

Biochemicals are chemical substances produced by living organisms. They range widely in size, from simple molecules such as formic acid and glucose to macromolecules such as proteins and nucleic acids. Their *in vitro* chemical synthesis is often impossibly difficult, and in such cases they are available (if at all) only as commercial tissue extracts which have been subjected to purification procedures of widely varying stringency. The desired chemical may be, initially, only a minor constituent of the source tissue which may vary considerably in its composition and complexity. Recent advances in molecular biology have made it possible to produce substantial amounts of biological materials, which are present in nature in extremely small amounts, by recombinant DNA technology and expression in bacteria, yeast, insect and mammalian cells. The genes for these substances can be engineered such that the gene products, e.g. polypeptides or proteins, can be readily obtained in very high states of purity, and in large amounts if necessary. However, many such products, which are still obtained from the original natural sources, are available commercially and may require further purification.

As a preliminary step the tissue might be separated into phases [e.g. whole egg into white and yolk, blood into plasma (or serum) and red cells], and the desired phase may be homogenised. Subsequent treatment usually comprises filtration, solvent extraction, salt fractionation, ultracentrifugation, chromatographic purification, gel filtration and dialysis. Fractional precipitation with ammonium sulfate gives crude protein species. Purification is finally judged by the formation of a single band of macromolecule (e.g. protein, DNA) on electrophoresis and/or analytical ultracentrifugation. Although these generally provide good evidence of high purity, nonetheless it does not follow that one band under one set of experimental conditions is an absolute indication of homogeneity [D.S. Vodopich and R. Moore, *Biology Laboratory Manual*, McGraw-Hill, 2007, ISBN 9780072995220].

During the past 20 or 30 years a wide range of methods for purifying substances of biological origin have become available. For small molecules (including many sugars and amino acids) reference should be made to Chapters 1 and 2. The more important methods used for large molecules, polypeptides and proteins in particular, comprise:

1. *Centrifugation*. In addition to centrifugation for sedimenting proteins after ammonium sulfate precipitation in dilute aqueous buffer, the technique has been used for fractionation of large molecules in a denser medium or a medium of varying density. By layering sugar solutions of increasing densities in a centrifuge tube, proteins can be separated in a sugar-density gradient by centrifugation. Smaller DNA molecules (e.g. plasmid DNA) can be separated from RNA or nuclear DNA by centrifugation in aqueous cesium chloride (*ca* 0.975g/ml of buffer) for a long time (e.g. 40 hours at 40,000 x g). The plasmid DNA band appears at about the middle of the centrifuge tube and is revealed by the fluorescent pink band formed by the binding of DNA to ethidium bromide which is added to the CsCl buffer. *Microfuges* are routinely used for centrifugation in Eppendorf tubes (1.2-2ml) and can run up to speeds of 12,000 x g or more. *Analytical centrifugation*, which is performed under specific conditions in an analytical ultracentrifuge is very useful for determining purity, aggregation of protein subunits and the molecular weights of macromolecules. [D. Rickwood, T.C. Ford and J. Steensgaard (Eds), *Centrifugation: Essential Data Series*, J Wiley & Sons, 1994, ISBN 9780471942719; L.L. Regel and W.R. Wilcox, *Processing by Centrifugation*, Springer, 2001, ISBN 9780306466546; J.M. Graham and D. Rickwood, *Biological Centrifugation*, Springer, 2001, 9781859960370; A. Records and K. Sutherland, *Decanter Centrifugation Handbook*, Elsevier, 2001, ISBN 1856173690].

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2. *Gel filtration* with polyacrylamide (mol wt exclusion limit from 3000 to 300,000) and agarose gel (mol wt exclusion limit 0.5 to 150×10^6) is useful for separating macromolecules. In this technique high-molecular-weight substances are too large to fit into the gel microapertures and pass rapidly through the matrix (with the void volume), whereas low-molecular-weight species enter these apertures and are held there for longer periods of time, being retarded by the column material in the equilibria, relative to the larger molecules. This method is also used for desalting solutions of macromolecules.

Dry gels and *crushed beads* are also useful in the gel filtration process. Selective retention of water and inorganic salts by the gels or beads (e.g. Sephadex G-25) results in increased concentration and purity of the protein fraction which moves with the void volume. (See also section on 'Gel filtration' in Chapter 1.)

3. *Ion-exchange matrices* are microreticular polymers containing carboxylic acid (e.g. Bio-Rad 70) or phosphoric acid (Pharmacia, Amersham Biosciences, Mono-P) exchange functional groups for weak acidic cation exchangers, sulfonic acid groups (Dowex 50W) for strong acidic cation exchangers, diethylaminoethyl (DEAE) groups for weakly basic anion exchangers and quaternary ammonium (QEAE) groups for strong anion exchangers. The old cellulose matrices for ion exchangers have been replaced by Sephadex, Sepharose or Fractogel which have more even particle sizes with faster and more reproducible flow rates. Some can be obtained in fine, medium or coarse grades depending on particle size. These have been used extensively for the fractionation of peptides, proteins and enzymes. The use of pH buffers control the strength with which the large molecules are bound to the support in the chromatographic process. Careful standardisation of experimental conditions and similarly the very uniform size distribution of Mono beads have led to high resolution in the purification of protein solutions. MonoQ is a useful strong anion exchanger, and MonoS is a useful strong cation exchanger, whereas MonoP is a weak cation exchanger (check with commercial sources, see Chapter 1). These have been successful with medium pressure column chromatography (HPLC, see below in 7). Chelex 100 binds strongly and removes metal ions from macromolecules. [See sections on 'HPLC', 'Ion-exchange Resins' and 'Ion-exchange Celluloses and Sephadex' in Chapter 1.]
4. *Hydroxylapatite (hydroxyapatite)* is used for the later stages of purification of enzymes. It consists essentially of hydrated calcium phosphate which has been precipitated in a specific manner. It combines the characteristics of gel and ionic chromatography. Crystalline hydroxylapatite is a structurally organised, highly polar material which, in aqueous solution (in buffers), strongly adsorbs macromolecules such as proteins and nucleic acids, permitting their separation by virtue of the interaction with charged phosphate groups and calcium ions, as well as by physical adsorption. The procedure therefore is not entirely ion-exchange in nature. Chromatographic separations of singly and doubly stranded DNA are readily achievable, whereas there is negligible adsorption of low-molecular-weight species. [Tiselius et al. *Arch Biochem Biophys* **65** 132 1916, DOI: 10.1016/0003-9861(56)90183-7; Siegelman et al. *Anal Biochem* **13** 402 1965, DOI: 10.1016/0003-2697(65)90332-5.]
5. *Affinity chromatography* is a chromatographic technique whereby the adsorbant has a particular and specific affinity for one of the components of the mixture to be purified. For example the adsorbant can be prepared by chemically binding an inhibitor of a specific enzyme (which is present in the crude complex mixture) to a matrix (e.g. Sepharose). When the mixture of impure enzyme is passed through the column containing the adsorbant, only the specific enzyme binds to the column. After adequate washing, the pure enzyme can be released from the column by either increasing the salt concentration (e.g. NaCl) in the eluting buffer or adding the inhibitor to the eluting buffer. The salt or inhibitor can then be removed by dialysis, gel filtration (above) or ultrafiltration (see below). [See W.H. Scouten, *Affinity Chromatography: Bioselective Adsorption on Inert Matrices*, J.Wiley & Sons, NY, 1981, ISBN 0471026492; H. Schott, *Affinity Chromatography: Template Chromatography of Nucleic Acids and Proteins*, Marcel Dekker, NY, 1984, ISBN 0824771117; P. Matejtschuk Ed. *Affinity Separations* Oxford University Press 1997 ISBN 0199635501 (paperback); M.A. Vijayalakshmi, *Biochromatography, Theory and Practice*, Taylor & Francis Publ, 2002, ISBN 0415269032; and the section on 'Other Types of Chromatography' in Chapter 1.]
6. In the *Isoelectric focusing* of large charged molecules on polyacrylamide or agarose gels; slabs of these are prepared in buffer mixtures (e.g. ampholines) which have various pH ranges along the length of the gel. When a voltage is applied for some time, the buffers arrange themselves on the slabs in respective areas according to their pH ranges (prefocusing). Then the macromolecules are applied near the middle of the slab and allowed to migrate in the electric field until they reach the pH area similar to their isoelectric points and focus at that position. This technique can also be used in a chromatographic mode, *chromatofocusing*, whereby a gel in a column is run (also under HPLC conditions) in the presence of ampholines (narrow or wide pH ranges as required) and the macromolecules are then run through in a buffer. *Capillary electrophoresis* systems in which a current is applied to set the gradient are available in which the columns are fine capillaries and are used for qualitative and quantitative purposes [See R. Kuhn and S. Hoffstetter-Kuhn, *Capillary Electrophoresis: Principles and Practice*, Springer-Verlag Inc, NY, 1993; P. Camilleri Ed. *Capillary Electrophoresis-Theory and*

Practice, CRC Press, Boca Raton, Florida, 1993; D.R. Baker, *Capillary Electrophoresis*, J Wiley & Sons, NY, 1995; P.G. Righetti, A. Stoyanov and M. Zhukov, *The Proteome Revisited, Isoelectric Focusing*; *J. Chromatography Library Vol 63* 2001, Elsevier, ISBN 0444505261, P. Schmitt-Kopplin, *Capillary Electrophoresis: Methods and Protocols*, Humana, 2007, ISBN 9781588295392; J.P. Landers, *Handbook of Capillary & Microchip Electrophoresis and Associated Microtechniques*, CRC Press, Boca Raton, Florida, 2007, ISBN 9780849333293, C. Henry, *Microchip Capillary Electrophoresis*, Humana, 2006, ISBN 9781588292933.] The bands are eluted according to their isoelectric points. Isoelectric focusing standards are available which can be used in a preliminary run in order to calibrate the effluent from the column, or alternatively the pH of the effluent is recorded using a glass electrode designed for the purpose. Several types of efficient commercial equipment are available for separating proteins on a preparative and semi-preparative scale.

7. *High performance liquid chromatography* (HPLC) is liquid chromatography in which the eluting liquid is sent through the column containing the packing (materials as in paragraphs 2. to 6. above, which can withstand higher than atmospheric pressures) under pressure. On a routine basis this has been found useful for purifying proteins (including enzymes) and polypeptides after enzymic digestion of proteins or chemical cleavage (e.g. with CNBr) prior to sequencing (using reverse-phase columns such as μ -Bondapak C18). Moderate pressures (50-300psi) have been found most satisfactory for large molecules (FPLC). [See *Scopes Anal Biochem* **114** 8 1981, DOI: 10.1016/0003-2697(81)90443-7; B.A. Bidlingmeyer *Practical HPLC Methodology and Applications*, J Wiley & Sons, NY 1991; L.R. Snyder, J.L. Glajch and J.J. Kirkland *Practical HPLC Method Development*, J Wiley & Sons, NY 1988; ISBN 0471627828; R.W.A. Oliver, *HPLC of Macromolecules: A Practical Approach*, 2nd Edn, Oxford University Press, 1998, T. Hanai, *HPLC: A Practical Guide*, Royal Society of Chemistry (UK), 1999, ISBN 084045155; P. Millner *High Resolution Chromatography*, Oxford University Press, 1999 ISBN 0199636486; see also Chapter 1, Bibliography.]
8. *Ultrafiltration* (UF) using a filter (e.g. Millipore) can remove water and low-molecular-weight substances without the application of heat. Filters with a variety of molecular-weight exclusion limits not only allow the concentration of a particular macromolecule to be determined, but also the removal (by washing during filtration) of smaller molecular-weight contaminants (e.g. salts, inhibitors or cofactors). This procedure has been useful for changing the buffer in which the macromolecule is present (e.g. from Tris-Cl to ammonium carbonate), and for desalting. Ultrafiltration can be carried out in a stirrer cell (Amicon) in which the buffer containing the macromolecule (particularly protein) is pressed through the filter, with stirring, under argon or nitrogen gas pressure (e.g. 20-60psi). During this filtration process the buffer can be changed. This is rapid (e.g. 2L of solution can be concentrated to a few mls in 1 to 2 hours depending on pressure and filter). A similar application uses a filter in a specially designed tube (Centricon tubes, Amicon) and filtration occurs under centrifugal force in a centrifuge (4-6000rpm at 0°/40min). The macromolecule (usually DNA) then rests on the filter and can be washed on the filter also by centrifugation. The macromolecule is recovered by inverting the filter, placing a conical receiver tube on the same side where the macromolecule rests, filling the other side of the filter tube with eluting solution (usually a very small volume e.g. 100 μ L), and during further centrifugation this solution passes through the filter and collects the macromolecule from the underside into the conical receiver tube. With the development of polymeric and ceramic nanofilters use can be made of nanofiltration (NF) in which particles or molecules of less than 2nm can be held back. This is to be compared with UF where the size limit is between 2nm and 0.1 μ m (see Chapter 7).
9. *Partial precipitation* of a protein in solution can often be achieved by controlled addition of a strong salt solution, e.g. ammonium sulfate. This is commonly the first step in the purification process. Its simplicity is offset by possible denaturation of the desired protein and the (sometimes gross) contamination with other proteins. It should therefore be carried out by careful addition of small aliquots of the powdered salt or concentrated solution (below 4°, with gentle stirring) and allowing the salt to be evenly distributed in the solution before adding another small aliquot. Under carefully controlled conditions and using almost pure protein, it is sometimes possible to obtain the protein in crystalline form suitable for X-ray analysis (see below).
10. *Dialysis*. This is a process by which small molecules, e.g. ammonium sulfate, sodium chloride, are removed from a solution containing the protein or DNA using a membrane which is porous to small molecules. The solution (e.g. 10ml) is placed in a dialysis bag or tube tied at both ends, and stirred in a large excess of dialysing solution (e.g. 1.5 to 2 L), usually a weak buffer at ca 4°. The dialysing buffer is replaced with fresh buffer several times, e.g. four times in 24 hours. This procedure is similar to ultrafiltration (above) and allows the replacement of buffer in which the protein, or DNA, is dissolved. It is also possible to concentrate the solutions by placing the dialysis tube or bag in Sephadex G25 which allows the passage of water and salts from the inside of the bag thus concentrating the protein (or DNA) solution. Dialysis tubing is available from various distributors, but 'Spectra/por' tubing (from Spectrum Medical Industries, Inc, LA) is particularly effective because it retains macromolecules and allows small molecules to dialyse out very rapidly, thus reducing dialysis-

sing time considerably. This procedure is used when the buffer has to be changed so as to be compatible with the next purification or storage step, e.g. when the protein (or DNA) needs to be stored frozen in a particular buffer over extended periods. UF and NF can also serve this purpose whereby the solvent can be completely replaced by washing with an alternative solvent.

11. *Gel Electrophoresis*. This is becoming a more commonly used procedure for purifying proteins, nucleic acids, nucleoproteins, polysaccharides and carbohydrates. The gels can be 'electroblotted' onto membranes, and the modern procedures of identifying, sequencing (proteins and nucleic acids) and amplifying (nucleic acids) on sub-micro scales have made this technique of separation a very important one. See below for polyacrylamide gel electrophoresis (PAGE), [D. Patel *Gel Electrophoresis*, J.Wiley-Liss, Inc., 1994; P. Jones and D. Rickwood, *Gel Electrophoresis: Nucleic Acids*, J. Wiley and Sons, NY 1999 (paperback) ISBN 0471960438; D.M. Gersten and D. Gersten, *Gel Electrophoresis: Proteins*, J. Wiley and Sons, NY, 1996, ISBN 0471962651; R. Westermeier *Electrophoresis in Practice*, 4th Edn, Wiley-VCH Publishing, 2004 ISBN 9783527311811].
12. *Crystallisation*. The ultimate in purification of proteins or nucleic acids is crystallisation. This involves very specialised procedures and techniques and is best left to the experts in the field of X-ray crystallography who can provide a complete picture of the structure of these large molecules. [A. Ducruix and R. Giegé Eds, *Crystallisation of Nucleic Acids and Proteins: A Practical Approach*, 2nd Edition, 2000, Oxford University Press, ISBN 0199636788 (paperback); T.L. Blundell and L.N. Johnson *Protein Crystallisation*, Academic Press, NY, 1976; A. McPherson *Preparation and Analysis of Protein Crystals*, J.Wiley & Sons, NY, 1982; A. McPherson, *Crystallisation of Biological Macromolecules*, Cold Spring Harbour Laboratory Press, 2001 ISBN 0879696176, see also Bibliography in Chapter 1.]

Other details of the above can be found in Chapters 1 and 2 which also contain relevant references.

Several illustrations of the usefulness of the above methods are given in the *Methods Enzymol* series (Academic Press) in which 1000-fold purifications or more have been readily achieved. In applying these sensitive methods to macromolecules, reagent purity is essential. It is disconcerting, therefore, to find that some commercial samples of the widely used affinity chromatography ligand Cibacron Blue F3GA contained this dye only as a minor constituent. The major component appeared to be the dichlorotriazinyl precursor of this dye. Commercial samples of Procion Blue and Procion Blue MX-R were also highly heterogeneous [Hanggi and Carr, *Anal Biochem* **149** 91 1985, DOI: 10.1016/0003-2697(85)90480-4]. Variations in composition of sample dyes can well account for differences in results reported by different workers. The purity of substances of biological origin should therefore be checked by one or more of the methods given above. Water of high purity should be used in all operations. Double glass distilled water or water purified by a MilliQ filtration system (see Chapter 2) is most satisfactory.

Brief general procedures for the purification of polypeptides and proteins.

Polypeptides of molecular weights up to *ca* 1-2000 (10-20 amino acid residues) are best purified by reverse phase HPLC. The desired fractions that are collected are either precipitated from solution with EtOH or lyophilised. The purity can be checked by HPLC and identified by microsequencing (1-30 picomoles required) to ascertain that the correct polypeptide is in hand. Polypeptides larger than these are sometimes classified as proteins and are purified by one or more of the procedures described above. The purification of enzymes and functional proteins which can be identified by specific interactions is generally easier to follow because enzyme activities or specific protein interactions can be checked (by assaying) after each purification step. The commonly used procedures for purifying soluble proteins involve the isolation of an aqueous extract from homogenised tissues or extracts from ruptured cells from microorganisms or specifically cultured cells, for example, by sonication, freeze shocking or passage through a small orifice under pressure. Contaminating nucleic acids are removed by precipitation with a basic protein, e.g. protamine sulfate. The soluble supernatant is then subjected to fractionation with increasing concentrations of ammonium sulfate. The required fractions are then further purified by the procedures described in sections 2-9 above. If an affinity adsorbant has been identified, then affinity chromatography can provide an almost pure protein in one step sometimes even from the crude extract. The rule of thumb is that a solution with a protein concentration of 1mg/ml has an absorbance A_{1cm} at 280nm of 1.0 absorbance unit. Membrane-bound proteins are usually insoluble in water or dilute aqueous buffer and are obtained from the insoluble fractions, e.g. the microsomal fractions from the >100,000 x g ultracentrifugation supernatant. These are solubilised in appropriate detergents, e.g. Mega-10 (nonionic), Triton X-100 (ionic) detergents, and purified by methods 2 to 8 (previous section) in the presence of detergent

in the buffer used. They are assayed also in the presence of detergent or membrane lipids.

The purity of proteins is best checked by *polyacrylamide gel electrophoresis* (PAGE). The gels are either made or purchased as pre-cast gels and can be with uniform or gradient gel composition. Proteins are applied onto the gels *via* wells set into the gels or by means of a comb, and travel along the gel surface by means of the current applied to the gel. When the buffer used contains sodium dodecylsulfate (SDS), the proteins are denatured and the denatured proteins (e.g. as protein subunits) separate on the gels mainly according to their molecular sizes. These can be identified by running marker proteins, with a range of molecular weights, simultaneously on a track alongside the proteins under study. The protein bands are visualised by fixing the gel (20% acetic acid) and staining with Coomassie blue followed by silver staining if higher sensitivity is required. Commercial 'Phast Gel Electrophoresis' apparatus, or related equipment, is very useful for rapid analysis of proteins. It uses small pre-cast polyacrylamide gels (two gels can be run simultaneously) with various uniform or gradient polyacrylamide concentrations as well as gels for isoelectric focusing. The gels are usually run for 0.5-1.5 hours and can be stained and developed (1-1.5 hours) in the same apparatus. The equipment can be used to 'electroblot' the protein bands onto a membrane from which the proteins can be isolated and sequenced or subjected to antibody or other identification procedures. It should be noted that all purification procedures are almost always carried out at *ca* 4° in order to avoid denaturation or inactivation of the protein being investigated.

There has been considerable necessity for, and interest in, the study of **Proteomics**. This involves the identification, quantitation and isolation of all the proteins produced by a cell or organism at a particular point in time. It provides information on the expression of **all** the proteins produced by particular cells at a desired stage of the cell's development, maturity, activation or condition. A sophisticated apparatus for this purpose is a flat bed polyacrylamide gel which is run electrophoretically in one direction according to the extent of polymerisation of the acrylamide, and then run at right angles along a pH gradient (isoelectric focusing). Hundreds of polypeptides and proteins are thus separated, collected and identified by various other techniques such as LC-MS-MS, capillary electrophoresis etc (T. Palzkill, *Proteomics*, Springer, 2001, ISBN 0792375653; T.D. Veenstra and R.D. Smith *Proteome Characterization and Proteomics*, Academic Press, 2003, ISBN 978079237565; R. Westermeier, T. Naven and H-R. Höpker, *Proteomics in Practice: A Guide to Successful Experimental Design*, J.Wiley & Sons, NY 2008, ISBN 9783527319411; J.M. Walker (ed) *The Proteomics Protocols Handbook* Springer, 2005, ISBN: 978-1-58829-343-5 Print, 978-1-59259-890-8 Online; the *Journal of Proteomics* (ISSN: 1874-3919), which is the official journal of *The European Proteomics Association* that is published by Elsevier, and has been running for several years, is but one of the many journals on Proteomics and Bioinformatics that are available and can be viewed on the internet; see also Bibliography in Chapter 1).

Another rapidly developing field is **metabolomics** where metabolites are screened and identified in the normal and diseased cell at specific time intervals. These can be identified from studies of *genomics*, *transcriptomics* or *proteomics*. Such studies are now possible because of the highly improved power of HPLC, GC, MS, NMR and the interfacing of these instruments with each other, and with laptop computers which may drive them, store the data and compare it built-in libraries of substances. Thus *metabolones* can be mapped for various biological systems (plant and animal). Publications that are available include the *Metabolomics Journal* published by SpringerLink and started in 2005, and *Journal of Metabolomics and Systems Biology* (JMSB), published by Academic Journals which started in 2011. They publish original papers, reviews and conference reports.

Anyone contemplating the purification of a protein is referred to: Professor R.K. Scopes's monograph *Protein Purification*, 3rd Edn, Springer-Verlag, New York, 1994, ISBN 0387940723; M.L. Ladisch Ed. *Protein Purification - from Molecular Mechanisms to Large-scale Processes*, American Chemical Society, Washington DC, 1990; E.L.V. Harris and S. Angal, *Protein Purification Applications - A Practical Approach*, IRL Press, Oxford, 1990; J.C. Janson and L. Rydén, *Protein Purification - Principles, High Resolution Methods and Applications*, VCH Publ. Inc., 1989; ISBN 0895731223, Satinder Ahja *Handbook of Bioseparations*, Academic Press, 2000, ISBN 0120455404; S.M. Wheelwright, *Protein Purification: Design and Scale up of Downstream Processing*, J Wiley & Sons, NY 1994; references in the bibliography in Chapter 1, and selected volumes of *Methods Enzymol*, e.g. M.P. Deutscher (Ed), *Guide to Protein Purification*, *Methods Enzymol*, Academic Press, Vol **597** 2017, ISBN 9780128114698, Electronic ISBN: 9780128114704; M.A. Vijayalakshmi, *Biochromatography, Theory and Practice*, Taylor & Francis Publ, 2002, ISBN 0415269032; J.S. Davies, *Amino Acids, Peptides and Proteins Vol 32* 2006, RSC Publishing, Royal Society of Chemistry, ISBN

0854042326, DOI: 10.1039/9781847552778-FX001; S. Roe, *Protein Purification Techniques: A Practical Approach*, 2nd Edn, Oxford University Press, 2001, ISBN 0199636737; J.M. Walker (Ed) *The Protein Protocols Handbook*, 3rd Edn, (Springer Protocols Handbooks) Humana Press, 2009, ISBN 978-1-60327-474-6, e-ISBN 978-1-59745-198-7; T. Palmer, *Enzymes, Biochemistry, Biotechnology, Clinical Chemistry*, Horwood Publishing, 2001, ISBN 1898563780. For a comprehensive treatise of many volumes see *Springer Handbook of Enzymes* D. Schonburg & I. Schonburg Eds (A. Chang co-Ed) Springer-Verlag, Berlin, Heidelberg, 2003-onwards (with 39 volumes in 2011) <<http://www.springer.de>>.

The point should be made that advances in liquid chromatography (HPLC, coupled or uncoupled to MS or MS-MS) are ever increasing. This is not only in the improvements of the hardware but also in the variety of column materials that can now be purchased. Thus after the first precipitations of proteins, the rest of the purification can be carried out by HPLC equipment. Also in the syntheses of polypeptides, small proteins, RNAs and DNAs the final purifications are invariably done using HPLC procedures.

Considerable advances have been made in recent years with the rapid development and applications of the **microchip**. This in a sense is a **laboratory on a chip** (refer to sections on *Advances in Physical Techniques used in Purification* and *Advances in Chemical Techniques Used in Purification* at the ends of Chapters 1 and 2 respectively for applications.) To mention one example, complete sequencing of DNA can be performed using microchips no larger 15-30mm square. Throughput of fluids (reagents, solvents etc) is considerably faster and uses very small volumes, hence the synonym *microfluidics*. No wonder that complete sequencing of genomes can be performed in a matter of a few days. These procedures require purification at various stages; all being carried out on the same or accompanying microchips. The use of very small volumes and amounts of reagents and nucleic acid materials, results in shorter turnover times and rapid and quite accurate results. The downside is that very expensive equipment is required. However, once this cost is overcome, which can be recuperated in a short period, the running costs are not high considering the value of the results.

Protein Function

A very brief description of the function of proteins is made here. Proteins, which are chains of assorted amino acids perform a variety of functions. Depending of their amino acids and the sequences in which they are linked, they invariably, but not always, fold in specific ways as they are synthesised from the messenger RNA. The folding process is sometimes assisted by other specific proteins or polypeptides called *chaperones*. The folding tends to dictate the function of the protein. The folding is usually complex. The process not only folds segments of the amino acid sequences to produce α -helices and/or β -sheets which run parallel or anti-parallel to each other, but can also allow folds that cause the monomeric protein to form dimers, trimers, tetramers etc. Thus, if the folding is such that many charged amino acids (e.g. lysine, histidines for basic residues, and aspartate and glutamate for acidic residues) are on the exposed surfaces of the protein then the protein is water soluble depending on the pH of the solution. At pH values which cause the positive charges of the exposed acid residues to neutralise the exposed basic residues, e.g. at the PI (isoelectric point), the protein may precipitate out of solution. On the other hand if hydrophobic residues e.g. phenylalanine, leucine, valine, are on the exposed surface of the protein then its solubility in H₂O would be limited or extremely poor, and the protein would confine itself in a lipid bilayer and become membrane bound. Proteins capable of catalysing chemical reactions are called **enzymes**, others which are embedded in a cellular membrane may function as **pores** which allow ions or small molecules to flow into or out of a cell. These may or may not be **gated**, i.e. controlling the movement of particles in or out of the cell, as in **neurons** (nerve cells). Other lipid bound proteins function in **signalling** and are generally associated with protein **receptors**. Among the smaller polypeptides are the **activators**, **inhibitors**, **hormones**, and **immunopeptides**.

Enzymes: These proteins are folded in such a specific manner that a **pocket** is formed. This pocket is flanked by residues (basic, acidic or hydrophobic) which would assist catalysis. Unlike the catalytic process of chemical reactions (which following first, second or higher order kinetics), enzymic reactions follow **Michaelis-Menten kinetics**, also known as **saturation kinetics**. The reactive pocket is generally known as the **Active Site** and may or may not bind a small molecule (**cofactor**) or two to assist the enzyme in the catalytic process by entering into the reaction. This may be involved in reversible chemical bond formation or electron movement between the enzyme substrate(s) or product(s). Enzymic reactions are very highly stereospecific reactions with very careful steric control. Stereospecificity is not surprising considering that all the amino acids, including those at the active sites, are pure chiral entities. The enzyme can be activated or deactivated, e.g. by the binding of another molecule at a site away from the active pocket which alters the conformation

at the pocket so as to assist or desist the reaction rate. This site of binding is called the *allosteric site*. A plethora of enzymes and enzymic reactions are known with a great spectrum of reactions and mechanisms. Hydrophilic enzymes are easier to study, as this can be carried out in aqueous buffers. Hydrophobic enzymes are more difficult to study because they are generally active in the presence of the lipid membrane in which they are embedded. The kinetics are therefore studied in the presence of detergents (otherwise they may not be active or be dissolved in the aqueous medium), which may not mimic the natural state of the enzyme. A very rough measurement of activity is C_{50} which is the concentration of substrate that causes the enzyme to reach 50% activity. However, thorough kinetic studies are usually made, and two enzyme parameters are determined and are as characteristic of individual enzymes/substrates as are the melting and boiling points of organic compounds. These parameters are the Michaelis Constant (K_m) and the maximum velocity V_{max} of the particular enzyme with a stated substrate under specified conditions (i.e. temperature, buffer pH, ionic strength, etc.). The K_m is vaguely associated with the reversible binding of the substrate at the active site, and the V_{max} is the ultimate steady state velocity when the active site is saturated with the substrate. V_{max} also depends on the concentration of the enzyme used. A better parameter than V_{max} is the *turnover number* or k_{cat} which is related to the maximum velocity but takes into account the molecular mass of the enzyme in which case the enzyme needs to be in a high state of purity. k_{cat} has the dimension of min^{-1} (or sec^{-1}) and is the number of catalytic events taking place per min (or sec), the number of μmoles of substrate converted to product(s) in $(\mu\text{mol enzyme})^{-1}\text{min}^{-1}$. The activities of enzymes are known to be inhibited, and these too can be simple or complex. **IC_{50} values**, the concentration of inhibitor which will reduce the maximum activity to 50%, is a very rough guide, but it says little about the type of inhibition. Various types of inhibition are known, such as *competitive inhibition*, *uncompetitive inhibition* or *non-competitive inhibition* depending on how the inhibitor molecule competes with the substrate at the active site. Similarly with *activator*, as well as the possibility that as the concentration of products increases they can reach values when the products can inhibit enzyme activity. Most of these interactions are *reversible*. On the other hand, the inhibitor can also be designed such that it possess a reactive chemical group which will react *irreversibly* with amino acid residues at the active site and *kill* the enzyme. Such inhibition is known as *suicide inhibition*.

It was found necessary to identify enzymes, in a similar way that the Chemical Abstract Service identifies chemical compounds by their CASRegistry Numbers (see information at the beginning of this book, and the CASRNumber Index). The International Enzyme Commission has thus identified enzymes by EC numbers which consist of four numbers each separated by a full stop e.g. **EC 1.6.99.7**. [See section on 'Proteins, Enzymes, RNA and DNA' in this Chapter, and in the General Index]. This particular EC number is for ***Dihydropteridine Reductase*** where **1.** refers to the **Group number** which is for *oxidoreductase* enzymes, **6.** refers to the cofactor being NADH-NAD⁺ acceptor, **99.** refers to a second acceptor — *quinonoid-dihydrobiopterin* in this case, and **7.** is for it being the seventh enzyme classified in this group. The Groups are: EC 1 for *Oxidoreductases*, EC 2 for *Transferases*, EC 3 for *Hydrolases*, EC 4 for *Lyases*, EC 5 for *Isomerases*, and EC 6 for *Ligases* [See the comprehensive list in https://en.wikipedia.org/wiki/Enzyme_Commission_number, www.chem.qmul.ac.uk/iubmb/enzyme, and for details of, and references for, enzymes see M. Dixon & E.C. Webb [assisted by C.J.R. Thorne & K.F. Tipton *The Enzymes* (3rd Edn) Longman Group, London 1979, ISBN: 0-12-218358-4, <https://www.researchgate.net/publication/247011955>; and references above]. It should be noted that this classification is for *enzyme function*, i.e. enzymes with similar function which could have different sequences, or come from different sources, would have the same EC number, except perhaps the last digit(s). [Bibliography: A. Fersht *Enzyme Structure and Mechanism* (2nd edn) Freeman & Co 1985, ASIN: B010WF331A; A. Fersht *Structure and mechanism in protein science: a guide to enzyme catalysis and protein folding* W.H. Freeman San Francisco: 1999. ISBN 0-7167-3268-8; K.G. Scrimgeour *Chemistry and Control of Enzyme Reactions* Academic Press 1977, ISBN: 0126341508; J. Tze-Fei Wong *Kinetics of Enzyme Mechanisms* Academic Press 1975, ISBN: 012762250; https://en.wikipedia.org/wiki/Enzyme_kinetics].

Transport Proteins: These are at least of two types. Those that bind to and assist molecules, e.g. small molecules, proteins, lipids or carbohydrates, to enter or leave the cell; and those that are embedded in the cell membrane and allow the desired *molecular traffic* to occur. They also include *pore proteins* which are embedded in the cell wall and allow, in a gated or non-gated process, for ions such as Na⁺, K⁺ and Ca²⁺ to move in and out of cells that require this. They are plentiful in the *synapses* of neurons which allow movement of ions, and small molecules, across them, causing electrical conductance. Special *pore proteins* in synapses may also cause small excitatory and inhibitory molecules to move across them (e.g. *neurotransmitters* like γ -aminobutyric acid GABA), or cause disturbances between synapses.

Receptor Proteins: These are lipophilic proteins that embed themselves into the lipid membranes of cells. Their sequences dictate their folding and dictate their specific action. They commonly have a small protein **anchor**, or a series of smaller proteins which are activated when a molecule (e.g. a drug, a hormone, a **cytokine**) binds to the outer surface of the receptor protein. The binding transfers a signal to the **anchor proteins** beneath (e.g. G-proteins, phosphorylating **tyrosine kinases**, Jak's) that are bound to the inner side (cell lumen) of the receptor. The signal can activate mechanisms further inside of the cell which carry out the intended function of the drug, cytokine etc. Among these mechanisms, for example, the signal molecule such as a phosphorylated protein at a tyrosine residue, can now enter the cell nucleus and alter gene production. These **receptor proteins** can be very complicated e.g. the string of their amino acids can traverse the lipid bilayer membrane several times, and occasionally can form a protein crown. In some cases the receptor proteins in the lipid bilayer are caused to come together (e.g. dimerise) under the influence of a bound **cytokine** before activating the signalling of the anchor proteins inside the cell. As in the above case with enzymes, the agonistic or antagonistic effects of the binding drug (**cytokine**, hormone etc) can be roughly quantitated by IC_{50} values, the concentrations that cause 50% of the desired physiological effects. [[https://en.wikipedia.org/wiki/Receptor_\(biochemistry\)](https://en.wikipedia.org/wiki/Receptor_(biochemistry)); https://en.wikipedia.org/wiki/Cell_surface_receptor; <https://www.ibiblio.org/virtualcell/textbook/chapter3/cmf3.htm>.]

Brief general procedures for purifying DNA.

Oligo-deoxyribonucleotides (up to *ca* 60-mers) are conveniently purified by HPLC (e.g. using a Bio-Rad MA7Q anion exchange column and a Rainin Instrument Co, Madison, Dynamax-300A C₈ matrix column) and used for a variety of molecular biology experiments. Plasmid and chromosomal DNA can be isolated by centrifugation in **cesium chloride** buffer (see paragraph 1. *centrifugation* above), and then re-precipitated with 70% ethanol at -70° (18 hours), collected by centrifugation (microfuge) and dried in air before dissolving in TE (10mM TrisHCl, 1mM EDTA pH 8.0). The DNA is identified on an Agarose gel slab (0.5 to 1.0% DNA grade in 45mM Tris-borate + 1mM EDTA or 40mM Tris-acetate + 1mM EDTA pH 8.0 buffers) containing ethidium bromide which binds to the DNA and under UV light causes it to be visualised as pink fluorescent bands. Marker DNA (from λ phage DNA cut with the restriction enzymes Hind III and/or EcoRI) with bands running from 72 to 353 base-pairs (bp) are run in a parallel track in order to estimate the size of the unknown DNA. Various other DNA markers are commercially available such as the step ladder ranging from 50bp to 800bp with bands at 50bp intervals, and the step ladder with bands ranging from 100bp to 4000bp with bands at 200bp intervals. The DNA can be isolated from the bands on the gel by transfer onto nitro-acetate paper (e.g. NA 45) electrophoretically, by binding to silica or an ion-exchange resin, then extracted from the adsorbent paper and precipitated with ethanol. The DNA pellet is then dissolved in TE buffer and its concentration determined. A solution of duplex DNA (or RNA) of 50 μ g/ml gives an absorbance of 1.0 unit at 260nm/1cm cuvette (single-stranded DNA or RNA gives a value of 1.3 absorbance units). DNA obtained in this way is suitable for molecular cloning.

Recombinant and chemically synthesised DNA and RNA are now routinely separated and purified by HPLC, and their structures are confirmed by sequencing an aliquot. A variety of commercially available HPLC systems are now available, and a desired system can be selected from them.

Brief mention must be made of the tremendous advances that have been made in recent years in the fields of DNA, RNA gene sequencing and synthesis. The development of instrumentation and analysers by the Illumina –Company [www.illumina.com/] and the '**Ion torrent**' semiconductor sequencing of DNA and RNA, using an ion PGM (personal genome machine) sequencer (see; <http://lifetech-it.hosted.jivesoftware.com/index.jspa>) have made it possible to sequence complete genomes in a matter of weeks or less.

For experimental details on the isolation, purification and manipulation of DNA and RNA the reader is referred to: J. Sambrook, E.F. Fritsch and T. Maniatis, *Molecular Cloning-A Laboratory Manual*, 2nd Edn, (3 volumes), Cold Spring Harbor Laboratory Press, (CSHL Press) NY, 1989, ISBN 0879693096 (paperback); P.D. Darbre, *Basic Molecular Biology: Essential Techniques*, J. Wiley and Sons, NY 1998, ISBN 0471977055; J. Sambrook and D.W. Russell, *Molecular Cloning-A Laboratory Manual*, 3rd Edn, (3 volumes), Cold Spring Harbor Laboratory Press, NY, 2001, ISBN 0079695773, ISBN 9780879695774 (paperback), ISBN 0079695765 (cloth bound); J. Sambrook and D.W. Russell, *The Condensed Protocols for Molecular Cloning: A Laboratory Manual*, CSHL Press, 2006, ISBN 9780879697716, also available on line; M.A. Vijayalakshmi, *Biochromatography, Theory and Practice*, Taylor & Francis Publ, 2002, ISBN 0415269032; A. Travers and M. Buckle, *DNA-Protein Interactions: A Practical Approach*, Oxford University Press, 2000, ISBN 0199636915

(paperback); R. Rapley and D.L. Manning Eds *RNA: Isolation and Characterisation Protocols*, Humana Press 1998 ISBN 0896034941; R. Rapley, *The Nucleic Acid Protocols Handbook*, Humana Press 2000 ISBN 0896038416 (paperback).

This chapter lists some representative examples of biochemicals and their origins, a brief indication of key techniques used in their purification, and literature references where further details may be found. Simpler low-molecular-weight organic compounds, particularly those that may have been prepared by chemical syntheses, e.g. phenylacetic acid, will be found in Chapter 3. Only a small number of enzymes and proteins are included because of space limitations. The purification of the ones that have been included has been described only briefly. The reader is referred to comprehensive texts such as the *Methods in Enzymology* (E-Book Series) series which currently runs to Vol **597** 2017, ISBN 9780128114698, Electronic ISBN: 9780128114704; and *The Enzymes* (3rd Edn, Academic Press) which ran to more than 28 volumes in 2010, for methods of preparation and purification of proteins and enzymes. Leading references on proteins will be found in *Advances in Protein Chemistry* which was incorporated with *Advances in Structural Biology* (84 volumes (2011), Elsevier Inc), and enzymes will be found in *Advances in Enzymology* which then became *Advances in Enzymology and Related Areas of Molecular Biology*, J Wiley & Sons, NY (up to volume 78 in 2011). The *Annual Reviews of Biochemistry* (Annual Reviews Inc. Patlo Alto California) also are an excellent source of key references to the up-to-date information on known and new natural compounds, from small molecules, e.g. enzyme cofactors, to proteins and nucleic acids. See also the *Springer Handbook of Enzymes* cited above.

Abbreviations of titles of periodical are generally defined as in the Chemical Abstracts Service Source Index (CASSI) but without punctuation. References to Fieser & Fieser's *Reagents for Organic Synthesis* will be shortened to Fieser throughout, e.g. Fieser **2** 254, **11** 88, etc. All temperatures are in degrees Centigrade unless otherwise stated. Other abbreviations are self evident.

Ionisation constants of ionisable compounds are given as **pK** values (published from the literature) and refer to the **pKa** values at room temperature (~ 15°C to 25°C). The values at other temperatures are given as superscripts, e.g. **pK**²⁵ for 25°C. Estimated values are entered as **pK**_{Est(1)} ~ (see section on 'Ionisation Constants' in Chapter 1 for further information).

Benzene, which has been used as a solvent successfully and extensively in the past for reactions and purification by chromatography and crystallisation is now considered a **very dangerous substance**, so it has to be used with extreme care. It is important that an alternative solvent to benzene (e.g. toluene, toluene-petroleum ether, or a petroleum ether to name a few) should be used first. However, if benzene has to be used then all operations have to be performed in well-ventilated fumehoods and precautions taken to avoid inhalation and contact with skin and eyes. Whenever benzene is mentioned in the text, an asterisk e.g. *C₆H₆ or *benzene, is inserted to remind the user that special precaution should be adopted.

Selected Amino acids and peptides, Proteins, Enzymes DNA and RNA, Carotenoids, Carbohydrates, Steroids, and Physiologically Active Compounds (including miscellaneous low-molecular-weight bioactive substances, drugs, antibiotics, coenzymes, vitamins, lipids, phospholipids, nucleosides, nucleotides, polynucleotides and useful reagents) are collected in the following separate respective sections of this chapter.

AMINO ACIDS and PEPTIDES

This section includes amino acid derivatives and related compounds.

N-Acetyl-L-alaninamide [15962-47-7] $C_5H_{10}N_2O_2$, **M 130.2, m 162°**. Crystallise the amide repeatedly from EtOH/diethyl ether. The (\pm)-*isomer* crystallises from H_2O and has **m 157-158°**. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 1838 1961, de Jong *Recl Trav Chim Pays-Bas* **19** 288 1900, DOI: 10.1002/recl.19000190902; *Beilstein* **4** H 295.]

N-Acetyl-β-alanine [3025-95-4] $C_5H_9NO_3$, **M 127.2, m 78.3-80.3°, pK²⁵ 4.45**. The β-alanine crystallises from acetone. [King & King *J Am Chem Soc* **78** 1089 1956, DOI: 10.1021/ja01587a005; *Beilstein* **4** IV 2526, 2548.]

N-Acetyl-L-alanyl-L-alaninamide [30802-37-0] $C_8H_{15}N_3O_3$, **M 201.2, m 250-251°**. Crystallise the dipeptide derivative repeatedly from EtOH/diethyl ether. [Puliti & Mattia *Acta Cryst Section C* **2** 51 1995, DOI: 10.1107/S0108270194008577.]

N-Acetyl-L-alanyl-L-alanyl-L-alaninamide [29428-34-0] $C_{11}H_{20}N_4O_4$, **M 272.3, m 295-300°**. Crystallise the tripeptide derivative from MeOH/diethyl ether.

N-Acetyl-L-alanylglycinamide [76571-64-7] $C_7H_{13}N_3O_3$, **M 187.2, m 148-149°, d²⁵ 1.191g/cm³**. Crystallise the dipeptide derivative repeatedly from EtOH/diethyl ether.

Acetyl-α-amino-n-butyric acid [34271-24-4] $C_6H_{11}NO_3$, **M 145.2, pK²⁵ 3.72**. Crystallise the acid twice from water (charcoal) and dry it in air [King & King *J Am Chem Soc* **78** 1089 1956, DOI: 10.1021/ja01587a005; Gördes et al. *Adv Synth & Catalysis* **345** 510 2003, DOI: 10.1002/adsc.200390059].

O-Acetylcarnitine chloride (2-acetoxy-3-carboxy-N,N,N-trimethylpropanamine HCl) [*S*(D+)- 5080-50-2, *R*(L-)- 5061-35-8, *RS* 2504-11-2] $C_9H_{17}NO_4 \cdot HCl$, **M 239.7, m 181°, 187°(corr, dec), 197°(dec), [α]_D²⁵ -28 (c 2, H₂O) for *S*-isomer, pK²⁵ 3.6**. Recrystallise the chloride from isopropanol. Dry it over P₂O₅ under high vacuum. The *S*-betaine crystallises from EtOH/Et₂O with **m 145°(dec)** and is *hygroscopic*; it has **[α]_D²⁰ -19.5 (c 6, H₂O)**. It is an endogenous mitochondrial metabolite that transports acetyl groups across the mitochondrial membrane. [Poorthuis *et al* 'Determination of acylcarnitines in urine of patients with inborn errors of metabolism using HPLC after derivatisation with 4'-bromophenacyl bromide' *Clin Chim Acta* **216** 53 1993, DOI:10.1016/0009-8981(93)90138-T, PMID: 8222273; Krimberg & Wittandt *Biochem Z* **251** 231 1932, Strack et al. *Z Physiol Chem* **238** 191 1936, *Beilstein* **4** III 1630, 1632.]

R(-)-N-Acetyl-L-cysteine methyl ester [7652-46-2] $C_6H_{11}NO_3S_6$, **M 177.2, m 71-78°, 80°, [α]_D²⁰ -24.0 (c 1, MeOH)**. The ester is purified by converting into the cuprous mercaptide which is decomposed by dilute H₂SO₄, extracted into Et₂O, dried (Na₂SO₄), filtered, evaporated and the residue is recrystallised from H₂O containing a little AcOH. The crystals are dried in a vacuum. These operations should be carried out in an inert atmosphere (N₂ or argon) to avoid oxidation to the disulfide cystin ester. *Note* that the cuprous salt is only stable when it is dry, but is readily oxidised when wet. It has been used as a sulfur transfer agent [Gilman & Spero *Tetrahedron Lett* **34** 1751 1993, DOI:10.1016/S0040-4039(00)60769-4]. [Pirie *Biochem J* **25** 614 1931, DOI: 10.1042/bj0250614; *Beilstein* **4** III 1607.]

N-Acetylglutamic acid [1188-37-0] $C_7H_{11}NO_5$, **M 189.2, m 185° (RS), 201° (S), [α]_D²⁵ -16.6 (in H₂O), [α]_D²⁵ -15.6 (c 4, MeOH) for *S*-enantiomer, pK_{Est}(1) ~3.4, pK_{Est}(2) ~4.3**. A likely impurity is glutamic acid. Crystallise it from boiling water. It inhibits *N*-acetyl-L-glutamate synthase. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 1948 1961, Shigesada & Tatibana *Eur J Biochem* **84** 285 1978, DOI: 10.1111/j.1432-1033.1978.tb12167.x; Coudé et al. *Biochem Biophys Res Commun* **102** 1016 1981, DOI: 10.1016/0006-291X(81)91639-9; *Beilstein* **4** IV 3047.]

N-Acetylglucinamide [2620-63-5] $\text{C}_4\text{H}_8\text{N}_2\text{O}_2$, **M 116.1, m 139-139.5°**. Crystallise the amide repeatedly from EtOH/Et₂O. Dry it in a vacuum desiccator over KOH. [Davis & Levy *J Chem Soc* 3479 1951, DOI: 10.1039/JR9510003479; Fischer & Otto *Chem Ber* 36 2106 1903, DOI: 10.1002/cber.190303602128; *Beilstein* 4 IV 2401.]

N-Acetylglycine (Aceturic acid) [543-24-8] $\text{C}_4\text{H}_7\text{NO}_3$, **M 117.1, m 206-208°, 207-209°, pK₁²⁵ -1.92, pK₂²⁵ 3.69**. N-Acetylglycine is treated with acid-washed charcoal and recrystallised three times from water or EtOH/Et₂O and is dried *in vacuo* over KOH [King & King *J Am Chem Soc* 78 1089 1956, DOI: 10.1021/ja01587a005]. [*Beilstein* 4 IV 2399.]

N-Acetylglycyl-L-alaninamide [34017-20-4] $\text{C}_7\text{H}_{13}\text{N}_3\text{O}_3$, **M 187.2**. Crystallise the dipeptide derivative repeatedly from EtOH/Et₂O. Dry it in a vacuum desiccator over KOH. [Hedwig et al. *JCS Faraday Trans* 87 1751 1991, DOI: 10.1039/FT9918701751.]

N-Acetylglycylglycinamide [27440-00-2] $\text{C}_6\text{H}_{11}\text{N}_3\text{O}_3$, **M 173.2, m 207-208°**. Crystallise the dipeptide derivative repeatedly from EtOH/Et₂O. Dry it in a vacuum desiccator over KOH. [Hedwig et al. *JCS Faraday Trans* 87 1751 1991, DOI: 10.1039/FT9918701751.]

N-Acetylglycylglycylglycinamide [35455-24-4] $\text{C}_8\text{H}_{14}\text{N}_4\text{O}_4$, **M 230.2, m 253-255°**. Crystallise the tripeptide derivative repeatedly from EtOH/Et₂O. Dry it in a vacuum desiccator over KOH.

N-Acetylhistidine (H₂O) [39145-52-3] $\text{C}_8\text{H}_{11}\text{N}_3\text{O}_2$, **M 181.2, m 148° (RS), 169° (S), [α]_D²⁵ +46.8 (c 1, H₂O) for S-enantiomer**. A likely impurity is histidine. Crystallise it from water, then 4:1 acetone/water. [For Co complexing see Marshall et al. *J Am Chem Soc* 78 4636 1956, DOI: 10.1021/ja01599a030; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 p 1990 1961, *Beilstein* 25 IV 4359.]

N-Acetyl-RS-homocysteine thiolactone (Citolone) [1195-16-0] [17896-21-8 for ±] $\text{C}_6\text{H}_9\text{NO}_2\text{S}$, **M 159.2, m 110°, 109-111°, 111.5-112.5°**. Dry Citolone in a vacuum desiccator. It recrystallises from toluene as needles. It is a ninhydrin -ve substance which gives a 'slow' nitroprusside test. It has λ_{max} at 238nm (ε 4,400 M⁻¹cm⁻¹); and ν_{max} (nujol) 1789s and 851ms cm⁻¹. [Benesch & Benesch *J Am Chem Soc* 78 1597 1956, DOI: 10.1021/ja01589a025; cf. Laliberté et al. *J Chem Soc* 2756 1963, DOI: 10.1039/JR9630002756.]

N-Acetyl-L-leucinamide (2-acetamido-4-methylvaleramide) [30130-37-1, 28529-34-2] $\text{C}_8\text{H}_{16}\text{N}_2\text{O}_2$, **M 172.2, m 202°**. Recrystallise it from aqueous EtOH or CHCl₃/petroleum ether (b 40-60°). [Gränacher *Helv Chim Acta* 8 216 1925, DOI: 10.1002/hlca.19250080135; *Beilstein* 4 II 864, for L, and Bergmann et al. *Justus Liebigs Ann Chem* 449 301 1926, DOI: 10.1002/jlac.19264490116; *Beilstein* 4 II 877 for DL.]

N-Acetyl-L-methionine [65-82-7] $\text{C}_7\text{H}_{13}\text{NO}_3\text{S}$, **M 191.3, m 103.5-104.5°, 104°, [α]_D²⁵ -24.5 (c 1, in H₂O), pK_{Est} ~3.4**. Crystallise N-acetyl-L-methionine from Me₂CO, H₂O or EtOAc. Dry it in a vacuum over P₂O₅. Its solubility at 25° in H₂O is 30.7%, and in Me₂CO it is 29.5%. [Mitzi & Schueter *Biochim Biophys Acta* 27 168 1958, DOI: 10.1016/0006-3002(58)90305-6; Birnbaum et al. *J Biol Chem* 194 455 1952, www.jbc.org/content/194/1/455; *Beilstein* 4 IV 3206.]

Acetylmethionine nitrile [538-14-7] $\text{C}_7\text{H}_{12}\text{N}_2\text{OS}$, **M 174.3, m 44-46°**. Crystallise the nitrile from diethyl ether. [Catch et al. *J Chem Soc* 1609 1947, DOI: 10.1039/JR9470001609; *Beilstein* 4 III 1654.]

N-Acetyl-N¹-methyl-L-alaninamide [19701-83-8] $\text{C}_6\text{H}_{12}\text{N}_2\text{O}_2$, **M 144.2, m 181.2-182° (sealed tube), [α]_D²⁵ -51.1 (c 2, EtOH)**. Crystallise the amide from EtOAc/Et₂O, then from EtOH and Et₂O. Also recrystallise it twice by dissolving ~2.5g in hot 200ml of toluene and cooling. It sublimes at ~170°, so its m is measured in a sealed tube. [Applewhite & Niemann *J Am Chem Soc* 81 2208 1959, DOI: 10.1021/ja01518a050; *Beilstein* 4 IV 2500.]

N-Acetyl-N'-methylglycinamide [7606-79-3] $C_5H_{10}N_2O_2$, **M 130.2, m 157.5-158°**. Recrystallise the amide from EtOH/Et₂O mixture. Also recrystallise it twice from EtOAc/EtOH (16:1) and once from EtOAc. [Applewhite & Niemann *J Am Chem Soc* **81** 2208 1959, DOI: 10.1021/ja01518a050.]

N-Acetyl-N'-methyl-L-leucine amide [32483-15-1] $C_9H_{18}N_2O_2$, **M 186.3, m 165.3-166.8° (sealed tube), $[\alpha]_D^{25}$ -33.9 (c 1, H₂O)**. Recrystallise the amide from EtOAc, EtOH/hexane mixture or toluene/hexane mixture. It sublimes above ~160°. [Applewhite & Niemann *J Am Chem Soc* **81** 2208 1959, DOI: 10.1021/ja01518a050.]

N-Acetyl-L-phenylalanine [2018-61-3] $C_{11}H_{13}NO_3$, **M 207.2, m 170-171°, 174-175°, $[\alpha]_D^{25}$ +47.5 (c 4, EtOH), +52.5 (c 2, EtOH), (DL) m 152.5-153°, pK_{Est} ~3.5**. N-Acetyl-L-phenylalanine is recrystallised from H₂O, 20% MeOH/H₂O, or CHCl₃; dry and store it at 4°. The **(DL)-mixture** crystallises from H₂O, Me₂CO, EtOAc, or CHCl₃ with **m 152-154°** and the solubilities in w% at 25° are 0.73 (H₂O), 4.3 (Me₂CO), 0.79 (EtOAc) and 0.34 (CHCl₃) [Kerr & Niemann *J Org Chem* **23** 893 1958 DOI: 10.1021/jo01100a601, Overby & Ingersoll *J Am Chem Soc* **73** 3363 1951, DOI: 10.1021/ja01151a110; L-form: Fu et al. *J Am Chem Soc* **76** 6054 1954, DOI: 10.1021/ja01652a057; Bender & Glasson *J Am Chem Soc* **81** 1590 1959, DOI: 10.1021/ja01516a020]. [*Beilstein* **14** I 238, **14** IV 1575.]

N-Acetyl-L-phenylalanine ethyl ester [2361-96-8] $C_{11}H_{17}NO_3$, **M 235.3, m 93-94°**. Crystallise the ester from aqueous EtOH or H₂O. [Izumiya & Fruton *J Biol Chem* **218** 59 1956, <http://www.jbc.org/content/218/1/59>, PMID: 13278315.]

N-Acetyltryptophan [87-32-1] $C_{13}H_{14}N_2O_3$, **M 246.3, m 206°, 207-208° (RS), pK_{Est} ~3.8, [1218-34-4] m 188°, 189.5-190.5° (S), $[\alpha]_D^{25}$ +30.1 (aqueous NaOH), +71.5° (dioxane/aqueous HCl)**. A likely impurity is tryptophan. Crystallise it from EtOH by adding water. [Cowgill *Biochim Biophys Acta* **200** 18 1970, DOI: 10.1016/0005-2795(70)90038-3, DL: Berg *J Biol Chem* **100** 79 1933, <http://www.jbc.org/content/100/1/79>; *Beilstein* **22/14** V 40-50.]

N-Acetyl-L-valine amide [37933-88-3] $C_7H_{14}N_2O_2$, **M 158.3, m 275°**. Recrystallise the amide from CH₃OH/Et₂O. [den Tweel, Harder & Buitelaar *Stability and Stabilization of Enzymes* Elsevier Sci Publ, Amsterdam 1993, ISBN: 0444893725.]

α-Alanine (RS) [302-72-7] $C_3H_7NO_2$, **M 89.1, m 295-296°, (S) [56-41-7] m 297°(dec), $[\alpha]_D^{15}$ +14.7 (in 1M HCl), (R) [338-69-2] m 289-291°(dec), $[\alpha]_D^{15}$ -14.1 (c 0.9, 1M HCl), pK_1^{25} 2.34, pK_2^{25} 9.87**. Crystallise alanine from H₂O or aqueous EtOH, i.e. crystallise it from 25% EtOH in water, or recrystallise it from 62.5% EtOH, wash it with EtOH and dry it to constant weight *in vacuo* over P₂O₅. **RS-α-alanineamide** [20108-77-4] has **m 62°** (from CHCl₃), **pKa 8.02**, the **hydrochloride** [80222-96-4] has **m 173°** (from EtOH) and the **acetate salt** has **m 136-137°**. **S-α-alanine methyl ester hydrochloride** [2491-20-5] has **m 109-111°, $[\alpha]_D^{15}$ +8.0 (c 1.6, MeOH)**, and **S-α-alanine N-methylamide** [7324-05-2] has **m 78°** (hygroscopic, also **m 72°** reported, from CHCl₃). [Gutter & Kegeles *J Am Chem Soc* **75** 3893 1953, DOI: 10.1021/ja01112a006; Minireview: Walsh *J Biol Chem* **264** 2393 1989, www.jbc.org/content/264/5/2393.full.pdf.] **2,2'-Iminodipropionic acid** is a likely impurity. [*Beilstein* RS: **4** H 387, **4** I 491, **4** II 814, **4** III 1222, **4** IV 2481; R: **4** H 385, **4** I 491, **4** II 812, **4** III 1219, **4** IV 2480; S: **4** H 381, **4** I 489, **4** II 809, **4** III 1208, **4** IV 2480.]

β-Alanine [107-95-9] $C_3H_7NO_2$, **M 89.1, m 197-198°(dec), 205°(dec), 205.5°(dec), 207°(dec, rapid heating), pK_1^{25} 3.55 (3.60, CO₂H), pK_2^{25} 10.24 (10.36, NH₃⁺)**. Crystallise β-alanine by dissolving it in a hot saturated aqueous solution, filtering, adding four volumes of absolute EtOH and cooling in an ice-bath. Recrystallise it in the same way and then finally, crystallise it from a warm saturated solution in 50% EtOH and adding four volumes of absolute EtOH with cooling in an ice-bath. The crystals are dried in a vacuum desiccator over P₂O₅. It also crystallises from H₂O, and sublimes at 170-180°/0.3mm. The **hydrochloride** [6057-90-5] forms plates with **m 123°**, its **methyl ester** has **b 69°/58mm**, the **methyl ester hydrochloride** [3196-73-4] has **m 107°** (from EtOH/Et₂O), the **amide** has **m 41°** and the **amide hydrochloride** has **m 149°**. **N-Methyl β-alanine** crystallises from EtOH, the **monohydrate** forms plates with **m 99-100°** and its **hydrochloride** [2679-14-3] has **m 105°**. **N-Methyl β-alanine amide** [4874-17-3] M 102.1 is a liquid with **d₄²⁵ 1.052, n_D²⁰ 1.458**, and is an antibacterial

[Altamura et al. *J Med Chem* **38** 4244 1955, DOI: 10.1021/jm00021a013]. [Donovan & Kegeles *J Am Chem Soc* **83** 255 1961, DOI: 10.1021/ja01463a001; for pKa see Albert *Biochem J* **47** 531 1950, DOI: 10.1042/bj0470531; *Beilstein* **4** H 401, **4** I 499, **4** II 827, **4** III 1258, **4** IV 2526.]

S-Alaninol [*S*-2-aminopropan-1-ol] [2749-11-3] C_3H_9NO , **M 75.1**, **b** 72-73°/11mm, **167-169°/760mm**, **d**₄²⁰ **0.961**, **d**₂₅²⁵ **0.925g/ml**, **n**_D²⁰ **1.456 (1.4498)**, **[α]_D²⁰ +26.0** (c 2, EtOH), **[α]_D²⁰ +18** (neat), **pK₂₅ 9.43**. Purify it as for *S*-2-amino-3-methylbutan-1-ol below. [*Beilstein* **4** IV 1615.]

D-Allothreonine [*2R,3R*(-)-isomer] [24830-94-2] $C_4H_9NO_3$, **M 119.1**, **m** 272-273°(dec), **276°(dec)**, **[α]_D²⁵ -9.1** (c 3.9, H₂O), **pK₁²⁵ 2.11**, **pK₂²⁵ 9.10**. Recrystallise D-allothreonine from aqueous EtOH or 50% EtOH. L-Allothreonine has **[α]_D²⁵ +9.8** (c 4 H₂O) [Elliot *J Chem Soc* 62 1950, DOI: 10.1039/JR9500000062; Birnbaum et al. *J Biol Chem* **194** 455 1952, <http://www.jbc.org/content/194/1/455>, PMID: 14927637; IR: Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** 1961, *Beilstein* **4** IV 3170.]

RS-α-Allylglycine (2-aminopent-4-enoic acid) [7685-44-1] $C_5H_9NO_2$, **M 115.1**, **m** 250-255°(dec), **258-260°(dec)**, **pK_{Est(1)} ~2.3**, **pK_{Est(2)} ~9.6**. Dissolve it in absolute EtOH and precipitate it with pyridine, then recrystallise it from aqueous EtOH [*R_F* on paper in BuOH/EtOH/NH₃/H₂O (4:4:1:1) is 0.37]. Store at -20°. The **hydrobromide** has **m** **136-140°** (from EtOAc) and the **phenylureido derivative** has **m** **159-161°**. [Schögl *Monatsh Chem* **89** 377 1958, DOI: 10.1007/BF00898759; *Beilstein* **4** IV 2852.]

Aminoacetic acid see **Glycine** below.

α-Amino acids. All the α-amino acids with the '*natural*' configuration [*S* (L), except for cysteine which is *R*(L)] at the α- carbon atom are available commercially in a very high state of purity. Many of the '*non-natural*' α-amino acids with the [*R*(D)] configuration as well as racemic mixtures are also available, and generally none require further purification before use unless they are of '*Technical Grade*' or were stored for a very long period. The *R* or *S* enantiomers are optically active except for glycine which has two hydrogen atoms on the α- carbon atom, but these are *pro*-chiral and enzymes or proteins *do* distinguish between them, e.g. serine hydroxymethyltransferase successfully replaces the *pro*-α- hydrogen atom of glycine with CH₂OH (from formaldehyde) to make *S*-serine. The twenty common natural α-amino acids are: **amino acid**, three-letter abbreviation, **one-letter abbreviation**, pK (-COOH) and pK (-NH₃⁺): **Alanine**, Ala, **A**, 2.34, 9.69; **Arginine**, Arg, **R**, 2.17, 9.04; **Asparagine**, Asn, **N**, 2.01, 8.80; **Aspartic acid**, Asp, **D**, 1.89, 9.60; **Cysteine**, Cys, **C**, 1.96, 8.18; **Glutamine**, Gln, **Q**, 2.17, 9.13; **Glutamic acid**, Glu, **E**, 2.19, 9.67; **Glycine**, Gly, **G**, 2.34, 9.60; **Histidine**, His, **H**, 1.8, 9.17; **Isoleucine**, Ile, **I**, 2.35, 9.68; **Leucine**, Leu, **L**, 2.36, 9.60; **Lysine**, Lys, **K**, 2.18, 8.95; **Methionine**, Met, **M**, 2.28, 9.20; **Phenylalanine**, Phe, **F**, 1.83, 9.12; **Proline**, Pro, **P**, 1.99, 10.96; **Serine**, Ser, **S**, 2.21, 9.15; **Threonine**, Thr, **T**, 2.11, 9.62; **Tryptophan**, Trp, **W**, 2.38, 9.39; **Tyrosine**, Tyr, **Y**, 2.2, 9.11, **Valine**, Val, **V**, 2.32, 9.61, respectively. Technical grade amino acids can be purified on ion-exchange resins (e.g. Dowex 50W and eluting with a gradient of HCl or AcOH), and the purity can be checked by TLC in two dimensions and stained with ninhydrin. [J.P.Greenstein & M.Winitz, *Chemistry of the Amino Acids* (3 Volumes), J.Wiley & Sons, NY, 1961; C.Cooper, N.Packer and K.Williams, *Amino Acid Analysis Protocols*, Humana Press, 2001, ISBN 0896036561.] Recently codons for a further two amino acids have been discovered which are involved in ribosome-mediated protein synthesis giving proteins containing these amino acids. The amino acids are *R*(L)-selenocysteine [Stadtman *Ann Rev Biochem* **65** 83 1996, PMID: 8811175] and pyrrolysine [(4*R*, 5*R*)-4-substituted (with Me, NH₂ or OH) pyrroline-5-carboxylic acid] [Srinivasan, James & Krzycki *Science* **296** 1459 2002, PMID: 12029131.] They are, however, rare at present and only found in a few microorganisms.

dl-α-Aminoadipic acid (hydrate) (2-aminoheptanoic acid) [542-32-5] $C_6H_{11}NO_4$, **M 161.2**, **m** **196-198°**, **204°**, **205-206°**, **pK_{Est(1)} ~2.0**, **pK_{Est(2)} ~4.5**, **pK_{Est(3)} ~9.8**. Crystallise the acid from H₂O. Alternatively, purify it by precipitating the Cu salt and decomposing the Cu salt suspended in H₂O by bubbling H₂S, filtering off the CuS, evaporating, and recrystallising the residue from H₂O. Note that prolonged refluxing of an aqueous solution converts the acid to the lactone: **piperid-2-one-6-carboxylic acid** which has **m** **177-178°**. [Linstead & Wang *J Chem Soc* 810, 811 1937, DOI: 10.1039/JR9370000807; Waalkes et al. *J Am Chem Soc* **72** 5760 1950,

DOI: 10.1021/ja01168a515; Greenstein & Winitz *The Chemistry of the Amino Acid* J. Wiley, **Vol 3** p. 2408 1961, *Beilstein* **4** III 1555, **4** IV 3070.] It is found in rat retina and is a biomarker for diabetes in humans [Wang et al. *J Clin Invest* **123** 4309 2013, DOI:10.1172/JCI64801, PMID: 24091325].

N-(p-Aminobenzoyl)-L-glutamic acid [4271-30-1] $C_{12}H_{14}N_2O_5$, **M 266.3**, **m 173°**, **174-175° (L-form)**, $[\alpha]_{546}^{25}$ **-17.5** (c 2, 0.1M HCl); **197° (DL)**, pK_1^{25} **2.61**, pK_2^{25} **3.76**, pK_3^{25} **4.83**. Crystallise the acid from H_2O . Also purify it by dissolving 2.7g in H_2O (130ml), adding aqueous NaOH to pH 5.5 and adding portionwise a solution of 0.5M $CuSO_4$ to complete precipitation of the Cu salt. This salt is filtered off, suspended in H_2O and H_2S is bubbled through to precipitate CuS, filter, evaporate and recrystallise the residue from H_2O . It has λ_{max} (H_2O) at 273nm. [Backer & Houtman *Recl Trav Chim Pays-Bas* **70** 738 1951, DOI: 10.1002/recl.19510700902; *Beilstein* **14** IV 1153.]

RS-2-Aminobutyric acid [2835-81-6] $C_4H_9NO_2$, **M 103.1**, **m 283-285°(dec)**, **287-288°(dec)**, **291°(dec)**, **303°(dec)**, **303°(dec, sealed tube)**, pK_1^{25} **2.29**, pK_2^{25} **9.83**. Crystallise the acid from water. [Stiles & Finkbeiner *J Am Chem Soc* **81** 505 1959, DOI: 10.1021/ja01511a067; Fe complexing: Perrin *J Chem Soc* 3125 1958, DOI: 10.1039/JR9580003125; *Beilstein* **4** IV 2584.]

S-2-Aminobutyric acid (Butyrine) [1492-24-6] $C_4H_9NO_2$, **M 103.1**, **m 292°(dec)**, $[\alpha]_D^{25}$ **+ 20.6** (c 2, 2.5N HCl), pK_1^{25} **2.55**, pK_2^{25} **9.60**. Crystallise butyrine from aqueous EtOH, and the melting point depends on heating rate but has **m 303°** in a sealed tube. **R-2-Aminobutyric acid (D-)** [2623-91-8] $C_4H_9NO_2$, **M 103.1**, **m 292°(dec)**, is the *enantiomer* and has $[\alpha]_D^{25}$ **- 20.6** (c 2, 2.5N HCl), $[\alpha]_D^{20}$ **-7.94** (c 4, H_2O). [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 2399 IR: 2401 1961, *Beilstein* **4** III 1294, **4** IV 2584.]

RS-3-Aminobutyric acid (BABA) [541-48-0, 2835-82-7] $C_4H_9NO_2$, **M 103.1**, **m 193-194°**, $pK_{Est(1)}$ **~3.5**, $pK_{Est(2)}$ **~10.3**. Crystallise the acid from aqueous EtOH or MeOH/Et₂O. Also crystallise it by heating a slightly diluted EtOH solution and adding Me₂CO. It gives a purple spot with R_F 0.89 on paper chromatography using 80% aqueous phenol (the α -amino acid has R_F 0.74). [Zukha & Rivlin *J Org Chem* **23** 94 1958, DOI: 10.1021/jo01095a604; Bruylants *Bull Soc Chim Belg* **32** 259 1923, *Beilstein* **4** IV 2595.] It induces plant disease resistance, and increases resistance to abiotic stress [Cohen *Plant Disease* **86** 448 2002, DOI:10.1094/PDIS.2002.86.5.448].

S-3-Aminobutyric acid [3775-72-2] $C_4H_9NO_2$, **M 103.1**, **m 212°**, **210-212°**, $[\alpha]_D^{18}$ **+ 38.8** (c 0.5, H_2O). Purify the acid by recrystallisation from absolute EtOH. It has also been crystallised from MeOH or MeOH/Et₂O and dried in a vacuum. [Synthesis from S-alanine: Balenovic et al. *J Chem Soc* 3316 1952, DOI: 10.1039/JR9520003313; Bruylants *Bull Soc Chim Belg* **32** 259 1923.] **Optical resolution** of the (\pm)-**methylester** was achieved *via* recrystallisation of the *d*-camphorsulfonic acid salts which, after decomposition of the salts and hydrolysis of the methyl esters gave pure *R*- and *S*- **3-aminobutyric acids** [Fischer & Scheibler *Justus Liebigs Ann Chem* **383** 337 1911, DOI: 10.1002/jlac.19113830206; *Beilstein* **4** IV 2595.]

4-Aminobutyric acid (GABA) [56-12-2] $C_4H_9NO_2$, **M 103.1**, **m 202°(dec)**, **203°(dec)**, pK_1^{25} **4.14**, pK_2^{25} **10.55**. Crystallise GABA from aqueous EtOH or MeOH/Et₂O. Also crystallise it by dissolving it in the least volume of H_2O and adding 5-7 volumes of absolute EtOH. It is a neurotransmitter. [Sherman *Biochemical Preparations* **4** 91 1955; de Witt *Org Synth Coll Vol* **2** 25 1943, DOI: 10.15227/orgsyn.017.0004; *Beilstein* **4** III 1316, **4** IV 2600.]

1-Amino-1-cyclopentanecarboxylic acid (cycloleucine) [52-52-8] $C_6H_{11}NO_2$, **M 129.2**, **m 328-335°(dec)**, **328-329°**, **330°(dec)**, pK_1^{20} **2.4**, pK_2^{20} **10.3**. Any Cl^- or other anions are removed by stirring with a strong cation exchange resin (Amberlite IR-120), filtering, and washing with distilled H_2O until the filtrate is free from the anion. The resin is then stirred overnight with 6N NH_4OH , filtered, the filtrate is decolourised (charcoal) and evaporated to dryness in a vacuum. The residue is recrystallised from H_2O /EtOH. Also crystallise it from aqueous EtOH. The **hydrochloride** has **m 222-224°(dec)**. [Neelakantan & Hartung *J Org Chem* **23** 964 1958, DOI: 10.1021/jo01101a007; Connors & Ross *J Chem Soc* 2119 1960, DOI: 10.1039/JR9600002119; O'Donnell et al. *Synthesis* 127 1984, DOI: 10.1055/s-1984-30749; *Beilstein* **14** IV 974.]

trans*-(1*RS*,4*RS*)-4-Aminoethylcyclohexane-1-carboxylic acid** (t*-AMCHA**, **Tranexamic acid**, **Tranex**, **Transamin**, **Trasamlon**, **Ugurol**, **Frenolyse**, **Hexapromin**, **Abvitoff** among other names) [1197-18-8] $\text{C}_6\text{H}_{11}\text{NO}_2$, **M 157.2**, **m 386-392°** (dec), and ***cis*-(1*RS*,4*SR*)-4-aminoethylcyclohexane-1-carboxylic acid** [1197-17-7] **M 157.2**, **m 236-238°** (dec); ***cis*- and *trans*- acids have the same pK_1^{25} 4.51 (CO_2H), pK_2^{25} 10.72 (NH_2)**. A mixture of *cis*- and *trans*- acids (2:1) is obtained by catalytic reduction of 4-acetamidomethylbenzoic acid (3.9g, 20mmol, from acetylation of 4-aminomethylbenzoic acid see [56-91-7]) in aqueous NaOH (0.8g, 20mmol in 15ml of H_2O) in the presence of Raney Ni (3ml of aqueous suspension) are shaken with H_2 in a bomb (100ml) at 170° and 82atm/cm² (1205psi). Reduction is complete after 2 hours, the catalyst is filtered off, the filtrate is acidified with 4N H_2SO_4 and evaporated to dryness *in vacuo*; the residue is extracted with Me_2CO , filtered and evaporated to dryness *in vacuo* to give crude 4-acetamidomethylcyclohexane-1-carboxylic acid (~3.9g). The acid is then refluxed with 20% HCl (20ml) for 3 hours (in an oil bath at 150°), evaporated to dryness *in vacuo*; the residue is dissolved in H_2O (20ml) and freed from HCl by passage through a column of ‘Amberlite IR-4B’ (15ml, in OH^- form) and eluted with H_2O . Evaporation of the eluate *in vacuo* and crystallisation of the residue from aqueous Me_2CO gives a **2:1 mixture of *cis*- and *trans*- 4-aminomethylcyclohexane-1-carboxylic acid** (2.24g, ~71%, **m ~232-236°, dec**). Both acids are symmetrical, i.e. mirror images are superimposable.

The mixture (10g, 64mmol) is **separated** by refluxing with $\text{Cu}(\text{CO}_3)_2$ (9.15g, 38mmol) in H_2O (100ml) for 1 hour (turning deep blue in colour), cooled and the blue precipitate is filtered off, dissolved into 8% aqueous NH_3 (100ml), filtered from a little solid, and passed through a column of ‘Diaion SK#1 (NH_4^+ form)’ and washed with H_2O . The eluted solution is then passed through an ‘Amberlite IR-4B’ (OH^- form) column and the effluent is evaporated to dryness. The residue (6.75g, **m ~221-223°, dec**.) is repeatedly recrystallised from H_2O - Me_2CO to give the pure ***cis*-amino acid** with **m 236-238°, dec**, and IR (KBr) peaks at ν_{max} 2940, 2660, 1639 (1640), 1560 (1563), 1509 (1515), 1408 (1403), 1305 (1308), 930 and 904 cm^{-1} , value in *italics* are from Meyer see below). The combined filtrates from the recrystallisation are evaporated, the residue is dissolved in H_2O and similarly de-ionised through the same columns. The final residue is recrystallised from H_2O - Me_2CO to give the pure ***trans*-4-aminoethylcyclohexane-1-carboxylic acid** (3.37g) with **m 286-292°, dec**, and IR (KBr) peaks at ν_{max} 2940, 2610, 1637, 1535 (1528), 1383 (1381) (1325) and 920 cm^{-1} , value in *italics* are from Meyer see below). On TLC (Silica gel G, eluted with ascending *n*-PrOH/ H_2O , 65:35) the R_F of the *cis*-isomer was always 1.2 times larger than that of the *trans*- isomer. [Note that the melting points of the isomers measured in the usual way vary somewhat, and by using a Du Pont Model 900 differential thermal analyzer the *cis*-acid had **m 252°**, and the *trans*-acid had **m 295-300°** as endotherms, Meyer *J Med Chem* **9** 641 1966, DOI: 10.1021/jm00322a059].

The ***cis*-hydrochloride** has **m 195-197°, dec** (prisms from H_2O - Me_2CO), the ***trans*-hydrochloride** has **m 238-241.5°, dec** (needles from H_2O - Me_2CO); the ***cis*-hydrobromide** has **m 205-208°, dec** (plates from Me_2CO), the ***trans*-hydrobromide** has **m 227-229°, dec** (plates from H_2O); the ***cis*-*p*-toluenesulfonic acid salt** has **m 177-178°** (plates from *n*-PrOH- Et_2O), the ***trans*-*p*-toluenesulfonic acid salt** has **m 262-264°** (plates from H_2O), the ***cis*-HCl-AuCl₃ salt** has **m 178-180°, dec** (yellow needles from H_2O), the ***trans*-HCl-AuCl₃ salt** has **m 205-206°, dec** (yellow prisms from H_2O), the ***cis*-HCl-PtCl₄ salt** has **m 233°, dec** (yellow needles from H_2O), the ***trans*-HCl-PtCl₄ salt** has **m 254-255°, dec** (yellow-orange plates from H_2O); the ***cis*-*N*-acetamide** has **m 189-190°** (prisms from EtOH), the ***trans*-*N*-acetamide** has **m 154-155°** (prisms from Me_2CO); the ***cis*-*N*-benzamide** has **m 157-158°** (plates from EtOH/* C_6H_6) and the ***trans*-*N*-benzamide** has **m 177-178°** (needles from EtOH/ H_2O).

The *cis*- and *trans*- acids are also readily separated from the mixture by recrystallisation of their ***p*-toluenesulfonic acid salts** which have very different solubility properties. **Isomerisation:** When a solution of the *cis*-acid (2g) in 0.5N NaOH (26ml) is heated in a silver vessel in an autoclave at 200° for 6 hours, and the cooled solution is passed through an ‘Amberlite IR-120’ (NH_4^+ form) column and washed with H_2O , evaporation of the eluate *in vacuo* and four recrystallisations from aqueous MeOH gave the *trans*-acid (0.8g, 40%), **m 384-390° dec**. [Naito et al. *Chem Pharm Bull Jpn* **16** 728 1968, Daiichi Seiyak *Dutch Pat* 6,414,942 1965, *Chem Abstr* **64** 3379 1966.] The width-at-half height of the peaks from the cyclohexane protons in the ^1H NMR spectra are larger from the *trans*-acid than for the corresponding peaks from the *cis*-acid. The structure of the *cis*-acid was confirmed by conversion to a cyclic **lactam** on fusion. It has **m 104°** after recrystallisation from hexane followed by sublimation at 100°/2.5 x 10⁻²mm. [Note that this melting point was recorded at the point where, under crossed Nicol prisms, birefringence was lost. The crystal form, however, was only slowly lost thereafter over a wide range of temperatures]. The **lactam** has IR (KBr) peaks at ν_{max} 1661 (amide I),

1421, 1325 and 1205 cm^{-1} [Meyer *J Med Chem* **9** 641 1966, DOI: 10.1021/jm00322a059].

These amino acids are haemostatic with antiplasmic activity. The *trans*-acid has potent antiplasmic activity (inhibiting the fibrinolytic enzyme system), being 50 times more active than the *cis*-acid, and 5-7 times more potent than ϵ -aminocaproic acid [Naito et al. *Chem Pharm Bull Jpn* **16** 357 and 728 1968.] The antifibrolitic activity is due to blocking of the lysine binding sites of plasminogen. *t*-AMCHA has been used as a lysine analogue to characterise binding sites in plasminogens. [Brockway & Castellino *Arch Biochem Biophys* **151** 194 1972, DOI:10.1016/0003-9861(72)90488-2; Hoover et al. *Biochemistry* **32** 10936 1993, DOI: 10.1021/bi00092a002; Marshall et al. *Biochemistry* **33** 3599 1994, DOI: 10.1021/bi00178a017.]

4-Amino hippuric acid (*N*-*p*-aminobenzoylglycine) [61-78-9] $\text{C}_9\text{H}_{10}\text{N}_2\text{O}_2$, **M 194.2, m 198-199°, 200-202°, $\text{pK}_{\text{Est}(1)} \sim 1.7(\text{NH}_2)$, $\text{pK}_{\text{Est}(2)} \sim 3.4(\text{CO}_2\text{H})$.** Crystallise the acid from H_2O . It is soluble in organic solvents. [Cohen & McGilvery *J Biol Chem* **169** 119 1947, <http://www.jbc.org/content/169/1/119>, PMID: 20240544; Cohen & McGilvery *J Biol Chem* **171** 121 1947, <http://www.jbc.org/content/171/1/121> PMID: 20240544; Beilstein **14** III 1069, **14** IV 1152.] Used for determining renal function [Schumann & Wüstenberg *Clin Nephrol* **33** 35 1990, PMID: 2302868.]

***dl*-4-Amino-3-hydroxybutyric acid** [924-49-2] $\text{C}_4\text{H}_9\text{NO}_3$, **M 119.1, m 218°(dec), 223°(dec), 225°(dec), $\text{pK}_{\text{I}}^{25} \sim 3.80(\text{CO}_2\text{H})$, $\text{pK}_{\text{Est}(2)} \sim 9.3$.** Crystallise the acid from H_2O or aqueous EtOH. Recrystallise it by dissolving it in H_2O and adding MeOH or EtOH. It is not very soluble in CHCl_3 or EtOAc. [Renaud & Seebach *Synthesis* 424 1986, DOI: 10.1055/s-1986-31665; Beilstein **4** II 938, **4** IV 3187.]

***R* (L)-4-Amino-3-hydroxybutyric acid (GABOB)** [352-21-6] $\text{C}_4\text{H}_9\text{NO}_3$, **M 119.1, m 212°(dec), 213-214°(dec), 216-217°(dec), $[\alpha]_{\text{D}}^{25} -20.5$ (c 1.75, H_2O), and the *S*-(+)- isomer has [7013-05-0] m 207-212°, $[\alpha]_{\text{D}}^{25} +21$ (c 1.7, H_2O).** Purify GABOB through a Dowex 50Wx8 resin, eluting with 1.3N NH_4OH , evaporating and crystallising the residue by dissolving it in H_2O and adding EtOH. It is an anticonvulsant. [Renaud & Seebach *Synthesis* 424 1986, DOI: 10.1055/s-1986-31665; Fukase et al. *Tetrahedron Lett* **29** 795 1988, DOI:10.1016/S0040-4039(00)80212-9 Beilstein **4** IV 3187.]

α -Amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid hydrate (AMPA) [*R*(+)- 83654-131; *RS*(\pm)- 74341-63-2; *S*(-)- 83643-88-3] $\text{C}_7\text{H}_{10}\text{N}_2\text{O}_4$, **M 204.2 (H_2O), *RS* m 252°(dec).** AMPA is an analogue of glutamic acid designed and studied by Hansen and Krogsgaard-Larsen [*JCS Perkin Trans 1* 1829 1980, DOI: 10.1039/P19800001826; Lauridsen et al. *J Med Chem* **28** 668 1985, DOI: 10.1021/jm50001a022] as a possible agonist for the **AMPA subgroup of ionotropic glutamate receptors**. The *racemic* amino acid has been synthesised in four steps from ethyl 3-methoxy-5-methylisoxazol-4-carboxylate \rightarrow 4-hydroxymethyl-3-methoxy-5-methylisoxazole \rightarrow 4-chloromethyl-3-methoxy-5-methylisoxazole \rightarrow diethyl acetamido-(3-methoxy-5-methylisoxazol-4-ylmethyl)malonate which was deprotected [de-acetylated, 3-MeO- \rightarrow 3-HO, ($=\text{COOEt}$)₂ \rightarrow ($=\text{COOH}$)₂ \rightarrow -COOH] to give *RS*(\pm)-AMPA. The last step was achieved by treating the diethyl *N*-Ac-OMe-AMPA with 48% aqueous HBr under a blanket of N_2 and refluxed (oil bath at 140°) for 15-20 minutes, cooled rapidly, evaporated *in vacuo*, dissolved in H_2O and re-evaporated. The residue was dried *in vacuo* over KOH and P_2O_5 , and recrystallised twice from *iso*PrOH/Et₂O (**m 206-209° dec**; 80% yield), and finally from *iso*-PrOH to give an analytical sample of the **hydrobromide (m 218-220° dec)**: IR (KBr) had ν_{max} at 3000br, 1740, 1660m, 1535, 1500, 1255 and 1215 cm^{-1} , and ^1H NMR (D_2O with MeCN internal standard at 2.02) with δ at 2.25(s, 3H), 2.95(d, 2H) and 4.25(t, 1H). The **free *RS*-amino acid (\pm)-AMPA [77521-29-0]** was obtained by neutralising the **hydrobromide** in EtOH with Me_3N and recrystallising from H_2O to provide pure **amino acid monohydrate** (86% yield) with (**m 252° dec**) and IR with ν_{max} at 3550—2450(several bands), 1655, 1625, 1585, 1535, 1515, 1400, 1335 and 1255 cm^{-1} . [Hansen and Krogsgaard-Larsen *JCS Chem Commun* 87 1979, DOI: 10.1039/C39790000087.]

The enzyme used for the **Enzymic optical resolution** was a gel-bound aminoacylase (Enzygel aminoacylase available from Boehringer) well known to hydrolyse *N*-acetyl *S*-amino acids at a rate of *ca* 10,000 times faster than the enantiomeric *N*-acetyl *R*-amino acids. This also established the absolute configuration of the products. It was found that ***RS*-*N*-Ac-3-OMe-AMPA** was a better substrate to use than *RS*-*N*-Ac-AMPA, and was prepared as follows: The above diethyl acetamido-(3-methoxy-5-methylisoxazol-4-ylmethyl)malonate was hydrolysed and monodecarboxylated without demethylating the 3-OMe group by refluxing with 1M aqueous

HCl for 8 hours, evaporating to dryness, twice dissolving in H₂O and re-evaporating; and to the residue was added H₂O (1part), Et₃N (1.0 mol) in EtOH (3parts) until the pH was ~6. The precipitated **RS-3-OMe-AMPA** [83643-90-7] had **m** 224-225°(dec), IR (KBr) had ν_{\max} at 2950br, 1660, 1510, 1475, 1405 and 1345 cm⁻¹, and ¹H NMR (D₂O-CF₃COOD) with δ at 2.29(s, 3H), 3.01(d, 2H), 3.97(s, 3H) and 4.28(t, 1H). This product (0.5mmol) was acetylated with Ac₂O (9.8mmol) in AcOH by refluxing for 70 minutes, adding H₂O, evaporating to dryness *in vacuo*, dissolving in toluene, re-evaporating twice, and finally the residue was recrystallised from EtOAc to give **RS-N-Ac-3-OMe-AMPA** [84751-72-4] (45% yield) which had **m** 161° and IR (KBr) with ν_{\max} at 3370, 2300—2950(several bands), 1725, 1610, 1520, 1465, 1410, 1345, 1230, 1200 and 1135 cm⁻¹, and ¹H NMR (CD₃OD; also as internal standard at δ 3.36) with δ at 1.19(s, 3H), 2.31(s, 3H), 2.84(m, 2H), 3.99(s, 3H) and 4.63(dd, 1H).

Enzymic optical resolution: A solution of **RS-N-Ac-3-OMe-AMPA** (28.6mg, 0.118mmol) in H₂O (19.7ml) containing Enzygel aminoacylase (212mgm ~40 IU) adjusted to pH 7.1, by dropwise addition of aqueous LiOH solution, was stirred gently to keep the enzyme gel in suspension at 30° and covered with Parafilm to 186 hours avoid evaporation, and sampled at time intervals to estimate the amount of deacetylated amino acid. After 50% of deacetylated amino acid was formed. The Enzygel was removed by centrifugation and washed with H₂O (3 x 10ml) also by centrifugation. The combined original supernatant and washings was placed on a column (~0.7 x 10cm) of Amberlite IR-120 (~4ml, in H⁺ form) and eluted with H₂O (30ml at ~0.5ml/min) to give a weakly acidic eluate followed by a neutral eluate. The column was then eluted with 0.1M aqueous NH₃ to give a basic eluate followed by a neutral eluate. (a) The combined first weakly acidic and neutral eluates were evaporated *in vacuo*, suspended in MeOH, filtered and the filtrate evaporated *in vacuo*, to give an oil which was extracted with hot EtOAc (3 x 5ml), filtered again an evaporated to provide pure (by TLC on silica gel 60 F₂₆₄, EtOAc-MeOH 9:1 + 1% HCOOH; visualised with I₂) unhydrolysed **R-N-Ac-3-OMe-AMPA** [84799-53-1] as an oil (11.7mg, >82% yield of 50% of *RS-material*) with $[\alpha]_D^{20}$ -25.2 (c 1.17, EtOH), $[\alpha]_{436}^{20}$ -56 (c 1.17, EtOH). (b) The combined basic and following neutral eluates were evaporated to dryness *in vacuo*, twice dissolved in H₂O and evaporated *in vacuo*, to give a white crystalline residue which was recrystallised from H₂O/EtOH to afford **pure** (by TLC on silica gel 60 F₂₆₄, EtOH/25% aqueous NH₃ 7:3; visualised with ninhydrin spray) **S-3-OMe-AMPA** [84799-54-2] (9.3mg, >79% yield of *RS-material*) with $[\alpha]_D^{20}$ -8.6 (c 0.47, H₂O), $[\alpha]_{436}^{20}$ -14.5 (c 0.47, H₂O). The Enzygel can still be reused efficiently after washing by centrifugation. The preceding **R-N-Ac-3-OMe-AMPA** (19.1mg 0.079mmol from two combined preparations) was deacetylated and demethylated by heating (oil bath temperature 140°), under reflux and a blanket of N₂ with 48% aqueous HBr (5ml) for 23 minutes (see above for *RS*-compound), cooled, evaporated *in vacuo*, washed with H₂O twice and re-evaporated *in vacuo*, dried also *in vacuo* over KOH and P₂O₅ and recrystallised from EtOH-petroleum ether (b 40-60°) to give **R-AMPA.HBr** [84799-50-8] (14.1g, 67% yield) with $[\alpha]_D^{25}$ -14.2 (c 0.71, EtOH), $[\alpha]_{436}^{25}$ -31.6 (c 0.71, EtOH). When this salt was dissolved in EtOH (0.3ml) and treated with 10v/v% of Et₃N in EtOH to pH 5-6 the amino acid precipitated; which upon recrystallisation from H₂O-EtOH gave pure (by TLC) **R-AMPA** [83654-13-1 anhydrous; 84799-49-5 monohydrate] (9.3mg), that melted with gradual dec > ~200°, and had elemental analysis for the **monohydrate** with $[\alpha]_D^{27}$ +19 (c 0.18, H₂O), and IR (KBr) broad bands with ν_{\max} at 3420, 3000, 1625, 1490, 1400, 1335, 1245 and 1200 cm⁻¹, and ¹H NMR (D₂O, MeCN internal standard at δ 2.02) with δ at 2.21(s, 3H), 2.83(d, 2H) and 2.88(t, 1H). The above **S-3-OMe-AMPA** (14.8mg, 0.074mmol, pooled from two enzymic reactions) was demethylated by reaction with 43% HBr in AcOH (5ml) at 22° and was complete after 77 hours (as shown by TLC). After evaporation *in vacuo*, treatment with H₂O twice (charcoal) and re-evaporation, the residue was dried (KOH and P₂O₅ *in vacuo*) and recrystallisation from EtOH-petroleum ether (b 40-60°) provided pure **S-AMPA.HBr** [84799-52-0] (12.3mg, 62%) with $[\alpha]_D^{25}$ +14.6 (c 0.62, EtOH), $[\alpha]_{436}^{25}$ -31.5 (c 0.62, EtOH). When this salt was dissolved in EtOH (0.3ml) and treated with 10v/v% of Et₃N in EtOH to pH 5-6 the **enantiomeric** amino acid precipitated out. Recrystallisation of this enantiomer from H₂O-EtOH gave **pure** (by TLC) **S-AMPA** [83643-88-3 anhydrous; 84799-51-9 monohydrate] (7.8mg), which had identical physical and spectral properties as its *R*-enantiomer except for the optical rotation which was $[\alpha]_D^{28}$ -21 (c 0.19, H₂O).

Receptor binding studies in rat brain membranes showed that **S-AMPA** was the active isomer. The following IC₅₀ values were determined: *RS*-AMPA = 0.8±0.3 μM; *RS*-5-OMe-AMPA = >100 μM; **S-AMPA** = 0.4±0.1 μM; *R*-AMPA = 4.8±0.8 μM; *S*-Glu = 1.3±0.6 and *R*-Glu = ~100 μM. [Hansen et al *J Med Chem* 26 901 1983, DOI: 10.1021/jm00360a021; for synthesis and activities of heteroaryl analogues of AMPA see Falch et al. *J Med Chem* 41 2513 1998, DOI: 10.1021/jm9801206; for glutamate agonist activity see Krosgaard et al. *Nature*

284 64 1980, DOI:10.1038/284064a0; for binding of ^3H AMPA to rat brain see Honoré et al. *J Neurochem* **38** 173 1982, DOI: 10.1111/j.1471-4159.1982.tb10868.x.]

α -Aminoisobutyric acid (2-amino-2-methylpropionic acid) [62-57-7] $\text{C}_4\text{H}_9\text{NO}_2$, **M 103.1**, **m** sublimes at **280-281°, 335°** (sealed tube), **pK₁²⁵ 2.36**, **pK₂²⁵ 10.21**. Crystallise the acid from aqueous EtOH and dry it at 110°. [Clarke & Dean *Org Synth Coll Vol* **2** 29 1943, DOI: 10.15227/orgsyn.011.0004; Zelinski & Stadnikoff *Chem Ber* **39** 1722 1906, DOI: 10.1002/cber.190603902108; *Beilstein* **4** IV 2616.]

***RS*- β -Aminoisobutyric acid** [144-90-1, 10569-72-9] $\text{C}_4\text{H}_9\text{NO}_2$, **M 103.1**, **m** **176-178°, 178-180°, 181-182°, *R*-(α -methyl- β -alanine isomer** has **m 183°, [α]_D²⁵ -21 (c 0.43, H₂O)**, **pK_{Est(1)} ~3.7**, **pK_{Est(2)} ~10.2**. *RS*- β -Amino-isobutyric acid forms colourless prisms by crystallisation from hot H₂O which are powdered and dried *in vacuo*. The purity is checked by paper chromatography (Whatman 1) using ninhydrin spray to visualise the amino acid; R_F values in 95% MeOH and *n*-PrOH/5N HCOOH (8:2) are 0.36 and 0.50 respectively. [Kupiecki & Coon *Biochemical Preparations* **7** 20 1960, Pollack *J Am Chem Soc* **65** 1335 1943, DOI: 10.1021/ja01247a021.] The ***R*-enantiomer**, isolated from iris bulbs or human urine, crystallises from H₂O and sublimes *in vacuo* [Asen et al. *J Biol Chem* **234** 343 1959, <http://www.jbc.org/content/234/2/343>; *R*(-)-absolute configuration: Balenovic & Bregant *Tetrahedron* **5** 44 1959, DOI: 10.1016/0040-4020(59)80069-7]. The ***RS*-hydrochloride** crystallises from EtOH/Et₂O with **m 128-129°** (also **130°**) [Böhme et al. *Chem Ber* **92** 1258 1959, DOI: 10.1002/cber.19590920604]. [*Beilstein* **4** III 1330.]

5-Aminolaevulinic acid hydrochloride (ALA-HCl, δ -aminolaevulinic acid HCl) [5451-09-2] $\text{C}_5\text{H}_9\text{NO}_3$, **HCl**, **M 167.6**, **m** **148°(dec), 150-151°(dec), 156-158°(dec)**, **pK₁²² 4.05**, **pK₂²² 8.90**. Dry ALA-HCl in a vacuum desiccator over P₂O₅ overnight, then crystallise it by dissolving it in cold EtOH and adding dry Et₂O. Also crystallise it by dissolving in the minimum volume of MeOH, and placing in a desiccator containing dry Et₂O (clamp the desiccator). During several days the Et₂O slowly distils into the MeOH causing the hydrochloride to separate as long needles. Filter them off and dry them in a Fischer pistol. [Neuberger & Scott *J Chem Soc* 1820, 1924 1954, DOI: 10.1039/JR9540001820; Wynn & Corwin *J Org Chem* **15** 203, 207 1950, DOI: 10.1021/jo01147a031; *Beilstein* **4** IV 3265.] It is a photosensitiser and is antineoplastic [Peng et al. *Cancer* **79** 2282 1997, DOI: 10.1002/(SICI)1097-0142(19970615)79:12<2282>.

4-Aminomethylbenzoic acid (α -amino-*p*-toluic acid) [56-91-7] $\text{C}_8\text{H}_9\text{NO}_2$, **M 151.2**, **m** **273-274°, 294-295°, >300°, 345°(dec, sealed tube), 347-350° (sealed tube)**, **pK₁²⁰ 3.59 (CO₂H)**, **pK₂²⁰ 9.64 (NH₂)**. This acid has been prepared in two different ways from *p*-cyanobenzoic acid [Levine & Sedlecky *J Org Chem* **24** 115 1959, DOI: 10.1021/jo01083a608; Albert & Magrath *J Chem Soc* 678 1944, DOI: 10.1039/JR9440000677], and by reduction of *p*-carboxybenzaldehyde oxime [Nair & Baugh *J Org Chem* **38** 2185 1973, DOI: 10.1021/jo00952a016]. A mixture of *p*-cyanobenzoic acid (14g, 619-65-8), Raney cobalt (2g, W-6 or W-7), 28% aqueous NH₃ (40ml) and H₂O (150ml) are shaken in a Parr hydrogenator at 3 atm (initial pressure) and 25° for ~3 hours when the theoretical volume of H₂ is absorbed. The catalyst is filtered off, the filtrate is boiled to remove NH₃ and the solid that separated is collected, and recrystallised from 18 parts of boiling H₂O (charcoal) to give the amino acid (m 347-350°) in 80% yield. It is soluble in 70 parts of H₂O. Alternatively, the oxime of *p*-carboxybenzaldehyde (1g, 619-66-9) in 95% EtOH (100ml) containing 5% Pd/C (100mg) is shaken with H₂ at 30 psi for 18 hours at 25°. The catalyst is filtered off and washed with hot glacial AcOH (2 x 20ml); the combined washings and filtrate are evaporated to dryness and the residue is recrystallised from H₂O (charcoal) to give the white crystalline amino acid (850mg, m 294-295°). The melting point appears to vary with heating rate. Its UV has λ_{max} at 234nm (H₂O), and the ¹H NMR [TFA] has δ at 4.15 (q, *J* = 6Hz, benzylic H), 7.3 (d, *J* = 8Hz, two protons adjacent to H₂NCH₂-) and 7.9 (d, *J* = 8Hz, two protons adjacent to CO₂H). The ***N*-acetyl derivative** crystallises from a large volume of xylene and has **m 199-200°** (lit: Levine & Sedlecky state **m 199-120°**). [For pKa see Goldacre *Nature* **154** 796 1944, DOI: 10.1038/154796b0; *Beilstein* **14** H 487, **14** III 1212, **14** IV 1362.]

***S*-2-Amino-3-methyl-1-butanol (*S*-valinol)** [2026-48-4] $\text{C}_5\text{H}_{13}\text{NO}$, **M 103.2**, **m** **31-32°, 35-36°, b 81°/8mm, 88°/11mm**, **d₄²⁵ 0.926g/ml**, **[α]_D²⁵ + 16.5 (c 6.32, *l* = 2 H₂O)**, **[α]_D²⁵ + 15.6 (EtOH)**, **pK_{Est} ~10.4**. Purify *S*-valinol by vacuum distillation using a short Vigreux column. Alternatively, it is purified by steam distillation. The steam distillate is acidified with HCl; the aqueous layer is collected and evaporated. The residue is dissolv-

ed in butan-1-ol, filtered and dry Et₂O added to crystallise the hydrochloride salt (*hygroscopic*), **m 113°**. The *free base* can be obtained by suspending the salt in Et₂O and adding small volumes of saturated aqueous K₂CO₃ until effervescence is complete and the mixture is distinctly alkaline. At this stage the aqueous layer should appear as a white sludge. The mixture is heated to boiling and refluxed for 30 minutes (more Et₂O is added if necessary). The Et₂O layer is decanted off from the white sludge, the sludge is extracted twice with Et₂O (by boiling for a few minutes), the combined organic layers are dried (KOH pellets), evaporated and the residue is distilled in a vacuum. The *R*-(*-*)-*enantiomer* has [4276-09-9] $[\alpha]_{\text{D}}^{20}$ -16 (c 10, EtOH); and the *racemate RS-valinol* [16369-05-4] has **b 75-77°/8mm, d²⁵ 0.936g/ml, n_D²⁰ 1.4543**. [Nagao et al. *J Org Chem* **55** 1148 1990, DOI: 10.1021/jo00291a012; *Beilstein* **4** III 805, *Fieser* **12** 563, **13** 341, **16** 380.]

S-(+)-3-Aminopentanoic acid [14389-77-6] and *R*-(*-*)-3-aminopentanoic acid [131347-76-7] C₅H₉NO₂, **M 115.1, m (175°), 185°, $[\alpha]_{\text{D}}^{20}$ (+) and (-) 43 (c 0.5, H₂O), pK₁²⁵ 3.54, pK₂²⁵ 10.25**. Crystallise the amino acids from EtOH/Et₂O. [*Beilstein* **4** II 843, **4** III 1342, **4** IV 2635.]

α -Aminothiophene-2-acetic acid [2-(2-thienyl)glycine] [*R*(+) 65058-23-3, *S*(-) 4052-59-9, (-)- 43189-45-3, *RS*(\pm) 21124-40-3] C₆H₇NO₂S, **M 157.2, m 236-237° (*R*), 235-236° (*S*), 208-210°, 223-224° (dec)(*RS*), $[\alpha]_{\text{D}}^{20}$ (+) and (-) 84 (c 1, 1% aqueous HCl), $[\alpha]_{\text{D}}^{25}$ (+) and (-) 71 (c 1 H₂O), pK_{Est(1)}~ 1.5, pK_{Est(2)}~ 8.0**. Recrystallise 2-(2-thienyl)glycine by dissolving it in H₂O (1g in 3 ml), adjusting the pH to 5.5 with aqueous NH₃, diluting with MeOH (20 ml), stirring, adjusting the pH to 5.5 and cooling to 0°. Also recrystallise it from small volumes of H₂O. [*R*-isomer: Nishimura et al. *Nippon Kagaku Zasshi* **82** 1688 1961, *S*-isomer: Johnson & Panetta *Chem Abstr* **63** 14869 1965, Johnson & Hardcastle *Chem Abstr* **66** 10930 1967, *RS*-isomer: LiBassi et al. *Gazz Chim Ital* **107** 253 1977.] The (\pm) *N*-acetyl derivative has **m 191°** (from H₂O), and the (\pm)-methylester hydrochloride has **m 180°** (from MeOH/Et₂O). [Schouteenten et al. *Bull Soc Chim Fr* II 248, II 252 1978]. [*Beilstein* **18** V/12 184.]

5-Amino-*n*-valeric acid (5-aminopentanoic acid, homopiperidinic acid) [660-88-8] C₅H₁₁NO₂, **M 117.2, m 157-158°, 158-161°, pK₁²⁵ 4.25, pK₂²⁵ 10.66**. The acid was isolated from **Kafir Bran 1**, recrystallised from H₂O/EtOH, and dried *in vacuo* over H₂SO₄. When heated above its melting point, it is converted to 2-piperidone with **m 200°**. [Wood & Colver *J Am Chem Soc* **67** 654 1945, DOI: 10.1021/ja01220a042; *Beilstein* **4** IV 2636.] The hydrochloride has [627-95-2] **M 153.6 and m 95-97°** (from EtOH/Et₂O, 1:2; see below); and on heating above its melting point gives it 2-piperidone hydrochloride **m 182°**. [*Beilstein* **4** H 418.] The aminovaleric acid, synthesised from cyclopentanone oxime, was reacted with aqueous potassium cyanate (foaming), evaporated to dryness at 100°, dissolved in hot H₂O, acidified with HCl, boiled with charcoal, filtered and cooled to give δ -ureido valeric acid with **m 178°** (also 177-178° dec, reported). δ -Carbomethoxyaminovaleric acid has **m 71-72°** (from EtOAc, 3.5g/20ml). [Schniepp & Marvel *J Am Chem Soc* **57** 1557 1935, DOI: 10.1021/ja01312a014].

5-Amino-*n*-valeric acid hydrochloride [627-95-2] C₅H₁₁NO₂·HCl, **M 153.6, m 92-94°, m 95-97°, 103-104°**. Crystallise the salt from CHCl₃. Otherwise dissolve it in EtOH and add 2 volumes of Et₂O and chill. [Schniepp & Marvel *J Am Chem Soc* **57** 1557 1935, DOI: 10.1021/ja01312a014]; Woods & Colver *J Am Chem Soc* **67** 654 1945, DOI: 10.1021/ja01220a042; *Beilstein* **4** III 1343, **4** IV 2636.]

Anserine [*N*, β -alanyl-1-methyl-*S*-histidine] [584-85-0] C₁₀H₁₆N₄O₃, **M 240.3, m 238-239°, $[\alpha]_{\text{D}}^{23}$ +12.3 (c 5, H₂O), pK₁²⁵ 2.64, pK₂²⁵ 7.04, pK₃²⁵ 9.49**. Crystallise anserine from aqueous EtOH. It is *hygroscopic* and is best stored as the nitrate salt (see below). Purify it by shaking the nitrate salt with Dowex 3 (x4 free base) and washing with H₂O, evaporating the filtrate and removing H₂O by 3 distillations with 10ml of propan-2-ol. Dissolve the crystals in MeOH and add H₂O dropwise until one phase is obtained and cool. Dry the crystals at 60° over P₂O₅ in a vacuum. The picrate has **m 145°** (from H₂O). [Rinderknecht et al. *J Org Chem* **29** 1968 1964, DOI: 10.1021/jo01030a075; *Beilstein* **25** II 408, **25** IV 4383.] This dipeptide occurs in vertebrate skeletal muscle as in birds and other animals, but apparently not in dog, cow, horse and man (see carnosine below).

S-Anserine nitrate [5937-77-9] C₁₀H₁₆N₄O₃·HNO₃, **M 303.3, m 225°(dec), 226-228°(dec), $[\alpha]_{\text{D}}^{30}$ +12.2**. Likely impurities are 1-methylimidazole-5-alanine and histidine. Crystallise the nitrate from aqueous MeOH or EtOH (needles). Also dissolve ~20g in 25ml of MeOH, add 2-propanol (150-200ml) and store the mixture at 5°

overnight to give shiny needles. Recrystallise it by heating 12g of the nitrate in MeOH (300ml) and adding H₂O (50-60ml) until one phase is obtained and refrigerating overnight. Filter and dry it at 60°/P₂O₅ in a vacuum. [Rinderknecht et al. *J Org Chem* **29** 1968 1964, DOI: 10.1021/jo01030a075; Behrens & duVigneaud *J Biol Chem* **120** 517 1937, <http://www.jbc.org/content/120/2/517>.]

S-Arginine (S-2-amino-5-guanidinopentanoic acid) [74-79-3] C₆H₁₄N₄O₂, M 174.2, m 205°(dec, anhydrous), 207°(dec, 2 H₂O), [α]_D +26.5 (c 5, in 5M HCl), [α]₅₄₆ +32 (c 5, in 5M HCl), pK₁²⁵ 2.18, pK₂²⁵ 9.36, pK₃²⁵ 11.5. S-Arginine crystallises from H₂O as the *dihydrate* and as plates from EtOH. It also crystallises from 66% EtOH. Its solubility in H₂O is 15% at 21°. Its isoelectric point is at pH 10.76. The *racemate* [7200-25-1] has m 228-233° (dec). [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 p 1841 1961, *Beilstein* 4 IV 817.] Natural precursor substrate for nitric oxide synthase which converts it to citrulline and NO. This induces the release of insulin by a nitric oxide dependent mechanism which also has other important physiological effects [Andrew & Mayer 'Enzymatic function of nitric oxide synthases' (review), *Cardiovascular Research* **43** (3) 521 1999, DOI: 10.1016/S0008-6363(99)00115-7].

S-Arginine hydrochloride [1119-34-2] C₆H₁₄N₄O₂·HCl, M 210.7, m 217°(dec), 222°(dec), [α]_D²⁰ +26.9 (c 6, M HCl). A likely impurity is ornithine. Crystallise the salt from H₂O at pH 5-7, by adding EtOH to 80% (v/v). [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 p 1841 1961, *Beilstein* 4 IV 2649.]

S-Argininosuccinic acid {N-(4S-4-amino-4-carboxybutylamino)iminomethyl-S-aspartic acid} [2387-71-5] C₁₀H₁₈N₄O₆, M 290.3, [α]_D²⁴ +16.4 (H₂O). A likely impurity is fumaric acid. In neutral or alkaline solution it readily undergoes ring closure to the '*anhydride*' (see below). Crystallise it from water by adding 1.5 volumes of EtOH. The barium salt is stable at 0-5° if dry. [Westfall *Biochem J* **77** 135 1960, DOI: 10.1042/bj0770135; Ratner & Kunkemueller *Biochemistry* **5** 1821 1966, DOI: 10.1021/bi00870a007.]

S-Argininosuccinic anhydride [28643-94-9] C₁₀H₁₆N₄O₅, M 272.3, [α]_D²³ -10 (H₂O for anhydride formed at neutral pH). Crystallise the anhydride from H₂O by adding two volumes of EtOH. An *isomeric anhydride* is formed if the free acid is allowed to stand at acid pH. In solution, the mixture of anhydride and free acid is formed [see above entry, Ratner & Kunkemueller *Biochemistry* **5** 1821 1966, DOI: 10.1021/bi00870a007; Kowalsky & Ratner *Biochemistry* **8** 899 1969, DOI: 10.1021/bi00831a020].

S-Asparagine (2-aminosuccinic acid 4-amide) [70-47-3] C₄H₈N₂O₃, M 132.1, m 234-235° (monohydrate) [5794-13-8] [α]_D +32.6 (0.1M HCl), pK₁²⁵ 1.98, pK₂²⁵ 8.84. Likely impurities are aspartic acid and tyrosine. Crystallise it from H₂O or aqueous EtOH. It slowly effloresces in dry air. The *R(-)-enantiomer* has [2058-58-4] and optical rotation of opposite sign. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 p 1856 1961, *Beilstein* 4 IV 3005.]

Aspartic acid (2-aminosuccinic acid) C₄H₇NO₄, M 133.1, m 338-339° (*RS*, [617-45-8]), m 271° (*S*, requires heating in a sealed tube [56-84-8]), [α]_D²⁵ +25.4 (3M HCl), pK₁²⁵ 1.99, pK₂²⁵ 3.90. Likely impurities are glutamic acid, cystine and asparagine. Crystallise the acid from water by adding 4 volumes of EtOH and dry it at 110°. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 p 1856 1961, *Beilstein* 4 IV 2998, 3000.]

L-Aspartic acid β-methyl ester hydrochloride [16856-13-6] C₅H₁₀ClNO₄, M 183.6, m 191-193°, 194°, pK₂₅²⁵ 8.62. Recrystallise it from MeOH by adding anhydrous Et₂O [Bach et al. *Biochemical Preparations* **13** 20 1971].

DL-Aspartic acid dimethyl ester hydrochloride [*RS*- 14358-33-9, *S*- 32213-95-9] C₆H₁₂ClNO₄, M 197.7, 116-117°. Crystallise it from absolute MeOH. [Kovach et al. *J Am Chem Soc* **107** 7360 1985, DOI: 10.1021/ja00311a024.] The *diethyl ester* has pK₂₅²⁵ 6.4.

Azaserine (O-diazoacetyl-S-serine) [115-02-6] C₅H₇N₃O₄, M 173.1, m 146-162°(dec), [α]_D^{27.5} -0.5 (c 8.5, H₂O, pH 5.2), pK_{Est(1)} ~4.53, pK_{Est(2)} ~5.40. Crystallise azaserine from 90% EtOH. Also dissolve it in H₂O,

filter it through Supercel and add EtOH to give azaserine as pale yellow crystals. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 1** pp 75-76 1961, Curphey & Daniel *J Org Chem* **43** 4666 1978, DOI: 10.1021/jo00418a033; *Beilstein* **4** IV 3124.] This antibiotic from *Streptomyces* sp, or by synthesis, has antifungal activity, and because it retards growth it is antineoplastic.

Benzoyl glycine (hippuric acid) [495-69-2] $C_9H_9NO_3$, **M 179.2**, **m 188°**, **pK²⁰ 3.81, 3.59. pK⁴⁰ 3.59**. Crystallise the acid from boiling H₂O. Dry it over P₂O₅. Also purify it by dissolving 135-140g in 2L of boiling H₂O, filtering through a steam-heated funnel and allowing to crystallise at ~20° (yield 115-122g first crop, **m 186-187°**). [Ingersoll & Babcock *Org Synth Coll Vol* **2** 328 1943, DOI: 10.15227/orgsyn.012.0040; *Beilstein* **9** 225, I 100.]

N-Benzoyloxycarbonylglycyl-L-alaninamide [17331-79-2] $C_{13}H_{17}N_3O_4$, **M 279.3**, **m dec >200°**. Recrystallise the dipeptide derivative from EtOH/Et₂O.

N-Benzoyloxycarbonyl-N'-methyl-L-alaninamide [33628-84-1] $C_{12}H_{16}N_2O_3$, **M 236.3**, **m dec >200°**. Recrystallise the amide from EtOAc.

Betaine (1-carboxymethyl-N,N,N-trimethylammonium zwitterion) [107-43-7 (anhydrous), 590-47-6, 17146-86-0 (monohydrate)] $C_5H_{11}NO_2$, **M 117.1**, **m 294-294°(dec) (monohydrate?) 301-305°(dec) (anhydrous), ~319°(dec)**, **pK²⁵ 1.83**. Crystallise betaine from aqueous EtOH or EtOH/Et₂O. The *monohydrate* loses H₂O above 100°. Betaine undergoes internal alkylation to methyl dimethylaminoacetate above its melting point. It is also prepared by treating the hydrochloride (below) with silver oxide and recrystallising from EtOH/Et₂O. [Raman spectra: Edsall *J Am Chem Soc* **65** 1767 1943, DOI: 10.1021/ja01249a029; IR: Leifer & Lippincott *J Am Chem Soc* **79** 5098 1957, DOI: 10.1021/ja01576a006; for pK see Grob et al. *Chem Ind (London)* 1222 1955, *Beilstein* **4** III 1127, **4** IV 2369.]

Betaine hydrochloride [590-46-5] $C_5H_{12}ClNO_2$, **M153.6**, **m 227-228°(dec), 232°(dec), 241-242°, 246-247°(dec)**. Recrystallise the salt from EtOH. Its solubility at 25° is 65% in H₂O, and 5% in EtOH. [Edsall *J Am Chem Soc* **66** 1767 1943, DOI: 10.1021/ja01249a029; Kuhn & Ruelius *Chem Ber* **83** 420 1950, DOI: 10.1002/cber.19500830504; *Beilstein* **4** III 1127, IV 2369.]

Bis-N-tert-butyloxycarbonyl-L-cystine [10389-65-8] $C_{16}H_{28}N_2O_8S_2$, **M 440.5**, **m 144.5-145°, [α]_D²⁰ -133.2 (c 1, MeOH), [α]_D²⁰ -120 (c 2, AcOH), pK_{Est} ~2.9**. Crystallise the cystine derivative from EtOAc by adding hexane [Ferraro *Biochemical Preparations* **13** 39 1971].

Bombesin (2-L-glutamine-3,6-L-asparaginealtesin, a tetradecapeptide) [31362-50-2] $C_{71}H_{110}N_{24}O_{18}S$, **M 1619.9**. Purify Bombesin by gel filtration on a small column of Sephadex G-10 and elute with 0.01 M AcOH. This procedure removes lower molecular weight contaminants which are retarded on the column. The procedure should be repeated twice, and the material should now be homogeneous on electrophoresis; and on chromatography it gives a single active spot which is negative to ninhydrin but positive to Cl₂ and iodoplatinate reagents. R_F on paper chromatography (*n*-BuOH/pyridine/AcOH/H₂O :: 37.5: 25: 7.5: 30) is 0.55 for Bombesin and 0.65 for Alytin. [Bernardi et al. *Experientia* **Part 1** **27** 166 1971, Anastasi et al. **Part 2** **27** 873 1971.] The *hydrochloride* has **m 185°(dec)** (from EtOH) **[α]_D²⁴ -20.6** [c 0.65, Me₂NCHO/(Me₂N)₃PO (8:2)]. [For the stimulation of inositol phosphate see Lloyd et al. *Biochem J* **260** 813 1989, DOI: 10.1042/bj2600813.] This polypeptide was first isolated from the skin of the European fire-bellied toad (*Bombina orientalis*) and has two known homologues in mammals known to cause the release of gastric secretions from G cells. [Gonzalez et al. 'Bombesin-related peptides and their receptors: recent advances in their role in physiology and disease states' *Current Opinion in Endocrinology, Diabetes and Obesity* **15** 58 2008, DOI:10.1097/ MED.0b013e3282f3709b, PMID: 18185064.]

Bradykinin [ArgProProGlyPheSerProPheArg] [5979-11-3] $C_{50}H_{73}N_{15}O_{11}$, **M_r 1,060.2**. Purify Bradykinin by ion-exchange chromatography on CMC (*O*-carboxymethyl cellulose) and partition chromatography on Sephadex G-25. The purity is checked by paper chromatography using BuOH/AcOH/H₂O

(4:1:5) as eluent. [Park et al. *Can J Biochem* **56** 92 1978, DOI: 10.1139/o78-015; ORD and CD: Bodanszky et al. *Experientia* **26** 948 1970, activity: Regoli & Barabé *Pharmacol Rev* **32** 1 1980, *Beilstein* **22** III/IV 91.] Bradykinin is an inflammatory peptide modulator which dilates blood vessels and causes blood pressure to decrease. [Dendorfer et al. 'Pathways of bradykinin degradation in blood and plasma of normotensive and hypertensive rats' *Am J Physiol Heart Circ Physiol* **280** H2182 2001, PMID: 11299220.]

S-Canavanine [2-amino-4-(guanidinooxy)butyric acid] [543-38-4] **M 176.2, m 184°**, $[\alpha]_D^{17} +19.4$ (c 2, H₂O), $[\alpha]_D^{20} +7.9$ (c 3.2, EtOH), 184°, $d^{25} 1.61\text{g/cm}^3$, $pK_1^{25} 2.43$, $pK_2^{25} 6.60$, $pK_3^{25} 9.25$. Crystallise S-canavanine from absolute EtOH or aqueous EtOH. [Tomiyaama *J Biol Chem* **111** 45 1935, <http://www.jbc.org/content/111/1/45>, gave $pK_1^{25} 9.25$ (pK_a COOH), $pK_2^{25} 7.40$ (pK_b guanidinium), $pK_3^{25} 11.50$ (pK_b NH₄⁺), Gulland & Morris *J Chem Soc* 763 1935, DOI: 10.1039/JR9350000763; (±) Frankel et al. *J Chem Soc* 3127 1963, DOI: 10.1039/JR9630003127; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2622-2628 1961, *Beilstein* **4** III 1636, **4** IV 3188.] It is a plant amino acid which can mimic S-arginine and be incorporated into proteins with unusual properties; and differs from arginine in having a CH₂ replaced by O. [Rosenthal et al. 'Aberrant, canavanyl protein formation and the ability to tolerate or utilize L-canavanine'. *Experientia* **43** (5) 558 1987, DOI: 10.1007/BF02143585. PMID: 3582574.]

S-Canavanine sulfate (from jackbean, *O*-guanidino-L-homoserine) [2219-31-0] **C₅H₁₂N₄O₃ · H₂SO₄, M 274.3, m 160-165°(dec), 172°(dec)**, $[\alpha]_D^{18.5} +19.8$ (c 7, H₂O), $pK_1^{25} 2.50$ (CO₂H), $pK_2^{25} 6.60$ (α-NH₂), $pK_3^{25} 9.25$ (guanidinooxy). Recrystallise the sulfate by dissolving (~1g) in H₂O (10ml), and adding with stirring 0.5 to 1.0 volumes of 95% EtOH whereby crystals separate. These are collected, washed with Me₂CO/EtOH (1:1) and dried over P₂O₅ in a vacuum. [Hunt & Thompson *Biochemical Preparations* **13** 416 1971, Fearon & Bell *Biochem J* **59** 221 1955, DOI: 10.1042/bj0590221; *Beilstein* **4** III 1636, **4** IV 3188.]

N-Carbamoylglycine (hydantoic acid, *N*-carboxymethylurea, ureidoacetic acid) [462-60-2] **C₃H₆N₂O₃, M 118.0, m 169-170°(dec), m 173-175°(dec), 178-180°(dec)**, pK_a (H₂O) **3.91 (5°), 3.80 (20.3°), 3.87 (32.1°) and 3.89 (50°)**. *Hydantoic* acid is prepared by reaction of potassium cyanate (KNCO) with glycine in H₂O, and is recrystallised from H₂O or EtOH. [Dakin *J Chem Soc* **107** 434 1915, DOI: 10.1039/CT9150700434; King *J Am Chem Soc* **78** 6020 1956, DOI: 10.1021/ja01604a017; Inouye & Watanabe *JCS Perkin Trans I* 1911 1977, DOI: 10.1039/P19770001911.] It has UV with λ_{max} (ε) at 215 (2800) and 240 (165)nm (neutral species: EtOH); 215 (2600) and 240 (105)nm (cation: EtOH + H⁺); 215 (2400) and 240 (80)nm (anion: EtOH + OH⁻) [Crombie & Hooper *J Chem Soc* 3010 1955, DOI: 10.1039/JR9550003010]. The *ethyl ester* [6293-20-5] **M 146.0**, crystallises from H₂O as needles with **m 135°**; and the *amide* [3530-79-8] **M 117.1**, crystallises from H₂O as prisms with **m 204° (180° has also been reported)**. [*Beilstein* **4** H 359, **4** I 477, **4** II 792, **4** III 1163, **4** IV 2411.]

Carnitine see **Vitamin B₇** in 'Physiologically Active....' compounds in this Chapter.

L-Carnosine (β-alanyl-L-histidine) [305-84-0] **C₉H₁₄N₄O₃, M 226.2, m 253°(dec), 258-260°(dec), 260°(capillary tube), 262°(dec)**, $[\alpha]_D^{25} +20.5$ (c 1.5, H₂O), $pK_1^{25} 2.64$, $pK_2^{25} 6.83$, $pK_3^{25} 9.51$. Likely impurities are histidine and β-alanine. Crystallise L-carnosine from water by adding EtOH in excess. Recrystallise it from aqueous EtOH by slow addition of EtOH to a strong aqueous solution of the dipeptide. Its solubility in H₂O is 33.3% at 25°. [Vinick & Jung *J Org Chem* **48** 392 1983, DOI: 10.1021/jo00151a026; Turner *J Am Chem Soc* **75** 2388 1953, DOI: 10.1021/ja01106a032; *Beilstein* **25** H 516, **25** I 717, **25** II 408, **25** IV 4381.]

S-Citrulline (2-amino-5-ureidopentanoic acid) [372-75-8] **C₆H₁₃N₃O₃, M 175.2, m 222°**, $[\alpha]_D^{20} +24.2$ (in 5M HCl), $pK_1^{25} 2.43$, $pK_2^{25} 9.41$. Likely impurities are arginine and ornithine. Crystallise S-citrulline from water by adding 5 volumes of EtOH. Also crystallise it from water by addition of MeOH. [Ellenbogen *J Am Chem Soc* **74** 5198 1952, DOI: 10.1021/ja01140a065; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2491-2494 1961, *Beilstein* **4** IV 2647.] Intermediate in the urea cycle and in a pathway by which mammals excrete ammonia.

Corticotropin [92307-52-3] **polypeptide from bovine hypothalamus M_r ~4697**. The extract is purified by

ion-exchange on CM-cellulose, desalted, evaporated and lyophilised. Then separated from impurities by gel filtration through Sephadex G-50. [Lande et al. *Biochemical Preparations* **13** 45 1971, Esch et al. *Biochem Biophys Res Commun* **122** 899 1984, DOI:10.1016/0006-291X(84)91175-6].

Creatine (N-guanidino-N-methylglycine) [6020-87-7 (*monohydrate*), 57-00-1 (*anhydrous*)] $C_4H_9N_3O_2$, **M 131.1 (anhydrous), 149.1 (hydrate) m 303°**, pK_1^{25} **2.63**, pK_2^{25} **14.3**. Likely impurities are creatinine and other guanidino compounds. It crystallises from the minimum volume of boiling H_2O as the *monohydrate*. The *hydrate* is also obtained by dissolving in H_2O and adding Me_2CO . Drying under vacuum over P_2O_5 or drying at 100° gives the *anhydrous* base. The *anhydrous base* can be obtained also by dissolving the hydrate in H_2O , seeding with the anhydrous base and cooling in ice. A **m** of **258-268°(dec)** was reported. The *picrate* crystallises from 17 parts of H_2O with **m** of **218-220°(dec)**. [King *J Chem Soc* 2377 1930, DOI: 10.1039/JR9300002374; anhydrous: Huffman et al. *J Am Chem Soc* **58** 1728 1936, DOI: 10.1021/ja01300a066; Mendel & Hodgkin *Acta Cryst* **7** 443 1954, DOI: 10.1107/S0365110X54001326; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 2750 1961, *Beilstein* **4** III 1170, **4** IV 2425.]

Creatine phosphate di Na, 4H₂O salt (phosphocreatine) [922-32-7] **M 327.1**, pK_1^{27} **2.7**, pK_2^{27} **4.58**, pK_3^{27} **~12**. To 3-4g of the salt in H_2O (220ml) is added 4 volumes of EtOH with thorough stirring and allowing to stand at 20° for 12 hours (this temperature is critical as crystals do not readily form at 23° or 25°). The salt first appears as oily droplets which slowly settle and crystallise. After 12 hours the supernatant is clear. Stirring and scratching the flask containing the filtrate brings out additional crystals (0.3-1g) if the salt is kept at 20° for 12 hours. Filter it off at room temperature, wash with 3 x 5ml of ice-cold 90% EtOH, then 5ml of absolute EtOH and dry it in a vacuum desiccator (Drierite or $CaCl_2$) for 16-30 hours. The *hexahydrate* (plates) is converted to the *tetrahydrate* salt (needles) in a vacuum at -10°. [Ennor & Stocken *Biochemical Preparations* **5** 9 1957, *Biochem J* **43** 190 1948, DOI: 10.1042/bj0430190; *Beilstein* **4** III 1170, **4** IV 2425.]

Creatinine [2-amino-1-methylimidazolidin-4-one, 2-imino-1-methyl-4(3H)-oxoimidazolidine] [60-27-5] $C_4H_7N_3O$, **M 113.1**, **m 295°(dec), ~305°(dec)**, pK_1^{25} **4.80**, pK_2^{25} **9.2**. Likely impurities are creatine and ammonium chloride. Dissolve it in dilute HCl, then neutralise with ammonia. Recrystallise it from H_2O by adding excess of Me_2CO . The *picrate* crystallises from 23 volumes of boiling H_2O and has **m 220-221°(dec)**. [King *J Chem Soc* 2377 1930, DOI: 10.1039/JR9300002374; *Beilstein* **25** III/IV 3543; see p 1007.]

Cycloserine [2-amino-3-isoxazolidone; D-(R-natural) and L-(S-non-natural)] [*R*- 68-41-7 and *S*- 339-72-0] $C_3H_6N_2O_2$, **M 102.1**, **m 145-150° (dec), 154-155°, 155-156° (dec), 156° (dec)**, $[\alpha]_D^{25}$ **(+) and (-) 137 (c 5, 2N NaOH)**, pK_1^{10} **4.5**, pK_2^{10} **7.74**, pK_1^{25} **4.50**, pK_2^{25} **7.43**, pK_1^{50} **4.44**, pK_2^{50} **7.20**. Purify cycloserine by recrystallisation from aqueous EtOH or MeOH or aqueous NH_3 /EtOH or isoPrOH. Also recrystallise it from aqueous ammoniacal solution at pH 10.5 (100mg/ml) by diluting with 5 volumes of isopropanol and then adjusting to pH 6 with acetic acid. An aqueous solution, buffered to pH 10 with Na_2CO_3 , can be stored in a refrigerator for 1 week without decomposition. Its UV has λ_{max} at 226nm ($A_{1cm}^{1\%}$ 4.02). The *tartrate salt* has **m 165-166° (dec), 166-168° (dec)**, and $[\alpha]_D^{24}$ **-41 (c 0.7, H₂O)**. [Stammer et al. *J Am Chem Soc* **79** 3236 1957, DOI: 10.1021/ja01569a065; UV: Kuehl *J Am Chem Soc* **77** 2344 1955, DOI: 10.1021/ja01613a105; *Beilstein* **27** III/IV 5549.] The *R-isomer* interferes with D-Ala transport and D-ala to L-Ala bond formation and interconversion of these, hence its bacteriostatic action; and is an antibiotic for Gram-positive bacteria. The *S-isomer* inhibits ketosphinganine synthase and blocks sphingosine biosynthesis.

Cystamine dihydrochloride [2,2'-diaminodiethylene disulfide dihydrochloride, 2,3'-dithio-bis(ethylamine) dihydrochloride] [56-17-7] $C_4H_{12}N_2S_2 \cdot 2HCl$, **M 225.2**, **m 217-220°(dec), 219-220°(dec)**, pK_1^{30} **8.82**, pK_2^{30} **9.58**. Recrystallise the salt by dissolving in EtOH containing a few drops of dry EtOH/HCl, filtering and adding dry Et_2O . The solid is dried in a vacuum and stored in a dry and dark atmosphere. It has been recrystallised from EtOH (solubility: 1g in 60ml of boiling EtOH) or MeOH (plates). The *free base* has **b 90-100°/0.001mm, 106-108°/5mm and 135-136°/760mm**, d_4^{20} 1.1559, n_D^{20} 1.5720. [Verly & Koch *Biochem J* **58** 663 1954, DOI: 10.1042/bj0580663; Gonick et al. *J Am Chem Soc* **76** 4671 1954, DOI: 10.1021/ja01647a055; Jackson & Block *J Biol Chem* **113** 135 1936, <http://www.jbc.org/content/113/1/135>.] The *dihydrobromide* has **m 238-239° (from EtOH/Et₂O)** [Viscontini *Helv Chim Acta* **36** 835 1953, DOI: 10.1002/hlca.19530360412]. [*Beilstein* **4** H 287, **4** IV 1578.] It is an -SH modifying reagent and a *heparin* antagonist.

S,S-(L,L)-Cystathionine (**S-2-amino-2-carboxyethyl-L-homocysteine, L-2-amino-4[(2-amino-2-carboxyethyl)thio]butyric acid**) [56-88-2] $C_7H_{14}N_2OS_2$, M 222.3, m >300°, dec at 312° with darkening at 270°, $[\alpha]_D^{20} +23.9$ (c 1, M HCl). S,S-Cystathionine is purified by converting it to the **HCl** salt in 20% HCl and carefully basifying with aqueous NH_3 until separation is complete. Filter it off and dry it in a vacuum. It forms prisms from H_2O . The **dibenzoyl** derivative has m 229° (from EtOH). [IR: Greenstein & Winitz *Chemistry of the Amino Acids* (J Wiley) Vol 3 p2690 1961 and Tallan et al. *J Biol Chem* 230 707 1958, <http://www.jbc.org/content/230/2/707>, PMID: 13525388; Synthesis: du Vigneaud et al. *J Biol Chem* 143 59 1942, <http://www.jbc.org/content/143/1/59>; Anslow et al. *J Biol Chem* 166 35 1946, <http://www.jbc.org/content/166/1/35>, PMID: 20273671.] [Prepn: Weiss & Stekol *J Am Chem Soc* 73 2497 1951, DOI: 10.1021/ja01150a026; see also Greenberg *Methods Enzymol* 5 936 1962, DOI: 10.1016/S0076-6879(62)05339-2; Beilstein 4 IV 3197.]

Cysteamine (**2-aminoethanethiol, 2-mercaptoethylamine**) [60-23-1] C_2H_7NOS , M 77.2, m 97-98.5°, 98-99°, 99-100°, pK_1^0 9.15, pK_2^0 11.93, pK_1^{30} 8.42, pK_2^{30} 10.83. It is soluble in H_2O giving an alkaline reaction, and it has a disagreeable odour. A likely impurity is the disulfide **cystamine** which is not soluble in alkaline solution. Under a N_2 atmosphere dissolve it in EtOH, evaporate to dryness and wash the white residue with dry petroleum ether, then sublime it at 0.1mm and store it under N_2 at 0-10° in the dark. Its **HgCl₂ (2:3) complex** has m 181-182° (from H_2O), and its **picrate** has m 125-126°. [Mills & Bogert *J Am Chem Soc* 62 1173 1940, DOI: 10.1021/ja01862a053; Baddiley & Thain *J Chem Soc* 800 1952, DOI: 10.1039/JR9520000800; Shirley *Preparation of Organic Intermediates* (J. Wiley) Vol 3 189 1951, Barkowski & Hedberg *J Am Chem Soc* 109 6989 1987, DOI: 10.1021/ja00257a014; Beilstein 4 IV 1570.]

Cysteamine hydrochloride [156-57-0] $C_2H_7NOS \cdot HCl$, M 113.6, m 70.2-70.7°, 70-72°. Purify the salt by recrystallisation from EtOH. It is freely soluble in H_2O and should be stored in a dry atmosphere. [Mills & Bogert *J Am Chem Soc* 62 1173 1940, DOI: 10.1021/ja01862a053.] The **picrate** has m 125-126°; see previous entry for **free base**. [Beilstein 4 IV 1570.]

(±)-Cysteic acid (**3-sulfoalanine, 1-amino-3-sulfopropionic acid**) [*RS*-13100-82-8, 3024-83-7, *R(L)*- 498-40-8] $C_3H_7NO_5S$, M 169.2, m 245°(± form, dec), 260°(+ form, dec). Likely impurities are cystine and oxides of cysteine. Crystallise the acid from water by adding 2 volumes of EtOH. It crystallises from H_2O as the **monohydrate**. The **anhydrous** acid has m ~260°(dec). [Chapeville & Formageot *Biochim Biophys Acta* 26 538 1957, DOI: 10.1016/0006-3002(57)90102-6; Gortner & Hoffman *J Biol Chem* 72 433 1927, <http://www.jbc.org/content/72/1/433>; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 p1908 1961.]

R(L)-Cysteic acid (H₂O) (*R*-abs config) [498-40-8, 23537-25-9] $C_3H_7NO_5S$, M 187.2, m 275-280°(dec), 289°, **L-Cysteic acid** $[\alpha]_D^{20} +8.66$ (c 7.4, H_2O , pH 1) and +1.54 (H_2O , pH 13), pK_1^{25} 1.89 (SO_3H), pK_2^{25} 8.7 (CO_2H), pK_3^{25} 12.7 (? NH_2). Likely impurities are cystine and oxides of cysteine. Crystallise it from water by adding 2 volumes of EtOH. When recrystallised from aqueous MeOH it has m 264-266°, and the **anhydrous** acid has m ~260°(dec). [Chapeville & Formageot *Biochim Biophys Acta* 26 538 1957, DOI: 10.1016/0006-3002(57)90102-6; Riordan & Giese *Methods Enzymol* 47 31 1977, DOI: 10.1016/0076-6879(77)47005-8; Beilstein 4 IV 3296.] It is a precursor of **taurine**.

D-(S)- and L-(R natural enantiomer)- Cysteine (**S- and R-2-amino-3-mercaptopropionic acid**) [*S*(+) 921-01-7, *R*(-) 52-90-4] $C_3H_7NO_2S$, M 121.2, m 230°, 240° (dec), $[\alpha]_D^{20} +$ and -7.6 (c 2, M HCl), + and -10.1 (c 2, H_2O , pH 10), pK_1^{25} 1.92 (CO_2), pK_2^{25} 8.35 (NH_2), pK_3^{25} 10.46 (SH). Purify it by recrystallisation from H_2O (free from metal ions) and dry it in a vacuum. It is soluble in H_2O , EtOH, Me_2CO , EtOAc, AcOH, C_6H_6 and CS_2 . Acidic solutions can be stored under N_2 for a few days without deterioration. [For synthesis and spectra see Greenstein & Winitz *Chemistry of the Amino Acids* (J. Wiley) Vol 3 p1879 1961, Beilstein 4 III 1618, 4 IV 3144.]

L-Cysteine hydrochloride (H₂O) [52-89-1 (anhydrous), 7048-04-6 (monohydrate)] $C_3H_7NO_2S \cdot HCl$, M

175.6, m 175-178° (dec), $[\alpha]_D^{25} +6.53$ (5M HCl). Likely impurities are cystine and tyrosine. Crystallise the salt from MeOH by adding diethyl ether, or from hot 20% HCl. Dry it under vacuum over P₂O₅. **Hygroscopic.** [Beilstein 4 III 1580, 1600.] It is an NMDA (*N*-methyl-D-aspartate) glutamatergic receptor agonist.

(±)-Cysteine hydrochloride [10318-18-0 (anhydrous), 116797-51-5 (monohydrate)] **C₃H₇NO₂S. HCl, M 157.6, m 140-141.5° (dec), pK₂²⁰ 8.36 (NH₂), pK₂²⁰ 10.28 (SH).** Crystallise the salt from hot 20% HCl and dry it under vacuum over P₂O₅. It also crystallises from EtOH with **m 175°** (hydrate?). When crystallised from absolute EtOH or EtOH/Et₂O, it has **m 140-141.5°** (anhydrous?). [Turner & Voitle *J Am Chem Soc* 72 628 1950, DOI: 10.1021/ja01157a510; cf. Albert *Biochem J* 50 690 1952, DOI: 10.1042/bj0500690; Beilstein 4 IV 3145.]

L-(R,R)-Cystine [56-89-3] **C₆H₁₂N₂O₄S₂, M 240.3, $[\alpha]_D^{18.5} -229$ (c 0.92 in M HCl), pK₁²⁵ 1.04 (1.65), pK₂²⁵ 2.05 (2.76), pK₃²⁵ 8.00 (7.85), pK₄²⁵ 10.25 (8.7, 9.85).** Cystine disulfoxide impurity is removed by treating an aqueous suspension with H₂S. The cystine is filtered off, washed with distilled water and dried at 100° under a vacuum over P₂O₅. Crystallise it by dissolving in 1.5M HCl, then adjusting to neutral pH with ammonia. Likely impurities are D-cystine, meso-cystine and tyrosine. Also purify it by dissolving it in 10% NH₃ and adding gradually dilute AcOH until the point of precipitation and cooling slowly [Oughton & Harrison *Acta Cryst* 12 396 1959, DOI: 10.1107/S0365110X59001177]. Alternatively dissolve it in 6N NH₄OH and evaporate it at room temperature for crystallisation to occur. [Chaney & Steinrauf *Acta Cryst B* 30 711 1974, DOI: 10.1107/S0567740874003566; Beilstein 4 IV 3155.]

meso-2,6-Diaminopimelic acid (2R,6S) [583-93-7] **C₇H₁₄N₂O₄, M 190.2, m 295°(dec), 313-315°(dec), pK₁²⁵ 1.04 (1.65, 1.8), pK₂²⁵ 2.05 (2.2, 2.76), pK₃²⁵ 8.00 (7.85, 8.8), pK₄²⁵ 10.25 (8.7, 9.85, 9.9), pI ~5.5.** Crystallise the acid from H₂O or aqueous EtOH. Also purify it by dissolving it in hot H₂O and adding 5 volumes of EtOH, filter after 12 hours at -10°. The acid has been recrystallised from 35% aqueous EtOH. The preparation and separation of *meso*- from *dl*- 2,6-diaminopimelic acid and the enzymic resolution of the latter and some of its derivatives as well as IR spectra were described by Greenstein and coworkers [Izumiya et al. *J Am Chem Soc* 79 648, 651 1957, DOI: 10.1021/ja01560a038]. [Beilstein 4 IV 3081.] Present in the cell wall of some bacteria.

L(S)-2,3-Diaminopropionic acid monohydrochloride (3-amino-L-alanine hydrochloride) [1482-97-9] **C₃H₈N₂O₂. HCl, M 140.6, m 132-133°(dec), 237°(dec), $[\alpha]_D^{25} +26.1$ (c 5.8, M HCl), pK₁²⁵ 1.30, pK₂²⁵ 6.79, pK₃²⁵ 9.51.** It forms needles from H₂O and can be recrystallised from aqueous EtOH. [IR: Koegel et al. *J Am Chem Soc* 77 5708 1977, DOI: 10.1021/ja01626a073; Bisht et al, *J Biol Chem* 287 20369 2012, DOI: 10.1074/jbc.M112.351809; Beilstein 4 IV 2501.] The *laevo* **D(R)-enantiomer** has [6018-56-0], and the **racemic salt** has [54897-59-5] and **m 232°(dec)** (see above)

meso-2,3-Diaminosuccinic acid [23220-52-2, 921-52-8] **C₄H₈N₂O₄, M 148.1, m 305-306°(dec, and sublimes), pK_{Est(1)} ~3.6, pK_{Est(2)} ~9.8.** Crystallise the acid from water. Also, dissolve it in dilute NaOH and add AcOH to pH 5-6 and allow it to crystallise (**m 304° dec**). Alternatively, dissolve the acid in aqueous NH₃ and boil; when the NH₃ has evaporated, the acid separates, filter it off and dry it at room temperature in a vacuum. In another procedure 1g of acid is dissolved in 10ml of concentrated HCl + 15ml of H₂O at 80°, filter immediately, dilute with 20ml of H₂O and allow to stand for 24 hours. When the **monohydrochloride** (0.7g, **m 175-156° dec**) crystallises out, filter and dry it. It has also been purified by dissolving it in the minimum volume of 10% HCl, filtering, and diluting with 5 volumes of H₂O when the crystals separate slowly on standing. The acid is filtered off after 24 hours and dried (**m 306-306° dec**). Similar procedures were used for the *dl*-isomer. [Wenner *J Org Chem* 13 28 1948, DOI: 10.1021/jo01159a004; McKennis & Yard *J Org Chem* 23 980 1958, DOI: 10.1021/jo01101a010; Beilstein 4 III 1528, 4 IV 3025.]

6-Diazo-5-oxo-L-norleucine [157-03-9] **C₆H₉N₃O₃, M 171.2, m 140-150°(dec), 145-155°(dec), $[\alpha]_D^{20} +21$ (c 5, EtOH), pK₁ 2.1, pK₂ 8.95.** Crystallise it from EtOH, H₂O/EtOH, MeOH, 95% aqueous MeOH or H₂O/Me₂CO. Its UV has λ_{max} at 274nm (E1%1cm: 683) and 244nm (E1%1cm: 276) at pH 7 in phosphate buffer. It is a tumour inhibitor. [DeWald & Moor *J Am Chem Soc* 80 3941 1958, DOI: 10.1021/ja01548a036;

Dion et al. *J Am Chem Soc* **78** 3075 1956, DOI: 10.1021/ja01594a036; *Beilstein* **4** IV 3278.]

Diglycyl glycine (triglycine) [556-33-2] $C_6H_{11}N_3O_4$, **M 189.2**, **m 246°(dec)**, **pK₁²⁵ 3.30**, **pK₂²⁵ 7.96**. Crystallise triglycine from H₂O or H₂O/EtOH and dry it at 110°. *Alternatively*, crystallise the tripeptide (103mg) by dissolving it in H₂O (0.4ml) and adding absolute EtOH (0.4ml) to give 64mg (68%) of Glyglygly which can be converted to *N*-*o*-nitrophenoxycetylGlyglygly with **m 212-217°**. [Yakel & Hughes *Acta Cryst* **5** 847 1952, DOI: 10.1107/S0365110X52002318; Hughes & Moore *Acta Cryst* **3** 313 1952, DOI: 10.1107/S0365110X50000793; Holley & Holley *J Am Chem Soc* **74** 3069 1952, DOI: 10.1021/ja01132a035; *Beilstein* **4** III 1198, **4** IV 2469.]

***N,N*-Di-(2-hydroxyethyl)glycine (BICINE, *N,N*-bis-(2-hydroxyethyl)glycine)** [150-25-4] $C_6H_{13}NO_4$, **M 163.2**, **m 193°(dec)**, **193-195°(dec)**, **pK₁²⁵ 2.50**, **pK₂²⁵ 8.11**. Dissolve bicine in a small volume of hot water and precipitate it with EtOH, twice. Repeat once more but treat the aqueous solution with charcoal and filter before adding EtOH. Also crystallise it from concentrated aqueous solutions. [Torn & Kolthoff *J Am Chem Soc* **77** 2061 1955, DOI: 10.1021/ja01613a008; Chaberek et al. *J Am Chem Soc* **75** 2185 1953, DOI: 10.1021/ja01105a049; *Beilstein* **4** IV 2390.] It is a useful buffer and a spacer on Sephadex for plasma protein fractionation.

3-(3,4-Dihydroxyphenyl)-L-alanine (DOPA, LEVODOPA, EUODOPA) [L- 59-92-7, 5796-17-8; DL- 63-84-3] $C_9H_{11}NO_4$, **M 197.2**, **m 275°(dec)**, **267-268°(dec)**, **284-286°(dec)**, **~295°(dec)**, **[α]_D¹⁵ -13.1° (c 5.12, N HCl)**, **pK₁²⁵ 2.32 (CO₂H)**, **pK₂²⁵ 8.72 (NH₂)**, **pK₃²⁵ 9.96 (OH)**, **pK₄²⁵ 11.79 (OH)**. Likely impurities are vanillin, hippuric acid, 3-methoxytyrosine and 3-aminotyrosine. DOPA recrystallises from large volumes of H₂O forming colourless white needles; its solubility in H₂O is 0.165%, but it is insoluble in EtOH, *C₆H₆, CHCl₃, and EtOAc. Also crystallise it by dissolving it in dilute HCl and adding dilute ammonia to give pH 5, under N₂. *Alternatively*, crystallise it from dilute aqueous EtOH. It is rapidly oxidised in air when moist, and darkens, particularly in alkaline solution. Dry it *in vacuo* at 70° in the dark, and store it in a dark container preferably under N₂. It has λ_{max} at 220.5nm (log ε 3.79) and 280nm (log ε 3.42) in 0.001N HCl. [Yamada et al. *Chem Pharm Bull Jpn* **10** 693 1962, Bretschneider et al. *Helv Chim Acta* **56** 2857 1973, DOI: 10.1002/hlca.19730560821; NMR: Jardetzky & Jardetzky *J Biol Chem* **233** 383 1958, <http://www.jbc.org/content/233/2/383>; *Beilstein* **4** IV 2492, 2493.]

3-(3,4-Dihydroxyphenyl)-2-methyl-L-alanine [methyldopa, 2-amino-3-(3,4-dihydroxy-phenyl)-2-methylpropionic acid] [555-30-6, 41372-08-1 (*sesquihydrate*)] $C_{10}H_{13}NO_4$, **M 211.2(anhydr)**, **m >300°**, **300-301°(dec)**, **pK₁²⁵ 2.2**, **pK₂²⁵ 9.2**, **pK₃²⁵ 10.6**, **pK₄²⁵ 12.0**. Recrystallise methyldopa from H₂O. [Reinhold et al. *J Org Chem* **33** 1209 1968, DOI: 10.1021/jo01267a059.] The **L-isomer** forms a *sesquihydrate* from H₂O **m 302-304° (dec)**, and the *anhydrous* crystals are *hygroscopic*, **[α]_D²³ -4.0 (c 1, 0.1N HCl)**, **[α]₅₄₆ +154.5 (c 5, CuSO₄ solution)**. It has λ_{max} at 281nm (ε 2780). Its solubility in H₂O at 25° is ~10mg/ml and the pH of an aqueous solution is ~5.0. It is insoluble in most organic solvents. The **3-methoxy-racemate** has [15073-80-8], separates when an ammoniacal solution of it is adjusted to pH 6.0, the aminoacid is then filtered off, washed with H₂O, dried *in vacuo*, and has **m 218-220°(dec)**. [Stein et al. *J Am Chem Soc* **77** 700 1955, DOI: 10.1021/ja01608a046; *Beilstein* **4** IV 2505.]

3,5-Diiodo-L-thyronine (3,5-diiodo-4-[4-hydroxyphenoxy]-1-phenylalanine) [1041-01-6] $C_{15}H_{13}I_2NO_4$, **M 525.1**, **m 255°(dec)**, **255-257°(dec)**, **[α]_D²⁵ +26 [2N HCl-EtOH (1:2)]**, **pK₁²⁰ 3.25**, **pK₂²⁰ 5.32**, **pK₃²⁰ 9.48**. The amino-acid is purified by dissolving in EtOH containing a little concentrated HCl, decolorising the solution with charcoal, diluting with H₂O, and precipitating it by adding boiling NaOAc solution. If the solution is slightly violet in colour due to some free iodine, aqueous sodium metabisulfite solution should be added dropwise until the colour is discharged as the free iodine is reduced to iodide ions. Collect the solid, wash well with cold H₂O, dry *in vacuo* (over H₂SO₄) and store it dry in the dark. Also, it can be recrystallise from EtOH after decolorising. [Chambers et al. *J Chem Soc* 3424 1949, DOI: 10.1039/JR9490003424; *Beilstein* **14** III 1565, **14** IV 2372.]

3,5-Diiodo-L-tyrosine dihydrate [300-39-0, 18835-59-0] $C_9H_9I_2NO_3 \cdot 2H_2O$, **M 469.0**, **m 199-210°**, **202°(dec)**, **204°(dec)**, **[α]_D²⁰ +2.89 (c 4.9, 4% HCl)**, **pK₁²⁵ 2.12**, **pK₂²⁵ 6.48**, **pK₃²⁵ 7.82**. It forms crystals from

H₂O [solubility (g/L): 0.204 at 0°, 1.86 at 50°, 5.6 at 75° and 17.0 at 100°]. Also recrystallise it from 50% or 70% EtOH. When boiled in EtOH the crystals swell, and on further boiling a gelatinous precipitate is formed [Harrington *Biochem J* **22** 1429 1928, DOI: 10.1042/bj0221436; Jurd *J Am Chem Soc* **77** 5747 1955, DOI: 10.1021/ja01626a091]. It also crystallises from cold dilute ammonia on adding acetic acid to pH 6. Dry under a vacuum. [Beilstein **14** IV 2370.] It is a substrate for assaying thyroid hormone aminotransferase, and an intermediate in an alternative biosynthetic pathways of metabolism of thyroid hormones.

dl-4-Dimethylamino-2,2-diphenylvaleramide (Dimevamide, Aminopentamide) [60-46-8] C₁₉H₂₄N₂O, M **296.4**, m **183-184°**, pK_{Est} ~**9.8**. Crystallise dimevamide from aqueous EtOH. The *hydrochloride* forms leaflets from EtOH/Et₂O with m **190-191°** and is *deliquescent*. The *picrate* has m **210-211°**. It is an antispasmodic. [Moffett et al. *J Am Chem Soc* **79** 4451 1957, DOI: 10.1021/ja01573a056; Beckett & Casy *J Chem Soc* 3076 1957, DOI: 10.1021/ja01573a056, Beilstein **14** III 1363, **14** IV 1865.]

(-)-L-4-Dimethylamino-2,2-diphenylvaleramide [6078-64-4] C₁₉H₂₄N₂O, M **296.4**, m **134.5-135.5°**, **136.5-137.5°**, [α]_D²⁰ **-112** (c **0.87**, EtOH), **-84.1** (c **0.9**, **0.04N** HCl), pK_{Est(1)} **8.3**. Crystallise the amide from petroleum ether, EtOH or as needles from aqueous EtOH. It is an analgesic. [Beckett & Casy *J Chem Soc* 3076 1957, DOI: 10.1021/ja01573a056.]

N,N-Dimethylglycinehydrazide hydrochloride [539-64-0] C₄H₁₂ClN₃O, M **153.6**, m **181°**. Crystallise the salt by adding EtOH to a concentrated aqueous solution. [Viscontini & Meier *Helv Chim Acta* **33** 1773 1950, DOI: 10.1002/hlca.19500330646; Beilstein **4** III 1127, **4** IV 2368.] The *dihydrochloride*, C₄H₁₁N₃O. 2HCl, M **190.1**, has m **214.5°**, and is used as a reagent for aldehydes and ketones.

Djenkolic acid (S,S'-methylene-bis-L-cysteine) [498-59-9] C₇H₁₄N₂O₄S₂, M **254.3**, m **300-350°(dec)**, [α]_D²⁰ **-65** (c **2**, HCl) [See pK of S-methyl-L-cysteine]. Crystallise djenkolic acid from a large volume of water (solubility is 1g/L at 30°). [du Vigneaud & Patterson *J Biol Chem* **114** 533 1936, <http://www.jbc.org/content/114/2/533>; Armstrong & du Vigneaud *J Biol Chem* **168** 373 1947, PMID: 20291097; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2682, 2687 1961, Beilstein **4** III 1591.] The *N,N'-dibenzoyl derivative* crystallises with 1H₂O from aqueous EtOH with m **87.5-89°** [Beilstein **9** III 1171.] The acid is nephrotoxic to humans.

S-Ethionine (S-Ethyl-L-homocysteine, S-2-amino-4-ethylsulfanylbutyric acid) [13073-35-3] C₆H₁₃NO₂S, M **163.2**, m **282°(dec)**, [α]_D²⁵ **+23.7** (in **5M** HCl), pK₂₅ **9.02** (for RS). Likely impurities are *N*-acetyl-(*R* and *S*)-ethionine, *S*-methionine, and *R*-ethionine. Crystallise it from water by adding 4 volumes of EtOH or 85% aqueous EtOH. It sublimes at 196-216°/0.3mm with 99.1% recovery and *unracemised* [Gross & Grodsky *J Am Chem Soc* **77** 1678 1955, DOI: 10.1021/ja01611a085]. [Stekol et al. *J Am Chem Soc* **72** 2309 1950, DOI: 10.1021/ja01161a540; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2658, 2659 1961, Beilstein **4** IV 3194.] It is an antimetabolite and methionine antagonist, highly toxic and a potent carcinogen [Shivapurkar et al. *Carcinogenesis* **5** 989 1984, DOI: 10.1093/carcin/5.8.989, PMID: 6744518].

Ethylene N,N'-bis[(*o*-hydroxyphenyl)glycine] (EHPG) [1170-02-1, 6021-71-2] C₁₈H₂₀N₂O₆, M **360.4**, m **249°(dec)**, pK_{Est(1)} ~**1.8**, pK_{Est(2)} ~**4.8**, pK_{Est(3)} ~**9.0**. Purify it by extensive Soxhlet extraction with acetone. Commercial sample is a 1:1 mixture or *R,R/S,S* pair and the *meso-RS* form. [Vanadium complexes: Bonadies & Carrano *J Am Chem Soc* **108** 4088 1986, DOI: 10.1021/ja00274a038; pKa Schroder & Sheik *Talanta* **21** 250 1974, DOI: 10.1016/0039-9140(74)80119-0; for metal chelation tendencies see Freedman et al. *Nature* **179** 1020 1957, DOI: 10.1038/1791020b0].

2-Fluorophenylalanine [*R*(+) 97731-02-7, *S*(-) 19883-78-4, *RS*(±) 2629-55-2] C₉H₁₀FN₂O₂, M **183.2**, m **226-232°**, **231-234°**, [α]_D²⁵ **(+)** and **(-)** **15** (c **2**, H₂O pH **5.5**), pK₁²⁴ **2.12**, pK₂²⁴ **9.01**. Recrystallise 2-fluorophenylalanine from aqueous EtOH. The *hydrochloride* has m **226-231°(dec)**, and the *N-acetyl derivative* has m **147-149°** (aqueous EtOH). [Bennett & Nieman *J Am Chem Soc* **72** 1800 1950, DOI: 10.1021/

ja01160a110; *Beilstein* **14** III 1268, **14** IV 1671.]

4-Fluorophenylalanine [*R*(+) 18125-46-7, *S*(-) 1132-68-9, *RS*(±) 51-65-0] $C_9H_{10}FNO_2$, **M 183.2**, **m 227-232°**, $[\alpha]_D^{25}$ + and -24 (c 2, H_2O), pK_1^{24} 2.13, pK_2^{24} 9.05. It is recrystallised from aqueous EtOH. The (*R*)-*N*-acetyl derivative has **m 142-145°**, $[\alpha]_D^{25}$ -38.6 (c 8, EtOH). The (*R*)-hydrochloride [122839-52-5] **M 219.6**, has **m 245-251°(dec)**, $[\alpha]_D^{25}$ +11 (c 1, H_2O); the (*S*)-hydrochloride [64231-54-5] has same **M** and **m** but opposite optical rotation. [Bennett & Nieman *J Am Chem Soc* **72** 1800 1950, DOI: 10.1021/ja01160a110; *Beilstein* **14** III 1268.]

L-5-Fluorotryptophan monohydrate [16626-02-1; 154-08-5] $C_{11}H_{11}FN_2O_2$, **M 240.2(H_2O)**, **m reported for the L-enantiomer 158-163°(dec)**, (±)-isomer >250°(dec), 264-265°(also 238-239° dec reported), $[\alpha]_D^{20}$ +5.5 (c 1, 0.1N HCl), $pK_{Est(1)} \sim 2.5$ (CO_2H), $pK_{Est(2)} \sim 9.4$ (NH_2), $pK_{Est(3)} \sim 16$ (indole-NH). Recrystallise it from EtOH, aqueous EtOH or AcOH. Also, purify it by passage through a Dowex AG1x2 (acetate form) column and recrystallise the L-enantiomer (from enzymic enrichment) from $H_2O/EtOH$, **m 158-163°(dec)**, $[\alpha]_D^{23}$ -8.3 (c 2.5, N NaOH). [Coy et al. *Biochemistry* **13** 3550 1974, DOI: 10.1021/bi00714a022; *Beilstein* **22/14** V 116.] It can be incorporated in place of Trp into proteins, and can sometimes lead to enhanced enzyme activity as in glutathione transferase [see Parsons et al. *Biochemistry* **37** 6286 1998, DOI: 10.1021/bi980219e, PMID: 9572843].

L(*S*)-Glutamic acid [56-86-0] $C_5H_9NO_4$, **M 147.1**, **m 205°(dec)**, 224-225°(dec), $[\alpha]_D^{25}$ +31.4 (c 5, 5M HCl), pK_1^{20} 2.06, pK_2^{20} 4.35, pK_3^{20} 9.85. Crystallise L-glutamic acid from H_2O acidified to pH 3.2 by adding 4 volumes of EtOH, and drying at 110°. Likely impurities are aspartic acid and cysteine. It sublimes at 170-175°/10mm. It melts at 160° with cyclisation to *L-pyrrolidone carboxylic acid*. [King & Kidd *J Chem Soc* 3315 1949, DOI: 10.1039/JR9490003315; Parikh et al. *J Am Chem Soc* **80** 953 1958, DOI: 10.1021/ja01537a052; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 1929-1952 1961, *Beilstein* **4** III 1530, **4** IV 3028.] The *S*-enantiomer is an excitatory neurotransmitter and an agonist to all subtypes of glutamate receptors. The non-natural *R*-enantiomer [6893-26-1] has similar properties except for opposite optical rotation. The racemic *RS*-glutamic acid [617-65-2, 19285-83-7 (H_2O)], which has physiological activity due to its *S* component, crystallises from H_2O [solubility is 1.2%(25°), 5%(50°), (25°), 12%(75°) and 28%(100°)], and has **m 225-227°(dec)** [199°(dec) and 201°(open tube) also reported], **b 94-95°/3mm**.

L-Glutamic acid- γ -benzyl ester [1676-73-9] $C_{12}H_{15}NO_4$, **M 237.3**, **m 179-181°**, 181-182°, $[\alpha]_D^{20}$ +19.3 (c 1, AcOH), pK_1^{25} 2.17, pK_2^{25} 9.00. Recrystallise the ester from H_2O and store it at 0°. [Estrin *Biochemical Preparations* **13** 25 1971, *Beilstein* **6** IV 2538.]

L-Glutamine [56-85-9] $C_{12}H_{15}NO_4$, **M 146.2**, **m 184-185°**, 187°, $[\alpha]_D^{25}$ +31.8 (M HCl), $[\alpha]_D^{22}$ +5.6 (H_2O), pK_1^{25} 2.17, pK_2^{25} 9.13. Likely impurities are glutamic acid, ammonium pyroglutamate, tyrosine, asparagine, isoglutamine, arginine. Crystallise it from water [solubility is 1.2%(0°), 2.6%(50°), (18°), 4.8%(30°)] or aqueous EtOH. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 1929-1925 1961, *Beilstein* **4** IV 3038.] The racemic (*RS*)-glutamine [585-21-7] crystallises in prisms from aqueous Me_2CO and has **m 173-173.5°** [Kline & Cox *J Org Chem* **26** 1854 1861, DOI: 10.1021/jo01065a040] and glistening prisims from H_2O/Me_2CO with reported **m 185-186°** [King & Kidd *J Chem Soc* 3315 1949, DOI: 10.1039/JR9490003315].

L-Glutathione (GSH, reduced form, γ -L-glutamyl-L-cysteinyl-glycine) [70-18-8] $C_{10}H_{17}N_3O_6S$, **M 307.3**, **m 188-190°(dec)**, 195°(dec), $[\alpha]_D^{20}$ -20.1 (c 1, H_2O), pK_1^{25} 2.12 (CO_2H), pK_2^{25} 3.59 (CO_2H), pK_3^{25} 8.75 (NH_2), pK_4^{25} 9.65 (10.0, SH). Crystallise L-glutathione from 50% aqueous EtOH, dry it in a vacuum and store it below 5°. Alternatively, recrystallise it from aqueous EtOH under N_2 , and store it dry in a sealed container below 4°. It is freely soluble in H_2O . It has been isolated via its Zn Hg or Cu complexes. [Weygand & Geiger *Chem Ber* **90** 634 1957, DOI: 10.1002/cber.19570900428; Zn complex: Li et al. *J Am Chem Soc* **76** 225 1954, DOI: 10.1021/ja01630a058; Martin & Edsall *Bull Soc Chim Fr* **40** 1763 1958; Berse et al. *Can J Chem* **37** 1733

1959, DOI: 10.1139/v59-251; du Vigneaud & Miller *Biochemical Preparations* **2** 87 1952, Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 2** p 1523 1961, *Beilstein* **4** IV 3165.] A 5-10mM solution will elute glutathione S-transferase (GST) from a glutathione agarose affinity column.

L-Glutathione (oxidised) [27025-41-8] $C_{20}H_{32}N_6O_{12}S_2$, **M 612.6, m 175-195°, 195°, $[\alpha]_D^{20}$ -98 (c 2, H₂O), pK₁ 3.15, pK₂ 4.03, pK₃ 8.75.** Purify it by recrystallisation from 50% aqueous EtOH. Its solubility in H₂O is 5%. Store it at 4°. [Zn complex: Li et al. *J Am Chem Soc* **76** 225 1954, DOI: 10.1021/ja01630a058; Berse et al. *Can J Chem* **37** 1733 1959, DOI: 10.1139/v59-251; *Beilstein* **4** IV 3168.]

Glycinamide hydrochloride [1668-10-6] $C_2H_6N_2O_2 \cdot HCl$, **M 110.5, m 186-189°, 203-205°, 207-208°, pK₁²⁵ -6.10, pK₂²⁵ -1.78, pK₃²⁵ 7.95.** Crystallise the salt from EtOH, EtOH/H₂O or MeOH. Generally dissolve the salt in the minimum volume of H₂O, add a three-fold volume of EtOH (or MeOH) and allow to crystallise over 3 days at -5° (yield >85%). [Karmas & Spoerri *J Am Chem Soc* **74** 1580 1952, DOI: 10.1021/ja01126a070; *Beilstein* **4** IV 2358.] Useful buffer at pH ~7, and for synthesising pyrazine.

Glycine (aminoacetic acid, glyocol) [56-40-6] $C_2H_5NO_2$, **M 75.1, m 262° (dec, goes brown at 226°, sublimes at 200°/0.1mm), d₄²⁵ 1.1607g/ml, pK₁²⁵ 2.35 (CO₂H), pK₂²⁵ 9.78 (α-NH₂).** Crystallise glycine from distilled water by dissolving at 90-95°, filtering, cooling to about -5°, and draining the crystals centrifugally. Alternatively, crystallise it from distilled water by addition of MeOH or EtOH (e.g. 50g dissolved in 100ml of warm water, and 400ml of MeOH is added). The crystals are washed with MeOH or EtOH, then with diethyl ether and dried *in vacuo*. Likely impurities are ammonium glycinate, iminodiacetic acid, nitrilotriacetic acid or/and ammonium chloride. It is **polymorphic** — has three forms. Its solubility in H₂O (g/100ml) is 25 (25°), 54.4 (75°) and 67 (100°); it is almost insoluble in EtOH (0.06%), and has some solubility in pyridine (0.61% w/v). [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 1955 1961, *Beilstein* **4** IV 2349; Fieser **1** 412.] Glycine tastes sweet and is an essential amino acid of proteins. It is an inhibitory neurotransmitter in the spinal chord, and is an allosteric regulator of NMDA (*N-methyl-D-aspartate*) receptors [Betz & Harvey *Glycine as a Neurotransmitter* eLS (Encyclopedia of Life Sciences), Wiley & Sons, 2001, DOI: 10.1038/npg.els.0000140.]

Tetraglycine Hydroperiodide [7097-60-1] $C_{16}H_{42}I_7N_8O_{16}$, with the composition of $[(H_2NCH_2COOH)_4 \cdot HI \cdot 5/4I_2]$, **M 1490.9, decomposes between 162° and 167°.** When **diglycine monohydroiodide** (cf. the corresponding hydrochloride below) (139g) in H₂O (1L) is placed in a wide-mouthed glass-stoppered flask with glycine (225g) and I₂ (80g), and stirred vigorously at 70°, a dark brown precipitate separates within a few minutes. After slow cooling, filtering and drying at ~25° (or slightly above to sublime any free elemental I₂), large flat needle (or blade) shaped crystals having a brassy-bronze metallic lustre in reflected light are obtained. They should be stored in the dark. [Frost & Eddy *J Am Chem Soc* **74** 1346 1952, DOI: 10.1021/ja01125a509]. It is soluble in H₂O (38g/100ml at ~25°). It is sold as tablets (20mg hyperiodide, 90mg disodium dihydrogen pyrophosphate, 5mg talc) which will dissolve in less than 1 minute in 1L of H₂O at ~25°, for disinfecting drinking H₂O by liberating 8mg of I₂. The water is palpable and if the tablets are packaged properly they should be stable under adverse climatic conditions for extended periods. [Morris et al. *Ind Eng Chem* **45** 1013 1953, DOI: 10.1021/ie50521a043.] It is also used for the treatment of hyperthyroidism and exposure to radiation.

Glycine anhydride (2,5-diketopiperazine) [106-57-0] $C_4H_6N_2O_2$, **M 114.1, m 309-310°, 311-312°(dec), ~315°(dec), pK₁ -4.45, pK₂ -2.16 (pK₂ -1.94 in AcOH).** Recrystallise glycine anhydride from H₂O (plates), and it can be sublimed (slowly) at 260° or at 140-170°/0.5mm. The **dihydrochloride** has **m 129-130°** and is prepared by dissolving it in concentrated HCl and adding EtOH to crystallisation point; dry it in a vacuum. The **bis-1-naphthylurethane** has **m 232°(dec)**, and the **diperchlorate** has **m 117° (hygroscopic)**. [MS: Johnstone *JCS Perkin Trans 1* 1297 1975, DOI: 10.1039/P19750001297; NMR: Bláha & Samek *Coll Czech Chem Commun* **32** 3780 1967, <http://dx.doi.org/10.1135/cccc19673780>; X-ray Cryst: Corey *J Am Chem Soc* **60** 1599 1938, DOI: 10.1021/ja01274a023; *Beilstein* **24** IV 1070.]

Glycine ethyl ester hydrochloride [623-33-6] $C_4H_9NO_2 \cdot HCl$, **M 136.9, m 145-146°, pK₂₅ 7.69.** Crystallise it from absolute EtOH or EtOH/Et₂O. [Marvel *Org Synth Coll Vol 2* 310 1943, DOI: 10.15227/orgsyn.014.0046; *Beilstein* **4** II 780, **4** III 3 75.]

Glycine *tert*-butyl ester (glycine 1,1-methylethyl ester) [6456-74-2] $C_6H_{13}NO_2$, M 131.2, b 29-31°/2mm, 65-67°/20mm, d_D^{20} 1.4237, n_D^{25} 1.424, $pK_{Est} \sim 7.6$. The ester, prepared from *tert*-butyl azidoacetate by catalytic reduction (5% Pd/C and H_2), has been purified *via* the phosphite salt. To the ester (23.6g, 0.18mole) in MeOH (150ml) is added phosphorous acid (15g, 0.18mole), the mixture is gently warmed to dissolve the latter and after cooling to 25°, Et_2O (150ml) is added slowly and the stirred mixture is cooled at 0° for 12 hours. The **phosphite salt** is collected, filtered off, washed with Et_2O and dried in an oven at 70° (32g, 82%, m 144-147° dec). After recrystallisation from MeOH-isopropyl ether, the phosphite salt has m 154-157° (dec). The phosphite salt (32g, 0.15mole) is added with stirring into aqueous 6N-sodium hydroxide solution (50ml) until all the solid has dissolved. The mixture is extracted with Et_2O (2 x 20ml), the extract is dried (Na_2SO_4), filtered, evaporated and the residue is distilled under vacuum to give the ***tert*-butyl ester** as an oil (14g, 72%). The versatile *tert*-butyl group is labile under acidic conditions which do not affect a blocked amino grouping. **Glycine *tert*-butyl ester hydrochloride** [27532-96-3] M 167.6, has m 143° (EtOH/ Et_2O). [Vollmar & Dunn *J Org Chem* **25** 387 1960, DOI: 10.1021/jo01073a020; More and Rydon *Org Synth Coll Vol* **5** 586 1973, DOI: 10.15227/orgsyn.045.0047; *Beilstein* **6** IV 2489.]

Glycine hydrochloride [6000-43-7] $C_2H_5NO_2 \cdot HCl$, M 111.5, m 176-178°, 185°, 187°. Crystallise the salt from absolute EtOH or 80% EtOH. **Monoglycine hydrochloride** has m 176-177°, but **diglycine monohydrochloride** has m 187°. Thus when using equivalent quantities of glycine and HCl in H_2O , the former salt is formed, but when **glycine monohydrochloride** (22.3g) and glycine (15g) are dissolved in H_2O (30ml) and heated to almost boiling, colourless crystals (24g after drying) of **diglycine monohydrochloride** (m 186-187°) separate on cooling. [Frost *J Am Chem Soc* **64** 1286 1942, DOI: 10.1021/ja01258a014; *Beilstein* **4** III 1111, **4** IV 2353.]

Glycine methyl ester hydrochloride [5680-79-5] $C_3H_8NO_2 \cdot HCl$, M 125.6, m 174°(dec), 177°(corrected), pK^{25} 7.66. Crystallise the ester salt from MeOH. [Werbin & Spoerri *J Am Chem Soc* **69** 1682 1947, DOI: 10.1021/ja01199a034; *Beilstein* **4** H 340, **4** III 1116.]

Glycine *p*-nitrophenyl ester hydrobromide [7413-60-7] $C_8H_8N_2O_4 \cdot HBr$, M 277.1, m 214° (dec). Recrystallise the ester salt from MeOH by adding diethyl ether. [Alners et al. *Biochemical Preparations* **13** 22 1971].

Glycocyanine (N-guanylglycine) [352-97-6] $C_3H_7N_3O_2$, M 117.1, m 280-284°(dec), >300°, pK^{25} 2.86 (NH_3^+). Recrystallise it from 15 parts of hot H_2O , or by dissolving it in slightly more than the calculated amount of 2N HCl and precipitating it by adding an equivalent of 2N NaOH, filtering, washing with cold H_2O and drying first *in vacuo*, then at 60° *in vacuo*. The **hydrochloride** has m 200°(dec) after recrystallisation from aqueous HCl as plates. The **picrate** forms needles from hot H_2O with m 210°(dec). [Brand & Brand *Org Synth Coll Vol* **3** 440 1955, DOI: 10.15227/orgsyn.022.0059; Failey & Brand *J Biol Chem* **102** 768 1933, <http://www.jbc.org/content/102/2/767>; King *J Chem Soc* 2374 1930, DOI: 10.1039/JR9300002374; *Beilstein* **4** H 359, **4** I 477, **4** II 793, **4** III 1165.]

N-Glycylanilide [555-48-6] $C_8H_{10}N_2O$, M 150.2, m 62°, $pK_{Est} \sim 8.0$. N-Glycylanilide crystallises from water as needles (**dihydrate**) and is soluble in Et_2O . [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol **3** pp1915-1970 1961, *Beilstein* **4** H 343.]

Glycylglycine [556-50-3] $C_4H_8N_2O_3$, M 132.1, m 260-262°(dec), pK^{20} 8.40, pK^{30} 8.04, pK_1^{15} 3.19, pK_2^{15} 8.40. Crystallise glycylglycine from aqueous 50% EtOH or water at 50-60° by addition of EtOH. Its solubility in H_2O is 13.2 w/v% at 20°. Dry it at 110°. It sublimes at 190-200°/0.3mm with 30% recovery [Gross & Gradsky *J Am Chem Soc* **77** 1678 1955, DOI: 10.1021/ja01611a085; King *J Am Chem Soc* **79** 6153 1957, DOI: 10.1021/ja01580a013]. [*Beilstein* **4** IV 2459.]

Glycylglycine hydrochloride [13059-60-4, 23273-91-8] $C_4H_8N_2O_3 \cdot HCl$, M 168.6, m 215-220°, 235-236°, 260-262°, pK_1^{25} 3.12, pK_2^{25} 8.17. Crystallise the salt twice from 95% EtOH. Single crystals are formed by slow evaporation of an aqueous solution. [Mellon & Hoover *J Am Chem Soc* **73** 3879 1951, DOI: 10.1021/ja01152a095; for Raman spectrum see Garfinkel & Edsall *J Am Chem Soc* **80** 3818 1958, DOI:

10.1021/ja01548a003; *Beilstein* **4** IV 2469.]

Glycyl-L-proline [704-15-4] $\text{C}_7\text{H}_{12}\text{N}_2\text{O}_3$, **M 172.2**, **m 184°(dec)**, **185°**, **204°**, **pK₁²⁵ 2.81**, **pK₂²⁵ 8.65**. Crystallise glycyl-L-proline from water at 50-60° by addition of EtOH. [UV: Saidel *J Am Chem Soc* **77** 3892 1955, DOI: 10.1021/ja01619a063; Bergmann et al. *Z Physiol Chem* **212** 79 1932, *Beilstein* **22** IV 49.] Exhibits anti-ischemic effects on neuroactive amino acids.

dl-Glycylserine [687-38-7] $\text{C}_5\text{H}_{10}\text{N}_2\text{O}_4$, **M 162.2**, **m 197-199°(dec, sealed tube)**, **207°(dec)**, **pK₁²⁵ 2.92**, **pK₂²⁵ 8.10**. Crystallise it from H₂O (charcoal) by addition of EtOH. [Fölsch *Acta Chem Scand* **12** 561 1958, DOI: 10.3891/acta.chem.scand.12-0561; NSR: Bovey & Tiers *J Am Chem Soc* **81** 2870 1959, DOI: 10.1021/ja01520a063; *Beilstein* **4** III 1572, **4** IV 3140.]

Gramicidin A (a linear pentadecapeptide from *Bacillus brevis*) [11029-61-1] **m ~229-230°(dec)**. Purify gramicidin A by countercurrent distribution from $\text{C}_6\text{H}_6/\text{CHCl}_3$, MeOH/H₂O (15:15:23:7) with 5000 tubes. Fractions are examined by UV (280nm) of small aliquots. Separation from gramicidin C and other material occurred after 999 transfers. Collectively called *Gramicidin D* [1405-97-6] $\text{C}_{99}\text{H}_{140}\text{N}_{20}\text{O}_{17}$, **M 1882.3**. [Gramicidin A, B and C, prep: Gross & Witkop *Biochemistry* **4** 2495 1965, DOI: 10.1021/bi00887a032; synth: Bauer et al. *Biochemistry* **11** 3266 1972, DOI: 10.1021/bi00767a022.] Purify it finally by recrystallisation from EtOH/H₂O and dry it at 100°/10⁻²mm over KOH. It forms platelets **m 229-230°**. It is almost insoluble in H₂O (0.6%) but soluble in lower alcohols, dry Me₂CO, dioxane, acetic acid and pyridine. The commercial material is more difficult to crystallise than the synthetic compound. [*seco*-Gramicidin: Sarges & Witkop *J Am Chem Soc* **86** 1861 1964, DOI: 10.1021/ja01063a048; Gramicidin A, structure: **87** 2011 1965, DOI: 10.1021/ja01087a027; Gramicidin A, synth: **87** 2020 1965, DOI: 10.1021/ja01087a028; Gramicidin B, structure: **87** 2027 1965, DOI: 10.1021/ja01087a029] It has characteristic $[\alpha]_D^{20}$ +27.3 (c 1.3, MeOH) and UV with λ_{max} at 282nm (ϵ 22,100). The *N-carbamoyldeformyl gramicidine A* precipitates from EtOAc/petroleum ether (b 40-60°). [*Beilstein* **26** III/IV 4273.] The difference between A, B, and C is in residue 11 which is Trp, Phe and Tyr respectively.

Gramicidin C (a linear pentadecapeptide from *Bacillus brevis*) [9062-61-7]. Purify as for Gramicidin A since they are isolated together and separated. [Structure: Sarges & Witkop *Biochemistry* **4** 2491 1965, DOI: 10.1021/bi00887a031; Hunter & Schwartz 'Gramicidins' in *Antibiotics I* (Gotlieb and Shaw Eds) Springer-Verlag, NY, p.642 1967, as well as references above for Gramicidin A.]

Gramicidin S [113-73-5] $\text{C}_{60}\text{H}_{92}\text{N}_{12}\text{O}_{10}$, **M 1120.0**, **m 268-270°**, $[\alpha]_D^{25}$ -290 (c 0.5, EtOH + 30mM aqueous HCl {7:3}). Gramicidin S crystallises from EtOH. The *di-HCl* [15207-30-4] crystallises from EtOH (+ few drops of HCl) with **m 277-278°** (see below). [¹³C NMR in MeOH and in Me₂SO: Gibbons et al. *Nature* **227** 840 1970, DOI:10.1038/227840a0; *Beilstein* **26** III/IV 4273.] Unlike Gramicidins A, B and C, this antibiotic is a cyclic decapeptide with two ornithine residues which have free amino groups, hence it forms salts readily (see below). Potentially harmful, protect eyes.

Gramicidin S 2HCl (from *Bacillus brevis* Nagano) [15207-30-4] $\text{C}_{60}\text{H}_{92}\text{N}_{12}\text{O}_{10} \cdot 2\text{HCl}$, **M 1192.9**, **m 277-278°(dec)**, $[\alpha]_D^{24}$ -289 (c 0.4, 70% H₂O+EtOH). It crystallises in prisms from EtOH+aqueous HCl.

N-Guanyltiramine hydrochloride [60-20-8] $\text{C}_9\text{H}_{13}\text{N}_3\text{O} \cdot \text{HCl}$, **M 215.7**, **m 218°**, **pK₁ 10.2 (phenolic OH)**, **pK₂ 12.4 (guanidino N)**. Purify the salt on a phosphocellulose column and elute with a gradient of aqueous NH₃ (0-10%). The second major peak has the characteristic tryptamine spectrum and is collected, and lyophilised to give white crystals of the *dihydrate* which dehydrate at 100°. It has UV with λ_{max} at 274.5nm (ϵ 1,310) in 0.1N NaOH, and 274.5nm (ϵ 1,330) at pH 7.0. Excitation λ_{max} is at 280nm and emission λ_{max} is at 330nm. [For use of ¹²⁵I N-guanyltiramine in assaying ADP-ribosyltransferase activity in cholera toxin see Mekalanos et al. *J Biol Chem* **254** 5849 1979, <http://www.jbc.org/content/254/13/5849>, PMID: 447682.]

S-Histidine [71-00-1] $\text{C}_6\text{H}_9\text{N}_3\text{O}_2$, **M 155.2**, **m 287°(dec)**, $[\alpha]_D^{25}$ -39.7 (c 1, H₂O), **+13.0 (6M HCl)**, **pK₁²⁵ 1.96**, **pK₂²⁵ 6.12**, **pK₃²⁵ 9.17**. A likely impurity is arginine. S-Histidine is adsorbed from aqueous solution onto

a Dowex 50-H⁺ ion-exchange resin, washed with 1.5M HCl (to remove other amino acids), then eluted with 4M HCl as the *dihydrochloride*. This purified *dihydrochloride* (see below) is finally dissolved in water, the pH adjusted to 7.0, and the *free zwitterionic base* crystallises out on addition of EtOH. Its solubility in H₂O is 4.2% at 25°. **RS-Histidine** [4998-57-6] has **m 273°(dec)**. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 pp 1971-1993 1961, *Beilstein* 25 III/IV 4346-4348.]

S-Histidine dihydrochloride [1007-42-7] C₆H₉N₃O₂. 2HCl, **M 242.1, m 245°, [α]_D²⁰ +47.5 (c 2, H₂O)**. The dihydrochloride crystallises from water or aqueous EtOH and is washed with acetone, then diethyl ether. Alternatively, convert it to the histidine *di-(3,4-dichlorobenzenesulfonate)* salt by dissolving 3,4-dichlorobenzenesulfonic acid (1.5g/10ml) in the aqueous histidine solution with warming, and then the solution is cooled in ice. The resulting crystals (**m 280° dec**) can be recrystallised from 5% aqueous 3,4-dichlorobenzenesulfonic acid, then dried over CaCl₂ under vacuum, and washed with diethyl ether to remove excess reagent. The dihydrochloride can be regenerated by passing the solution through a Dowex-1 (Cl⁻ form) ion-exchange column. The solid is obtained by evaporating the solution on a steam bath or better in a vacuum. [Greenstein & Winitz, *The Amino Acids* Vol 3 p 1976 1961, *Beilstein* 25/16 V 366.]

S-Histidine monohydrochloride (H₂O) [5934-29-2 (H₂O), 7048-02-4] C₆H₉N₃O₂. HCl, **M 209.6(H₂O), m 80° monohydrate, 254°(dec, anhydrous), [α]_D²⁵ +13.0 (6M HCl)**. Crystallise the mono-hydrochloride from aqueous EtOH or 60% aqueous EtOH (**m 259°dec**). Alternatively, dissolve 10g in 50ml of H₂O, decolourise with Norite, filter, evaporate it in a vacuum to a syrup, cool to room temperature, add 95% EtOH with stirring until slightly turbid, scratch the sides of the vessel until crystals form, then add slowly 40ml of EtOH and keep at 0° overnight, filter the solid off, wash it several times with EtOH and dry it in a vacuum. [Rose & Cox *J Biol Chem* 68 217 1926, <http://www.jbc.org/content/68/1/217>; Cox et al. *J Biol Chem* 81 755 1929, <http://www.jbc.org/content/81/3/755>; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 pp 1972, 2098 1961, *Beilstein* 25 II 407, 25 III/IV 4346.]

L-Homocysteine (2-amino-4-mercaptobutyric acid) [6027-13-0] C₄H₉NO₂S, **M 135.2, m 232-233°, [α]_D²⁵ +153 (c 13, 5N HCl), pK₁²⁵ 2.22 (CO₂H), pK₂²⁵ 8.87 (NH₃⁺), pK₃²⁵ 10.86 (SH)**. Crystallise L-homocysteine from aqueous EtOH. All operations should be carried out under N₂ as the thiol readily oxidises in air. The acid (3g) is dissolved in freshly boiled H₂O (30ml) under N₂ cooled under N₂ (all operations should be under N₂), add absolute EtOH (100ml), the acid is filtered off, and a second crop is obtained by diluting the filtrate to 500ml with absolute EtOH, keeping overnight in a refrigerator, filtering, washing with EtOH and drying in a vacuum. Store it under N₂ or argon. The *S-benzyl derivative* is repeatedly crystallised from H₂O, or by dissolving it in HCl followed by slow addition of ammonia. It has **m 240-241°, [α]_D²⁵ +27 (c 13, 5N HCl)**. [Riedel & du Vigneaud *J Biol Chem* 112 149 1935, <http://www.jbc.org/content/112/1/149>; du Vigneaud & Patterson *J Biol Chem* 109 97 1935, <http://www.jbc.org/content/109/1/97>; du Vigneaud et al. *J Biol Chem* 126 217 1938, <http://www.jbc.org/content/126/1/217>; du Vigneaud & Brown *Biochemical Preparations* 5 93, 95 1975, Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 pp 2667-2670 1961, *Beilstein* 4 IV 3189, IR: Koegel et al. *J Am Chem Soc* 77 5708 1955I, DOI: 10.1021/ja01626a073.]

dl-Homocysteine (2-amino-4-mercaptobutyric acid) [454-29-5] C₄H₉NO₂S, **M 135.2, m 234-235°(corr, dec)**. Purify it as for the L-isomer under N₂. [Allen & Steinmann *J Am Chem Soc* 74 3932 1952, DOI: 10.1021/ja01135a502; and references for the L-isomer above, *Beilstein* 4 IV 3189.]

dl-Homocystine [462-10-2, 870-93-9 (±)] C₈H₁₆N₂O₄S₂, **M 268.4, m 263-265°(dec), pK₁²⁵ 1.59 (CO₂H), pK₂²⁵ 2.54 (CO₂H), pK₃²⁵ 8.52 (NH₃⁺), pK₄²⁵ 9.44 (NH₃⁺)**. dl-Homocystine crystallises in platelets from water with 1H₂O and **m 258-260°(dec)**, all operations should be carried out under N₂. [Sudo *J Chem Soc Jpn (Pure Chem Sect)* 79 81, 86, 87 1958, *Beilstein* 4 IV 3199.]

L(S,S)-Homocystine [626-72-2] C₈H₁₆N₂O₄S₂, **M 268.4, m 281-284°(dec), [α]_D²⁶ +79 (c 1, M HCl), [α]_D²¹ -16 (c 0.06, H₂O), pK (see above)**. The acid (3g) is dissolved in freshly boiled H₂O (30ml) under N₂, cooled under N₂ (all operations should be under N₂), absolute EtOH (100ml) is added, the acid is filtered off, and a second crop is obtained by diluting the filtrate to 500ml with absolute EtOH, kept overnight in a refrigerator, filtered, washed with EtOH and dried in a vacuum. The **D(R,R)-form** has similar properties but is -ve in M HCl

and +ve in H₂O. [du Vigneaud & Patterson *J Biol Chem* **109** 97 1935, <http://www.jbc.org/content/109/1/97>; du Vigneaud & Brown *Biochemical Preparations* **5** 93, 95 1975, Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2667-2670 1961, *Beilstein* **4** III 1643, **4** IV 3199; IR: Koegel et al. *J Am Chem Soc* **77** 5708 1955, DOI: 10.1021/ja01626a073.]

L-Homoserine (2-amino-4-hydroxybutyric acid) [672-15-1] C₄H₉NO₃, M 119.1, m 203°, [α]_D²⁶ +18.3 (in 2M HCl), pK_{Est(1)} ~2.1, pK_{Est(2)} ~9.3. Likely impurities are *N*-chloroacetyl-L-homoserine, *N*-chloroacetyl-D-homoserine, L-homoserine, homoserine lactone, homoserine anhydride (formed in strong solutions of homoserine if slightly acidic). It crystallises from water by adding 9 volumes of EtOH. It cyclises to the **lactone** in strongly acidic solution — on standing at room temperature for 8 hours, the optical rotation of a solution in HCl decreases to almost zero due to the formation of the **laevo-rotating** lactone. On refluxing a 2N HCl solution for 2 hours and cooling, **L-homoserine-γ-lactone hydrochloride** [2185-03-7] crystallises out and has m 210-220°(dec), C₄H₇NO₂·HCl, M 137.6, [α]_D²⁶ -27.0 (c 5, 2M HCl) and [α]_D²⁰ -27.8 (c 1, H₂O). [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2612-2616 1961, *Beilstein* **4** IV 3187.]

erythro-3-Hydroxy-RS-aspartic acid [6532-76-9] C₄H₇NO₅, M 149.1, pK₁²⁵ 1.91, pK₂²⁵ 3.51, pK₃²⁵ 9.11. When prepared from sodium glyoxalate and copper glycinate in N NaOH at 5° overnight and removing Na ions by stirring with Dowex 50 resin, filtering, concentrating *in vacuo* to a small volume then applied to a Dowex 1 (formate form) column, followed by washing with H₂O to remove glycine, then eluting with N formic acid and the pooled fractions lyophilised, a crude mixture of **erythro** and **threo** 3-hydroxyaspartic acid (44:56) is obtained. These are separated on a Dowex 1 (formate form), whereby the **erythro**- isomer elutes first as shown by being identical with the **erythro** hydroxyamino acid obtained in the transaminase system from sheep brain. The **threo**- isomer is inactive in this enzymic system. Their solubilities in H₂O at 20° are 2.1g/100ml for **erythro** and 0.2g/100ml for **threo**, hence the former is recrystallised from H₂O/EtOH (or small volumes of H₂O), and the latter from a larger volume of H₂O. Likely impurities are 3-chloromalic acid, ammonium chloride, **threo**-3-hydroxyaspartic acid. [Kornguth & Shallach *Arch Biochem Biophys* **91** 39 1960, DOI: 10.1016/0003-9861(60)90451-3.] On heating either diastereomer in 6N HCl at 120° for 96 hours, equilibration to a 5:2 mixture of **erythro** to **threo** occurs. However, when heated in H₂O at 120° some deamination with formation of glycine occurs, and after 50 hours the **erythro/threo** ratio is 1:2. [Kornguth & Shallach *Arch Biochem Biophys* **104** 79 1964, DOI: 10.1016/S0003-9861(64)80037-0; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 1** p 214, **Vol 3** p 2416 1961.]

β-Hydroxy-L-glutamic acid [533-62-0] C₅H₉NO₅, M 163.1, m 100°(hydrate, dec), 135°(anhydr, dec), [α]_D²⁰ +17.6 (c 2, 6M HCl) and [α]_D²⁰ +1.2 (c 2, H₂O). pK₁²⁵ 2.27, pK₂²⁵ 4.29, pK₃²⁵ 9.66. Crystallise the acid from water (prisms). The **racemic** acid also is recrystallised from H₂O (needles) and has m 198°(dec). [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 1** pp 211-213, **Vol 3** p 2422 1961.]

5(R)-Hydroxy-L(2S)-lysine dihydrochloride monohydrate (2S-5R-2,6-diamino-5-hydroxycaproic acid 2HCl, H₂O) [32685-69-1; 13204-98-3 (DL & Allo)] C₆H₁₄N₂O₃·2HCl·H₂O, M 253.1, m 225°(dec), [α]_D²⁵ +17.8 (c 1, 6M HCl), pK₂²⁵ 8.85, pK₃²⁵ 9.83. Likely impurities are 5-*allo*-hydroxy-(D and L)-lysine, histidine, lysine, ornithine. Crystallise the hydrochloride from water by adding 2-9 volumes of EtOH stepwise. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 2000-2009 1961.]

DL-erythro-3-Hydroxynorvaline (2-amino-3-hydroxypentanoic acid) [34042-00-7] M 133.2, m 257-259°(dec), 263°(dec), pK₁²⁰ 2.32, pK₂²⁰ 9.12. Prepared by amination of α-bromo-β-hydroxy-*n*-valeric acid Purify it by recrystallisation from aqueous EtOH. The **Cu salt** has m 255-256° (dec), the **benzoyl** derivative has m 181°, the ***N*-phenylcarbamoyl** derivative has m 164°, and the ***N*-naphthylcarbamoyl** derivative has m 179°, [Buston et al. *J Biol Chem* **204** 665 1953, <http://www.jbc.org/content/204/2/665>, PMID: 13117840; *Beilstein* **4** IV 3220.]

***N*-(*p*-Hydroxyphenyl)glycine** [122-87-2] C₈H₉NO₃, M 167.2, m 244°(dec). Purify by recrystallising from a large volume of H₂O. It is sparingly soluble in most organic solvents but is soluble in aqueous acid or alkali. It

is used in photography (developing agent) and as an acid indicator in bacteriology. With FeCl_3 it produces a methylene blue colour (oxidation to the *p*-quinone-imine?). [Galatis *Helv Chim Acta* **4** 574 1921, DOI: 10.1002/hlca.19210040162; *Beilstein* **13** IV 1210.]

***p*-Hydroxy-D-phenylglycine** [D-2-(4-hydroxyphenyl)glycine] [22818-40-2] $\text{C}_8\text{H}_9\text{NO}_3$, **M 167.2**, **m** >240°(dec), $[\alpha]_{\text{D}}^{20}$ -156 (c 1, M HCl), $\text{pK}_{\text{Est}(1)} \sim 2$, $\text{pK}_{\text{Est}(2)} \sim 4.5$, $\text{pK}_{\text{Est}(3)} \sim 10.3$. Crystallise it from water and dry it *in vacuo*. ***p*-Hydroxy-L-phenylglycine** [32462-30-9] has $[\alpha]_{\text{D}}^{20}$ +158 (c 1, M HCl). [*Beilstein* **14** I 659.]

***trans*-L-4-Hydroxyproline** (2*S*,4*R*-4-hydroxypyrrolidine-2-carboxylic acid) [51-35-4] $\text{C}_5\text{H}_9\text{NO}_3$, **M 131.1**, **m** 273°(dec), 274°, $[\alpha]_{\text{D}}^{20}$ -76.0 (c 5, H_2O), pK_1^{25} 1.86, pK_2^{25} 9.79. Crystallise it from MeOH/EtOH (1:1). Separation from normal *allo*-isomer can be achieved by crystallisation of the copper salts [see Levine *Biochemical Preparations* **8** 114 1961]. Separation from proline is achieved via the crystalline *picrate*, CdCl_2 , or *acid ammonium rhodanate* salts [see Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 2182 1961]. [*Beilstein* **22/5** V 7.]

5-Hydroxy-L-tryptophan [4350-09-8] $\text{C}_{11}\text{H}_{12}\text{N}_2\text{O}_3$, **M 220.2**, **m** 270°(dec), 273°(dec), $[\alpha]_{\text{D}}^{22}$ -32.5, $[\alpha]_{\text{D}}^{20}$ -73.5 (c 1, H_2O), $\text{pK}_{\text{Est}(1)} \sim 2.4$, $\text{pK}_{\text{Est}(2)} \sim 9.0$, $\text{pK}_{\text{Est}(3)} \sim 9.4$, $\text{pK}_{\text{Est}(4)}$ 16 (NH). Likely impurities are 5-hydroxy-D-tryptophan and 5-benzoyloxytryptophan. Crystallise 5-hydroxy-L-tryptophan under nitrogen from water by adding EtOH. Store it under nitrogen. Also dissolve it in the minimum volume of hot H_2O (~0.7g in 4ml) under nitrogen (charcoal) and allow it to crystallise at 5°. The *picrolonate* crystallises from H_2O with **m 184-186°(dec)**. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 2732-2737 1961, Morris & Armstrong *J Org Chem* **22** 306 1957, DOI: 10.1021/jo01354a022; *Beilstein* **22/14** V 278.]

(±)-Ibotenic acid monohydrate (α -[3-hydroxy-5-isoxazolyl]-glycine, α -amino-3-hydroxy-5-isoxazoleacetic acid) [2552-55-8] $\text{C}_5\text{H}_6\text{N}_2\text{O}_4$, **M 176.1**, **m** 144-146° (monohydrate), 151-152° (anhydrous), 148-151°, pK_1 2, pK_2 5.1, pK_3 8.2. It has been converted to the ammonium salt (**m** 121-123° dec) dissolved in H_2O , passed through an Amberlite IR 120 resin (H^+ form) and eluted with H_2O . The acidic fractions are collected, evaporated to dryness and the residue recrystallises from H_2O as the *monohydrate* (**m** 144-146°). The *anhydrous acid* is obtained by making a slurry with MeOH, decanting and evaporating to dryness, and repeating the process twice more (**m** 151-152°). Recrystallisation from H_2O gives back the *monohydrate*. [Improved synth: Nakamura *Chem Pharm Bull Jpn* **19** 46 1971, DOI: 10.1248/cpb.19.46.] The *ethyl ester* forms needles when recrystallised from a small volume of Et_2O and has **m** 78-79°, and IR (CHCl_3) with ν_{max} at 3500-2300 (OH), 1742 (ester C=O), 1628, 1528 cm^{-1} , and UV with λ_{max} at (EtOH) at 206nm (ϵ 7,080). The *hydrazide* has **m** 174-175° (from MeOH) with IR (KBr) 1656 (C=O) cm^{-1} . It occurs in muscarinic mushrooms, is hallucinogenic and is poisonous [Becker et al. *Psychopharmacology* **144** 333 1999, DOI:10.1007/s002130051015. PMID: 10435405].

Iminodiacetic acid [142-73-4] $\text{C}_4\text{H}_7\text{NO}_4$, **M 133.1**, **m** 225°(dec), 240°(dec), 247.5°(dec), **b** 126-127°/14mm, pK_1^{25} 2.50, pK_2^{25} 9.40. Crystallise the acid several times from water. The *N*-Methyl derivative **m** 215° is purified by dissolving it in an equal weight of warm H_2O and adding 3 volumes of MeOH [Kiematsu et al. *Org Synth Coll Vol* **2** 397 1943]. [Chaberek & Martell *J Am Chem Soc* **74** 5052 1952, DOI: 10.1021/ja01140a018; *Beilstein* **4** III 2428, **4** IV 1176.]

3-Iodo-L-tyrosine [70-78-0] $\text{C}_9\text{H}_{10}\text{NO}_3$, **M 307.1**, **m** 205-208°(dec), 210°(dec), $[\alpha]_{\text{D}}^{25}$ -4.4 (c 5, 1M HCl), $\text{pK}_{\text{Est}(2)} \sim 2.1$, $\text{pK}_{\text{Est}(3)} \sim 6.4$, pK_4^{25} 8.7. Likely impurities are tyrosine, diiodotyrosine and iodide. Crystallise it by dissolving it in concentrated ammonia (~200mg in ~20ml), evaporating to ~5ml, and NH_4Cl is added to pH 4.5-5.0. After a few hours at 0°, the amino acid crystallises in needles. It is filtered off, washed with a little ice-cold H_2O and dried in a vacuum. Alternatively, dissolve it in dilute ammonia at room temperature, then add dilute acetic acid to pH 6. Store it at 0°. Recrystallisation of ~250mg from H_2O (~5ml) removes any diiodotyrosine. It is an inhibitor of tyrosine hydroxylase with a K_i of ~500nM. [Harrington & Rivers *Biochem J* **38** 320 1944, DOI: 10.1042/bj0380320; Rivers *Chem Ind (London)* **21** 1956, *Beilstein* **14** III 1562, **14** IV 1562, **14** IV 2369.]

L-Isoleucine [73-32-5] $C_6H_{13}NO_2$, M 131.2, m 285-286°(dec), $[\alpha]_D^{20}$ +40.6 (6M HCl) pK_1^{25} 2.66, pK_2^{25} 9.69. Crystallise L-isoleucine from H_2O by addition of 4 volumes of EtOH or from aqueous MeOH. It sublimes at 170-181°/0.3mm with 99.7% recovery, unracemised [Gross & Gradsky *J Am Chem Soc* **77** 1678 1955, DOI: 10.1021/ja01611a085]. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 1 p 183-191, Vol 3 pp 2043-2073 1961, Huffman & Ingersoll *J Am Chem Soc* **73** 3366 1951, DOI: 10.1021/ja01151a111; Beilstein **4** IV 2775.]

DL-Isoserine (±-3-amino-2-hydroxypropionic acid) [565-71-9, 632-12-2] $C_3H_7NO_3$, M 105.1, m 235°(dec), 237°(dec), 245°(dec), 250-252°(dec), pK_1^{25} 2.78 (acidic), pK_2^{25} 9.27 (basic). Recrystallise it from H_2O or 50% aqueous EtOH. It has an isoelectric pH of 6.02. [Rinderknecht & Niemann *J Am Chem Soc* **75** 6322 1953, DOI: 10.1021/ja01120a523; Gundermann & Holtmann *Chem Ber* **91** 160 1958, DOI: 10.1002/cber.19580910128; dissociation constants: Emerson et al. *J Biol Chem* **92** 451 1931, <http://www.jbc.org/content/92/2/449>.] The *hydrobromide* has m 128-130° (from aqueous HBr) [Schöberl & Braun *Justus Liebigs Ann Chem* **542** 288 1939, DOI: 10.1002/jlac.19395420120]. [Beilstein **4** H 503, **4** IV 3116.]

L-Isovaline (2-amino-2-methylbutyric acid) [595-40-4] $C_5H_{11}NO_2$, M 117.2, m ca 300° (sublimes in vac), $[\alpha]_D^{25}$ +113.1 (c 5, H_2O), $[\alpha]_D^{25}$ +10 (5M HCl), $pK_{Est(1)} \sim 2.4$, $pK_{Est(2)} \sim 9.7$. Crystallise it from aqueous Me_2CO , or better dissolve in H_2O and add excess Me_2CO . [Baker et al. *J Am Chem Soc* **74** 4701 1952, DOI: 10.1021/ja01138a502; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 p 2573-2577 1961.]

L(S)-Leucine [61-90-5] $C_6H_{13}NO_2$, M 131.2, m 293-295°(dec), $[\alpha]_D^{25}$ +15.6 (5M HCl), pK_1^{25} 2.33, pK_2^{25} 9.74. Likely impurities are isoleucine, valine, and methionine. Crystallise L-leucine from water by adding 4 volumes of EtOH. It sublimes at 180-188°/0.3mm with 99.1% recovery, and *unracemised* [Gross & Gradsky *J Am Chem Soc* **77** 1678 1955, DOI: 10.1021/ja01611a085]. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 p 2075-2094 1961, Kameda et al. *J Pharm Soc Jpn* **78** 763 1958, Beilstein **4** IV 2738.]

L-Lysine [56-87-1] M 146.2, [39665-12-8 monohydrate] $C_5H_{14}N_2O_2$, M 164.2, m >210°(dec), $[\alpha]_D^{20}$ +25 (c 2, 6M HCl), pK_1 2.18, pK_2 8.95, pK_3 10.53. Crystallise this basic amino acid from aqueous EtOH. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 pp 2097-2122 1961, Kearley & Ingersoll *J Am Chem Soc* **73** 5783 1951, DOI: 10.1021/ja01156a089; Beilstein **4** IV 2717.]

L-Lysine dihydrochloride [657-26-1] $C_5H_{14}N_2O_2 \cdot 2HCl$, M 219.1, m 193°, 199-201°, 203-204°, $[\alpha]_D^{25}$ +25.9 (5M HCl). Crystallise it from MeOH, in the presence of excess HCl, by adding diethyl ether. [Yoneya *J Biochem(Tokyo)* **38** 343 1951, Kearley & Ingersoll *J Am Chem Soc* **73** 5783 1951, DOI: 10.1021/ja01156a089; Beilstein **4** IV 2717.]

L-Lysine monohydrochloride [657-27-2] $C_5H_{14}N_2O_2 \cdot HCl$, M 182.7, m 256°(dec), $[\alpha]_D^{25}$ +20.5 (c 5, 5M HCl). Likely impurities are arginine, D-lysine, 2,6-diaminoheptanedioic acid and glutamic acid. Crystallise the mono-hydrochloride from water at pH 4-6 by adding 4 volumes of EtOH. At above 60% relative humidity it forms a *dihydrate*. [Birhbaum et al. *J Biol Chem* **194** 455 1952, <http://www.jbc.org/content/194/1/455>, PMID: 14927637; Kearley & Ingersoll *J Am Chem Soc* **73** 5783 1951, DOI: 10.1021/ja01156a089; Beilstein **4** IV 2717.]

α-Melanotropin [581-05-5] (a tridecapeptide, α-MSH, melanocyte stimulating hormone), $C_{77}H_{109}N_{21}O_{19}S$, M 1664.9, $[\alpha]_D^{25}$ -58.5 (c 0.4, 10% aqueous AcOH). Its solubility in H_2O is 1mg/ml. It is separated from the extract by ion-exchange on carboxymethyl cellulose, desalted, evaporated and lyophilised, then chromatographed on Sephadex G-25. [Lande et al. *Biochemical Preparations* **13** 45 1971.] α-MSH is a melanocyte-stimulating hormone [Varga et al. *J Mol Neurosci* **50** 558 2013, DOI: 10.1007/s12031-013-9998-3. PMID: 23504281].

β-Melanotropin [9034-42-8] (octadeca to docosa peptides), amorphous. An extract of β-melanotropin is

purified by ion-exchange on carboxymethyl cellulose, desalted, evaporated and lyophilised, then chromatographed on Sephadex G-25. [Lande et al. *Biochemical Preparations* **13** 45 1971.] Increased plasma β -melanotropin (β -MSH) leads to hyperpigmentation [Sawin et al. *Arch Intern Med* **125** 708 1970, DOI:10.1001/archinte.1970.00310040132018].

Melphalan (4-[bis-(2-chloroethyl)amino]-L-phenylalanine, L-sarcosine) [148-82-3] $\text{C}_{13}\text{H}_{18}\text{Cl}_2\text{N}_2\text{O}_2$, **M** 305.2, **m** 182-183° (dec), 183-185°, $[\alpha]_{\text{D}}^{25}$ +7.5 (c 1.33, 1.0 N HCl), $[\alpha]_{\text{D}}^{20}$ -28 (-31.5) (c 0.8, MeOH), $\text{pK}_{\text{Est}} \sim 6.4$. Purify melphalan by recrystallisation from MeOH (colourless needles), and its solubility is 5% in 95% EtOH containing one drop of 6N HCl. It is soluble in EtOH and propylene glycol but is almost insoluble in H_2O . The **R-form** crystallises from MeOH (needles) with **m** 181.5-182° and $[\alpha]_{\text{D}}^{21}$ -7.5 (c 1.26, 1.0 N HCl). The **RS-form** has **m** 180-181° (very small needles from MeOH). [Bergel & Stock *J Chem Soc* 2409 1954, DOI: 10.1039/JR9540002409; *Beilstein* **14** IV 1689.] This *nitrogen mustard* is antineoplastic and a **CARCINOGEN**.

dl-Methionine (**RS-2-amino-4-methylthiobutyric acid**) [59-51-8] $\text{C}_5\text{H}_{11}\text{NO}_2\text{S}$, **M** 149.2, **m** 281°(dec), pK_1^{25} 2.28, pK_2^{25} 9.21. Crystallise it from hot water [solubility in H_2O (g/100ml) is 1.8 (0°), 3.4 (25°), 10.5 (75°) and 17.6 (100°)] or EtOH. Also purify it by dissolving it in H_2O and passing through an Amberlite IR-120 resin (NH_4^+ form). The eluate is concentrated and then passed through Amberlite IR-4B resin, and this eluate is evaporated to dryness. The residue is washed with EtOH, then Me_2CO , dried and recrystallised from aqueous EtOH (colourless plates) [Baddiley & Jamieson *J Chem Soc* 4280 1954, DOI: 10.1039/JR9540004280]. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 2125 1961, *Beilstein* **4** IV 3190.]

L-Methionine [63-68-3] $\text{C}_5\text{H}_{11}\text{NO}_2\text{S}$, **M** 149.2, **m** 277-279°(dec), 283°(dec), $[\alpha]_{\text{D}}^{25}$ -8.1 (c 0.8, H_2O), $[\alpha]_{\text{D}}^{25}$ +21.2 (0.2M HCl), $[\alpha]_{\text{D}}^{25}$ +23.1 (c 1, M HCl), [note signs of rotation], pK_1^{25} 2.13, pK_2^{25} 9.73. Crystallise L-methionine from aqueous EtOH. Also purify it by dissolving ~0.5g of amino acid in ~10ml of hot H_2O , filtering, adjusting the pH to 5.8 with 5N HCl, and collecting the solid after adding ~20ml of EtOH. It is recrystallised by dissolving in H_2O and adding EtOH. It sublimes at 197-208°/0.3mm with 99.8% recovery and **unracemised** [Gross & Gradsky *J Am Chem Soc* 77 1678 1955, DOI: 10.1021/ja01611a085]. [Baddiley & Jamieson *J Chem soc* 1085 1955, DOI: 10.1039/JR9550001085; Milne & Peng *J Am Chem Soc* **79** 645 1957, DOI: 10.1021/ja01560a037; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2125-2152 1961, *Beilstein* **4** IV 3189.] Methionine is a kidney acidifier, protects kidneys, and is used as an antidote in acetaminophen poisoning.

dl-Methionine sulfoxide [454-41-1, 62697-73-8] $\text{C}_5\text{H}_{11}\text{NO}_3\text{S}$, **M** 165.2, **m** >240°(dec), 241-242°(dec). Likely impurities are dl-methionine sulfone and dl-methionine. Crystallise the sulfoxide by dissolving it in hot H_2O and adding excess EtOH. [Lepp & Dunn *Biochemical Preparations* **4** 80 1955, Micheel & Schmitz *Chem Ber* **72** 518 1939, DOI: 10.1002/cber.19390720310; *Beilstein* **4** III 1650, **4** IV 3192.] **dl-Methionine sulfone** [820-10-0] $\text{C}_5\text{H}_{11}\text{NO}_4\text{S}$, **M** 181.2, has **m** ~250°(dec), and is excreted in the urine of rats when the *sulfoxide* is injected intraperitoneally [Wingo et al. *Arch Biochem Biophys* **47** 307 1953, DOI: 10.1016/0003-9861(53)90468-8 [Beilstein **4** IV 3193.]

S-Methyl-L-cysteine [1187-84-4] $\text{C}_4\text{H}_9\text{NO}_2\text{S}$, **M** 135.2, **m** 207-211°, ~240°(dec), 267-270°, $[\alpha]_{\text{D}}^{26}$ -32.0 (-34.5) (c 5, H_2O), pK_1^{25} 1.94 (COOH), pK_2^{25} 8.73 (NH_2 , 8.97). Likely impurities are cysteine and S-methyl-dl-cysteine. Crystallise it from H_2O by adding 4 volumes of EtOH. It also crystallises from MeOH with **m** 234-236°(dec), but after sublimation it has **m** 267-270° and $[\alpha]_{\text{D}}^{27}$ -31.6 (c 1, H_2O). [Rinderknecht et al. *Helv Chim Acta* **41** 1 1958, DOI: 10.1002/hlca.660410102; Theodoropoulos *Acta Chem Scand* **13** 383 1959, DOI: 10.3891/acta.chem.scand.13-0383; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 1904 1961, *Beilstein* **4** IV 3145.]

α -Methylmethionine [562-48-1] $\text{C}_6\text{H}_{13}\text{N}_2\text{OS}$, **M** 163.0, **m** 283-284°, $\text{pK}_{\text{Est}(1)}$ ~ 2.1, pK^{30} 9.45. Crystallise α -methylmethionine from aqueous EtOH or H_2O . Pfister et al. stated '**m** 283-284°' when crystallised from aqueous EtOH which gave **α -methylmethioninemethylsulfonium iodide** with MeI, and upon recrystallisation from 50%aqueous EtOH/absolute EtOH had **m** 95°(dec). Potts stated 'white needles' from H_2O with **m** 134°. [Pfister et al. *J Am Chem Soc* **77** 697 1955, DOI: 10.1021/ja01608a045; Potts *J Chem Soc* (in Notes starting on p 1626) 1632 1955, DOI: 10.1039/JR9550001626; Greenstein & Winitz *The Chemistry of the Amino Acids* J.

Wiley, Vol 3 p 2566 1961.]

S-Methyl-L-methionine chloride See Vitamin U in 'Physiologically Active...', this chapter.

N-Methyltryptophan (L-abrine) [526-31-8] $C_{12}H_{14}N_2O_2$, M 218.3, m 295°(dec with darkening and sintering), $[\alpha]_D^{21} +44.4$ (c 2.8, 0.5M HCl), $[\alpha]_D^{20} +65$ (c 1, 0.5N NaOH), pI 10.10, $pK_{Est(1)} \sim 2.3$, $pK_{Est(2)} \sim 9.7$. Crystallise L-abrine from H₂O or EtOH/H₂O mixture. Its solubility in MeOH is 1w/v% at ~25°. Dry it for 2 days at 60° in high vacuum; it has m 275-290°(dec with browning at 230°) and $[\alpha]_D^{21} +47.2$ (c 2, 0.5N HCl) [Peter et al. *Helv Chim Acta* **46** 577 1963, DOI: 10.1002/hlca.19630460217]. The **acetyl derivative**, $C_{14}H_{14}N_2O_3$, has m 176° and $[\alpha]_D^{25} -148$ (c 0.86, N NaOH). [Gregory & Morley *J Chem Soc C* 910 1968, DOI: 10.1039/J39680000910; for configuration and racemisation see Cahill & Jackson *J Biol Chem* **126** 29 1938, <http://www.jbc.org/content/126/1/29>; *Beilstein* **22/14** V 40.]

dl-5-Methyltryptophan [951-55-3] $C_{12}H_{14}N_2O_2$, M 218.3, m 275°(dec), 284-288° [pK see tryptophan]. Crystallise dl-5-methyltryptophan from aqueous EtOH after dissolving it in aqueous NaOH, precipitating with AcOH, filtering the solid off and drying for 24 hours at 50°. The purity can be determined by titrating an NaOH solution with perchloric acid [Jackman & Archer *J Am Chem Soc* **68** 2105 1946, DOI: 10.1021/ja01214a507; *Beilstein* **22** IV 6815.] The **picrate** crystallises from MeOH with m 202°(dec). The **N-phenylcarbamoyl derivative** crystallises from aqueous MeOH with m 202°. [Gordon & Jackson *J Biol Chem* **110** 151 1935, <http://www.jbc.org/content/110/1/151>.]

Nisin [1414-45-5] $C_{143}H_{230}N_{42}O_{37}S_7$, M 3354.2. This polypeptide from *S. lactis* is purified by crystallisation from 80% (v/v) EtOH and by countercurrent distribution. The synthetic polypeptide antibiotic can also be purified by preparative HPLC and assayed by HPLC on a Nucleosil 3007C₁₈ (6 x 250mm) column using a MeCN—0.01M HCl gradient (30-50%), at 2%/minute, and flow rate of 1.5ml/minute to give a retention time of 8.1 minutes; or MeCN—0.3M guanidine-HCl gradient (30-50%), at 2%/minute, and flow rate of 1.5ml/minute to give a retention time of 10.9 minutes. FAB-MS gave the *pseudomolecular ion* m/z at 3352.7 (M + H)⁺. It is soluble in dilute acid and is stable even on boiling. [Berridge et al. *Biochem J* **52** 529 1952, DOI: 10.1042/bj0520529; synthesis by Fukase et al. *Tetrahedron Lett* **29** 795 1988, DOI: 10.1016/S0040-4039(00)80212-9.] This polypeptide has 34 amino acids, 8 of which are unusual, having 5 sulfide bridges but no Trp residues [Gross & Morell *J Am Chem Soc* **93** 4634 1971, DOI: 10.1021/ja00747a073]. Nisin is used as a food preservative.

Norleucine (α-amino-n-caproic acid) [R(+) 327-56-0, S(-) 327-57-1] $C_6H_{13}NO_2$, M 131.2, m 301°(some dec), $[\alpha]_{546}^{20}$ (+) and (-) 28 (c 5, 5M HCl), (+) and (-) 6.26 (c 0.7, H₂O), [RS: 616-06-8] m 297-300° (sublimes partially at ~280°), pK_1^{25} 2.39 and pK_2^{25} 9.76 (for RS). Crystallise norleucine from water or aqueous MeOH. *Note* that the crystals (shiny leaflets from H₂O) of the (+)-form are sweet, whereas those of the (-)-form are bitter. The (RS)-racemate also crystallises as shiny leaflets from H₂O [solubility in (g/100ml) is 1.1 (25°), 2.9 (75°) and 5.2 (100°)], and is slightly soluble in EtOH (0.42 w/w% at 25°). [Huffman & Ingersoll *J Am Chem Soc* **73** 3366 1951, DOI: 10.1021/ja01151a111; *Beilstein* **4** III 1386, **4** IV 2628.]

Norvaline (α-amino-n-valeric acid, 2-aminopentanoic acid) [unnatural D (R-) 2031-12-9, natural L (S+) 6600-40-4] $C_5H_{11}NO_2$, M 117.2, m 305°(dec), $[\alpha]_{546}^{20}$ (+) and (-) 25 (c 10, 5M HCl), pI 6.04, pK_1^{25} 2.36, pK_2^{25} 9.87 (9.72). Crystallise R- or S+ norvaline from aqueous EtOH or water. Crystallise the RS-racemate from hot H₂O [solubility at 18° is 1g/10ml] in which it is freely soluble, but is insoluble in most organic solvents. It sublimes unchanged. [Finkbeiner & Stiles *J Am Chem Soc* **85** 616 1963, DOI: 10.1021/ja00888a031; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 pp 2390-2399 1961, *Beilstein* **4** III 1331-1333, **4** IV 128, 2629.]

L-Ornithine (2,5-diaminopentanoic acid) [70-26-8] $C_5H_{12}N_2O_2$, M 132.2, m 140°, $[\alpha]_D^{25} +16$ (c 0.5, H₂O), pK_1^{20} 2.11, pK_2^{20} 8.39, pK_3^{25} 10.59. Crystallise L-ornithine from water containing 1mM EDTA (to remove metal ions). An aqueous solution is alkaline since ornithine has an alkylamino chain which is strongly

basic. It is used in the treatment of hyperammonemia. [Perrin *J Chem Soc* 3125 1958, DOI: 10.1039/JR9580003125; Rivard *Biochemical Preparations* **3** 97 1955, Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2477-2491 1961, *Beilstein* **4** III 1346, **4** IV 2644.] The *racemate* [616-07-9] can be purified by dissolving in H₂O, and adding EtOH; its *monohydrochloride* [1069-31-4] has **m** ~233°(dec) (from MeOH/Et₂O).

L-Ornithine monohydrochloride [3184-13-2] C₅H₁₂N₂O₂·HCl, **M 168.6**, **m 230-232°(dec)**, **233°(dec)**, **236.5-237.5°(dec)**, [α]_D²⁵ +28.3 (**5M HCl**). Likely impurities are citrulline, arginine and D-ornithine. Crystallise the monohydrochloride from water by adding 4 volumes of EtOH and dry it in a vacuum desiccator over fused CaCl₂. [Rivard *Biochemical Preparations* **3** 98 1955.] The *dihydrochloride* [6211-16-1] has **m 202-203°** and [α]_D²⁰ +18.4 (c 2.3, 6N HCl) after recrystallisation from MeOH/Et₂O [Zaoral & Rudinger *Coll Czech Chem Commun* **24** 1993 1959, DOI: org/10.1135/ccccc19591993]. [*Beilstein* **4** IV 2644.]

Oxytocin [50-56-6] C₄₃H₆₆N₁₂O₁₂S₂, **M 1007.2**, **m dec on heating**, [α]_D²² -26.2° (c 0.53, N AcOH). It is a cyclic nonapeptide which is purified by countercurrent distribution between solvent and buffer. It is soluble in H₂O, *n*-BuOH and isoBuOH. [Bodanszky & du Vigneaud *J Am Chem Soc* **81** 2504 1959, DOI: 10.1021/ja01519a053; Sakakibara et al. *Bull Chem Soc Jpn* **38** 120 1965, DOI: 10.1246/bcsj.38.120; for solid phase synthesis see Bayer & Hagenmyer *Tetrahedron Lett* **9** 2037 1968, DOI: 10.1016/S0040-4039(00)89739-7.] It was also synthesised on a solid phase matrix and finally purified as follows: A Sephadex G-25 column is equilibrated with the aqueous phase of a mixture of 3.5% AcOH (containing 1.5% of pyridine)/*n*-BuOH/*C₆H₆ (2:1:1) and then the organic phase of this mixture is run through. A solution of oxytocin (100mg) in H₂O (2ml) is applied to the column which is then eluted with the organic layer of the above mixture. The fractions containing the major peak [as determined by the Folin-Lowry protein assay: Fryer et al. *Anal Biochem* **153** 262 1986, DOI: 10.1016/0003-2697(86)90090-4] are pooled, diluted with twice their volume of H₂O, evaporated to a small volume and lyophilised to give oxytocin as a pure white powder (20mg, 508 U/mg). [Ives *Can J Chem* **46** 2318 1968, DOI: 10.1139/v68-378; *Beilstein* **22** III/IV 82.] It is a pituitary hormone which is the main stimulant for uterine contraction in pregnancy and for lactation. This hormone elicits affectionate feelings.

dl-Phenylalanine [150-30-1] C₉H₁₁NO₂, **M 165.2**, **m 265-266°(capillary, dec)**, **271-273°(dec)**, **282-284°(dec)**, **pK₁²⁵ 2.58**, **pK₂²⁵ 9.24**. *dl*-Phenylalanine crystallises from H₂O or H₂O/EtOH in large plates and is dried under vacuum over P₂O₅. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2156-2175 1961, *Beilstein* **14** III 1229, **14** IV 1553.]

L(S)-Phenylalanine [63-91-2] C₉H₁₁NO₂, **M 165.2**, **m 280°(dec)**, **281-183°(dec)**, [α]_D²⁵ -34.0 (c 2, H₂O). Likely impurities are leucine, valine, methionine and tyrosine. Crystallise L-phenylalanine from water by adding 4 volumes of EtOH. Dry it *in vacuo* over P₂O₅. Also crystallise it from saturated refluxing aqueous solutions at neutral pH, or 1:1 (v/v) EtOH/water solution, or concentrated HCl. It sublimes at 176-184°/0.3mm with 98.7% recovery and *unracemised* [Gross & Grodsky *J Am Chem Soc* **77** 1678 1955, DOI: 10.1021/ja01611a085]. *S*-Phenylalanine *ethyl ester hydrochloride* [3182-93-2] has **m 156-158°** and [α]_D²⁰ -7.8 (c 2, H₂O) after crystallisation from EtOH/Et₂O [Billimoria & Cook *J Chem Soc* 2323 1949, DOI: 10.1039/JR9490002323; *Beilstein* **14** IV 1556]. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2156-2175 1961, *Beilstein* **14** IV 1552.]

L(S)-α-Phenylglycine [2935-35-5] C₈H₉NO₂, **M 151.2**, **m 305-310°, 305-308°(capillary, dec)**, [α]_D²⁰ +157 (c 1, N HCl), [α]_D²⁵ +188 (c 1, M HCl), **pK₁²⁵ 1.83**, **pK₂²⁵ 4.39** (for *dl*). Crystallise it from EtOH. [Kaneko *J Chem Soc Jpn* **60** 538 1939, Rudman et al. *J Am Chem Soc* **74** 551 1952, DOI: 10.1021/ja01122a083; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2694-2697 1961, *Beilstein* **14** III 1187, **14** IV 1317; Fieser **17** 278.] The **D(R)-enantiomer** [875-74-1] has similar properties but a negative optical rotation. The *racemate* [69-91-0] sublimes at ~255° without melting. [Steiger *Org Synth Coll Vol* **3** 84 1955, DOI: 10.15227/orgsyn.022.0023.]

Phenylglycine-*o*-carboxylic acid [*N*-(2-carboxyphenyl)glycine] [612-42-0, 64241-57-2 *Me Ester*, 67990-19-6 *mono-Na Salt*, 71807-57-3 *di-Na Salt*] C₉H₉NO₄, **M 195.2**, **m 206°, 208°, 218°(dec)**, **220°, pK₁²⁰ 5.44**

(CO₂H), pK₂²⁰ 6.96 (CO₂H) (in 50% aqueous dioxane). It is prepared by boiling under reflux a solution containing anthranlic acid (14g), chloroacetic acid (10g) and anhydrous Na₂CO₃ (20g) in H₂O (200ml) for 3 hours, then adjusted to pH~2 with concentrated HCl, and set aside overnight. The acid which precipitates is filtered off, washed well with cold H₂O, dried in air, recrystallised from hot H₂O (charcoal) and dried at 100° (yield 12g, m 208°). Crystallise the acid further from hot water (charcoal) if necessary. It forms complexes with Cu²⁺, Zn²⁺, Cd²⁺, Co²⁺ and Ni²⁺ in aqueous dioxane. [Vogel's Textbook of Practical Organic Chemistry, Third edition, 1961, p. 980; Roileanu et al. *Rev Roumaine Chim* **12** 105 1967, Krause et al. US Pat No 5,821,385A Oct 13 1998, www.google.com/patents/US5821385; *Aldrich library of ¹³C, ¹H FTNMR Spectra*, NMR **2** 1181A, *Beilstein* **14** H 348, **14** I 544, **14** II 225, **14** III 938.]

D-Pipecolinic acid (R-piperidine-2-carboxylic acid) [1723-00-8] C₆H₁₁NO₂, M 129.2, m 264°(dec), 267°(dec), ~280°(dec), [α]_D¹⁹ +26.2 (c 2, H₂O), [α]_D²⁵ +35.7 (H₂O), pK₁²⁰ 2.29 (CO₂H), pK₂²⁰ 10.77 (NH⁺). D-Pipecolinic acid recrystallises as platelets from EtOH and is soluble in H₂O. The *hydrochloride* has m 256-257°(dec) from H₂O and [α]_D²⁵ +10.8 (c 2, H₂O). [Lukés et al. *Coll Czech Chem Commun* **22** 286 1957, DOI: [org/10.1135/cccc19570286](https://doi.org/10.1135/cccc19570286); Bayerman *Recl Trav Chim Pays-Bas* **78** 134 1959, DOI: [10.1002/recl.19590780209](https://doi.org/10.1002/recl.19590780209); Asher et al. *Tetrahedron Lett* **22** 141 1981, DOI: [10.1016/0040-4039\(81\)80170-0](https://doi.org/10.1016/0040-4039(81)80170-0); *Beilstein* **22/1** V 220.]

L-Pipecolinic acid (S-piperidine-2-carboxylic acid, L-homoproline) [3105-95-1, 535-75-1 (RS)] C₆H₁₁NO₂, M 129.2, m 268°(dec), 271°(dec), ~280°(dec), [α]_D²⁰ -26 (c 4, H₂O), [α]_D²⁵ -34.9 (H₂O). Recrystallise L-pipecolinic acid from aqueous EtOH, and it sublimes as needles in a vacuum. It is sparingly soluble in absolute EtOH, Me₂CO or CHCl₃ but insoluble in Et₂O. The *hydrochloride* has m 258-259°(dec, cryst from MeOH) and [α]_D²⁵ -10.8 (c 10, H₂O). [Synthesis from L-lysine: Fuji & Myoshi *Bull Chem Soc Jpn* **48** 1341 1975, DOI: [10.1246/bcsj.48.1341](https://doi.org/10.1246/bcsj.48.1341); synthesis of racemate: Shuman *J Org Chem* **55** 738 1990, DOI: [10.1021/jo00289a058](https://doi.org/10.1021/jo00289a058); *Beilstein* **22/1** V 220.] This is a metabolite of lysine and is observed in pipecolic acidemia, a cerebro-hepato-renal syndrome in the onset of neonatal adrenoleukodystrophy which is an infantile Refsum disease (build-up of phytanic acid causing ataxia, scaly skin, difficulty in hearing and eye problems).

Piperidine-4-carboxylic acid (isonipecotic acid) [498-94-2] C₆H₁₁NO₂, M 129.2, m 336°(dec, darkens at ~300°), pK_{Est(1)}~ 4.3 (CO₂H), pK_{Est(2)}~ 10.6 (NH⁺). It crystallises from H₂O or EtOH as needles. The *hydrochloride* crystallises from H₂O or aqueous HCl with m 293°dec (also 298°dec, 300°dec). The *amide* [39546-32-2] C₆H₁₁N₂O, M 128.2, has m 145-148°. [Wibaut *Recl Trav Chim Pays-Bas* **63** 141 1944, DOI: [10.1002/recl.19440630704](https://doi.org/10.1002/recl.19440630704); IR: Zacharius et al. *J Am Chem Soc* **76** 2908 1954, DOI: [10.1021/ja01640a015](https://doi.org/10.1021/ja01640a015); *Beilstein* **22/1** V 244.]

Polypeptides. These are strings of α-amino acids usually with the *natural* S(L) [L-cysteine is an exception and has the *R* absolute configuration, because of the *sequence rule*] or sometimes '*unnatural*' R(D) configuration (much less common) at the α-carbon atom. They generally have less than ~100 amino acid residues. They can be naturally occurring or, because of their small size, can be synthesised chemically from the desired amino acids. Their properties can be very similar to those of small proteins. Many are commercially available, and can be custom made commercially or locally with a peptide synthesiser. They are purified by HPLC and can be used without further purification. Their purity can be checked as described under proteins (Introduction).

L-Proline [147-85-3] C₅H₉NO₂, M 115.1, m 215-220°(dec) (D-isomer), 220-222°(dec) (L-form), 205°(dec)(DL-isomer), [α]_D²⁰ -53 (c 0.6, 0.5N HCl), -93 (c 2.4, 6N KOH) for L-isomer, pI 6.3, pK₁²⁵ 1.95, pK₂²⁵ 10.64. A likely impurity is hydroxyproline. Purify L-proline *via* its *picrate* which is crystallised twice from water, then decomposed with 40% H₂SO₄. The picric acid is extracted with diethyl ether, the H₂SO₄ in solution is precipitated with Ba(OH)₂, and the filtrate is evaporated. The residue is then recrystallised from hot absolute EtOH [Mellon & Hoover *J Am Chem Soc* **73** 3879 1951, DOI: [10.1021/ja01152a095](https://doi.org/10.1021/ja01152a095)] or EtOH/Et₂O. Its solubility in H₂O is >100%. It sublimes at 182-187°/0.3mm with 99.4% recovery and *unracemised* [Gross & Gradsky *J Am Chem Soc* **77** 1678 1955, DOI: [10.1021/ja01611a085](https://doi.org/10.1021/ja01611a085)]. It is *hygroscopic* and is stored in a desiccator. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol **3** pp 2178-2199 1961, *Beilstein* **22** III/IV 8, **22/1** V 31.]

L-Prolylglycine [2578-57-6] $C_7H_{12}N_2O_3$, **M 172.2**, **m 236°**, $[\alpha]_D^{20} +21.1$ (c 4, H_2O), **pK₁²⁵ 3.19**, **pK₂²⁵ 8.97**. Recrystallise L-prolylglycine from water at 50-60° by addition of EtOH. [Appel et al. *Chem Ber* **108** 2680 1975, DOI: 10.1002/cber.19751080825; Rydon & Smith *J Chem Soc* 3642 1956, DOI: 10.1039/JR9560003642.]

L-Propargylglycine (S-2-aminopent-4-ynoic acid) [23235-01-0] $C_5H_7NO_2$, **M 113.1**, **m 230°**(dec starting at 210°), $[\alpha]_D^{20} -35$ (c 1, H_2O), -4° (c 5, 5N HCl), **pK_{Est(1)}~ 2.3** (CO_2H), **pK_{Est(2)}~ 9.8** (NH_2). The acid crystallises readily when ~4g in 50ml H_2O are treated with absolute EtOH at 4°/3 hours, and is collected, washed with cold absolute EtOH and Et_2O and dried in a vacuum. Also, it recrystallises from aqueous Me_2CO , R_F on SiO_2 TLC plates with n -BuOH/ H_2O /AcOH (4:1:1) is 0.26. The **racemate** has **m 238-240°**. [Leukart et al. *Helv Chim Acta* **59** 2181 1976, DOI: 10.1002/hlca.19760590629; Eberle & Zeller *Helv Chim Acta* **68** 1880 1985, DOI: 10.1002/hlca.19850680711; Jansen et al. *Recl Trav Chim Pays-Bas* **88** 819 1969, DOI: 10.1002/recl.19690880707.] It is a suicide inhibitor of γ -cystathionase and other enzymes [Washtien & Abeles *Biochemistry* **16** 2485 1977, DOI: 10.1021/bi00630a026; Shinozjika et al. *Eur J Biochem* **124** 377 1982, DOI: 10.1111/j.1432-1033.1982.tb06603.x].

R-Pyroglutamic acid (5-oxo-D-proline, R-2-pyrrolidone-5-carboxylic acid) [4042-36-8] $C_5H_7NO_3$, **M 129.1**, **m 156-158°**, $[\alpha]_D^{20} +11.2$ (c 1, H_2O). Purify R-pyroglutamic acid by dissolving it in H_2O , filtering, passing the filtrate through Dowex 50 (H^+ form), washing with H_2O , pooling washings, evaporating, removing H_2O azeotropically with Me_2CO and $*C_6H_6$, washing the residue with Et_2O and recrystallising from EtOH/petroleum ether. [Pradeles et al. *Coll Czech Chem Commun* **42** 79 1977, DOI: org/10.1135/cccc19770079; Beilstein **22/6** V 7.]

S-Pyroglutamic acid (5-oxo-L-proline) [98-79-3] $C_5H_7NO_3$, **M 129.1**, **m 156-158°**, **162-164°**, $[\alpha]_{546}^{20} -11$ (c 5, H_2O), **pK²⁵ 12.7** (by electron spin resonance). Crystallise S-pyroglutamic acid by dissolving it in boiling EtOH (20g in 100ml), cooling and after a few minutes adding petroleum ether (b 40-60°, 120ml), then after 5 minutes adding a further 120ml, and cooling to room temperature with 90% recovery. This has **m 155.5-157.5°** and $[\alpha]_D^{20} -11.4$ (c 4.4, H_2O) [Hardy *Synthesis* 290 1978, DOI: 10.1055/s-1978-24726; Pellegata et al. *Synthesis* 614 1978, DOI: 10.1055/s-1978-24834]. The NH_4 salt has **m 184-186°** (from EtOH). [Beilstein **22/6** V 7.] The **racemate** [149-87-1], purified in the same way, has **m 183-185°**.

Quisqualic acid (Quis, 3-[3,5-dioxo-1,2,4-oxadiazolin-2-yl]-L-alanine) [52809-07-1] $C_5H_7N_3O_5$, **M 189.1**, **m 190-191°**, $[\alpha]_D^{20} +17$ (c 2, 6M HCl), **pK_{Est(1)}~ 2.1** (CO_2H), **pK_{Est(2)}~ 8.9** (NH_2). It has been purified by ion-exchange chromatography on Dowex 50W (x 8, H^+ form); the desired fractions are lyophilised and recrystallised from H_2O /EtOH. It has IR (KBr) with ν_{max} at 3400—2750br, 1830s, 1775s, 1745s and 1605s cm^{-1} ; and 1H NMR ($NaOD/D_2O$, pH 13) δ : 3.55-3.57 (1H m, X of ABX, H-2), 3.72-3.85 (2H, AB of ABX, H-3), ^{13}C NMR (D_2O) δ : 50.1(t), 53.4(d), 154.8(s), 159.7(s) and 171.3(s). [Baldwin et al. *JCS Chem Commun* 256 1985, DOI: 10.1039/C39850000256.] It is an excitatory amino acid agonist for the quisqualate receptor [Joels et al. *Proc Natl Acad Sci USA* **86** 3404 1989, PubMed ID2470102], acts on AMPA receptors and group I metabotropic glutamate receptors [Porter et al. *Br J Pharmacol* **106** 509 1992, PMCID: PMC1907545]; and sensitises neurons in the hippocampus (CA-1 pyramidal neurons) to depolarisation by L-AP6 (L-2-amino-6-phosphonohexanoic acid), the so called ‘quis’ effect [Schulte et al. *Brain Res* **649** 203 1994, DOI:10.1016/0006-8993(94)91065-0; Littman et al. *Neuropharmacology* **34** 829 1995, DOI:10.1016/0028-3908(95)00070-M].

Sarcosine (N-methylglycine) [107-97-1] $C_3H_7NO_2$, **M 89.1**, **m 2 12-213°**(dec), **pK₁²⁰ 2.12**, **pK₂²⁰ 10.19**. Crystallise sarcosine from absolute EtOH, 95% EtOH or H_2O . It sublimes at 180-185°/0.3mm with 99.1% recovery [Gross & Gradsky *J Am Chem Soc* **77** 1678 1955, DOI: 10.1021/ja01611a085]. [Cocker & Harris *J Chem Soc* 1290 1940, DOI: 10.1039/JR9400001290; Cocker & Lapworth *J Chem Soc* 1897 1931, DOI: 10.1039/JR9310001894; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** p 2750 1961, Beilstein **4** III 1121, **4** IV 2363.]

Sarcosine anhydride (1,4-dimethylpiperazin-2,5-dione) [5076-82-4] $C_6H_7N_2O_2$, **M 142.2**, **m 146-147°**,

148°, $pK_{\text{Est}(1)} \sim -4.2$, $pK_{\text{Est}(2)} \sim 1.9$. Crystallise the anhydride from H_2O , EtOH or EtOAc. Dry it in a vacuum at room temperature. [Karrer et al. *Helv Chim Acta* **5** 140 1922, DOI: 10.1002/hlca.19220050115; *Beilstein* **24** II 144, **24** IV 1072.]

Seleno-DL-methionine (± 2 -amino-4-methylselenanylbutyric acid) [1464-42-2, 2578-28-1 (\pm)] $\text{C}_5\text{H}_{11}\text{NO}_2\text{Se}$, **M 196.1**, **m 265°(dec)**, **267-269°(dec)**, **270°** (see **pKs of methionine**). It crystallises in hexagonal plates from MeOH and H_2O . [Klosterman & Painter *J Am Chem Soc* **69** 2009 1949, DOI: 10.1021/ja01200a054.] The **L-isomer** [3211-76-5] is purified by dissolving it in H_2O , adjusting the pH to 5.5 with aqueous NH_3 , evaporating to near-dryness, and the residue is washed several times with absolute EtOH till a solid is formed and then recrystallise from Me_2CO . It has **m 266-268°(dec)** [also **275°(dec)**], and $[\alpha]_{\text{D}}^{25} +18.1$ (c 1, N HCl), $[\alpha]_{\text{D}}^{22} +21.6$ (c 0.5, 2N HCl). [Pande et al. *J Org Chem* **35** 1440 1970, DOI: 10.1021/jo00830a040; *Beilstein* **4** IV 3216.]

L(S)-Serine [56-45-1] $\text{C}_3\text{H}_7\text{NO}_3$, **M 105.1**, **m 228°(dec)**, **233-235°(dec)**, $[\alpha]_{\text{D}}^{25} +14.5$ (1M HCl), $[\alpha]_{\text{D}}^{20} +16$ (c 5, 5M HCl), $pK_1^{25} 2.15$, $pK_2^{25} 9.21$. It is biologically formed from glycine and formaldehyde by the enzyme serinehydroxymethyl transferase. A likely impurity is glycine. Crystallise L-serine from H_2O by adding 4 volumes of EtOH. Dry and store it in a desiccator. It sublimes at 160-170°/0.3mm with 99.7% recovery, and **unracemised** [Gross & Gradsky *J Am Chem Soc* **77** 1678 1955, DOI: 10.1021/ja01611a085]. [Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 3** pp 2202-2235 1961, *Beilstein* **4** IV 3118; Fieser **12** 430, **14** 282.] **D(R)-Serine** [312-84-5] is physiologically active as a mimic at the strychnine-intensive glycine binding site associated with the NMDA (*N-methyl-D-aspartate*) receptor site and the inhibitory post-synaptic glycine receptor.

Somatostatin [38916-34-6] $\text{C}_{76}\text{H}_{104}\text{N}_{18}\text{O}_{19}\text{S}_2$, **M 1637.9**, $[\alpha]_{\text{D}}^{25} -36$ (c 0.57, 1% AcOH). Somatostatin is a tetradecapeptide which is purified by gel filtration on Sephadex G-25, eluting with 2N AcOH, and then by liquid partition chromatography on Sephadex G-25 using *n*-BuOH/AcOH/ H_2O (4:1:5) and has $R_F = 0.4$. It is a brain growth hormone releasing-inhibiting factor which has also been synthesised. [Burgus et al. *Proc Natl Acad Sci USA* **70** 684 1973, DOI: 10.1073/pnas.70.3.684; Sarantakis & McKinley *Biochem Biophys Res Commun* **54** 234 1973, DOI: 10.1016/0006-291X(73)90913-3; Hartrodt et al. *Pharmazie* **37** 403 1982, PMID: 6126893.]

L-Threonine (2*S*,3*R*-2-amino-3-hydroxybutyric acid) [72-19-5] $\text{C}_4\text{H}_9\text{NO}_3$, **M 119.1**, **m 251-253°**, **254°(dec)**, **262-263°(dec)**, $[\alpha]_{\text{D}}^{26} -28.4$ (H_2O), $pK_1^{25} 2.17$, $pK_2^{25} 9.00$. Likely impurities are *allo*-threonine and glycine. Crystallise L-threonine from H_2O by adding 4 volumes of EtOH. Dry and store it in a desiccator. It also crystallises from 80% EtOH to give hexagonal plates **m 262-263°(dec)**. It sublimes at 200-226°/0.3mm with 99.6% recovery and **unracemised** [Gross & Gradsky *J Am Chem Soc* **77** 1678 1955, DOI: 10.1021/ja01611a085]. [Elliot *J Chem Soc* 62 1950, Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, **Vol 1** pp 176-183, **Vol 3** pp 2238-2257 1961, *Beilstein* **4** IV 3171.] **D-Threonine** (2*R*,3*S*-2-amino-3-hydroxybutyric acid) [632-20-2] has the same properties but with opposite rotation, and the **racemate** [80-68-2] has a lower melting point, **m 244°(dec)**; whereas **L-*allo*-threonine** (2*S*,3*S*-2-amino-3-hydroxybutyric acid) [28954-12-3] has **m 277°(dec)** and is optically active with $[\alpha]_{\text{D}}^{22} +9$ (c 2, H_2O). [*Beilstein* **4** IV 3170.]

L-Thyroxine sodium salt (5*H*₂O) [6106-07-6] $\text{C}_{15}\text{H}_{10}\text{I}_4\text{NO}_4 \text{Na} \cdot 5\text{H}_2\text{O}$, **M 888.9**, $[\alpha]_{\text{D}}^{20} +20$ (c 2, 1M HCl + EtOH, 1:4). Crystallise the sodium salt from absolute EtOH and dry it for 8 hours at 30°/1mm. [Joel & Canepa *Acta Cryst* **4** 283 1951, DOI: 10.1107/S0365110X51000921; *Beilstein* **14** II 378, **14** III 1566, **14** IV 2374.]

D-Thyroxine {*O*-[3,5-diiodo-4-oxyphenyl]-3,5-diiodo-D-(-)-tyrosine, 3,3',5,5'-tetraiodo-D-thyronine} [51-49-0] $\text{C}_{15}\text{H}_{11}\text{I}_4\text{NO}_4$, **M 776.9**, **m 235°(dec)**, **235-236°(dec)**, **340°(dec)**, $[\alpha]_{\text{D}}^{20} +4.5$ (c 3, aqueous 0.2N NaOH in 70% EtOH), $[\alpha]_{\text{D}}^{20} -17$ (c 2, aqueous N HCl + EtOH 1:4), $pK_1^{25} 2.2$ (CO_2H), $pK_2^{25} 8.40$ (OH), $pK_3^{25} 10.1$ (NH_2). Recrystallise D-thyroxine from H_2O (needles) or from an ammonical solution by dilution with H_2O , MeOH or Me_2CO . It has also been purified by dissolving ~6.5 g in a mixture of MeOH (200ml) and 2N HCl (20ml), adding charcoal, filtering then adding NaOAc solution to pH 6. On standing the thyroxine separates, it is filtered off, washed with MeOH then Me_2CO and dried *in vacuo*. **N-Formyl-D-thyroxine** has

m 210° and $[\alpha]_{546}^{21}$ -26.9 (c 5, EtOH). (\pm)-*Thyroxine* [300-30-1] has **m 256° (231-233°** also reported) and is purified in the same way. [Siedel & Siedel *Chem Ber* **96** 1 1963, DOI: 10.1002/cber.19630960102; Harington & Salter *Biochem J* **24** 456 1930, DOI: 10.1042/bj0240456; *Beilstein* **14** I 671, **14** II 384, **14** III 1566, **14** IV 2374.]

L-Thyroxine (*O*-[3,5-diiodo-4-oxyphenyl]-3,5-diiodo-L-(+)-tyrosine, 3,3',5,5'-tetraiodo-L-thyronine, **T₄**) [51-48-9] **C₁₅H₁₁I₄NO₄**, **M 776.9**, **m 229-230°(dec)**, **~235°(dec)**, **237°(dec)**, $[\alpha]_{\text{D}}^{22}$ -5.1 (c 2, aqueous N NaOH + EtOH 1:2), $[\alpha]_{\text{D}}^{22}$ +15 (c 5, aqueous N HCl in 95% EtOH 1:2), $[\alpha]_{\text{D}}^{22}$ +26 (EtOH/1M aqueous HCl, 1:1) (**pK²⁵ 6.6**). Purification of this **natural L-enantiomer** is the same as for the D-isomer above. Likely impurities are tyrosine, iodotyrosine, iodothyroxines and iodide. Dissolve it in dilute ammonia at room temperature, then crystallise it by adding dilute acetic acid to pH 6. *N-Formyl-L-thyroxine* has **m 214°(dec)** and $[\alpha]_{546}^{21}$ +27.8 (c 5, EtOH). [Harington & Pitt Rivers *Biochem J* **39** 157 1945, DOI: 10.1042/bj0390157; Siedel & Siedel *Chem Ber* **96** 1 1963, DOI: 10.1002/cber.19630960102; Reineke & Turner *J Biol Chem* **161** 613 1945, <http://www.jbc.org/content/161/2/613>, PMID: 21006943; Chalmers et al. *J Chem Soc* 3424 1949, DOI: 10.1039/JR9490003424; *Beilstein* **14** II 378, **14** III 1566, **14** IV 2373.] This **tetra-iodo**-containing hormone **T₄**, and the related **tri-iodo**-containing hormone **T₃**, are produced from *thyroglobin* in the thyroid follicular cells. The regulation of growth and development, and the metabolic rate of these hormones seems to be due to their effects on DNA transcription, consequently on protein synthesis.

N-Tosyl-L-lysine chloromethyl ketone HCl (TLCK, 3S-1-chloro-3-tosylamino-7-amino-2-heptanone HCl) [4272-74-6] **C₁₄H₂₁ClN₂O₃S**. **HCl**, **M 369.3**, **m 150-153°(dec)**, **156-158°(dec)**, **~165°(dec)**, $[\alpha]_{\text{D}}^{20}$ -7.3 (c 2, **H₂O**), **pK_{Est} ~ 10.6 (7-NH₂)**. The hydrochloride slowly crystallises from a concentrated solution in absolute EtOH, thinned with EtOH/Et₂O for collection and dried *in vacuo*. TLCK is soluble in H₂O, and a 10mM stock solution is prepared in 1mM HCl, pH 3.0, or in a buffer at pH 6.0 (solutions are unstable above this pH at 25°, thus 48% will decompose in ~5 minutes at pH 9.0). Solutions should be prepared freshly, and effective concentrations are 10-100μM. Solutions of 5mg/100ml in EtOH, and 5mM in Me₂SO, can also be prepared. It is a **suicide enzyme inhibitor of serine proteases**, e.g. trypsin and clostripain. [Matsuda et al. *Chem Pharm Bull Jpn* **30** 2512 1982, DOI: [org/10.1248/cpb.30.2512](https://doi.org/10.1248/cpb.30.2512); Shaw et al. *Biochemistry* **4** 2219 1965, DOI: 10.1021/bi00886a039; improved prep: Shaw & Glover *Arch Biochem Biophys* **139** 298 1970, DOI: 10.1016/0003-9861(70)90481-9].

Triglycyl glycine (tetraglycine) [637-84-3] **C₈H₁₄N₄O₅**, **M 246.2**, **m 270-275°(dec)**, **pK₁²⁵ 3.21(CO₂H)**, **pK₂²⁵ 7.94 (NH₃⁺)**. Crystallise it from H₂O (optionally, by the addition of EtOH). [Li et al. *J Am Chem Soc* **79** 5859 1957, DOI: 10.1021/ja01579a006; Rising et al. *J Am Chem Soc* **56** 1178 1934, DOI: 10.1021/ja01320a057; *Beilstein* **4** II 807, **4** III 1201, **4** IV 2472.]

Trigonellamide chloride (1-methylnicotinamide chloride) [1005-24-9] **C₇H₉ClN₂O**, **M 172.6**, **m 240°(dec)**. It crystallises from MeOH, and is dried *in vacuo*. It is prepared from nicotinamide and MeI in refluxing MeOH then shaking with AgCl [Karrer et al. *Helv Chim Acta* **19** 826 1936, DOI: 10.1002/hlca.193601901112]. It is soluble in organic solvents but moderately in H₂O. It is a metabolite of nicotinic acid in man, and was isolated from urine [Huff & Perlzweig *J Biol Chem* **150** 395 1943, <http://www.jbc.org/content/150/2/395>]. With ketones in aqueous alkali, it produces a green-blue fluorescence which turns blue on acidification and intensifies on heating. [*Beilstein* **22** III/IV 468, **22/2** V 80.]

3,3',5-Triiodo-S-thyronine [6893-02-3] **C₁₅H₁₂I₃NO₄**, **M 651.0**, **m 234-238°**, **236-237°(dec)**, $[\alpha]_{\text{D}}^{29.5}$ +21.5 (+23.0) (EtOH/1M aqueous HCl, 2:1, EtOH or 0.1M HCl), **pK₁²⁵ 6.48**, **pK₂²⁵ 7.62**, **pK₃²⁵ 7.82**. Likely impurities are as in *thyroxine*. Purify it by dissolving in dilute NH₃ at ~20°, then crystallise it by addition of dilute acetic acid to pH 6. Alternatively, 35g are purified by dissolving it in a mixture of EtOH (250ml) and 2N NaOH (100ml), then hot 2N HCl is added to the boiling solution until the pH is 4-5. After cooling for a few hours, the solid is filtered off and dried in a vacuum [**m 233-235°(dec)**]. [Chambers et al. *J Chem Soc* 2424 1949, DOI: 10.1039/JR9490003424; *Beilstein* **14** III 1566, **14** IV 2373.]

N,N,N-Trimethyl glycinehydrazide chloride (Girard Reagent T, 2-hydrazino-N,N,N-trimethyl-2-oxoethanaminium chloride) [123-46-6] **C₅H₁₄ClN₃O**, **M 167.6**, **m 188-192°(dec)**, **192°**. It is prepared by

reacting ethyl chloroacetate with Me_3N to form $(\text{Me}_3\text{NCH}_2\text{CO}_2\text{Et})^+ \text{Cl}^-$ followed by reaction with hydrazine. It is purified by crystallisation from absolute EtOH (slight decomposition) until it has only a slight odour. Store it in well-stoppered containers because it is very **hygroscopic**. It is very soluble in H_2O , AcOH and glycerol but slightly soluble in EtOH (0.66%). It forms water-soluble hydrazones with carbonyl compounds, and is used to purify them by allowing impurities in the aqueous solution to be extracted out. The carbonyl compounds are then recovered from the hydrazones. By replacing Me_3N by pyridine, the pyridine analogue *Girard P* is obtained which works in the same way. [Beilstein 4 III 1133; Fieser 1 410.]

N-Tris-(hydroxymethyl)methylglycine (TRICINE) [5704-04-1] $\text{C}_6\text{H}_{13}\text{NO}_5$, M 179.2, m 186-188°(dec), $\text{pK}_1^{20} \sim 2.3$, $\text{pK}_2^{20} 8.15$. Crystallise Tricine from EtOH and water. It is a good buffer in the pH range 7.4–8.8. [Good et al. *Methods Enzymol* 24B 53 1968, McGlothlin & Jordan *Analyt Lett* 9 245 1976, DOI: 10.1080/00032717608059100; Beilstein 18 III/IV 3454.]

L(S)-Tryptophan [73-22-3] $\text{C}_{11}\text{H}_{12}\text{N}_2\text{O}_2$, M 204.3, m 278°, 281-282°, 290°, $[\alpha]_D^{20} -33.4$ (EtOH), $[\alpha]_{546}^{20} -36$ (c 1, H_2O), $[\alpha]_D^{28} +28$ (c 2.1, 10% HCl), $\text{pK}_1^{25} -6.23$ (aqueous H_2SO_4), $\text{pK}_2^{25} 2.46$, $\text{pK}_3^{25} 9.41$, $\text{pK}_4^{25} 14.82$ (acidic NH, in aqueous NaOH). Crystallise L-tryptophan from H_2O /EtOH, wash it with anhydrous diethyl ether and dry it at room temperature in a vacuum over P_2O_5 . It sublimes at 220-230°/0.03mm with 99% recovery and **unracemised** [Gross & Gradsky *J Am Chem Soc* 77 1678 1955, DOI: 10.1021/ja01611a085]. [Cox & King *Org Synth Coll Vol* 2 612 1943, DOI: 10.1227/orgsyn.010.0100; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 pp 2316-2345 1961, Beilstein 22 IV 6765.] The non-natural **D(S)-enantiomer** [153-94-6] has m 282-285°(dec) [Beilstein 22/14 V14.]

Tyrocidine A (cyclic decapeptide antibiotic with two D-Phe amino acids) [1481-70-5] $\text{C}_{66}\text{H}_{57}\text{N}_{13}\text{O}_{13}$, M 1270.5, m 240°(dec), $[\alpha]_D^{25} -115$ (c 0.91, MeOH). Crystallise tyrocidine A as the **hydrochloride** from aqueous MeOH or EtOH/HCl, which has $[\alpha]_D^{25} -111$ (c 1.37, 1:1 EtOH/ H_2O). [Paladini & Craig *J Am Chem Soc* 76 688 1954, DOI: 10.1021/ja01632a015; King & Craig *J Am Chem Soc* 77 6624 1955, DOI: 10.1021/ja01629a063; Okamoto et al. *Bull Chem Soc Jpn* 50 231 1977, DOI: 10.1246/bcsj.50.231; for separation from tyrocidine B and C see Battersby & Craig *J Am Chem Soc* 74 4019 1952, DOI: 10.1021/ja01136a014; Beilstein 26 III/IV 4280.]

L-Tyrosine [60-18-4] $\text{C}_9\text{H}_{11}\text{NO}_3$, M 181.2, m 290-295°(dec), 294-300°(dec), $[\alpha]_D^{25} -10.0^\circ$ (5M HCl), $\text{pK}_1^{25} 2.18$ (CO_2H), $\text{pK}_2^{25} 9.21$ (OH), $\text{pK}_3^{25} 10.47$ (NH_3^+). Likely impurities are L-cysteine and the ammonium salt. L-Tyrosine is dissolved in dilute ammonia, then crystallised by adding dilute acetic acid to pH 5. Also, crystallise it from H_2O or EtOH/ H_2O , and dry it at room temperature in a vacuum over P_2O_5 . Its solubility in H_2O (g/100ml) is 0.02 (0°), 0.105 (50°), 0.244 (75°) and 0.056 (100°), but it is insoluble in common organic solvents. It sublimes at 235-240°/0.03mm with 99.2% recovery and **unracemised** [Gross & Gradsky *J Am Chem Soc* 77 1678 1955, DOI: 10.1021/ja01611a085]. It is the precursor of dopamine and catecholamines. [Albert *Biochem J* 50 690 1952, DOI: 10.1042/bj0500690; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 pp 2348-2366 1961, Beilstein 14 IV 2264.] The non-natural **D-enantiomer** has [556-02-5], and the **racemate** decomposes at ~316°, with similar solubilities as the optically active isomers.

L(S)-Valine [72-18-4] $\text{C}_5\text{H}_{11}\text{NO}_2$, M 117.2, m 305-308°(dec), 315°, $[\alpha]_D^{17} +28.4$ (c 2, 5M HCl), $\text{pK}_1^{20} 2.38$ (CO_2H), $\text{pK}_2^{20} 9.59$ (NH_3^+). Crystallise L-valine from water by addition of EtOH. Its solubility in H_2O (g/100ml) is 8.3 (0°), 9.2(50°) and 10.2 (65°), but is insoluble in common organic solvents. It sublimes at 178-188°/0.03mm with 99.3% recovery and **unracemised** [Gross & Gradsky *J Am Chem Soc* 77 1678 1955, DOI: 10.1021/ja01611a085]. [Perrin *J Chem Soc* 3125 1958, DOI: 10.1039/JR9580003125; Greenstein & Winitz *The Chemistry of the Amino Acids* J. Wiley, Vol 3 pp 2368-2377 1961, Beilstein 4 IV 2659.] The **racemate** sublimes without melting, but decomposes at ~298°(very rapid heating in a sealed capillary), and has similar solubilities as the optically active isomers.

PROTEINS, ENZYMES, DNA and RNA

Abrin A and C (agglutinins from *Abrus seeds*) [1393-62-0] M_r 63,000-67,000. These are toxic lectins (proteins) from seeds of *Abrus precatorius*. The yellow-white powder is purified by successive chromatography on DEAE-Sephadex A-50, carboxymethylcellulose, and DEAE-cellulose. Abrin A is more positively charged on the DEAE-cellulose column and has been crystallised from $(\text{NH}_4)_2\text{SO}_4$ by the free interface diffusion technique. Its molecular weight (by sedimentation equilibrium) is 60,000, whereas Abrin C has molecular weight of 63,800. Treatment of A with mercaptoethanol at 100°/2 hours followed by SDS-PAGE gave a main band with M_r 32,000 and two very weak bands, whereas C (which is more toxic) gave two intense bands with M_r 28,000 and 33,000. [Wei et al. *J Biol Chem* **249** 3061 1974, PMID: 4830236.] Abrin C has been crystallised for X-ray analysis by the free interface diffusion technique described by Salemme [*Arch Biochem Biophys* **151** 533 1972, DOI: 10.1016/0003-9861(72)90530-9]. The crystals were grown at 37° in Pyrex tubes (5 x 30 cm) by layering 50 μl of protein solution (22mg/ml) over 100 μl of unbuffered 70% saturated $(\text{NH}_4)_2\text{SO}_4$ [Wei & Einstein *J Biol Chem* **249** 2985 1974, PMID: 4828333.] [UV and CD: Herrmann & Behnke *Biochim Biophys Acta* **621** 43 1980, DOI: 10.1016/0005-2795(80)90060-4; for physical and chemical properties see Herrmann & Behnke *Biochim Biophys Acta* **667** 397 1981, DOI: 10.1016/0005-2795(81)90206-3; for the crystal structure of Arbin-A see Tahirov et al. *J Mol Biol* **250**(3) 354 1995, DOI: 10.1006/jmbi.1995.0382, PMID: 7608980; Beilstein **22** III/IV 6776.] The median toxic dose for humans is in the range 10 to 1000 $\mu\text{g/Kg}$ [Johnson et al. *J Analyt Toxicol* **33** 77 2009, <http://jat.oxfordjournals.org/content/33/2/77.full.pdf>; Gill *Microbiological Reviews* **46** (1): 86–94 1982, PMC 373212, PMID: 6806598].

Acetoin dehydrogenase [from beef liver, acetoin NAD oxidoreductase] [9028-49-3] M_r 76000, [EC 1.1.1.5]. Purify it *via* the acetone cake, then Ca-phosphate gel filtration (unabsorbed), lyophilised and then fractionated through a DEAE-22 cellulose column. The K_m for *diacetyl* is 40 μM , and for *NADH* it is 100 μM in phosphate buffer at pH 6.1. [Burgos & Martin *Biochim Biophys Acta* **268** 261 1972, DOI: 10.1016/0005-2744(72)90321-X; **289** 13 1972, DOI: 10.1016/0005-2744(72)90102-7.]

β -D-N-Acetylglucosaminidase [from *M sexta* insects] [9012-33-3] M_r ~61,000 [EC 3.2.1.52]. Purify it by chromatography on DEAE-Biogel, hydroxylapatite chromatography and gel filtration through Sephacryl S200. Two isoforms: a *hexosaminidase* EI with K_m 177 μM (V_{\max} 328 sec^{-1}) and EII a *chitinase* with K_m 160 μM (V_{\max} 103 sec^{-1}) with 4-nitrophenyl- β -acetylglucosamine as substrate. [Dziadik-Turner et al. *Arch Biochem Biophys* **212** 546 1981, DOI: 10.1016/0003-9861(81)90398-2.]

β -D-N-Acetylhexosaminidase A and B (from human placenta) [9012-33-3] M_r ~61,000, [EC 3.2.1.52]. Purify it by Sephadex G-200 filtration and DEAE-cellulose column chromatography. The hexosaminidase A is further purified by DEAE-cellulose column chromatography, followed by an ECTEOLA-cellulose column, Sephadex-200 filtration, electrofocusing and Sephadex G-200 filtration. Hexosaminidase B is purified by a CM-cellulose column, electrofocusing and Sephadex G-200 filtration. [Srivastava et al. *J Biol Chem* **249** purification 2043, <http://www.jbc.org/content/249/7/2043>, PMID: 4818822; for kinetics see p2049, <http://www.jbc.org/content/249/7/2049>, PMID: 4818822; for biochemical genetics see p2054 <http://www.jbc.org/content/249/7/2054>, PMID: 4206549; 1974.]

N-Acetyl neuraminic acid aldolase [from *Clostridium perfringens*, N-acetylneuraminic acid pyruvate lyase] [9027-60-5] M_r 32,000 [EC 4.1.3.3]. Purify the aldolase by extraction with H_2O , protamine precipitation, $(\text{NH}_4)_2\text{SO}_4$ fractionation, Me_2CO precipitation, acid treatment at pH 5.7 and precipitation at pH 4.5. The equilibrium constant for pyruvate + *n*-acetyl-D-mannosamine \rightleftharpoons N-acetylneuraminidate at 37° is 0.64. The K_m for N-acetylneuraminic acid is 3.9mM in phosphate at pH 7.2 and 37°. [Comb & Roseman *Methods Enzymol* **5** 391 1962, DOI: 10.1016/S0076-6879(62)05246-5.] The enzyme from hog kidney (cortex) has been purified 1700-fold by extraction with H_2O , protamine sulfate precipitation, $(\text{NH}_4)_2\text{SO}_4$ fractionation, heating between 60-80°, a second $(\text{NH}_4)_2\text{SO}_4$ fractionation and starch gel electrophoresis. The K_m for N-acetylneuraminic acid is 1.5mM. [Brunetti et al. *J Biol Chem* **237** 2447 1962, <http://www.jbc.org/content/237/8/2447>, PMID: 13874013.]

Acyl-coenzyme A Synthase [from beef liver] [9013-18-7] M_r 57,000 [EC 6.2.1.2]. Purify the synthase by extraction with sucrose/ HCO_3 buffer, protamine sulfate precipitation, $(\text{NH}_4)_2\text{SO}_4$ (66-65%) fractionation (pH 4.35) and a second $(\text{NH}_4)_2\text{SO}_4$ (35-60%) fractionation (pH 4.35). It has K_m 0.15mM (V_{rel} 1.0) for octanoate and 0.41mM (V_{rel} 2.37) for heptanoate. The K_m for ATP is 0.5mM, all at pH 9.0 in ethylene glycol buffer at 38°. [Mahler et al. *J Biol Chem* **204** 453 1953, <http://www.jbc.org/content/204/1/453>, PMID: 13874013; Jencks *Methods Enzymol* **5** 467 1962, DOI: 10.1016/S0076-6879(62)05259-3.]

Acyl-coenzyme A Synthase (from yeast) [9012-31-1] M_r ~151,000 [EC 6.2.1.1]. This enzyme has been purified by extraction into phosphate buffer pH 6.8-7.0 containing 2-mercaptoethanol and EDTA, protamine sulfate precipitation, polyethylene glycol fractionation, Alumina γ gel filtration, concentration by $(\text{NH}_4)_2\text{SO}_4$ precipitation, Bio-Gel A-0.5m chromatography and DEAE-cellulose gradient chromatography. It has K_m (apparent) 0.24mM (for acetate) and 0.035mM (for CoA); 1.2 mM for ATP and Mg^{2+} 4.0mM. [Frenkel & Kitchens *Methods Enzymol* **71** 317 1981. DOI: 10.1016/0076-6879(81)71040-1]

ADP-Ribosyl transferase (adenyllyl transferase, polynucleotide, from human placenta) [9026-30-6] M_r ~115,000 [EC 2.4.2.30]. Purify the transferase by making an affinity absorbent for ADP-ribosyltransferase by coupling 3-aminobenzamide to Sepharose 4B. [Burtscher et al. *Anal Biochem* **152** 285 1986, DOI: 10.1016/0003-2697(86)90410-0]

Agglutinin (from peanuts) [*Arachis hypogaea*] [1393-62-0] M_r 134,900 (tetramer). Agglutinin is purified by affinity chromatography on Sepharose- ζ -aminocaproyl- β -D-galactopyranosylamine. [Lotan et al. *J Biol Chem* **250** 8518 1975, <http://www.jbc.org/content/250/21/8518>, PMID: 811657.]

Albumin (bovine and human serum) [9048-46-8 (bovine), 70024-90-7 (human)] M_r ~67,000 (bovine), 69 000 (human), UV: $A_{280\text{nm}}^{1\%}$ 6.6 (bovine) and 5.3 (human) in H_2O , $[\alpha]_{546}^{25}$ -78.2° (H_2O). Albumin is purified by dissolving it in conductivity water and passage at 2-4° through two ion-exchange columns, each containing a 2:1 mixture of anionic and cationic resins (Amberlite IR-120, H-form, Amberlite IRA-400, OH-form). This treatment removes ions and lipid impurities. Care is taken to exclude CO_2 , and the solution is stored at -15°. [Möller et al. *Trans Faraday Soc* **57** 312 1961, DOI: 10.1039/TF9615700312.] More complete lipid removal is achieved by lyophilising the de-ionised solution, covering the dried albumin (human serum) with a mixture of 5% glacial acetic acid (v/v) in iso-octane (previously dried with Na_2SO_4) and allowing it to stand at 0° (without agitation) for upwards of 6 hours before decanting and discarding the extraction mixture, washing with iso-octane, re-extracting, and finally washing twice with iso-octane. The purified albumin is dried under vacuum for several hours, then dialyzed against water for 12-24 hours at room temperature, lyophilised, and stored at -10°C [Goodman *Science* **125** 1296 1957, DOI: 10.1126/science.125.3261.1296]. It has been recrystallised in high (35%) and in low (22%) EtOH solutions from Cohn's Fraction V.

The **high EtOH recrystallisation** is as follows: To 1kg of Fraction V albumin paste at -5° is added 300ml of 0.4 M pH (pH 5.5) acetate buffer in 35% EtOH pre-cooled to -10° and 430 ml of 0.1 M NaOAc in 25% EtOH also at -10°. Best results are obtained by adding all of the buffer and about half of the NaOAc and stirring slowly for 1 hour. The rest of the NaOAc is added when all the lumps have disintegrated. The mixture is set aside at -5° for several days to crystallise. 35% EtOH (1 L) is then added to dilute the crystalline suspension and lower the ionic strength prior to centrifugation at -5° (yield 80%). The crystals are further dissolved in 1.5 volumes of 15% EtOH/0.02M NaCl at -5° and clarified by filtration through washed, calcined diatomaceous earth. This solution may be recrystallised by re-adjusting to the conditions in the first crystallisation, or it may be recrystallised at 22% EtOH with the aid of a very small amount of decanol (enough to give a final concentration of 0.02%). **Note** that crystallisation from lower EtOH concentration gave better purification (i.e. by removing globulins and carbohydrates) and producing a more stable product.

The **low EtOH recrystallisation** is as follows: To 1kg of Fraction V at -10° to -15° is added 500ml of 15% EtOH at -5°, stirred slowly until a uniform suspension is formed. To the 15% EtOH (500ml) is added sufficient 0.2M NaHCO_3 solution (125-150ml) at 0° to bring the pH (1:10 dilution) to 5.3. Some temperature rise occurs, and care must be taken to keep the temperature < -5°. If the albumin is incompletely dissolved a small amount of H_2O is added (100ml at a time at 0°, allowing 15 minutes between additions). Undissolved albumin can be easily distinguished from small amounts of undissolved globulins, or as the last albumin dissolves, the appearance of the solution changes from milky white to hazy grey-green in colour. Keep the solution at -5° for

12 hours and filter by suspending in 15g of washed fine calcined diatomaceous earth, and filtering using a Büchner funnel precoated with coarser diatomaceous earth. The filtrate may require two or more similar filtrations to give a clear solution. To crystallise the filtrate, add through a capillary pipette, and with careful stirring, 1/100 volume of a solution containing 10% decanol and 60% EtOH (at -10°), and seed with the needle-like albumin crystals. After 2-3 days, crystallisation is complete. The crystals are centrifuged off. These are suspended with gentle mechanical stirring in one-third their weight of 0.005 M NaCl pre-cooled to 0° . With careful stirring, H_2O (at 0°) is added slowly in an amount equal to 1.7 times the weight of the crystals. At this stage there is about 7% EtOH, and the temperature cannot be made lower than -2.5° to -1° . Clarify, and collect as above. [Cohn et al. *J Am Chem Soc* **69** 1753 1947, DOI: 10.1021/ja01199a051.]

Human serum albumin has been purified similarly with 25% EtOH and 0.2% decanol. The isoelectric points of bovine and human serum albumins are 5.1 and 4.9, respectively.

Alamethicin (peptide antibiotic from *Trichoderma viridae*) [27061-78-5] $C_{92}H_{150}N_{22}O_{25}$, **M 1964.3**, **m 259-260 $^{\circ}$, 275-270 $^{\circ}$** , $[\alpha]_D^{22}$ -45 (c 1.2, EtOH), pK^{25} 6.04 (aqueous EtOH). Recrystallise alamethicin from MeOH. [MS, ^{13}C NMR and GC on optically active substrate confirms all L- amino acids: Panday et al. *J Am Chem Soc* **99** 8469 1977, DOI: 10.1021/ja00468a016.] The **acetate** [64918-47-4] $C_{94}H_{152}N_{22}O_{26}$, **M 2004**, has **m 195-180 $^{\circ}$** from MeOH/Et $_2$ O, the **methyl ester** has **m 175-176 $^{\circ}$ (240-242 $^{\circ}$)** from $CHCl_3$ /Et $_2$ O, and the **acetate-methyl ester** [64936-53-4] has **m 145-140 $^{\circ}$** from aqueous MeOH/Et $_2$ O. [Martin & Williams *Biochem J* **153** (2) 181 1976, DOI: 10.1042/bj1530181.]

Angiotensin (from rat brain) [70937-97-2] **M_r ~1524.8**. Angiotensin is purified using extraction, affinity chromatography and HPLC [Hermann et al. *Anal Biochem* **159** 295 1986, DOI: 10.1016/0003-2697(86)90346-5.]. It is obtained by the action of the ‘converting enzyme’ on **angiotensinogen** (see below). The following structures have been identified: **Angiotensin I** as Asp-Arg-Val-Tyr-Ile-His-Pro-Phe-His-Lue; **Angiotensin II** as Asp-Arg-Val-Tyr-Ile-His-Pro-Phe, and **Angiotensin III** as Arg-Val-Tyr-Ile-His-Pro-Phe [Ehlers & Riordan *Biochemistry* **28** 5311 1989, DOI: 10.1021/bi00439a001; history of discovery: Basso & Terragno *Hypertension* **38**(6) 1246 2001, DOI: 10.1161/hy1201.101214, PMID: 11751697].

Angiotensin-converting enzyme (ACE, angiotensinase, peptidyl peptide hydrolase) (from rabbit lung) [9015-82-1] **M 129,000 Dal (equilibrium sedimentation)**, **M_r ~140,000 (SDS-PAGE)** [EC 3.14.15.1]. Purify ACE by fractionation on DEAE-cellulose, Ca phosphate gel chromatography, elution from Sephadex G-200 and lectin affinity chromatography. The MW varied with glycosidation and is higher by gel filtration. It contains one atom of Zn/mol of protein and has K_m values for hydrolysis of hippurylhistidinylleucine and angiotensin I of 2.3 and 0.07 mM, and turnover of 15,430 and 792 mol/min/mol (min^{-1}) at 37° , respectively. The activity is inhibited by EDTA and increased amounts of Ca ions but required Ca ions. [Das & Soffer *J Biol Chem* **250** 6762 1975, <http://www.jbc.org/content/250/17/6762>, PMID: 169257; Reviewed by Ehlers & Riordan *Biochemistry* **28** 5311 1989, DOI: 10.1021/bi00439a001.]

Angiotensinogen (from porcine plasma) [64315-16-8] **M_r 59,400, 60,000, 62,600, 63,600 depending on sialic acid content**. This rennin substrate is purified 390-fold from the serum by chromatography on Blue Sepharose, phenyl sepharose, hydroxyapatite and finally by affinity chromatography on 5-hydroxytryptamine (5-HT)-sepharose to which it specifically binds to the 5-HT. It is applied to the latter column in 50mM sodium phosphate at pH 7 and after washing, it is eluted by increasing the ionic strength with 100mM sodium phosphate buffer containing 250mM NaCl. The multiple forms (with a tetradecapeptide having various degrees of glycosylation) are separated by SDS-PAGE and have pI 4.40-4.82. [Campbell et al. *Biochem J* **243** 121 1987, DOI: 10.1042/bj2430121; Beilstein **25** III/IV 4390 for angiotensin.]

Avidin (from egg white) [1405-69-2] **M_r ~70,000**. Avidin is purified by chromatography of an ammonium acetate solution on CM-cellulose [Green *Biochem J* **101** 774 1966, DOI: 10.1042/bj1010774]. It is also purified by affinity chromatography on 2-iminobiotin-6-aminoethyl-Sepharose 4B [Orr *J Biol Chem* **256** 761 1981, PMID: 6161128]. It is a biotin-binding protein.

Azurin (from *Pseudomonas aeruginosa*) [12284-43-4] **M_r 30,000**. Azurin with $A_{625/280} = 0.56$ is purified by gel chromatography on G-25 Sephadex with 5mM phosphate pH 7 buffer as eluent [Cho et al. *J Phys Chem* **91**

3690 1987, DOI: 10.1021/j100297a046]. It is a blue Cu protein used in biological electron transport, and its reduced form is obtained by adding a slight excess of $\text{Na}_2\text{S}_2\text{O}_4$. [See *Structure and Bonding* Springer Verlag, Berlin **23** 1 1975; for blue copper proteins see Rienzo et al. *Protein Sci* **9**(8) 1439 2000, DOI: 10.1110/ps.9.8.1439, PMID: 10975566.]

Bromelain (anti-inflammatory Ananase from pineapple) [37189-34-7] M_r ~33,000, [EC 3.4.33.4]. This protease has been purified *via* the acetone powder, G-75 Sephadex gel filtration and Bio-Rex 70 ion-exchange chromatography, and has $A_{1\text{cm}}^{1\%}$ 20.1 at 280nm. The protease from pineapple hydrolyses benzoyl glycine ethyl ester with a K_m (app) of 210mM and k_{cat} of 0.36 sec^{-1} . [Murachi *Methods Enzymol* **19** 273 1970, DOI: 10.1016/0076-6879(70)19021-5; Balls et al. *Ind Eng Chem* **33** 950 1941, DOI: 10.1021/ie50379a028.]

Carbonic anhydrase (carbonate hydrolase) [9001-03-0] M_r 31,000 [EC 4.2.1.1]. Purify carbonic anhydrase by hydroxylapatite and DEAE-cellulose chromatography [Tiselius et al. *Arch Biochem Biophys* **65** 132 1956, DOI: 10.1016/0003-9861(56)90183-7; Lindskog *Biochim Biophys Acta* **39** 218 1960, DOI: 10.1016/0006-3002(60)90156-6], and is then dialysed for crystallisation. A 0.5 to 1% solution of the enzyme in 0.05 M Tris-HCl pH 8.5 is dialysed against 1.75M solution of $(\text{NH}_4)_2\text{SO}_4$ in the same buffer, and this solution is slowly increased in salt concentration by periodic removal of small amounts of dialysate and replacing with an equal volume of 3.5M $(\text{NH}_4)_2\text{SO}_4$. The final salt concentration, in which the DEAE-cellulose fractions give beautiful birefringent suspensions of crystals, ranged from 2.4 to 2.7M and appeared first as fine crystals, then underwent transition to thin fragile plates. Carbonic anhydrase is a Zn enzyme which exists as several isoenzymes of varying degrees of activity [Funakoshi & Deutsch *J Biol Chem* **243** 6474 1968, PMID: 4973232; for crystal structure see Liljas et al. *Nature, New Biology* **235** 131 1972, DOI: 10.1038/newbio235131a0; see also P.D. Boyer Ed. *The Enzymes* Academic Press NY, pp 587-665 1971].

Carboxypeptidase A (from bovine pancreas, peptidyl-L-aminoacid lyase) [11075-17-5] M_r 34,600 [EC 3.4.17.1]. Carboxypeptidase A is purified by DEAE-cellulose chromatography, activation with trypsin and dialysed against 0.1M NaCl, yielding crystals. It is recrystallised by dissolving in 20 ml of M NaCl and dialysed for 24 hours each against the following salts present in 500ml of 0.02M sodium veronal pH 8.0, 0.5M NaCl, 0.2M NaCl and 0.15M NaCl. The last dialysate usually induces crystallisation. If it does not crystallise, then dialyse the last solution against 0.02M sodium veronal containing 0.10M NaCl. Only 2 or 3 recrystallisations are required to attain maximum activity. [Cox et al. *Biochemistry* **3** 44 1964, DOI: 10.1021/bi00889a008.] Enzyme activity is measured by hydrolysing hippuryl-L-phenylalanine (or phenylacetic acid) and observing the rate of change of optical density at 254nm (reaction extinction coefficient is $\sim 0.592 \text{ cm}^2/\mu\text{mole}$ at pH 7.5) [Bergmeyer *Methods in Enzymatic Analysis* (Academic Press) **1** 436 1974, ISBN 10: 0120913011 ISBN 13: 9780120913015].

Cathepsin B (from human liver) [9047-22-7] M_r 27,500 [EC 3.4.22.1]. Cathepsin B is purified by affinity chromatography on the semicarbazone of Gly-Phe-glycinal-linked to Sepharose 4B, with elution by 2,2'-dipyridyl disulfide [Rich et al. *Biochem J* **235** 731 1986, DOI: 10.1042/bj2350731; Barrett & Kirschke *Methods Enzymol* **80** 535 1981, DOI: 10.1016/S0076-6879(81)80043-2].

Cathepsin D (from bovine spleen) [9025-26-7] M_r 56,000 [EC 3.4.23.5]. Cathepsin D is purified on a CM column after $(\text{NH}_4)_2\text{SO}_4$ fractionation and dialysis, then starch-gel electrophoresis and by ultracentrifugal analysis. Finally chromatograph on a DEAE column [Press et al. *Biochem J* **74** 501 1960, DOI: 10.1042/bj0740501].

Ceruloplasmin (from human blood plasma) [9031-37-2] M_r 134,000. This blue protein is the principal Cu transporter (up to 90% of circulating Cu) and is purified by precipitation with polyethylene glycol 4000, batchwise adsorption and elution from QAE-Sephadex, and gradient elution from DEAE-Sepharose CL-6B. Ceruloplasmin is thus purified 1640-fold and is homogeneous on anionic polyacrylamide gel electrophoresis

(PAGE), SDS-PAGE, isoelectric focusing and low-speed equilibrium centrifugation. It has λ_{max} at 280, 260nm ($A_{1\text{cm}}^{1\%}$ 14.9, 0.68). [Oosthuizen *Anal Biochem* **146** 1 1985, DOI: 10.1016/0003-2697(85)90386-0; Cohn et al. *J Am Chem Soc* **68** 459 1946, DOI: 10.1021/ja01207a034.]

Chemokines. These are small proteins formed from longer precursors and are chemo-attractants for lymphocytes and lymphoid organs. They are characterised by having cysteine groups in specific relative positions. The two largest families are the α and β families that have four cysteine residues arranged (C-X-C) and (C-C) respectively. The mature chemokines have ~70 amino acids with internal cys S-S bonds and attract myeloid type cells *in vitro*. The γ -family (Lymphotactin) has only two cys residues. The δ -family (Neurotactin, Fractalkine) has the C-C-X-X-X-C sequence (*ca* 387 amino acids), binds to membrane and promotes adhesion of lymphocytes. It is the soluble domain of human Fractalkine ‘chemo-attract’ monocytes and T cells. Several chemokines are available commercially (some prepared by recombinant DNA techniques), including 6CKine/exodus/SLC which belongs to the β -family with 6 cysteines (110 amino acids, mature protein), as the name implies (C-C-C-C-X.....X-C-C) and homes lymphocytes to secondary lymphoid organs with lymphocyte adhesion antitumor properties. Other chemokines available are C10 (β CC) and Biotaxin. Several chemokine receptors and antibodies are available commercially and can generally be used without further purification. [Murphy ‘Molecular biology of lymphocyte chemo-attractant receptors’ in *Ann Rev Immunol* **12** 593 1994, DOI: 10.1146/annurev.iy.12.040194.003113.]

Chirazymes. These are commercially available enzymes, e.g. lipases, esterases, that can be used for the preparation of a variety of optically active carboxylic acids, alcohols and amines. They can cause regio and stereospecific hydrolysis and do not require cofactors. Some can be used also for esterification or transesterification in neat organic solvents. The proteases, amidases and oxidases are obtained from bacteria or fungi, whereas esterases are from pig liver and thermophilic bacteria. For preparative work the enzymes are covalently bound to a carrier and do not therefore contaminate the reaction products. Chirazymes are available from Roche Molecular Biochemicals or similar biochemicals suppliers, and are used without further purification.

α -Chymotrypsin [9004-07-3] M_r ~25000 [EC 3.4.21.1]. α -Chymotrypsin is crystallised twice from four-tenths saturated ammonium sulfate solution, then dissolved in 1mM HCl and dialysed against 1mM HCl at 2-4°. The solution is stored at 2° [Lang et al. *J Am Chem Soc* **80** 4923 1958, DOI: 10.1021/ja01551a041].

Citric acid cycle components (from rat heart mitochondria). These are resolved by anion-exchange chromatography [LaNoue et al. *J Biol Chem* **245** 102 1970, PMID: 4312474].

Clostripain [9028-00-6] M_r ~55,000, [EC 3.4.22.8]. Clostripain is isolated from *Clostridium histolyticum* collagenase by extraction in pH 6.7 buffer, followed by hydroxylapatite chromatography with a 0.1-0.2 M phosphate gradient, then Sephadex G-75 gel filtration with 0.05M phosphate pH 6.7, dialysis and a second hydroxylapatite chromatography (gradient elution with 0.1M \rightarrow 0.3M phosphate, pH 6.7) purification. It has proteinase and esterase activity and is assayed by hydrolysing *N*-benzoyl-L-arginine methyl ester. [Mitchell & Harrington *J Biol Chem* **243** 4683 1968, PMID: 4971659; *Methods Enzymol* **19** 635 1970, DOI: 10.1016/0076-6879(70)19050-1.]

Colicin E1 (from *E.coli*) [11032-88-5] M_r 56,000, pI 9.5. Colicin E1 is purified (8.6-fold to Specific Activity of 1.5×10^5 units/mg) from *E.coli* JC411 by salt extraction of extracellular-bound colicin followed by $(\text{NH}_4)_2\text{SO}_4$ (40-60% saturation) fractionation and ion-exchange chromatography on a DEAE-Sephadex A 50 column, and then by CM-Sephadex column chromatography [Schwartz & Helinski *J Biol Chem* **246** 6318 1971, PMID: 5001789].

Collagenase (from human polymorphonuclear leukocytes) [9001-12-1] M_r 68,000-125,000 [EC 3.4.24.3]. Collagenase is purified by using *N*-ethylmaleimide to activate the enzyme, and wheat germ agglutinin-agarose affinity chromatography [Callaway et al. *Biochemistry* **25** 4757 1986, DOI: 10.1021/bi00365a006].

Copper-zinc-superoxide dismutase (from blood cell haemolysis) [9054-89-1] M_r ~32,000 [EC 1.15.1.1].

The dismutase is purified by DEAE-Sepharose and copper chelate affinity chromatography. The preparation so achieved is homogeneous by SDS-PAGE, by analytical gel filtration chromatography and by isoelectric focusing [Weselake et al. *Anal Biochem* **155** 193 1986, DOI: 10.1016/0003-2697(86)90246-0; McCord & Fridovich *J Biol Chem* **244** 6049 1969, PMID: 5389100].

Cytochrome c_1 (from horse, beef or fishes' heart, or pigeon breast muscle) [9007-43-6] $M_r \sim 13,000$. Cytochrome c_1 is purified by chromatography on CM-cellulose (CM-52 Whatman) [Brautigan et al. *Methods Enzymol* **53** 128 1978, DOI: 10.1016/S0076-6879(78)53021-8]. It has a high PI (isoelectric point) and has been purified further by adsorption onto an acidic cation exchanger, e.g. Amberlite IRC-50 (polycarboxylic) or in ground form Amberlite XE-40 (100-200 mesh) or Decalco-F (aluminium silicate), where the non-cytochrome protein is not adsorbed and is readily removed. The cytochrome is eluted using a solution containing 0.25g ions/L of a univalent cation at pH 4.7 adsorbed onto the NH_4^+ salt of Amberlite IRC-50 at pH 7, washed with H_2O and then with 0.12M NH_4OAc to remove non-cytochrome protein. When the cytochrome begins to appear in the eluate, then the NH_4OAc concentration is increased to 0.25 M. The fractions with ca $\text{Fe} = 0.465-0.467$ are collected, dialysed against H_2O and adsorbed onto a small IRC-50 column and eluted with 0.5M NH_3 , then dialysed and lyophilised. (A second fraction II can be eluted from the first resin with 0.5M NH_3 but is discarded). [Keilin & Hartree *Biochemical Preparations* **1** 1 1952, Margoliash *Biochemical Preparations* **8** 33 1957.]

Cytochrome c has been recrystallised as follows: The above eluate (ca 100ml) is dialysed against H_2O (10 vols) at 4° for 24 hours (no more), then passed through an XE-40 column (2 x 1 cm above) which is equilibrated with 0.1M NH_4OAc pH 7.0. The column is washed with 0.1% $(\text{NH}_4)_2\text{SO}_4$ pH 8.0, and the dark red resin in the upper part of the column is collected and in 0.1% $(\text{NH}_4)_2\text{SO}_4$ pH 8.0 is transferred to another column (7mm diameter) and the cytochrome c is eluted with 5% $(\text{NH}_4)_2\text{SO}_4$ pH 8.0. More than 98% of the red colour is collected in a volume of ca 4ml in a weighed centrifuge tube. Add a drop of octanol and 0.43g of $(\text{NH}_4)_2\text{SO}_4$ /g of solution. When the salt has dissolved, ascorbic acid (5mg) is added as well as a few drops of 30% aqueous NH_3 , and it is kept at 10° for 10 minutes (turns lighter colour due to reduction). Then add finely powdered $(\text{NH}_4)_2\text{SO}_4$ in small portions (stir with a glass rod) until the solution becomes turbid. Stopper the tube tightly, and set aside at $15-25^\circ$ for 2 days while the cytochrome c separates as fine needles or rosettes. Further $(\text{NH}_4)_2\text{SO}_4$ (20mg) are added per ml of suspension and kept in the cold for a few days to complete the crystallisation. The crystals are collected by centrifugation (5000xg), suspended in saturated $(\text{NH}_4)_2\text{SO}_4$ (pH 8.0 at 10°), then centrifuged again. For recrystallisation the crystals are dissolved in the least volume of H_2O , one drop of ammonia and 1 mg of ascorbic acid are added and the above process is repeated. The yield of twice recrystallised cytochrome c from 2Kg of muscle is ca 200 mg, but this varies with the source and freshness of the muscle used. The crystals are stored as a solid after dialysis against 0.08M NaCl or 0.1M sodium buffer and lyophilising, or as a suspension in saturated $(\text{NH}_4)_2\text{SO}_4$ at 0° . [Hagihara et al. *Biochemical Preparations* **6** 1 1958.]

Purity of cytochrome c : This is checked by the ratio of the absorbance at 500nm (reduced form) to 280nm (oxidised form), i.e. $\epsilon_{500}/\epsilon_{280}$ should be between 1.1 and 1.28, although values of up to 1.4 have been obtained for pure preparations.

For the preparation of the **reduced form** see Margoliash *Biochemical Preparations* **5** 33 1957 and Yonetani *Biochemical Preparations* **11** 19 1966.

Cytochrome from *Rhodospirillum rubrum* ($\epsilon_{270}/\epsilon_{551}$ 0.967) is purified by chromatography on a column of CM-Whatman cellulose [Paleus & Tuppy *Acta Chem Scand* **13** 641 1959, DOI: 10.3891/acta.chem.scand.13-0641].

Cytochrome c oxidase (from bovine heart mitochondria) [9001-16-5] M_r 100,000/haeme [EC 1.9.3.1]. This oxidase is purified by selective solubilisation with Triton X-100 and subsequently with lauryl maltoside, and finally by sucrose gradient centrifugation [Li et al. *Biochem J* **242** 417 1987, DOI: 10.1042/bj2420417].

It has also been purified by extraction into 0.02 M phosphate buffer (pH 7.4) containing 2% of cholic acid (an inhibitor which stabilises as well as solubilises the enzyme) and fractionated with $(\text{NH}_4)_2\text{SO}_4$, collecting the 26-33% saturation cut and refractionating again and collecting the 26-33% saturation fraction. The pellet collected at 10,000xg appears as an oily paste. **The cholate needs to be removed to activate the enzyme as follows:** The precipitate is dissolved in 10ml of 0.1M phosphate buffer pH 7.4, containing 1% of Tween-80 and dialysed against 1L of 0.01 M PO_4 buffer (pH 7.4) containing 1% of Tween-80 for 10 hours at 0° and aliquoted. The enzyme is stable at 0° for 2 weeks and at -15° for several months. It is assayed for purity (see reference) by oxidation of reduced cytochrome c (K_m 10 μ M). [Yonetani *Biochemical Preparations* **11** 14 1966, *J Biol Chem*

236 1680 1961, PMID: 13787373.]

Cytokines See chemokines, interferons, interleukins.

Deoxyribonucleic acid (from plasmids). These are purified by two buoyant density ultracentrifugations using ethidium bromide-CsCl. The ethidium bromide is extracted with Et₂O, and the DNA is dialysed against buffered EDTA and lyophilised. [Marmur & Doty *J Mol Biol* **5** 109 1962, DOI: 10.1016/S0022-2836(62)80066-7; Guerry et al. *J Bacteriol* **116** 1064 1973, PMID: 4583233.] See ‘Introduction’ in this chapter.

Dermatan sulfate (condroitin sulfate B from pig skin) M_r 20,000-36,000 [54328-33-5 (Na salt)]. Dermatan sulfate is purified by digestion with papain and hyaluronidase, and fractionation using aqueous EtOH. [Gifonelli & Roden *Biochemical Preparations* **12** 1 1968.]

Dihydrofolate reductase (from *Mycobacterium phlei*) [9002-03-3] M_r ~18,000 [EC 1.5.1.3]. Dihydrofolate reductase is purified by (NH₄)₂SO₄ precipitation, then fractionation on a Sephadex G-75 column, applied to a Blue Sepharose column and eluted with 1mM dihydrofolate. [Al- Rubeai & Dale *Biochem J* **235** 301 1986, DOI: 10.1042/bj2350301.]

Dihydropteridine reductase (from sheep liver) [9074-11-7] M_r 52,000 [EC 1.6.99.7]. Dihydropteridine reductase is purified by fractionation with ammonium sulfate, dialysed against Tris buffer, adsorbed and eluted from hydroxylapatite gel. It is then run through a DEAE-cellulose column and also subjected to Sephadex G-100 filtration. [Craine et al. *J Biol Chem* **247** 6082 1972, PMID: 4405600.]

Dihydropteridine reductase (from human liver) [9074-11-7] M_r 52,000 [EC 1.6.99.7]. Dihydropteridine reductase is purified to homogeneity on a naphthoquinone affinity adsorbent, followed by DEAE-Sephadex and CM-Sephadex chromatography. [Firgaira, Cotton and Danks, *Biochem J* **197** 31 1981, DOI: 10.1042/bj1970031.] [For other dihydropteridine reductases see Armarego et al. *Med Res Rev* **4**(3) 267 1984, DOI: 10.1002/med.2610040302.]

3,4-Dihydroxyphenylalanine-containing proteins. Boronate affinity chromatography is used in the selective binding of proteins containing 3,4-dihydroxyphenylalanine to a *m*-phenylboronate agarose column and eluted with 1M NH₄OAc at pH 10. [Hawkins et al. *Anal Biochem* **159** 187 1986, DOI: 10.1016/0003-2697(86)90326-X.]

Dipeptidyl aminopeptidase (dipeptidyl peptidase IV, from rat brain) [9031-94-1, 54249-88-6] M_r 87,500 (monomer SDS-PAGE), (88,107 from nucleotide sequence), up to 4000,000 [EC 3.4.14.5]. The aminopeptidase is purified about 2000-fold by column chromatography on CM-cellulose, hydroxylapatite and Gly-Pro AH-Sepharose. [Imai et al. *J Biochem (Tokyo)* **93** 431 1983, PMID: 6341372; Schomburg & Schomburg *Springer Handbook of Enzymes* 2nd Edn vol **6** p 286 2002.]

DNA (deoxyribonucleic acids). The essential structures of chromosomes are DNA and contain the genetic ‘blueprint’ in the form of separate genes. They are made up of the four deoxyribonucleic acids (nucleotides): adenylic acid, guanylic acid, cytidylic acid and thymidylic acid (designated A, G, C, T respectively) linked together by their phosphate groups in ester bonds between the 3' and 5' hydroxy groups of the 2'-deoxy-D-ribose moiety of the nucleotides. The chains form a double-stranded spiral (helix) in which the two identical nucleotide sequences run antiparallel with the heterocyclic bases hydrogen bonded (A....T, G....C) forming the ‘ladder’ between the strands. Short sequences of DNA are available commercially, are commercially custom made or synthesised in a DNA synthesiser and purified by HPLC. Their purity can be checked by restriction enzyme cleavage followed by gel electrophoresis, or directly by gel electrophoresis or analytical HPLC. Commercial DNAs are usually pure enough for direct use but can be further purified using commercially available kits involving binding to silica or other matrices and eluting with Tris buffers. There are now rapid ‘throughput’ techniques for sequencing DNA which are very accurate.

Dopamine- β -hydroxylase (from bovine adrenal medulla) [9013-38-1] M_r ~290,000 [EC 1.14.17.1]. The Cu-containing glycoprotein enzyme has been isolated by two procedures. The **first** is an elaborate method requiring extraction, two $(\text{NH}_4)_2\text{SO}_4$ fractionations, calcium phosphate gel filtration, EtOH fractionation, DEAE-cellulose chromatography followed by two Sephadex-G200 gel filtrations giving enzyme with a specific activity of 65 Units/mg. [Friedman & Kaufman *J Biol Chem* **240** 4763 1965, PMID: 5846992; Rush et al. *Biochem Biophys Res Commun* **61** 38 1974, DOI: 10.1016/0006-291X(74)90530-0.] The **second** procedure is much gentler and provides good quality enzyme. Sedimented chromaffin vesicles are lysed in 10 volumes of 5mM K-phosphate buffer pH 6.5 using a loosely fitting Teflon-glass homogeniser. The mixture is centrifuged at 40,000xg/0.5 hours, and the supernatant is diluted with an equal volume of 100mM phosphate buffer (pH 6.5) containing 0.4M NaCl. This lysate is applied to a concanavalin A-Sepharose column (4 x 0.7cm) which has been equilibrated with 50 mM of phosphate buffer (pH 6.5 + 0.2M NaCl) with a flow rate of ~ 0.3 ml/minute. The column is washed thoroughly with the buffer until OD_{280nm} is 0.005. The enzyme is then eluted with the same buffer containing 10% α -methyl-D-mannoside (flow rate 0.1 ml/minute), and the enzyme is collected in 20-column volumes. The pooled eluate is concentrated by ultrafiltration in an Amicon Diaflo stirrer cell using an XM100A membrane. The concentrated enzyme is dialysed against 50mM phosphate buffer (pH 6.5) containing 0.1% NaCl. The enzyme gives one band (+ two very weak bands) on disc gel electrophoresis indicating better than 93% purity (67% fold purification) and has a specific activity of 5.4 Units/mg. [Rush et al. *Biochem Biophys Res Commun* **57** 1301 1974, DOI: 10.1016/0006-291X(74)90837-7; Stewart & Klinman *Ann Rev Biochem* **57** 551 1988, DOI: 10.1146/annurev.bi.57.070188.003003.]

Exonucleases. Like the endonucleases they are restriction enzymes which act at the 3' or 5' ends of linear DNA by hydrolysing off the nucleotides. Although they are highly specific for hydrolysing nucleotides at the 3' or 5' ends of linear DNA, the number of nucleotides cleaved is time dependent and usually has to be estimated from the time allocated for cleavage. Commercially available exonucleases are used without further purification.

Ferritin (from human placenta) [9007-73-2] M_r ~445,000 (**Fe free protein**). The purification of this major iron-binding protein is achieved by homogenisation in water and precipitation with ammonium sulfate, repeating the cycle of ultracentrifugation, and molecular sieve chromatography through a Sephadex 4B column. Isoelectric focusing reveals a broad spectrum of impurities which can be separated by ion-exchange chromatography on Sephadex A-25 and stepwise elution. [Konijn et al. *Anal Biochem* **144** 423 1985, DOI: 10.1016/0003-2697(85)90135-6.]

Fibrinogen (from human plasma) [9001-32-5] M_r **341,000**. This protein is made up of 2A α , 2B β and 2 γ subunits connected by disulfide bridges. A likely impurity is plasminogen. It is purified by glycine precipitation [Mosesson & Sherry *Biochemistry* **5** 2829 1966, DOI: 10.1021/bi00873a008] to obtain fractions 1-2, then further purified [Blombäck & Blombäck *Arkiv Kemi* **10** 415 1956] and contaminating plasminogen is removed by passage through a lysine-Sepharose column. Such preparations are at least 95% clottable as determined by Mosesson and Sherry's method (above ref.) in which the OD₂₈₀ is measured before and after clotting with 5 Units/ml of thrombin (> 3000U/mg). All fibrinogen preparations are treated with calf intestinal alkaline phosphatase to convert any fibrinogen peptide-AP to fibrinogen peptide-A by removing serine-bound phosphate. Solutions are then lyophilised and stored at -20°. [Higgins & Shafer *J Biol Chem* **256** 12013 1981, PMID: 7298640.] For the resolution of polypeptide chains of sulfitolysed fibrinogen by polyacrylamide gel electrophoresis (PAGE) see Takagi & Iwanaga [*Biochim Biophys Acta (Protein Structure)* **194** 594 1969, DOI: 10.1016/0005-2795(69)90121-4]. It is sparingly soluble in H₂O. Aqueous solutions are viscous with isoelectric point at pH 5.5. It is readily denatured by heating above 56° or by chemical agents, e.g. salicylaldehyde, naphthoquinone sulfonates, ninhydrin or alloxan. [Edsall et al. *J Am Chem Soc* **69** 2731 1947, DOI: 10.1021/ja01203a048; Purification: Cama et al. *Naturwissenschaften* **48** 574 1961, DOI: 10.1007/BF00589726; Lorand & Middlebrook *Science* **118** 515 1953, DOI: 10.1126/science.118.3070.515; cf. Fuller in *Methods Enzymol* **163** 474 1988, DOI: 10.1016/0076-6879(88)63044-8.]

For plasminogen-deficient fibrinogen from blood plasma, the anticoagulated blood is centrifuged and the plasma is frozen and washed with saline solution. It is treated with charcoal, freeze-thawed and dialysed *versus* Tris/NaCl buffer. [Maxwell & Nikel *Biochemical Preparations* **12** 16 1968.]

Fibronectin (from human plasma) [86088-83-7] M_r ~220,000. This glycoprotein contains 5-12% of carbohydrate. It has been purified by glycine fractionation and DEAE-cellulose chromatography. This material is dissolved in 0.25M Tris-phosphate buffer pH 7.0, diluted to 20% and glycine added gradually till 2.1M when the temperature falls to below 15°. The precipitate contains mainly fibrinogen. The supernatant is discarded, and the precipitate is treated with an equal volume of H₂O, cooled (to 0°) and precipitated by adding EtOH to 16% (v/v) at -4°. The precipitate contains some CI (Cold Insoluble) globulin, fibronectin and small quantities of other proteins. To remove these, the precipitate is dissolved in 0.25M Tris-phosphate buffer (pH 7.0) *ca* 0.5% and purified by DEAE-cellulose chromatography after diluting the buffer to 0.05M buffer. [Morrison et al. *J Am Chem Soc* **70** 3103 1948, DOI: 10.1021/ja01189a080; Mosesson & Umfleet *J Biol Chem* **245** 5728 1970, PMID: 4097343; Mosesson & Amrani *Blood* **56** 145 1980, Akiyama & Yamada *Adv Enzymol* **59** 51 1987.]

Follicle Stimulating Hormone (FSH, follitropin) [9002-68-0] M_r ~36,000. FSH is purified by Sephadex G100 gel filtration followed by carboxymethyl-cellulose with NH₄OAc pH 5.5. The latter separates luteinising hormone from FSH. Its solubility in H₂O is 0.5%. It has an isoelectric point of 4.5. A solution of 1mg in saline (100ml) can be kept at 60° for 0.5 hour. Activity is retained in a solution at pH 7-8 for 0.5 hour at 75°. The activity of a 50% aqueous EtOH solution is destroyed at 60° in 15 minutes. [Bloomfield et al. *Biochim Biophys Acta* **533** 371 1978, DOI: 10.1016/0005-2795(78)90383-5; Hartree *Biochem J* **100** 754 1966, DOI: 10.1042/bj1000754; Pierce & Parsons *Ann Rev Biochem* **50** 465 1981, DOI: 10.1146/annurev.bi.50.070181.002341.]

β-Galactosidase (from bovine testes) [9031-11-2] M_r 510,000 [EC 3.2.1.23]. It is purified 600-fold by (NH₄)₂SO₄ precipitation, acetone fractionation and affinity chromatography on agarose substituted with terminal thio-β-galactopyranosyl residues. [Distler & Jourdan *J Biol Chem* **248** 6772 1973, PMID: 4270451.]

Glucose oxidase (from *Aspergillus niger*) [9001-37-0] M_r 186,000 [EC 1.1.3.4]. The oxidase is purified by dialysis against deionized water at 6° for 48 hours and by molecular exclusion chromatography with Sephadex G-25 at room temperature. [Holt & Cotton *J Am Chem Soc* **109** 1841 1987, DOI: 10.1021/ja00240a039.]

Glucose-6-phosphate dehydrogenase [9001-40-5] M_r 128,000 (from Baker's yeast), 63,300 (from rat mammary gland) [EC 1.1.1.49]. The enzyme is useful for measuring pyridine nucleotides in enzyme recycling. The enzyme from Baker's yeast has been purified by (NH₄)₂SO₄ fractionation, Me₂CO precipitation, a second (NH₄)₂SO₄ fractionation, concentration by DEAE-SF chromatography, a third (NH₄)₂SO₄ fractionation and recrystallisation. Crystallisation is induced by addition of its coenzyme NADP, which in its presence causes rapid separation of crystals at (NH₄)₂SO₄ concentration much below that required to precipitate the amorphous enzyme. To recrystallise, the crystals are dissolved in 0.01M NADP (pH 7.3) with (NH₄)₂SO₄ at 0.55 saturation, and the crystals appear within 10 to 60 minutes. After standing for 2-3 days (at 4°) the (NH₄)₂SO₄ is increased to 0.60 of saturation, and more than 80% of the activity in the original crystals is recovered in the fresh crystals. [Noltmann et al. *J Biol Chem* **236** 1225 1961, PMID: 13729473]. Large amounts can be obtained from rat livers. The livers are extracted with 0.025M phosphate buffer (pH 7.5) and precipitated with 3M (NH₄)₂SO₄ (70% of activity). The precipitate is dissolved in 3 volumes of 0.025M phosphate (pH 7.5), dialysed against this buffer + 0.2mM EDTA at 4° for 5 hours, then diluted to 1% protein and the nucleic acids are precipitated by addition of 0.4 volumes of 1% protamine sulfate. (NH₄)₂SO₄ is added to a concentration of 2M (pH adjusted to 7.0 with NH₃), the precipitate is discarded and the supernatant is adjusted to 2.8M (NH₄)₂SO₄, dialysed, and the protein is adjusted to 1% and treated with Ca₃(PO₄)₂ gel. The gel is added in three steps (1.5ml of 0.4% gel/ml per step), and the gel is removed by centrifugation after each addition. The third gel adsorbed 50% of the activity. The gel is eluted with 0.2M phosphate buffer (pH 7.4, 40ml/g of gel; 60% recovery). The extract is precipitated in 3 volumes of (NH₄)₂SO₄ (adjusted to 4M) to give enzyme with an activity of 30μmoles/mg of protein per hour. [Lowry et al. *J Biol Chem* **236** 2746 1961, PMID: 14466981.] The Km values for the yeast enzyme are 20μM for G-6P and 2μM for NADP (Tris pH 8.0, 10⁻² M MgCl₂, 38°) [Noltmann & Kubly *The Enzymes* **VII** 223 1963].

Glutathione S-transferase (human liver) [50812-37-8] M_r 25,000 [EC 2.5.1.18]. It is purified by affinity chromatography using a column prepared by coupling glutathione to epoxy-saturated Sepharose. After washing

off contaminating proteins, the pure transferase is eluted with buffer containing reduced glutathione. The solution is then concentrated by ultrafiltration, dialysed against phosphate buffer at pH ~7 and stored in the presence of dithiothreitol (2mM) in aliquots at <-20°. [Simons & Vander Jagt *Anal Biochem* **82** 334 1977, DOI: 10.1016/0003-2697(77)90169-5.]

Glyceraldehyde-3-phosphate dehydrogenase [9001-50-7] M_r **144,000** [EC **1.2.1.12**]. Purify the dehydrogenase from rabbit muscle by extraction with 0.03N KOH and precipitate it with $(\text{NH}_4)_2\text{SO}_4$ (0.52 of saturation). The clear supernatant is adjusted to pH 7.5, and NH_3 is added dropwise to pH 8.2-8.4. Crystals appear sometimes even without seeding. The crystals are dissolved in H_2O , filtered to remove suspended material and 2 volumes of saturated $(\text{NH}_4)_2\text{SO}_4$ at pH 8.2-8.4 is added. After 1 hour the crystals appear. Recrystallise it in the same way. [Cori et al. *J Biol Chem* **173** 605 1948, PMID: 18910716; Furfine & Velick *J Biol Chem* **240** 844 1965, PMID: 14275144; *The Enzymes* **7** 243 1963, Lui & Huskey *Biochemistry* **31** 6898 1992, DOI: 10.1021/bi00145a005.] The K_m values are: NADH ($3.3\mu\text{M}$) and 1,3-diphosphoglycerate ($8 \times 10^{-7}\text{M}$) in pH 7.4 imidazole buffer at 26°, NAD ($13\mu\text{M}$), glyceraldehyde-3-P ($90\mu\text{M}$), P_i ($2.9 \times 10^{-4}\text{M}$), and arsenate ($69\mu\text{M}$) in 8.6 M NaHCO_3 buffer at 26°C. [Orsi & Cleland *Biochemistry* **11** 102 1972, DOI: 10.1021/bi00751a018.]

Glycerol kinase (from *Candida mycoderma*, *E coli*, rat or pigeon liver glycerokinase) [9030-66-4] M_r **251,000** [EC **2.7.1.30**]. Commercial enzyme has been dialysed against 2mM Hepes, 5mM dithiothreitol and 0.3mM EDTA, followed by several changes of 20mM Hepes and 5mM dithiothreitol prior to storage under N_2 at -20°. [Knight & Cleland *Biochemistry* **28** 5728 1989, DOI: 10.1021/bi00440a005.] The enzyme from pigeon liver is purified by acid-precipitation (acetate buffer at pH 5.1), $(\text{NH}_4)_2\text{SO}_4$ fractionation, heat treatment (60°/1 hour), calcium phosphate gel filtration, a second $(\text{NH}_4)_2\text{SO}_4$ fractionation, dialysis, elution of inert proteins and crystallisation. This is done by repeatedly extracting the precipitate from the last step with 0.05M sodium pyrophosphate (pH 7.5) containing 1mM EDTA, and 0.2M $(\text{NH}_4)_2\text{SO}_4$ is added. Careful addition of solid $(\text{NH}_4)_2\text{SO}_4$ to this solution leads to crystallisation of the enzyme. Recrystallisation is repeated. The enzyme is activated by Mg^{2+} and Mn^{2+} ions and is most stable in solutions in the pH 4.5-5.5 range. The stability is greatly increased in the presence of glycerol. It has K_m for glycerol is $60\mu\text{M}$, and for ATP is $9\mu\text{M}$ in glycine buffer pH 9.8 and 25°. [Kennedy *Methods Enzymol* **5** 476 1962, DOI: 10.1016/S0076-6879(62)05261-1.]

L-Glycerol-3-phosphate dehydrogenase (GDH, from rabbit muscle) [9075-65-4] M_r **78,000** [EC **1.1.1.8**]. The dehydrogenase is recrystallised by adding $(\text{NH}_4)_2\text{SO}_4$ to 0.45 saturation at pH 5.5 at 4°C, and the small amount of precipitate is removed, then a saturated solution of $(\text{NH}_4)_2\text{SO}_4$ is added dropwise from time to time over several days in the cold room. The crystals are collected and recrystallised until they have maximum activity. The enzyme is stable in half-saturated $(\text{NH}_4)_2\text{SO}_4$ for several weeks at 4°. The equilibrium $[\text{dihydroxyacetone}][\text{NADH}][\text{H}^+]/[\text{G-3-P}][\text{NAD}]$ is $1.0 \times 10^{-12}\text{M}$ in Tris buffer at 25°. It uses NAD ten times more efficiently than NADP. The K_m for G-3-P is $1.1 \times 10^{-4}\text{M}$, for NAD it is $3.8 \times 10^{-4}\text{M}$ and for dihydroxyacetone it is $4.6 \times 10^{-4}\text{M}$ in phosphate buffer pH 7.0 and at 23.3°. Dihydroxyacetone phosphate and fructose-1,6-diphosphate are inhibitors. [Baranowski *J Biol Chem* **180** 535 1949, PMID: 18135786; *The Enzymes* **7** 85 1963, Young & Pace *Arch Biochem Biophys* **75** 125 1958, DOI: 10.1016/0003-9861(58)90403-X; Walsh & Sallach *Biochemistry* **4** 1076 1965, DOI: 10.1021/bi00882a015.]

Glycogen synthase (from bovine heart) [9014-56-6] M_r **60,000** [EC **2.4.1.11**]. Purify the synthase by precipitation in the presence of added glycogen with polyethylene glycol, chromatography on DEAE-Sephacel and high-speed centrifugation through a sucrose-containing buffer. [Dickey-Dunkirk & Killilea *Anal Biochem* **146** 199 1985, DOI: 10.1016/0003-2697(85)90416-6.]

Haemoglobin A (from normal human blood) [9008-02-0] M_r **~64,500**, amorphous. Purify it from blood using CM-32 cellulose column chromatography. [Matsukawa et al. *J Am Chem Soc* **107** 1108 1985, DOI: 10.1021/ja00291a004.] For the purification of the α and β chains see Hill et al. *Biochemical Preparations* **10** 55 1963.

Histones (from S4A mouse lymphoma). The purification of histones uses a macroprocess column, heptafluorobutyric acid as solubilising and ion-pairing agent and an acetonitrile gradient. [McCroskey et al. *Anal Biochem* **163** 427 1987, DOI: 10.1016/0003-2697(87)90244-2.]

Hyaluronidase [9001-54-1, 37326-33-3] M_r **43,000 (bovine testes), 89,000 (bacterial)** [EC **3.2.1.35**]. Hyaluronidase is purified by chromatography on DEAE-cellulose prior to use. [Distler & Jourdain *J Biol Chem* **248** 6772 1973, PMID: 4270451.]

D-Hydantoinase [dihydropyrimidinase, also called **5,6-dihydropyrimidine amidohydrolase, from microorganisms e.g. Pseudomonas, Bacillus, Agrobacterium** as well as from mammalian and human tissues) [9030-74-4] M **51,720 (monomer from amino acid sequence, usually dimer or tetramer)** [EC **3.5.2.2**], pI **~6.5**. This cytosolic enzyme hydrolyses dihydropyrimidines and hydantoins to N-carbamoylamino acids, and with the appropriate substrates are useful for preparing D-amino acids. The enzyme from a recently isolated species of *Agrobacterium* was purified to homogeneity and found to possess hydantoinase activity that was free from dihydropyrimidinase activity. It had an estimated subunit molecular weight of ~66,500 and a theoretical molecular weight of 265,000. The preferred substrates were 5-mono-substituted hydantoins with aromatic groups as shown from the K_m values. 5,5-Dimethylhydantoin and thio analogues of 5-*p*-hydroxyphenylhydantoins were competitive inhibitors. [Runser & Meyer *Eur J Biochem* **213** 1315 1993, DOI: 10.1111/j.1432-1033.1993.tb17883.x]

A commercially available hydantoinase preparation from *Vigna angularis* from aduki bean with a minimum activity of 300U/g is commercially available. These enzymes are generally inhibited by N-carbamoylamino acids (reaction products), 8-hydroxyquinoline, EDTA, Sn^{2+} and Zn^{2+} , but are activated by uracil, 2-thiouracil, Co^{2+} , Fe^{2+} , Mg^{2+} , Mn^{2+} and Ni^{2+} , with turnovers of ~27,000min⁻¹ for hydantoin. Their pH-range is 6–9.5, and the temperature range is ~4–60°. D-Hydantoinase genes have been cloned and expressed in *E coli* and the enzymes from several sources have been crystallised.

A **unit** of enzyme activity is defined as the amount that catalyses the formation of 1mmole of N-carbamoylglycine from hydantoin per minute at pH 9.0 and 40°. [Morin *Enz Microbiol Technol* **15** 208 1993, *Springer Handbook of Enzymes* D. Schonburg & I. Schonburg Eds (A. Chang co-Ed) Springer-Verlag, Berlin, Heidelberg.]

3-Hydroxy butyrate dehydrogenase (from *Rhodopseudomonas spheroides*) [9028-38-0] M_r **~85,000** [EC **1.1.1.30**], **amorphous**. Purify the dehydrogenase by two sequential chromatography steps on two triazine dye-Sepharose matrices. [Scawen et al. *Biochem J* **203** 699 1982, DOI: 10.1042/bj2030699.]

Interferons [α IFN, β IFN and γ IFN]. Interferons are a family of glycosylated proteins and are cytokines which are produced a few hours after cells have been infected with a virus. Interferons protect cells from viral infections and have antiviral activities at very low concentrations (~3 x 10⁻⁴M; less than 50 molecules are apparently sufficient to protect a single cell). Double-stranded RNAs are very efficient inducers of IFNs. There are three main types of IFNs. The α IFNs are synthesised in lymphocytes, and the β IFNs are formed in infected fibroblasts. The α and β families are fairly similar, consisting of *ca* 166 to 169 amino acids. Although γ IFNs are also small glycosylated proteins (*ca* 146 amino acids), they are different because they are not synthesised after viral infections but are produced by lymphocytes when stimulated by **mitogens** (agents that induced cell division).

Several of these IFNs of mouse and human lymphocytes and fibroblasts are available commercially and have been best prepared in quantity by recombinant DNA procedures because they are produced in very small amounts by the cells. The commercial materials do not generally require further purification for their intended purposes. [Pestkas 'Interferons and Interferon standards and general abbreviations' *Methods Enzymol*, Wiley & Sons, **119** 1986, ISBN 012182019X; Lengyel 'Biochemistry of interferons and their actions' *Ann Rev Biochem* **51** 251-282 1982, DOI: 10.1146/annurev.bi.51.070182.001343; De Maeyer & De Maeyer-Guignard, Interferons in *The Cytokine Handbook*, 3rd Edn, Thomson et al. Eds, pp. 491-516 1998 Academic Press, San Diego, ISBN 0126896623.]

Interleukin (from human source). Purify these using lyophilisation and desalting on a Bio-Rad P-6DC desalting gel, then two steps of HPLC, first with hydroxylapatite, followed by a TSK-125 size exclusion column. [Köck & Luger *J Chromatogr* **296** 293 1984, DOI: 10.1016/S0021-9673(01)96423-4.]

Interleukin-2 (recombinant human) [94218-72-1] M_r ~15,000, amorphous. Purify it by reverse phase HPLC. [Weir & Sparks *Biochem J* **245** 85 1987, DOI: 10.1042/bj2450085; Robb et al. *Proc Natl Acad Sci USA* **81** 6486 1984, PMID: 6333684.]

Interleukins (IL-1, IL-2 – IL18). Interleukins are cytokines which cause a variety of effects including stimulation of cell growth and proliferation of specific cells, e.g. stem cells, mast cells, activated T cells, colony stimulating factors etc., as well as stimulating other ILs, prostaglandins release etc. They are small glycosylated proteins (*ca* 15 kD, 130-180 amino acids produced from longer precursors) and are sometimes referred to by other abbreviations, e.g. IL-2 as TCGF (T cell growth factor), IL-3 as multi-CSF (multilineage colony stimulating factor, also as BPA, HCSF, MCSF and PSF). They are produced in very small amounts and are commercially made by recombinant DNA techniques in bacteria or Sf21 insect cells. Interleukins for human (h-IL), mouse (m-IL) and rat (r-IL) are available, and up to IL-18 are available commercially in such purity that they can be used directly without further refinement, particularly those that have been obtained by recombinant DNA procedures which are specific. As well as the interleukins, a variety of antibodies for specific IL reactions are available for research or IL identification. [Symons et al. *Lymphokines and Interferons, A Practical Approach*, Clemens et al. Eds, p.272 1987, IRL Press, Oxford, ISBN 1852210354, 1852210362, Thomson et al. Eds, *The Cytokine Handbook*, 3rd Edn, 1998, Academic Press, San Diego, ISBN 0126896623.]

Lactate dehydrogenase (from dogfish, Beef muscle) [9001-60-9] M_r 140,000 [EC 1.1.1.27]. A forty-fold purification of the dehydrogenase is effected by affinity chromatography using Sepharose 4B coupled to 8-(6-aminohexyl)amino-5'-AMP or -NAD⁺. [Lees et al. *Arch Biochem Biophys* **163** 561 1974, DOI: 10.1016/0003-9861(74)90515-3; Pesce et al. *J Biol Chem* **239** 1753 1964, PMID: 14213346.]

Lactoferrin (from human whey) [55599-62-7, Fe Salt] M_r ~90,000. This iron-binding protein is purified by direct adsorption on cellulose phosphate by batch extraction, then eluted by a stepped salt and pH gradient. The Fe bound protein forms red crystals with λ_{\max} at 465nm (pH 8.2). [Foley & Bates *Anal Biochem* **162** 296 1987, DOI: 10.1016/0003-2697(87)90040-6.]

Lectins (proteins and/or glycoproteins of non-immune origin that agglutinate cells, e.g. from seeds of *Robinia pseudoacacia*), M_r ~100,000. Lectins are purified by precipitation with (NH₄)₂SO₄ and dialysed, then chromatographed on DE-52, DEAE-cellulose anion-exchanger, hydroxylapatite and Sephacryl S-200. [Wantyghem et al. *Biochem J* **237** 483 1986, DOI: 10.1042/bj2370483.]

Lectins are a group of proteins that are classed as sugar-binding proteins or glycoproteins of non-immune origin and which agglutinate cells and/or precipitate glyco-conjugates. They are present in plants (seeds, roots, leaves or bark) and some invertebrates (snails, clams, crabs) and have M_r values of 10,000-400,000. They may contain Mn²⁺ and/or Ca²⁺. Mono- or oligo- saccharides of appropriate specificity inhibit lectins. Some lectins are specific to human blood groups and induce mitosis in lymphocytes. [Goldstein et al. *Nature* **285** 66 1980 DOI: 10.1038/285066b0.]

Lipoprotein lipase (from bovine skimmed milk) [9004-02-8] M_r ~34,000 and 63,000 (SDSPAGE), 96,900 (sedimentation and diffusion), 100,000-120,000 (gel filtration) [EC 3.1.1.34]. Purify the lipase by affinity chromatography on heparin-Sepharose. It has K_i 0.026mM for very low density lipoprotein. It is inhibited by 2-mercaptoethanol, Cys, Ca, Hg, Mg and Mn ions. Protamine sulfate, 1mg of bovine serum albumin/ml or in 50% glycerol at -70°, stabilises the lipase for several days. 60% loss of activity occurs at 0°/hour in the presence of 1% of bovine serum albumin. [Kohji et al. *Biochim Biophys Acta* **665** 504 1981, DOI: 10.1016/0005-2760(81)90264-2.]

Lipoproteins (from human plasma). Individual human plasma lipid peaks are removed from plasma by ultra-

centrifugation; then they are separated and purified by agarose-column chromatography. Fractions are characterised immunologically, chemically, electrophoretically and by electron microscopy. [Rudel et al. *Biochem J* **139** 89 1974, DOI: 10.1042/bj1390089.]

Lipoteichoic acids (from gram-positive bacteria) [56411-57-5]. These acids, which are present in bacterial cell walls, are extracted by hot phenol/water from disrupted cells. Nucleic acids are also extracted and are removed by treatment with nucleases. Nucleic resistant acids, proteins, polysaccharides and teichoic acids are separated from lipoteichoic acids by anion-exchange chromatography on DEAE-Sephacel or by hydrophobic interaction on octyl-Sepharose [Fischer et al. *Eur J Biochem* **133** 523 1983, DOI: 10.1111/j.1432-1033.1983.tb07495.x].

Lysozyme (Muramidase, N-acetylmuramyl hydrolase, N-acetylmuramide, peptidoglycan N-acetylmuramoyl hydrolase, Globulin G₁) [human 174883-18-2, from human neutrophils 9001-63-2, from human milk 12671-19-1, from chicken egg white 126050-88-3, chloride from chicken egg white 9066-59-5] **M 14,400 ±100, E_{280nm} 2.65 (c 1mg/ml) Isoelectric Point (IP) 10.5-11.0, [EC 3.2.1.17]**. Lysozymes from human and bird sources are 129 amino acid enzymes that contain four disulfide bonds. Hen lysozyme was isolated in quantity from chicken egg white, crystallised, and its X-ray crystalline structure was determined [Phillips *Proc Natl Acad Sci USA* **57** 484 1967, Symposium]. Lysozyme occurs in many tissues of invertebrate and vertebrate animals. It is found in milk, blood serum and various secretions (saliva, nasal mucus and tears). It also occurs in some moulds and in the latex of certain plants. Lysozyme from human milk was studied in some detail and is very similar to the hen enzyme [Jollés & Jollés *Helv Chim Acta* **52** 2671 1969, DOI: 10.1002/hlca.19690520848 and **54** 2668 1971, DOI: 10.1002/hlca.19710540830]. Lysozyme is a glycosidase which dissolves various bacterial cell walls [particularly Gram-positive bacteria (which have surface lipoproteins) to give spheroblasts, and various Gram-negative bacteria in the presence of EDTA in hypotonic solutions, or non-ionic detergents]. Bacterial cell walls contain 1,4-β-N-acetylglucosaminyl oligosaccharides which are cleaved by the enzyme at the glycosidic C—O bond between the 4th and 5th sugar residues from the non-reducing end of the chain. The mechanism and kinetics of this hydrolysis have been studied extensively [cf. Fersht *Enzyme Structure and Mechanism*, 2nd edn, W.H. Freeman & Co, Reading 1985, ISBN 0716716151]. Lysozyme was an extremely useful antibacterial in the pre-antibiotic era. It is a basic protein with 20-22 basic residues and only 3-4 acidic groups (see their isoelectric points); it forms soluble salts and is stable up to 55°. It is purified from egg white by chromatography through Amberlite IRC50 at pH 7.18 in 0.2M phosphate buffer followed by recrystallisation at pH 9.5 by adding NaCl to a concentration of 5%. Alternatively, ~1L of homogenised egg white in the absence of air (from 3 dozen eggs) is added to a 10% suspension of Bentonite [1302-78-9] (150ml, a native hydrated aluminium silicate) in 1% aqueous KCl, and stirred vigorously enough to avoid excessive foaming for ~5 minutes to give a smooth suspension. Separate the clay by centrifugation, wash it twice with 0.5M phosphate buffer (300ml each, pH 7.5) and three times with 5% aqueous pyridine (300ml each) while decanting from the clay and discarding inactive supernatants. From the combined washed clay, lysozyme is extracted out twice with 300ml of 5% aqueous pyridine (adjusted to pH5 with H₂SO₄ using a glass electrode). The combined extracts are dialysed against running tap H₂O (at 12-15°) until free from pyridine odour and dialysed further for 24 hours. Essentially pure amorphous lysozyme is obtained by lyophilising the dialysate below 25°. *Crystalline isoelectric lysozyme* is obtained from the amorphous powder by adding 0.5g of NaCl to a 5% aqueous solution of the enzyme (10ml), adjusting the pH to 9.5-10 (glass electrode) with aqueous NaOH and storing at 4°, whereby it crystallises out. *Crystalline lysozyme carbonate* is obtained from the amorphous powder by adding 0.5g of NaHCO₃ to a 5% aqueous solution of the enzyme (10ml), giving a final pH of 8.0-8.5 (glass electrode) with aqueous NaOH and allowing to stand at room temperature, whereby the carbonate crystallises out. *Crystalline lysozyme chloride* is obtained from a 5% solution of amorphous powder by adjusting the pH to 4.5 with hydrochloric acid (glass electrode) followed by solid NaCl to 5%, and storing at 4°, whereby the *chloride* crystallises out in 4 to 5 days. [Alderton & Fevold *J Biol Chem* **164** 1 1946, PMID: 20989461; Fevold & Alderton *Biochemical Preparations I* 67 1949.] Other salts include the *L-ascorbate* [119189-24-1], the *lactate* [72497-48-4], the *phosphate* [72497-47-3] and the *2,3-dihydroxypropyl dihydrogen phosphate* [119189-23-0].

Dialysis and recrystallisation are simple and yield enzyme of high purity. Several forms of crystals are obtained depending on the pH of the crystallising solution. The activity of lysozyme is measured by the rate of decrease of

turbidity (at 570nm) as hydrolysis of acetone-dried cell walls of the Gram-positive bacterium *Micrococcus lysodeiktitus* (as substrate) occurs on addition of the enzyme [Hirs *Methods Enzymol* **I** 113 1968, DOI: 10.1016/0076-6879(55)01017-3].

At high concentrations or in the presence of albumin, lysozyme can be lyophilised or desalted without loss of activity. Freezing and thawing does not inactivate it, and Na^+ , Mg^{2+} , Ag^+ , Ca^{2+} , and Cu^{2+} exert an activating effect. It can be stored for several weeks at -20° without inactivation, and in the presence of 0.1% of bovine serum albumin it can be stored for months at -20° without inactivation. [*Springer Handbook of Enzymes* 2nd edn. Schonburg & Schonburg eds (A. Chang co-ed) Springer-Verlag Berlin, Heidelberg, **Volume 12** 2003 ISBN 3-540-00519-6.]

Agarose-bound lysozyme from hen egg (5,000-10,000 Units/g of solid), a lysozyme Biotin-caproyl solution (20,000 U/mg) and a carboxymethylated-maleylated lysozyme (Lysozyme-RCM, reduced form), and lysozyme of up to 100,000 U/mg protein are commercially available.

The chemical synthesis and high resolution X-ray structure of **human lysozyme** has been described by S.B.H. Kent and coworkers [Durek, Torbeev & Kent *Proc Natl Acad Sci* **104**(12) 4846 2007, DOI: 10.1073/pnas.0610630104, PMID: 17360367

A *T4 bacteriophage lysozyme* (from phage grown on *E.coli* B¹) [12585-29-4] was extracted and freed from DNA, cell debris, intact cells and acidic proteins by precipitation with Rivanol (6,9-diamino-2-ethoxyacridine *dl*-lactate). The filtrate was purified by concentration through an Amberlite IRC50 column which was thoroughly washed with 0.1M phosphate buffer pH 6.5 containing 10^{-3}M MgSO_4 and eluted with the phosphate buffer at pH 5.8 containing a 0 to 0.5M NaCl gradient. This was repeated twice, followed by gel filtration through Sephadex G-75 in pH 5.8 phosphate buffer and eluted with 0.5M NaCl. The eluate was dialysed against H_2O and lyophilised to give a 1500-fold purification with 40% recovery. Like the hen lysozyme it is a basic protein but with 164 amino acid residues, and it is unstable $>40^\circ$. [Tsugita et al. *J Biol Chem* **243** 391 1968, PMID: 4865643; Inouye et al. *J Biol Chem* **245** 3439, PMID: 5470815; 3479, PMID: 5470817 1970, Matthews *Biochim Biophys Acta* **405** 442 1975, DOI: 10.1016/0005-2795(75)90109-9].

Metallothionein (from rabbit liver) [9038-94-2]. Purify the cysteine-rich protein by precipitation to give Zn- and Cd-containing protein fractions and running it on a Sephadex G-75 column, then isoelectric focusing to give two protein peaks. It is made up of several isoforms with Mr 500 to 14000 Da. [Nordberg et al. *Biochem J* **126** 491 1972, DOI: 10.1042/bj1260491; Comeau et al. *Prep Biochem* **22** 151 1992, DOI: 10.1080/10826069208021365].

Myoglobin (from sperm whale muscle) [9047-17-0] **M_r ~17,000**. Myoglobin is purified by CM-cellulose chromatography and Sephadex G-50 followed by chromatography on Amberlite IRC-50 Type III or BioRex 70 (<400mesh). The crystalline product, as a paste in saturated $(\text{NH}_4)_2\text{SO}_4$ at pH 6.5-7.0, may be stored at 4° for at least 4 years unchanged, but must not be kept in a freezer. [Andres & Atassi *Biochemistry* **12** 942 1980, DOI: 10.1021/bi00729a024; Edmundson *Biochemical Preparations* **12** 41 1968.]

5'-Nucleotidase (from Electric ray, *Torpedo sp*) [9027-73-0] [EC 3.1.3.5], **amorphous**. Purify it by dissolving it in Triton X-100 and deoxycholate, and by affinity chromatography on concanavalin A-Sepharose and AMP-Sepharose [Grondal & Zimmerman *Biochem J* **245** 805 1987, DOI: 10.1042/bj2450805].

Orsomucoid (glycoprotein α_1 acid, from human plasma) [66455-27-4] **M_r 42000-44000, amorphous**. Purify the glycoprotein α_1 acid by passage through a carboxymethyl cellulose column, then through a Sephadex G-25 column, and lyophilise the protein fraction. [Aronson et al. *J Biol Chem* **243** 4564 1968, PMID: 5684010.]

Papain [9001-73-4] **M_r ~21,000** [EC 3.4.22.2], **amorphous**. A suspension of 50g of papain (freshly ground in a mortar) in 200ml of cold water is stirred at 4° for 4 hours, then filtered through a Whatman No 1 filter paper. The clear yellow filtrate is cooled in an ice-bath while a rapid stream of H_2S is passed through it for 3 hours, and

the suspension is centrifuged at 2000rpm for 20 minutes. Sufficient cold MeOH is added slowly with stirring to the supernatant to give a final MeOH concentration of 70 vol%. The precipitate, collected by centrifugation for 20 minutes at 2000rpm, is then dissolved in 200ml of cold water, the solution is saturated with H₂S, centrifuged, and the enzyme is again precipitated with MeOH. The process is repeated four times. [Bennett & Niemann *J Am Chem Soc* **72** 1798 1950, DOI: 10.1021/ja01160a109.] Papain has also been purified by affinity chromatography on a column of GlyGlyTyrArg-agarose [Stewart et al. *J Am Chem Soc* **108** 3480 1986, DOI: 10.1021/ja00272a052].

Pepsin [9001-75-6] **M_r 31,500(human), 6000(hog), [EC 3.4.23.1]**. Pepsin is re-chromatographed on a column of Amberlite CG-50 using a pH gradient prior to use. Crystallise it from EtOH. [Richmond et al. *Biochim Biophys Acta* **29** 453 1958, DOI: 10.1016/0006-3002(58)90220-8; Huang & Tang, *J Biol Chem* **244** 1085 1969, PMID: 4886180; **245** 2189 1970, PMID: 4909888.]

Pertussis toxin (from *Bordetella pertussis*) [70323-44-3] **M_r 117,000**. Purify the toxin by stepwise elution from 3 columns comprising Blue Sepharose, Phenyl Sepharose and hydroxylapatite, and the purity is checked by SDS-PAGE. [Svoboda et al. *Anal Biochem* **159** 402 1986, DOI: 10.1016/0003-2697(86)90360-X; Tamura et al *Biochemistry* **21** 5516 1982, DOI: 10.1021/bi00265a021.]

Phosphatase alkaline (alkaline phosphatase) [9001-78-9] **M_r ~40,000 (bovine liver), ~140,000 (bovine intestinal mucosa), 80,000 (*E.coli*), [EC 3.1.3.1]**. The *E.coli* supernatant in sucrose (20%, 33mM) in Tris-HCl pH 8.0 is purified through a DEAE-cellulose column and recrystallised. To the column eluates in 0.125M NaCl is added MgCl₂ (to 0.01M) and brought to 50% saturation in (NH₄)₂SO₄ by adding the solid (0.20g/ml). The mixture is centrifuged to remove bubbles and is adjusted to pH 8.0 (with 2N NaOH). Saturated (NH₄)₂SO₄ at pH 8.0 is added dropwise until the solution becomes faintly turbid (~61% saturation). It is set aside at ~25° for 1 hour (turbidity will increase). The mixture is placed in an ice bath for several minutes when turbidity disappears and a clear solution is obtained. It is then placed in a large ice bath at 0° (~5L) and allowed to warm slowly to room temperature in a dark room whereby crystals are formed appearing as a silky sheen. The crystals are collected by centrifugation at 25° if necessary. The crystalline solutions are stable at ~25° for many months. They can be stored at 0°, but are unstable when frozen. Cysteine at 10⁻³M and thioglycolic acid at 10⁻⁴M are inhibitory. This is reversed on addition of Zn²⁺ ions. Many organic phosphates are good substrates for this phosphatase. [Malamy & Horecker *Methods Enzymol* **9** 639 1966, DOI: 10.1016/0076-6879(66)09129-8; Torriani et al. *Methods Enzymol* **12b** 212 1968, DOI: 10.1016/0076-6879(67)12135-6; Engström *Biochim Biophys Acta* **92** 71 1964, DOI: 10.1016/0926-6569(64)90270-6.]

Alkaline phosphatase from rat osteosarcoma has been purified by Me₂CO precipitation and chromatography on DEAE-cellulose, Sephacryl S-200, and hydroxylapatite. [Nair et al. *Arch Biochem Biophys* **254** 18 1987, DOI: 10.1016/0003-9861(87)90076-2.]

Phosphoproteins (various). These are purified by adsorbing onto an iminodiacetic acid substituted agarose column to which are bound ferric ions. This chelate complex acts as a selective immobilised metal affinity adsorbent for phosphoproteins. [Muszynska et al. *Biochemistry* **25** 6850 1986, DOI: 10.1021/bi00370a018.]

5'-Phosphoribosyl pyrophosphate synthetase (from human erythrocytes, or pigeon or chicken liver) [9015-83-2] **M_r 60,000 [EC 2.7.6.1]**. It is purified 5100-fold by elution from DEAE-cellulose, fractionation with (NH₄)₂SO₄, filtration on Sepharose 4B and ultrafiltration. [Fox & Kelley *J Biol Chem* **246** 5739 1971, PMID: 4328836; Flaks *Methods Enzymol* **6** 158 1963, DOI: 10.1016/0076-6879(63)06158-9; Kornberg et al. *J Biol Chem* **215** 389 1955, PMID: 14392173.]

Pituitary Growth Factor (from human pituitary gland) [336096-71-0]. It is purified by heparin and copper affinity chromatography, followed by chromatography on carboxymethyl cellulose (Whatman 52). [Rowe et al. *Biochemistry* **25** 6421 1986, DOI: 10.1021/bi00369a012.]

Plasmids. These are circular lengths of *double-stranded* DNA which invade bacteria or other cells, e.g. insect cells, yeast cells, and have sequences which are necessary for their replication using enzymes and other ingredients, e.g. nucleotides, present in the cells. They contain engineered, or already have, genes which

produce enzymes that provide the cells with specific antibiotic resistance and are thus useful for selecting bacteria containing specific plasmids. Plasmids have been extremely useful in molecular biology since they can be very easily identified (from their size or the sizes of the DNA fragments derived from their restriction enzyme digests) and can be readily engineered *in vitro* (outside the cells). Genes coding for specific enzymes or other functional proteins can be inserted into these plasmids which have DNA sequences that allow the expression of large quantities of bacterial or non-bacterial (e.g. human) proteins. They have also been engineered in such a way as to produce ‘fusion proteins’ [in which the desired protein is fused with a specific ‘reporter, marker or carrier protein’ which will facilitate the isolation of the desired protein e.g. by binding strongly to a nickel support], and then the desired protein can be cleaved from the eluted fusion protein and obtained in very pure form. A large number of plasmids with a variety of sequences for specific purposes are commercially available in very pure form. They can be used to infect cells and can be isolated and purified from cell extracts in large amounts using a number of available procedures. These procedures generally involve lysis of the cells (e.g. with alkaline sodium dodecylsulfate, SDS), separation from nuclear DNA, precipitation of plasmid DNA from the cell debris, adsorbing it on columns which specifically bind DNA, and then eluting the DNA from the column (e.g. with specific Tris buffers as recommended by the suppliers) and precipitating it (e.g. with Tris buffer in 70% EtOH at -70°C). The purity is checked on agarose gels (containing ethidium bromide to visualise the DNA) by electrophoresis. A large number of plasmids are now commercially available (see Clontech GmbH, <http://www.clontech.com>, Invitrogen <http://www.invitrogen.com>, among other suppliers) and used as vectors for bacterial, mammalian, yeast and insect cells, and for baculovirus expression.

Protamine kinase (from rainbow trout testes) [37278-10-7] [EC 2.7.1.70]. Partial purification of the kinase is achieved by chromatography through hydroxylapatite followed by biospecific chromatography on nucleotide coupled Sepharose 4B [the nucleotide is 8-(6-aminoethyl)amine coupled cyclic-AMP]. It requires Mg ions specifically, but is inactivated by Ca, Mn, Cu, Zn ions and thiol compounds. [Jergil & Dixon *J Biol Chem* **245** 425 1970, PMID: 4312674; Jergil et al. *Biochem J* **139** 441 1974, DOI: 10.1042/bj1390441.]

Protamine sulfate (from herring sperm) [9007-31-2] [α] D^{22} -85.5 (saturated H₂O), pK²⁵ 7.4-8.0. Protamine sulfate is a strongly basic protein (white powder, see pK) used to precipitate nucleic acids from crude protein extracts. It dissolves to the extent of 1.25% in H₂O. It is freely soluble in hot H₂O but separates as an oil on cooling. It has been purified by chromatography on an IRA-400 ion-exchange resin in the SO₄²⁻ form and is washed with dilute H₂SO₄. Eluates are freeze-dried under high vacuum below 20°. This method is used to convert proteamine and protamine hydrochloride to the sulfate. [UV: Rasmussen *Hoppe Seyler's Z Physiol Chem* **224** 97 1934, Ando & Sawada *J Biochem (Tokyo)* **49** 252 1961, PMID: 13683326; Felix & Hashimoto *Hoppe Seyler's Z Physiol Chem* **330** 205 1963.] It reverses the anticoagulant effects of heparin by binding to it, and enhances lipid-mediated gene transfer [Sorgi et al. *Gene therapy* **4**(9) 961 1997, DOI: 10.1038/sj.gt.3300484].

Protease nexin (from cultured human fibroblasts) [148263-58-5]. This serine protease inhibitor is purified simply in two steps first by affinity binding to dextran sulfate-Sepharose, then purified to homogeneity involving a combination of DEAE-Sepharose in-line with dextran sulfate-Sepharose to simultaneously purify and concentrate the protein [Farrell et al. *Biochem J* **237** 907 1986, DOI: 10.1042/bj2370907]. [See also Farrell and Cunningham *Biochem J* **245** 543 1987, DOI: 10.1042/bj2450543; and for properties and purification see Scott et al. *J Biol Chem* **260** 7029 1985, PMID: 3858278]

Proteins. These are usually naturally occurring (or deliberately synthesised in microorganisms, e.g. bacteria, insect cells, or animal tissues), and are composed of a large number of α -S (L)amino acid residues (except for L-cysteine which has the *R* absolute configuration), selected from the 20 or so natural amino acids, in specific sequences and in which the α -amino group forms an amide (peptide) bond with the α -carboxyl group of the neighbouring amino acid. The number of residues is usually upwards of 100. Proteins with less than 100 amino acids are better referred to as **polypeptides**. Aqueous soluble proteins generally fold into ball-like structures mainly with hydrophilic residues on the outside of the ‘balls’ and hydrophobic residues on the inside. Proteins can exist singly or can form dimers, trimers, tetramers etc., consisting of similar or different protein subunits. [See ‘Introduction’ in this Chapter]. They are produced by cells for a large variety of functions, e.g. enzymology, reaction mediation as in regulation of DNA synthesis or chaperonins for aiding protein folding,

formation of pores in membranes for transport of ions or organic molecules, or for intra or intercellular signalling etc. The purity of proteins can be checked in denaturing (SDS, sodium dodecylsulfate) or non-denaturing polyacrylamide gels using electrophoresis (PAGE), and staining appropriately (e.g. with Coomassie Blue, followed by silver staining for higher sensitivity). If the protein is partly impure, then it should be purified further according to the specific literature procedures for the individual protein (see specific proteins in the *Methods Enzymol*, Wiley series).

Proteoglycans (from cultured human muscle cells). These are separated by ion-exchange HPLC using a Bio-gel TSK-DEAE 5-PW analytical column. [Harper et al. *Anal Biochem* **159** 150 1986, DOI: 10.1016/0003-2697(86)90320-9.]

Prothrombin (Factor II, from equine blood plasma) [9001-26-7] M_r 72,000. Prothrombin is purified by two absorptions on a barium citrate adsorbent, followed by decomposition of the adsorbents with a weak carboxylic cation-exchanger (Amberlite IRF-97), isoelectric precipitation (pH 4.7-4.9) and further purification by chromatography on Sephadex G-200 or IRC-50. Finally it is recrystallised from a 1% solution adjusted to pH 6.0-7.0 and partial lyophilisation to *ca* 1/5 to 1/10th volume and set aside at 2-5° to crystallise. Occasionally seeding is required. [Miller *Biochemical Preparations* **13** 49 1971.] For ‘Has the time arrived to replace the quick prothrombin time test for monitoring oral anticoagulant therapy?’ see Jackson & Esnouf [*Clin Chem* **51**(3) 483-5 2005, DOI: 10.1373/clinchem.2004.045393, PMID: 15738512].

Prymnesin (toxic protein from phytoflagellate *Prymnesium parvum*) [11025-94-8]. Prymnesin is purified by column chromatography, differential dissolution and precipitation in solvent mixtures and differential partition between diphasic mixtures. The product has at least 6 components as observed by TLC. [Ulitzur & Shilo *Biochim Biophys Acta* **201** 350 1970, DOI: 10.1016/0304-4165(70)90310-7.]

Pyruvate kinase isoenzymes (from *Salmonella typhimurium*) [9001-59-6] M_r 64,000 [EC 2.7.1.40], amorphous. These are purified by $(\text{NH}_4)_2\text{SO}_4$ fractionation and gel filtration, ion-exchange and affinity chromatography. [Garcia-Olalla & Garrido-Pertierra *Biochem J* **241** 573 1987, DOI: 10.1042/bj241057.]

Renal dipeptidase (from porcine kidney cortex) [9031-96-3] M_r 47,000 [EC 3.4.13.11]. The dipeptidase is purified 700-fold by homogenising the tissue, extracting with Triton X-100, elimination of insoluble material, and ion-exchange, size exclusion and affinity chromatography. [Hitchcock et al. *Anal Biochem* **163** 219 1987, DOI: 10.1016/0003-2697(87)90116-3.]

Restriction enzymes (endonucleases). These are enzymes which cleave double-stranded DNA (linear or circular) at specific nucleotide sequences within the DNA strands which are then used for cloning (by ligating bits of DNA sequences together) or for identifying particular DNA materials, e.g. plasmids, genes etc. A very large number of restriction enzymes are now available commercially and are extensively used in molecular biology. They are highly specific for particular nucleotide arrangements and are sensitive to the reaction conditions, e.g. composition of the medium, pH, salt concentration, temperature etc, which have to be strictly adhered to.

Endonucleases are essentially of two types. One type cleaves the DNA producing *blunt ends*, i.e. the nucleotides are paired at each end. The second type cleaves the DNA leaving *hanging ends*, i.e. the nucleotides at the ends of each strand (usually four nucleotides) of the double strands protrude ahead of the paired nucleotides. The latter enzymes particularly allow ligation of the cleaved sequence to occur in a desired direction.

The enzymes do not require further purification, and the reaction conditions are also provided by the suppliers from which the necessary reaction media can also be purchased (see commercial catalogues).

Reverse transcriptase (from avian or murine RNA tumour viruses) [9068-38-6] M_r ~170,000 [EC 2.7.7.49]. This enzyme produces the complementary DNA from the RNA (as template). These are purified by solubilising the virus with non-ionic detergents. Lysed virions are adsorbed on DEAE-cellulose or DEAE-Sephadex columns, and the enzymes are eluted with a salt gradient, then chromatographed on a phosphocellulose column and fractions with enzyme activity are eluted in a salt gradient. They are also purified

from other viral proteins by affinity chromatography on a pyran-Sepharose column. [Verma *Biochim Biophys Acta* **473** 1 1977, DOI: 10.1016/0304-419X(77)90005-1; Smith *Methods Enzymol* **65** 560 1980, DOI: 10.1016/S0076-6879(80)65060-5; see commercial catalogues for other transcriptases.]

Ribonuclease (from human plasma) [9001-99-4] M_r ~13,700 [EC 3.1.27.5], **amorphous**. Purify it by $(\text{NH}_4)_2\text{SO}_4$ fractionation, followed by PC cellulose chromatography and affinity chromatography (using Sepharose 4B to which $(\text{G})_n$ is covalently bonded). [Schmukler et al. *J Biol Chem* **250** 2206 1975, PMID: 234961.]

Ribozymes. These are ribonucleic acids which act like protein enzyme in catalysing the making and breaking of peptide bonds as well as catalysing reactions and cleavage of DNA and RNA molecules. The short RNAs are being intensively studied (see RNAi below). MicroRNAs (miRNA, see also below) are ubiquitous and are also genetically produced. They are involved in numerous reactions from splicing RNA, e.g. mRNA (messenger RNA) to controlling transcription of DNA to RNA, and translation of RNA to protein. Each miRNA is capable of being involved in a small number to hundreds of interactions with nucleic acids and with proteins.

Ricin (toxin from Castor bean *Ricinus communis*) [A chain 96638-28-7, B chain 96638-29-8] M_r ~30,000, **amorphous**. Crude ricin, obtained by aqueous extraction and $(\text{NH}_4)_2\text{SO}_4$ precipitation, is chromatographed on a galactosyl-Sepharose column with sequential elution of pure ricin. The second peak is due to ricin agglutinin. [Simmons & Russell *Anal Biochem* **146** 206 1985, DOI: 10.1016/0003-2697(85)90417-8.] It is an inhibitor of protein synthesis. **EXTREMELY DANGEROUS, USE EXTREME CARE [instructions accompany product].**

RNA (ribonucleic acids). Ribonucleic acids are like DNA except that the 2'-deoxy-D-ribose moiety is replaced by a D-ribose moiety and the fourth nucleotide thymidylic acid (T) is replaced by uridylic acid (U). RNA does not generally form complete duplex molecules like DNA, i.e. it is generally monomeric, except in certain viruses. The two main classes of RNA are **messenger-RNA** (mRNA) and **transfer-RNA** (tRNA). Pre-mRNAs are transcribed from the DNA genes, and non-coding segments (the introns) are spliced out to give the mRNAs which code for specific proteins. There are many different tRNAs, at least one of which is linked to a specific α -amino acid that binds to the mRNA *via* the ribosome (a set of proteins) to the RNA triplets (three nucleotides) which code for the particular α -amino acids. An enzyme then joins the α -amino acids of two adjacent tRNA- α -amino acid ribosome complexes bound to the mRNA to form a peptide bond. Thus peptide bonds and consequently polypeptides and proteins coded by the DNAs, *via* the respective mRNA, are produced. Martin et al. [*Biochem J* **89** 327 1963, DOI: 10.1042/bj0890327] purified RNA by dissolving it (5g) in 90ml of 0.1mM EDTA, then homogenised this with 90ml of 90% (w/v) phenol in water using a Teflon pestle. The suspension was stirred vigorously for 1 hour at ~20°, then centrifuged for 1 hour at 0° at 25000rpm. The lower (phenol) layer was extracted four times with 0.1mM EDTA, and the aqueous layers were combined, then made 2% (w/v) with respect to AcOK and 70% (v/v) with respect to EtOH. After standing overnight at -20°, the precipitate was centrifuged down, dissolved in 50ml of 0.1mM EDTA, made 0.3M in NaCl and kept for 3 days at 0°. The purified RNA was then centrifuged down at 10000xg for 30 minutes, dissolved in 100ml of 0.1mM EDTA, dialysed at 4° against water, and freeze-dried. It was stored at -20° in a desiccator. Michelson [*J Chem Soc* 1371 1959, DOI: 10.1039/JR9590001371] dissolved 10g of RNA in water, added 2M ammonia to adjust the pH to 7, then dialysed in *Visking* tubing against five volumes of water for 24 hours. The process was repeated three times; then the material after dialysis was treated with 2M HCl and EtOH to precipitate the RNA which was collected, washed with EtOH, ether and dried [see commercial catalogues for further examples]. RNAs are now routinely sequenced.

RNAi (interfering RNA). RNAi technology has exploded during the past few years and has become invaluable for exploring gene function, interaction and control. It has considerable potential in therapeutics. This came about from the discovery that a double-stranded RNA (**dsRNA**) complementary to the coding sequence of a muscle protein 'turned off' the gene instead of enhancing the effect. It seemed that the dsRNA activated the cellular machinery to degrade the target mRNA. This led to the development and the use of interfering RNA (RNAi) to 'silence' specific genes. This silencing of genes comes in various different forms. It can be brought about by **short interfering RNA (siRNA)**, **micro RNA (miRNA)** and **short-hairpin RNA (shRNA)**. RNA can

be notoriously unstable in biological systems, e.g. its half-life in serum is ~6 minutes, but recently methods have been devised (using nanoparticle delivery systems) where they are more stable and can be used to target specific cancer cells.

About 50% of the human genome is apparently transcribed into mRNA and translated into protein or used to produce small RNA fragments. miRNAs are small molecules of about 23 or less ribonucleotides which are cleaved from larger fragments of RNA transcripts encoded from within genes, or large tracts of DNA between the genes. miRNAs are a special group of non-coding RNAs which work by blocking (silencing) gene expression. More than 400 have been discovered in the human genome so far and some appear to regulate many genes, as in embryonic development, e.g. cell growth, apoptosis. A cancer-causing miRNA was identified and called *onco-miRNA*. miRNA expression in particular cancers can now be quantified.

Single stranded RNAs consisting of 100s of bases fold back on themselves and undergo complementary base-pairing to form double-stranded 'short-hairpin' RNA (shRNA). The loop of the hair-pin is excised by endonucleases, e.g. *Dicer*, to produce multiple functional miRNAs.

Many of the above RNAs for specific purposes are available commercially from several suppliers in high purity because they are synthetic. These firms will also synthesise particular RNAi's to suit one's requirements.

Subtilisin (from *Bacillus subtilis*) [9014-01-1] M_r 27,000 (sedimentation equilibrium) [EC 3.4.21.62].

This alkaline protease is purified 211-fold by affinity chromatography using 4-(4-aminophenylazo)phenylarsonic acid complex to activated CH-Sepharose 4B. It is inhibited by 2-phenylethane boronic acid, PMSF, 3,4-dichloroisocoumarin, acetone and benzamide. [Chandrasekaran & Dhar *Anal Biochem* **150** 141 1985, DOI: 10.1016/0003-2697(85)90452-X; Schomburg & Schomburg *Springer Handbook of Enzymes* 2nd Edn vol 7 p 286 2002.]

Synexin (from bovine liver) M 47,000 Da. This Ca binding protein is purified by $(\text{NH}_4)_2\text{SO}_4$ precipitation, then by a specific pH step elution from a chromatofocusing medium in the absence of ampholytes. The pI is 7.5. [Scott et al. *Anal Biochem* **149** 163 1985, DOI: 10.1016/0003-2697(85)90489-0.]

Thrombin (from bovine blood plasma) [9002-04-4] M_r 32,600 [EC 3.4.4.13]. Thrombin is purified by chromatography on a DEAE-cellulose column, while eluting with 0.1M NaCl, pH 7.0, followed by chromatography on Sephadex G-200. The final preparation is free from plasminogen and plasmin. [Yin & Wessler *J Biol Chem* **243** 112 1968, PMID: 5635939.] Thrombin from bovine blood is also purified by chromatography using *p*-chlorobenzylamino- ϵ -aminocaproyl agarose, and gel filtration through Sephadex G-25. [Thompson & Davie *Biochim Biophys Acta* **250** 210 1971, DOI: 10.1016/0005-2744(71)90136-7.] Thrombin from various species was purified initially by precipitation of impurities with **Rivanol** (see [6402-23-9]). [Miller *Nature* **184** 450 1959 DOI: 10.1038/184450a0.]

Tissue inhibitor of metalloproteins (TIMP, from human blood plasma), M_r ~30,000. These are purified by an [anti-human amniotic fluid-TIMP]-Sepharose immuno-affinity column and eluted with 50mM glycine/HCl pH 3.0 buffer that is 0.5M in NaCl, and followed by gel filtration [Cawston et al. *Biochem J* **238** 677 1986, DOI: 10.1042/bj2380677].

Transferrin (from human or bovine serum) [11096-37-0] M_r ~80,000. This transferrin is purified by affinity chromatography on phenyl-boronate agarose followed by DEAE-Sephacel chromatography. The product is free from haemopexin. [Cook et al. *Anal Biochem* **149** 349 1985, DOI: 10.1016/0003-2697(85)90581-0; Aisen & Listowsky *Ann Rev Biochem* **49** 357 1980, DOI: 10.1146/annurev.bi.49.070180.002041.]

Trehalase (from kidney cortex, α,α -trehalose glycohydrolase) [9025-52-9] M_r ~80,000 [EC 3.2.1.28]. This trehalase is purified by solubilising in Triton X-100 and sodium deoxycholate, and submitting to gel filtration, ion-exchange chromatography, conA-Sepharose chromatography, phenyl-Sepharose CL-4B hydrophobic interaction chromatography, Tris-Sepharose 6B affinity and hydrolyapatite chromatography. Activity is increased 3000-fold. [Yoneyama *Arch Biochem Biophys* **255** 168 1987, DOI: 10.1016/0003-9861(87)90307-9.]

T4-RNA ligase (from bacteriophage-infected *E.coli*) M_r 43,500 [EC 6.5.1.3 for RNA lyase]. This ligase is purified by differential centrifugation and separation on a Sephadex A-25 column, then through hydroxylapatite and DEAE-glycerol using Aff-Gel Blue to remove DNAase activity. (Greater than 90% of the protein in the enzyme preparation migrated as a single band on gradient polyacrylamide gels containing SDS during electrophoresis.) [McCoy et al. *Biochim Biophys Acta* **562** 149 1979 DOI: 10.1016/0005-2787(79)90134-5.]

Tyrosine kinases (PTKs) are a large group of *transferase* enzymes e.g. [EC 2.7.10.X] that transfer a phosphate group from ATP to a protein in living cells, which activate or deactivate cellular functions. As the name implies, the phosphate group is attached to the phenolic oxygen of a tyrosine residue in the protein. Cell signaling (transferring messages within the cell— *signal transduction*) is an important mechanism for communicating information from one part of the cell to another. It is carried out by phosphorylating proteins with tyrosine kinases. Other kinases, such as serine and threonine kinases, function in an analogous manner. There are two main families of tyrosine kinases: (A) the *transmembrane kinases (RTKs)* which are anchors for membrane receptors and function by signaling (transducing) for outside of the cell to the interior of the cell thus triggering a cascade of events which may ultimately reach the nucleus and effecting gene expression. (B) *cytoplasmic-non-receptor* tyrosine kinases which function (traffic) within various parts of the cell.

[Cox & Nelson *Lehninger: Principles of Biochemistry* (5th ed.). W H Freeman & Co 2008, ISBN 1-4292-2416-9; Dengjel et al. 'Receptor tyrosine kinase signaling: a view from quantitative proteomics' *Mol Biosyst* **5** 1112 2009, DOI: 10.1039/b909534a, PMID: 19756300; Hanks et al. 'The protein kinase family: conserved features and deduced phylogeny of the catalytic domains' *Science* **241** (4861) 42 1988, DOI: 10.1126/science.3291115, PMID: 3291115.]

Ubiquinol-cytochrome c reductase (from beef heart mitochondria) [9027-03-6] [EC 1.10.2.2]. This reductase is purified by solubilising the crude enzyme with Triton X-100, followed by hydroxylapatite and gel chromatography. The minimum unit contains nine polypeptide subunits of M_r 6000 - 49000 kD. [Engel et al. *Biochim Biophys Acta/Bioenergetics* **592** 211 1980, DOI: 10.1016/0005-2728(80)90182-6.]

Uridine 5'-diphosphoglucose pyrophosphorylase (from rabbit skeletal muscle) [9029-22-6] M_r 350,000, [EC 2.7.7.9]. The pyrophosphorylase is purified by two hydrophobic chromatographic steps and gel filtration. [Bergamini et al. *Anal Biochem* **143** 35 1984, DOI: 10.1016/0003-2697(84)90554-2.] It is also purified from calf liver by $(\text{NH}_4)_2\text{SO}_4$ (40-58%) precipitation, $\text{Ca}_3(\text{PO}_4)_2$ gel filtration, DEAE-cellulose chromatography and recrystallisation [by dialysis against concentrations of $(\text{NH}_4)_2\text{SO}_4$ (from 10%) in 0.02M TEA, at 2.5% increments, until 20% $(\text{NH}_4)_2\text{SO}_4$ when it crystallises out]. [Hansen et al. *Methods Enzymol* **8** 248 1966, DOI: 10.1016/0076-6879(66)08042-X.]

Urokinase (from human urine) [9039-53-6] M_r 53,000 [EC 3.4.21.31]. Crystallisation of this enzyme is induced at pH 5.0 to 5.3 (4°) by careful addition of NaCl with gentle stirring until the solution becomes turbid (silky sheen). The NaCl concentration is increased gradually (over several days) until 98% of saturation is achieved whereby urokinase crystallises out as colourless thin brittle plates. It can be similarly recrystallised to maximum specific activity [104K CTA units/mg of protein (Sherry et al. *J Lab Clin Med* **64** 145 1964, PMID: 14192559)]. [Lesuk et al. *Science* **147** 880 1965, DOI: 10.1126/science.147.3660.880; NMR: Bogusky et al. *Biochemistry* **28** 6728 1989, DOI: 10.1021/bi00442a028.] It is a plasminogen activator [Gold et al. *Biochem J* **262** 529 1989, DOI: 10.1042/bj2620529; de Bock & Wang *Med Res Rev* **24**(1) 13 2004, DOI: 10.1002/med.10054].

Xylanase (from *Streptomyces lividans*) [37278-89-0] M_r 43,000 [EC 3.2.1.8]. This xylanase is purified by anion-exchange chromatography on an Accell QMA column and finally by HPLC using a ProteinPak DEAE 5PW anion-exchange column. Solutions are stored frozen at -70°. [Morosoli et al. *Biochem J* **239** 587 1986, DOI: 10.1042/bj2390587; Wong et al. *Microbiol Rev* **52** 305 1988, PMID: 3141761.]

CARBOHYDRATES

This section includes natural and synthetic carbohydrates and glycosides.

Z-O-(2-Acetamido-2-deoxy-D-glycopyranosylideneamino)-N-phenylcarbamate

(PUGNAC) $C_{15}H_{19}N_3O_7$, [132063-05-9] **M 335.3, m 171-174° (dec), 174-180°(dec), $[\alpha]_D^{20} +67.5$ (c 0.2, MeOH).** Purify PUGNAC by flash chromatography (silica gel and elute with EtOAc/hexane 3:2), evaporate, and the foam is recrystallised from EtOAc/MeOH. The purity is checked by TLC on Merck SiO₂ gel 60 F₂₅₄ and the spots are detected by spraying with 0.025M I₂ in 10% aqueous H₂SO₄ and heating at 200°. It has R_F 0.21. The acetate is hydrolysed with NH₃/MeOH. [Beer & Vasella *Helv Chim Acta* **68** 2254 1985, DOI: 10.1002/hlca.19850680821; Beer et al. *Helv Chim Acta* **73** 1918 1990, DOI: 10.1002/hlca.19900730714.] It is an *in vitro* and *in vivo* inhibitor of the acetylglucosaminidase enzyme which removes *O*-linked *N*-acetylglucosamine from *O*-linked glycosylated proteins.

Acetobromo-α-D-galactose [3068-32-4] $C_{14}H_{19}BrO_9$, **M 411.2, m 87°, $[\alpha]_{546}^{20} +255$, $[\alpha]_D^{20} +210$ (c 3, CHCl₃).** Purify acetobromo-α-D-galactose as for the glucose analogue (next entry). If the compound melts lower than 87° or is highly coloured, then dissolve it in CHCl₃ (ca 3 volumes) and extract with H₂O (2 volumes), 5% aqueous NaHCO₃, and again with H₂O and dry it over Na₂SO₄. Filter and evaporate it in a vacuum. The partially crystalline solid or syrup is dissolved in dry Et₂O (must be very dry) and recrystallised by adding petroleum ether (b 40-60°) to give a white product. [McKellan & Horecker *Biochemical Preparations* **11** 111 1960, *Beilstein* **17/6** V 369.]

Acetobromo-α-D-glucose (α-acetobromoglucose 2,3,4,6-tetraacetyl-α-D-glucopyranosyl bromide) [572-09-8] $C_{14}H_{19}BrO_9$, **M 411.2, m 87-88°, 88-89°, $[\alpha]_{546}^{20} +230$, $[\alpha]_D^{20} +195$ (c 3, CHCl₃).** If nicely crystalline, recrystallise it from Et₂O/pentane or petroleum ether (b 40-60°). Alternatively, dissolve it in diisopropyl ether (dried over CaCl₂ for 24 hours, then over P₂O₅ for 24 hours) by shaking and warming (for as short a period as possible), and filter warm. Cool to ca 45°, then slowly to room temperature and finally at 5° for more than 2 hours. Collect the solid, wash it with cold dry diisopropyl ether and dry it in a vacuum over Ca(OH)₂ and NaOH. Store it dry in a desiccator in the dark. Solutions can be stabilised with 2% CaCO₃. [Redemann & Niemann *Org Synth Coll Vol* **3** 11 1955, DOI: 10.15227/orgsyn.022.0001; for the preparation of the α-2-cyanoethyl derivative from it see Glese et al. *Org Synth* **65** 236 1987, DOI: 10.15227/orgsyn.065.0236; *Beilstein* **17/6** V 368.]

α-Acetyldigitoxin [5511-98-8, 1111-39-3] $C_{43}H_{66}O_{15}$, **M 823.0, m 153-180° (monohydrate, dec), 199-201°, 217-221°, 258°(dec), $[\alpha]_D^{20} +5.0$ (c 0.7, pyridine), +24.5 (c 1, MeOH).** α-Acetyldigitoxin is obtained from the commercial mixture [α:β (2:1)] [25395-32-8]. The α-form is obtained from the β-form by heating in anhydrous or aqueous organic solvent (e.g. aqueous MeOH) at pH 3.5–8. It crystallises from MeOH as plates, CHCl₃/Et₂O or Me₂CO/Et₂O. At 20° 1g dissolves in 16ml of MeOH, 66ml of Me₂CO and ~880ml of EtOAc. Store at -20°. [Stoll & Kreis *Helv Chim Acta* **35** 1318, 1952, DOI: 10.1002/hlca.19520350431; Stoll et al. *Helv Chim Acta* **35** 1324 1952, DOI: 10.1002/hlca.19520350432; *Beilstein* **18** III/IV 1479.] It is a cardenolide.

β-Acetyldigitoxin [5355-48-6, 1264-51-3] $C_{43}H_{66}O_{15}$, **M 823.0, m 225°, 240°(dec), 271-273°(dec), $[\alpha]_D^{20} +16.2$ (c 0.7, pyridine), +26.7 (c 0.7, MeOH).** β-Acetyldigitoxin crystallises from MeOH as a methanolate which loses MeOH in a vacuum desiccator. The solubility at room temperature is: H₂O (0.0005%), EtOAc (0.4%), MeOH (0.6%), Me₂CO (1.6%) and CHCl₃(12%). Store at -20°. [Stoll & Renz *Helv Chim Acta* **35** 1310 1952, DOI: 10.1002/hlca.19520350430; Kreis US 2776963, 1957 to Sandoz; *Beilstein* **18** III/IV 1479]. It is a cardenolide.

N-Acetyl-D-lactosamine [LacNAc, 2-acetyl-amino-*O*-β-D-lactopyranosyl-2-deoxy-D-glucose] [32181-59-2] $C_{14}H_{25}NO_{11}$, **M 383.4, m 169-171°, 170-171°, $[\alpha]_D^{18} +51.5 \rightarrow +28.8$ (in 3hrs, c 1, H₂O).** Purify *N*-acetyl-D-lactosamine by recrystallisation from MeOH (with 1 mol of MeOH) or from H₂O. It is available commercially

as a solution of 0.5g/ml of H₂O. Store at -20°. [Zilliken et al. *J Biol Chem* **217** 79 1955, <http://www.jbc.org/content/217/1/79>, PMID: 13271370; Zilliken et al. *J Biol Chem* **208** 199 1954, <http://www.jbc.org/content/208/1/199>; PMID: 13174538; *Beilstein* **17** IV 3452.]

1-O-Acetyl-2,3,5-tri-O-benzoyl-β-D-ribofuranose [6974-32-9] C₂₈H₂₄O₉, M 504.5, m 128-130°, 130-131°, 131-132°, [α]_D²² +45.1 (c 1.32, CHCl₃). Recrystallise it from EtOH or isoPrOH (10 parts) and dry over P₂O₅. [Synthesis from D-ribose: Recondo & Rinderknecht *Helv Chim Acta* **42** 1171 1959, DOI: 10.1002/hlca.19590420409; NMR: Stevens & Fletcher *J Org Chem* **33** 1799 1968, DOI: 10.1021/jo01269a021; synthesis and IR of the DL *racemate* m 115-117°: Iwai et al. IR: *Chem Pharm Bull Jpn* **11** 188 1963, DOI: 10.1248/cpb.11.188; *Beilstein* **17/6** V 213.]

N-Acetyl muramic acid [NAM] See 'Physiologically Active....' in this Chapter.

Alginic acid [9005-32-7] (C₆H₈O₆)_n, M 48,000-186,000. The acid, found in kelp, a brown algae, is a straight-chain, hydrophilic, colloidal, polyuronic acid (mixed polymer of mann-uronic and gul-uronic acids) composed primarily of anhydro-β-D-mannuronic acid residues with 1→4 linkage. To 5g of acid in 550ml water containing 2.8g KHCO₃ are added 0.3ml of acetic acid and 5g potassium acetate. EtOH is added to make the solution to 25% (v/v) in EtOH, and any insoluble material is discarded. Further addition of EtOH, to 37% (v/v), precipitated alginic acid. Collect the acid and dry it *in vacuo* (3.6g). A striking property of alginic acid is that it is capable of absorbing 200-300 times its weight of H₂O and to a lesser extent salts (~60% of its weight). It liberates CO₂ from carbonates (titration point with phenolphthalein). **Sodium alginate (Algin)** [9005-38-3] has [α]_D²⁰ -133 (H₂O). [Prepn from *Macrosystis pyrifera*: Nelson & Cretcher *J Am Chem Soc* **51** 1914 1929, DOI: 10.1021/ja01381a045; metachromatic properties: Pal & Schubert *J Am Chem Soc* **84** 4384 1962, DOI: 10.1021/ja00882a004.]

Aloin [10-glucopyranosyl-1,8-dihydroxy-3-(hydroxymethyl)-9(10H)anthracenone, Barbaloin] [8015-61-0, 5133-19-7, 1415-73-2] C₂₁H₂₂O₉, M 418.4, m 148-148.5°, 148-150°. Aloin forms the *monohydrate* as lemon yellow crystals on crystallisation from H₂O (450g/1.5l) which has a lower m ~70-80° than the anhydrous substance. Dry *in vacuo* over P₂O₅ at 100°, but picks up lost H₂O on standing in air for 2 days. The *heptamethyl ether* (MeI.Ag₂O/Me₂CO/N₂/8hr reflux) crystallises from EtOH and has m 180-182°, [α]_D²⁰ -12.3 (c 1.46, CHCl₃). [For constitution see Cahn & Simonsen *J Chem Soc* 2573 1932, DOI: 10.1039/JR9320002573; Hay & Haynes *J Chem Soc* 3141 1956, DOI: 10.1039/JR9560003141.]

D(+)-Altrose [D- 1990-29-0, L- 1949-88-8] C₆H₁₂O₆, M 180.2, m 103-105°, 106-108°, [α]₅₄₆ +35 (c 7.6, H₂O). Crystallise D-altrose from aqueous EtOH. If it is obtained by hydrolysis of the acetate, then it may contain sodium and acetate ions. Ions are best removed by dissolving in H₂O, passing through suitable columns of ion-exchange resins, e.g. Amberlite IR-120 and Duolite A, and concentrating in a vacuum to a syrup. This is dissolved in MeOH, filtered and evaporated in a vacuum desiccator over granular CaCl₂. The thick syrup is inoculated with seed crystals, stirred, and before it sets to a magma of crystals, transfer the crystals with MeOH to a Büchner funnel. Recrystallise them in the same way. β-D-Altrofuranside has initial [α]_D²⁰ ~-69 (c 4, H₂O) which mutarotates to +33. [Richtmeyer *Methods in Carbohydrate Chemistry* **I** 107 Academic Press 1962, *Beilstein* **1** IV 4301, see Angyal *Adv Carbohydrate Chem Biochem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5, for ratio of anomers in solution.]

D-Amygdalin (D-mandelonitrile-β-gentiobioside) [29883-15-6] C₂₀H₂₇NO₁₁, M 457.4, m 214-216° (trihydrate), 220°(anhydrous), [α]_D²² -38 (c 1.2, H₂O). D-Amygdalin recrystallises from water as the *trihydrate*, or from EtOH. It is present in bitter almonds. After melting at ~220°, the melt solidifies and re-melts at 125-130°. [For ORD and sugar structure see Hudson *J Am Chem Soc* **46** 483 1924, DOI: 10.1021/ja01667a025; for the biose see Haworth & Wylam *J Chem Soc* **123** 3120 1923, DOI: 10.1039/CT9232303120; for catalytic racemisation see Smith *Chem Ber* **64** 1115 1931, DOI: 10.1002/cber.19310640527; *Beilstein* **17/8** V 188.]

Amylose [9005-82-7] (C₆H₁₀O₅)_n (for use in iodine complex formation). Amylopectin is removed from impure amylose by dispersing in aqueous 15% pyridine at 80-90° (concentration 0.6-0.7%) and allowing the

solution to stand at 44-45° for 7 days. The precipitate is re-dispersed and recrystallises during 5 days. After a further dispersion in 15% pyridine, it is cooled to 45°, allowed to stand at this temperature for 12 hours, then cooled to 25° and left for a further 10 hours. The combined precipitates are dispersed in warm water, precipitated with EtOH, washed with absolute EtOH, and dried *in vacuo* [Foster & Paschall *J Am Chem Soc* **75** 1181 1953, DOI: 10.1021/ja01101a051].

1,6-Anhydro-β-D-glucose (levoglucosan, β-glucosan, β-D-pyranose-form) [498-07-7] $C_6H_{10}O_5$, **M 162.1, m 177°, 179-180°, 182-184°, 184°, $[\alpha]_D -66.3$ (c 2, H₂O).** Levoglucosan is readily formed by pyrolysis (dry distillation) of starch, ash-free amylose, cellulose and β-D-glucose in a vacuum. Best results are obtained when the acidity of the distillate is least. The product is dissolved in MeOH, decolourised with charcoal and cooled to give the glucosan in good yields. Other β-D-glucosides such as Arbutin, Salicin and Phloridzin also yield levoglucosan, and is evidence for the β- linkage configuration because α-glucose and α-glucosides require more drastic conditions for pyrolysis, i.e. longer heating and higher temperatures since the α- orientation at C-1 needs to be inverted to the β- orientation [Karrer *Helv Chim Acta* **3** 258 1920, DOI: 10.1002/hlca.19200030125; Pictet & Cramer *Helv Chim Acta* **3** 640 1920, DOI: 10.1002/hlca.19200030159, Ruchel & Schuerch *J Org Chem* **31** 2233 1966, DOI: 10.1021/jo01345a035; Irvine & Oldham *J Chem Soc* 2903 1925, DOI: 10.1039/CT9252702903; Carvalho et al. *J Am Chem Soc* **81** 4054 1959, DOI: 10.1021/ja01524a060].

Another ready source of levoglucosan is β-phenylglucoside (2.0, see [1464-44-4]) which is dissolved 1.3N aqueous KOH (100ml), heated on a boiling water bath for 9 hours, while monitoring the specific optical rotation until it is constant (−64.4). The clear mixture is cooled, neutralised (to methyl orange, the slight colour of the cooled solution does not interfere) with 3N sulfuric acid, evaporated to dryness *in vacuo*, extracted from the K₂SO₄ residue with warm EtOH, evaporated and the residue is recrystallised from EtOH (10ml) to give **levoglucosan, m 179-180°, $[\alpha]_D -66.3$ (c 2, H₂O), in 88% yield.** Other β-phenylglyco-pyrano-sides give anhydro-glycosans, but the α- linked sugars require much longer heating (~20 hours or weeks) to give the anhydro-sugars. This alkaline reaction has been studied in detail, and similarly β-phenylgalactoside provided a 91% yield of **1,6-anhydro-β-D-galactose (anhydrogalactosan)** [644-76-8] **M 162.1, with m 226°** (plates from 50% aqueous EtOH) and **$[\alpha]_D -22.0$ (c 2, H₂O).** [Montgomery et al. *J Am Chem Soc* **65** 3 1943, DOI: 10.1021/ja01241a002].

Levoglucosan is one of the more important anhydro-sugars as it, and its 2-, 3- and 4- substituted derivatives, are readily hydrolysed by aqueous acid to give glucose and the respective derivatives [Peat *Adv Carbohydrate Chem* **2** 37 1946, DOI: 10.1016/S0096-5332(08)60006-5]. Also at elevated temperature, in the presence of zinc dust, ZnCl₂, or platinum black, levoglucosan polymerises to form a variety of oligomers described as **dextrans**, and these can be commonly 20,000 to 50,000 mers (but 300,000 mers in the extreme); and have found use in chromatographic purification of large molecules, e.g. proteins. The polymerisation process has been studied in detail [Carvalho et al. *J Am Chem Soc* **81** 4054 1959, DOI: 10.1021/ja01524a060; Irvine & Oldham *J Chem Soc* 2903 1925, DOI: 10.1039/CT9252702903; Ruchel & Schuerch *J Org Chem* **31** 2233 1966, DOI: 10.1021/jo01345a035; see also Pictet & Sarasin *Helv Chim Acta* **1** 87 1918, DOI: 10.1002/hlca.19180010109; and later papers]. The 2-, 3- and 4- OH groups are readily derivatised in the usual way, and the **triacetyl derivative** has [13242-55-2] **M 288.2, m 109.5-110.5°, $[\alpha]_D^{25} -50.8$ (CHCl₃), the trifluoroacetyl derivative** has **M 450.2, m 63.5-65.0°, $[\alpha]_D^{25} -39.9$ (c 2.1, CHCl₃), the tribenzoyl derivative** has [13567-05-7] **M 474.4, m 202-203°, $[\alpha]_D^{20} -36.4$ (CHCl₃), the tri-O-methyl derivative** has **M 204.2, m 57.5-58.5°, $[\alpha]_D^{25} -63.7$ (c 2, H₂O), the tri-O-ethyl derivative** has **M 246.3, b 90°/0.60mm, $d^{28}_{20} 1.081$, $[\alpha]_D^{25} -51.5$ (c 2.8, CHCl₃), the tri-O-benzyl derivative** has **M 432.5, m 89.5-90.5°, $[\alpha]_D^{25} -30.8$ (c 2.7, CHCl₃), and the tri-O-methylsilyl derivative** has **M 378.7, b 87-87.5°/0.05mm, $[\alpha]_D^{30} -33.8$ (c 3.6, CHCl₃); all of which have been subjected to the polymerisation process with the ‘ether’ derivatives being more successful than the ‘ester’ derivatives in producing high polymers. [Ruckel & Scheurch *J Org Chem* **31** 2233 1966, DOI: 10.1021/jo01345a035]. The ¹H NMR of levoglucosan and its derivatives in DMSO-*d*₆ and D₂O have been reported in detail [Wollwage & Seib *J Chem Soc (C)* 3143 1971, DOI: 10.1039/J39710003143]. [Beilstein **19** H 894, **19** I 452, **19** III/V 1181. **19/3** V 498.]**

β-L-(+)-Arabinose (natural) [87-72-9, 5328-37-0] $C_5H_{10}O_5$, **M 150.1, m 158°, 160-163°, 164-165°, $[\alpha]_D +104$ (c 4, H₂O after 24hr), pK¹⁷ 12.4.** β-L-(+)-Arabinose is recrystallised slowly, twice, from 80% aqueous EtOH, then dried under vacuum over P₂O₅. It can also be purified by heating the arabinose (200g) with glacial acetic acid (300ml) on a boiling water bath for 45 minutes, cooling, filtering, washing with 95% EtOH (500ml)

in four portions and drying at 56-60° over P₂O₅. It has been recrystallised from 5 times its weight of 76% EtOH using charcoal (10g) to yield 127g, **m 155-157°**, $[\alpha]_{\text{D}}^{20} +190.6$ and mutarotating to +104 (c 4, H₂O). [Anderson & Sands *Org Synth Coll Vol* **1** 67 1941, DOI: 10.15227/orgsyn.008.0018; Wolfrom & Christian *J Am Chem Soc* **48** 3172 1926, DOI: 10.1021/ja01691a025; *Beilstein* **1** IV 4217.]

D-(-)-Arabinose [10323-20-3, 28697-53-2 (pyranoside) C₅H₁₀O₅, **M 150.1**, **m 164°**, **164-165°**, $[\alpha]_{\text{D}}^{20} -104.5$ (c 1, H₂O), $[\alpha]_{546} -123$ (c 10, H₂O after 24 hours), **pK₂₅ 12.54**. Crystallise D-(-)-arabinose three times from EtOH, dry it *in vacuo* at 60° for 24 hours and store it in a vacuum desiccator. It also crystallises from a mixture of H₂O/MeOH/EtOH with **m 158-160°**, or H₂O with **m 160-161°**. [Whistler & Schweiger *J Am Chem Soc* **81** 5190 1959, DOI: 10.1021/ja01528a042; Fletcher et al. *J Am Chem Soc* **72** 4546 1950, DOI: 10.1021/ja01166a058; *Beilstein* **1** IV 4215, see Angyal *Adv Carbohydrate Chem Biochem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5; for ratio of anomers in solution.]

D-Arabitol (D-arabinitol) [488-82-4] C₅H₁₂O₅, **M 152.2**, **m 103°**, **103-104°**, $[\alpha]_{546}^{20} +13$ (c 5, 8% borax solution). This pentol, which occurs in lichens and fungi, is purified by recrystallisation from 90% EtOH or MeOH. [Ashina & Yanagita *Chem Ber* **67** 799 1934, DOI: 10.1002/cber.19340670521; derivatives: Nakagawa et al. *Bull Chem Soc Jpn* **40** 2150 1967, DOI: org/10.1246/bcsj.40.2150; Prince & Reichstein *Helv Chim Acta* **20** 101 1937, DOI: 10.1002/hlca.19370200113; Hough & Theobald *Methods in Carbohydrate Chemistry* **I** 94 1962, Academic Press, *Beilstein* **1** IV 2832.]

L-Arabitol (L-arabinitol) [7643-75-6] C₅H₁₂O₅, **M 152.2**, **m 102°**, **103°**, $[\alpha]_{546} -13$ (c 5, 8% borax solution). This pentol, which occurs in the urine of pentosuric subjects, is purified by recrystallisation from 90% EtOH or MeOH. It has a higher rotation in the presence of molybdate: $[\alpha]_{\text{D}}^{20} -130$ (c 0.16, acidified molybdate) [Richtmyer & Hudson *J Am Chem Soc* **73** 2249 1957, DOI: 10.1021/ja01149a095]. [Gätzi & Reichstein *Helv Chim Acta* **21** 195 1938, DOI: 10.1002/hlca.19380210132; *Beilstein* **1** IV 2832.]

DL-Arabitol (DL-arabinitol) [2152-56-9] C₅H₁₂O₅, **M 152.2**, **m 105-106°**. This synthetic arabitol is purified by recrystallisation from 90% EtOH, MeOH or EtOH/Me₂CO. [Raphael *J Chem Soc, Supplement* 44 1949, DOI: 10.1039/JR9490000S44; Ashina & Yamagita *Chem Ber* **67** 799 1934, DOI: 10.1002/cber.19340670521; *Beilstein* **1** IV 2832.]

p-Arbutin (p-hydroquinone-O-β-D-glucopyranoside) [497-76-7] C₁₂H₁₆O₇, **M 272.3**, **m 195-198°**, **199°**, **200°** (sintering at 163-164°), $[\alpha]_{\text{D}}^{20} -65$ (c 2, H₂O), **pK_{Est} ~10.0**. The glycoside from *Protea exima* is purified by recrystallisation from H₂O or moist EtOAc (as *monohydrate*), after chromatography through silica Gel using EtOAc/MeOH. Crystallisation from EtOH/CHCl₃ gives crystals **m 199-200°** with intermediate melting at 164° and resolidifying. Its solubility in H₂O is 5g/100ml at ~25°. The *pentaacetate* crystallises from EtOH in fine needles with **m 145-146°**, $[\alpha]_{\text{D}}^{20} -28.2$ (c 2, Me₂CO). [Robinson & Waters *J Chem Soc* 2729 1930, DOI: 10.1039/JR9300002729; IR, NMR, MS: Perold et al. *JCS Perkin Trans 1* 239 1979, DOI: 10.1039/P19790000239; *Beilstein* **17/7** V 110.]

L-(+)-Ascorbic acid [Vitamin C, 5R-5-(1S-1,2-dihydroxyethyl)-3,4-dihydroxyfuran-2(5H)-one] [50-81-7] C₆H₈O₆, **M 176.1**, **m 193°**(dec), $[\alpha]_{546} +23$ (c 10, H₂O), **pK₁²⁵ 4.04**, **pK₂²⁵ 11.34**. Crystallise it from MeOH/Et₂O/petroleum ether. Its solubility in g/100ml at ~25° is 33 (40 at 45°, 80 at 100°) in H₂O, 2.0 in EtOH, 5.0 in propylene glycol, and 1.0 in glycerol. [Herbert et al. *J Chem Soc* 1270 1933, DOI: 10.1039/JR9330001270]. It is an important vitamin, is anti-ascorbitic (deficiency causes Pellagra), an antimicrobial, a reducing agent and an antioxidant used also as a food additive. [Davies et al. *Vitamin C: Its Chemistry and Biochemistry*, The Royal Society of Chemistry Publication, p. 48 1991, ISBN 0-85186-333-7; *Beilstein* **18/5** V 26.]

6-Bromo-2-naphthyl-α-D-galactopyranoside [α- 25997-59-5; β- 15548-61-5] C₁₆H₁₇BrO₆, **M 385.2**, **m 178-180°**, **224-226°**, **225°**, $[\alpha]_{\text{D}}^{28} +60$ (c 1.2, pyridine). It is prepared from penta-O-acetyl-D-galactoside, 6-bromo-2-naphthol and ZnCl₂. The resulting tetra-acetate (2g) is hydrolysed by dissolving in 0.3N KOH (100ml) and heating until the solution is clear, then filtering and cooling to give colourless crystals of the

α -isomer which are collected and recrystallised twice from hot MeOH. The high specific rotation is characteristic of the α -isomer. The **tetraacetate** has **m 155-156°**, $[\alpha]_{\text{D}}^{20} +60$ (c 1, CHCl₃) [Dey & Pridham *Biochem J* **115** 47 1969, DOI: 10.1042/bj1150047] [reported **m 75-85°**, $[\alpha]_{\text{D}}^{24} +94$ (c 1.3, dioxane), Monis et al. *J Histochem Cytochem* **11** 653 1963, DOI: 10.1177/11.5.653]. [Beilstein **17** IV 2972.]

Carminic acid (7- α -D-glucopyranosyl-9,10-dihydro-3,5,6,8-tetrahydroxy-1-methyl-9,10-dioxo-2-anthracene carboxylic acid, Neutral Red 4: CI 75470, hydroxyanthrapurin glucoside) [1260-17-9] C₂₂H₂₀O₁₃, M 492.4, m 120°(dec), $[\alpha]_{\text{D}}^{15} +51.6$ (H₂O), $[\alpha]_{\text{D}}^{20} +3.1$ (c 1, H₂O). (several phenolic pKs). Carminic acid, the colourant principle of cochineal, forms red prisms from EtOH. It gives a red colour in Ac₂O and yellow to violet in acidic solution. Its UV has λ_{max} (H₂O) at 500nm (ϵ 6,800); (0.02N HCl) 490-500nm (ϵ 5,800) and (0.0001N NaOH) 540nm (ϵ 3,450). Its IR has ν_{max} (Nujol) at 1708s, 1693s, 1677m, 1648m, 1632m, 1606s, 1566s, 1509 cm⁻¹. Periodate oxidation is complete after 4 hours at 0° with the consumption of 6.2 mols. The **tetra-O-methyl carminate** has **m 186-188°** (yellow needles from *C₆H₆/petroleum ether). [For IR see Ali & Haynes *J Chem Soc* 1033 1959, DOI: 10.1039/JR9590001033; for synthesis see Allevi et al. *JCS Perkin I* 575 1998, DOI: 10.1039/A705145J; and Davis & Smith *Biochemical Preparations* **4** 38 1955, Beilstein **18** III/IV 6697.] A red dye (e.g. for clothes) found in insects (e.g. cochineal).

λ -Carrageenan [9064-57-7 (κ), 9000-07-1 (κ + little of λ)]. This D-galactose-anhydro-D or L-galactoside polysaccharide is precipitated from 4g of Carrageenan in 600ml of water containing 12g of KOAc by addition of EtOH. Collect the fractions that precipitate between 30 and 45% (v/v) of EtOH and dry them *in vacuo*. It is a metachromatic compound, cf. Alginic acid above and Heparin below [Pal & Schubert *J Am Chem Soc* **84** 4384 1962, DOI: 10.1021/ja00882a004].

Colchicoside [7S-7N-[3-(β -D-glucopyranosyloxy)-1,2,10-tetramethoxy-9-oxobenzo[a]-heptalen-7-yl-(7S)]-acetamide, 7-glucosylcolchicine] [477-29-2] C₂₇H₃₃NO₁₁, M 547.5, m 194-197°, 216-218° (block), d²⁵ 1.44g/cm³, $[\alpha]_{\text{D}}^{15} -128.5$ (c 1, CHCl₃), $[\alpha]_{\text{D}}^{15} -231$ (c 1, MeOH), $[\alpha]_{\text{D}}^{15} -355$ (c 1, H₂O). Purify the pale yellow colchicoside by chromatography through alumina and eluting with CHCl₃, then recrystallising from Me₂CO, m 275-280° (after releasing solvent at 180°). It also crystallises from EtOH; see the aglycon colchicine [64-86-8] in 'Physiologically Active.....' in this Chapter. Store in a refrigerator. The **tetraacetate** has **m 175-177°**, $[\alpha]_{\text{D}}^{15} -57$ (c 1, CHCl₃). [Bellet et al. *Ann Pharm Fr* **10** 241 1952, *Chem Abstr* **47** 3323 1953, Beilstein **14** IV 946 for colchicine.]

Convallatoxin (α -cardenolide mannoside) [508-75-8] C₂₉H₄₂O₁₀, M 550.6, m 238-239°, 238-241° (dec), d²⁵ 1.41g/cm³, $[\alpha]_{\text{D}}^{20} -9.4$ (c 0.7, dioxane), $[\alpha]_{\text{D}}^{16} -1.0 \pm 3$ (c 0.7, EtOH). Crystallise convallatoxin from EtOAc, CHCl₃/EtOH (9:1) or MeOH/Et₂O. The **tetraacetate** has **m 238-242°** (from MeOH/Et₂O), $[\alpha]_{\text{D}}^{25} -5$ (CHCl₃). [Reyle et al. *Helv Chim Acta* **33** 1541 1950, DOI: 10.1002/hlca.19500330621; Fieser & Jacobson *J Am Chem Soc* **59** 2335 1937, DOI: 10.1021/ja01290a070; for isolation from *lily of the valley* blossoms (*Convallaria majalis* L) see Karrer *Helv Chim Acta* **12** 506 1929, DOI: 10.1002/hlca.19290120154; Beilstein **18** III/IV 3142.] It is a most potent cardiotoxin.

α -Cyclodextrin (H₂O) [10016-20-3] C₃₆H₆₀O₃₀, M 972.9, m >280°(dec), $[\alpha]_{\text{D}}^{546} +175$ (c 10, H₂O). Recrystallise α -cyclodextrin from 60% aqueous EtOH, then twice from water, and dry it for 12 hours in a vacuum at 80°. It is also purified by precipitation from water with 1,1,2-trichloroethylene. The precipitate is collected, washed and resuspended in water (solubility ~50mg/ml at ~25°). This is boiled to steam distil the trichloroethylene. The solution is then freeze-dried to recover the cyclodextrin. [Armstrong et al. *J Am Chem Soc* **108** 1418 1986, DOI: 10.1021/ja00267a010]. [Beilstein **19/12** V 789.]

β -Cyclodextrin (H₂O) [7585-39-9, 68168-23-0] C₄₂H₇₀O₃₅, M 1135.0, m >300°(dec), $[\alpha]_{\text{D}}^{546} +170$ (c 10, H₂O). Recrystallise β -cyclodextrin from water and dry it for 12 hours in a vacuum at 110°, or 24 hours in a vacuum at 70°. The purity is assessed by TLC on cellulose containing a fluorescent indicator. [Taguchi, *J Am Chem Soc* **108** 2705 1986, DOI: 10.1021/ja00270a032; Tabushi et al. *J Am Chem Soc* **108** 4514 1986, DOI: 10.1021/ja00275a043; for absorption and fluorescence of indole inclusion complex see Orstan & Ross *J Phys Chem* **91** 2739 1987, DOI: 10.1021/j100295a019.] [Beilstein **19** IV 6287, **19/12** V 801.]

***N*-Decanoyl-*N*-methylglucamine** (Mega-10, *N*-D-glucidyl-*N*-methyl deconamide) [85261-20-7] $C_{17}H_{35}NO_6$, M 349.5, m 91-93°, 92°. Possible impurities are decanoic acid and *N*-methylglucamine. The former is removed by grinding the solid with Et₂O and then with petroleum ether and drying over P₂O₅. It is twice recrystallised from MeOH/Et₂O (1:9, v/v; 10ml/g) by dissolving in the minimum volume of MeOH, adding Et₂O and drying in a vacuum. To remove the glycamine, the solid (800mg) is dissolved in hot H₂O (10ml) and set aside. Mega-10 crystallises as colourless needles. These are filtered off and dried in a vacuum to constant weight. It is a good non-ionic non-hygroscopic detergent with a critical micelle concentration (CMC) of 7.4mM (0.26%) in 0.1M Tris-HCl pH 7.4 at 25°. [Hildreth *Biochem J* **207** 363 1982, DOI: 10.1042/bj2070363; for detergent activity in solubilising microsomes see Taguchi & Armarego *Med Res Rev* **18** 43 1998, DOI: 10.1002/(SICI)1098-1128(199801)18:1<43.] The ***N*-Methyl-D-glucamine [(2*R*,3*R*,4*R*,5*S*)-6-(methylamino)hexane-1,2,3,4,5-pentol]** [6284-40-8] $C_7H_{17}NO_5$, M 195.2, has m 128-129°, 126-129°, 129-131.5°, $[\alpha]_D^{20}$ -18.5 (c 2, H₂O). Recrystallise it from MeOH. Its solubility in g/1ml is 1.0 in H₂O (25°) and 0.012 in EtOH (70°). Aqueous solutions are alkaline and its sulfonate salts are detergents. [Karrer & Herkenrath *Helv Chim Acta* **20** 83 1937, DOI: 10.1002/hlca.19370200109; *Beilstein* **4** IV 1914.] The antimonite salt [133-51-7] is anti-protozoal (leishmanial) in dogs [Chapman et al. *Am J Vet Res* **45** 1028 1984, PMID: 6732008].

Dehydro-L(+)-ascorbic acid {DHA, (5*R*)-5-[(1*S*)-1,2-dihydroxyethyl]furan-2,3,4(5*H*)-trione} [490-83-5] $C_6H_6O_6$, M 174.1, m 196°(dec), 228°(dec), $[\alpha]_D^{20}$ +42.5 (c 1, H₂O), pK_a 3.90. This is the oxidised form of *ascorbic* acid. Crystallise dehydro-L(+)-ascorbic acid from MeOH. The *anhydrous acid* is formed by heating it in a vacuum at 100°/1 hour to give a crisp glassy product which when shaken with absolute EtOH and then kept at 0° for 2 days gives microcrystals of the anhydrous acid. This is then washed with absolute EtOH and dried in a vacuum. It has m 225°(dec) and is stable in acidic solution but decomposes rapidly in alkaline solution. A 1% solution of the anhydrous acid when dissolved in phthalate/HCl buffer pH 3.5 at 60° and cooled to 20° has $[\alpha]_D^{20}$ +56(0 minutes), +53.5(2 hours), +19(3 days), -2(5 days) and -6(6 days); then it becomes orange in colour. A freshly prepared 1% solution in H₂O has $[\alpha]_D^{20}$ +50(0 minutes), +44(2 hours), +16(3 days) and 0(5 days). [Herbert et al. *J Chem Soc* 1270 1933, DOI: 10.1039/JR9330001270; Kenyon et al. *J Chem Soc* 158 1948, DOI: 10.1039/JR9480000158; *Beilstein* **18/5** V 411.]

2-Deoxy-D-allose (2-deoxy-D-ribo-hexose) [6605-21-6] $C_6H_{12}O_5$, M 164.2, m 140-142°, $[\alpha]_D^{24}$ +57.5 (c 1.2, H₂O). Purify 2-deoxy-D-allose by two recrystallisations from absolute EtOH. The *p*-nitrophenylhydrazone has m 61-62°, $[\alpha]_D^{16.5}$ -55 (MeOH). An equilibrium solution at 31° in D₂O contains 15% α-pyranose, 58% β-pyranose, 12% α-furanose and 15% β-furanose forms as estimated by ¹HNMR spectroscopy. [see Angyal *Adv Carbohydrate Chem Biochem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5; for ratio of anomers in solution, DOI: 10.1016/S0065-2318(08)60122-5; Zorbach & Ollapally *J Org Chem* **29** 1790 1964, DOI: 10.1021/jo01030a030.] [*Beilstein* **1** IV 4283.]

2-Deoxy-β-D-galactose (2-deoxy-D-lyxo-hexose) [1949-89-9] $C_6H_{12}O_5$, M 164.2, m 110°, 120-121°, (126-128°), $[\alpha]_D^{20}$ +60 (c 2, H₂O, 2 hours). Crystallise 2-deoxy-β-D-galactose from MeOH or diethyl ether. The *aniline derivative* has m 142-143°, $[\alpha]_D^{16.5}$ -149 (c 0.8, pyridine). [Overend et al. *J Chem Soc* 671 1950, DOI: 10.1039/JR9500000671; Overend et al. *J Chem Soc* 992 1951, DOI: 10.1039/JR9510000992.] A 30% equilibrium solution at 31° in D₂O contains 40% α-pyranose, 44% β-pyranose, 8% α-furanose and 8% β-furanose forms as estimated by ¹HNMR spectroscopy [Angyal & Pickles *Aust J Chem* **25** 1711 1972, DOI: 10.1071/CH9721711]. [*Beilstein* **1** IV 4283.]

2-Deoxy-α-D-glucose (2-deoxy-D-arabino-hexose) [154-17-6] $C_6H_{12}O_5$, M 164.2, m 146-147°, 148-151°, $[\alpha]_D^{20}$ +46 (c 0.5, H₂O after 45 hours). Crystallise 2-deoxy-α-D-glucose from MeOH/Me₂CO, Me₂CO or butanone to give a mixture of α- and β- *anomers*, m 142-144°, $[\alpha]_D^{18}$ +38 (35 minutes) to +46 (c 0.5, H₂O). Recrystallisation from *iso*PrOH gives mainly the *α-anomer* m 134-136°, $[\alpha]_D^{18}$ +156 to +103 (c 0.9, pyridine). ¹H NMR studies showed that at 44° in D₂O the solution contained 36% of α-pyranose and 64% of β-pyranose sugars, but furanose structures were undetectable. [Snowden & Fischer *J Am Chem Soc* **69** 1048 1947, DOI: 10.1021/ja01197a022; derivatives: Bollinger & Schmidt *Helv Chim Acta* **34** 989 1951, DOI: 10.1002/hlca.19510340333; see Angyal & Pickles *Aust J Chem* **25** 1711 1972 for ratio of isomers in solution, DOI: 10.1071/CH9721711; *Beilstein* **1** IV 4282.]

6-Deoxy-D-glucose (D-quinovose) [7658-08-4] $C_6H_{12}O_5$, M 164.2, m 146°, $[\alpha]_D^{20}$ +73 (after 5 minutes) and +30° (final, after 3 hours) (c 8.3, H_2O). 6-Deoxy-D-glucose is purified by recrystallisation from EtOAc and is soluble in H_2O , EtOH but insoluble in Et_2O and Me_2CO . [Srivastava & Lerner *Carbohydr Res* **64** 263 1978, DOI: 10.1016/S0008-6215(00)83707-6; NMR: Angyal & Pickles *Aust J Chem* **25** 1711 1972, DOI: 10.1071/CH9721711; *Beilstein* **1** IV 4260.]

2-Deoxy-β-L-ribose [18546-37-7] $C_6H_{10}O_4$, M 134.1, m 77°, 80°, $[\alpha]_D^{25}$ +91.7 (c 7, pyridine, +40 final). Crystallise 2-deoxy-β-L-ribose from diethyl ether. It can also be purified by dissolving the ribose (7.3g) in EtOAc (3L) by reflux, decanting from any insoluble material and evaporating at 50°/vacuum to 2.2L, setting aside for 1-2 hours, filtering, and concentrating to 840ml. The ribose separates as compact nodules during 3-5 days at 0° and has m 87-93°, and after repeated recrystallisations it has m 92-95°. Mutarotation is as follows: $[\alpha]_D^{16.5}$ +80(0 minutes), +71(6 minutes), +59(21 minutes) and +59(41 minutes) (c 1.14, H_2O); $[\alpha]_D^{20}$ +105(0 minutes), +71(9.5 minutes), +67(12.5 minutes), +49(81 minutes), +49(136 minutes) (c 0.9, MeOH). [Deriaz et al. *J Chem Soc* 1879 1949, DOI: 10.1039/JR9490001879; *Beilstein* **1** IV 4181.]

2-Deoxy-β-D-ribose [533-67-5] $C_6H_{10}O_4$, M 134.1, m 86-87°, 87-90°, $[\alpha]_D^{20}$ -56 (c 1, H_2O after 24 hours). Dissolve 2-deoxy-β-D-ribose in a little H_2O , evaporate to a syrup (in a vacuum), and seed to crystallise. Triturate the crystals with a little EtOAc containing 5% MeOH, decant and dry in vacuum over P_2O_5 . It is best purified via the **anilide** which separates from a mixture of the ribose (100-125g) in MeOH (100ml) and redistilled aniline (40ml) in a few minutes. After standing for 20 hours at room temperature, it is cooled to 0°, filtered, washed with 50% aqueous MeOH and Et_2O followed by recrystallisation from ethylene glycol monomethyl ether. The **anilide** has m 172-173°, $[\alpha]_D^{25}$ +46 (equilibrium in pyridine). The anilide (5g), benzaldehyde (5ml) and benzoic acid (0.5g) in H_2O (150ml) are shaken mechanically for 20-24 hours. The aqueous phase is extracted with Et_2O (3x), decolourised with a little charcoal and evaporated in a vacuum to a syrup. This is dried over P_2O_5 in high vacuum. The syrupy sugar weighs 3.1g and crystallises in a few days, but more rapidly on seeding. Triturate it with a little EtOAc containing 5% MeOH, decant and dry it over P_2O_5 . At this stage it has m 78-82°, $[\alpha]_D^{25}$ -57 (c 1, H_2O final). This is a mixture of α- and β- anomers. Pure **β-anomer** is obtained by recrystallisation from EtOAc. The β-anomer when recrystallised from EtOAc and isoPrOH has m 96-98°, $[\alpha]_D^{25}$ -55 (c 0.5, H_2O final). [Sowden *Biochemical Preparations* **5** 75 1957.] The mutarotation is as follows: $[\alpha]_D^{20.5}$ +96.3(0 minutes), -76(33 minutes), -56(24 hours) (c 5.8 MeOH). It is moderately hygroscopic and should be kept in a well stoppered bottle. It also crystallises from diethyl ether. [Deriaz et al. *J Chem Soc* 1879 1949, DOI: 10.1039/JR9490001879; *Beilstein* **1** IV 4181, Hauske & Rapoport *J Org Chem* **44** 2472 1979, DOI: 10.1021/jo01328a029.]

The **1,3,4-tribenzoate (α- and β-mixture)**, obtained by benzylation, has m 127° (after crystallisation from EtOH), $[\alpha]_D^{23}$ -65 (c 1, $CHCl_3$). The crude syrupy mixture in $*C_6H_6$ is applied to an acid-washed Alorco Al_2O_3 column. Elution with $*C_6H_6$ /hexane (1:1) affords (after crystallisation from MeOH), **β-1,3,4-tri-O-benzoyl-2-deoxy-D-ribose**, m 159-169°, $[\alpha]_D^{20}$ -196 (c 1, $CHCl_3$). Further elution with $*C_6H_6$ gives, after recrystallisation from MeOH pure **α-1,3,4-tri-O-benzoyl-2-deoxy-D-ribose**, m 151-152°, $[\alpha]_D^{20}$ +41.6 (c 0.83, $CHCl_3$). [Pedersen et al. *J Am Chem Soc* **82** 3425 1960, DOI: 10.1021/ja01498a047; see Angyal *Adv Carbohydrate Chem Biochem* **42** 15 1984, for ratio of anomers and ring forms, DOI: 10.1016/S0065-2318(08)60122-5.]

Dextran [9004-54-0] $H(C_6H_{10}O_5)_xOH$ M_r 6,000-220,000. Solutions of dextran keep indefinitely at room temperature if 0.2ml of Roccal (10% alkyl dimethylbenzylammonium chloride) or of 2mg phenyl mercuric acetate are added per 100ml solution. This inhibits mould growth. [See Levoglucosan above; Evans & Hibbert *Adv Carbohydr Chem* **2** 203 1946, DOI: 10.1016/S0096-5332(08)60011-9; for Dextran: Structure & Synthesis see Neely *Adv Carbohydr Chem* **15** 341 1961, DOI: 10.1016/S0096-5332(08)60191-5.]

Diacetone-D-glucose (1,2:5,6-di-O-isopropylidene-α-D-glucofuranoside) [582-52-5] $C_{12}H_{20}O_6$, M 260.3, m 107-110°, 110.5°, 111-113°, 112°, $[\alpha]_D^{15}$ -18.4 (c 1, H_2O). Diacetone-D-glucose crystallises from Et_2O , (needles), petroleum ether or $*C_6H_6$ and sublimes *in vacuo*. It is soluble in 7 volumes of H_2O and 200 volumes of petroleum ether at their boiling points. The solubility in H_2O at 17.5° is 4.3%. It precipitates from aqueous solutions on basification with NaOH. [Schmid & Karrer *Helv Chim Acta* **32** 1371 1949, DOI: 10.1002/hlca.19490320504; Fischer & Rund *Chem Ber* **49** 88 1916, DOI: 10.1002/cber.19160490111; IR: Kuhn

Anal Chem **22** 276 1950, DOI: 10.1021/ac60038a015; *Beilstein* **19/12** V 318.]

***N,N'*-Diacetylchitobiose** (2-acetyl-*O*⁴-[2-acetylamino-2-deoxy- β -D-glucopyranosyl]-2-deoxy-D-glucose) [35061-50-8] $C_{16}H_{22}N_2O_{11}$, **M 424.4**, **m 245-247°(dec)**, **251.5-252.5°**, **260-262°**, $[\alpha]_D^{25} +39.5$ (extrapolated) $\rightarrow +18.5$ (after 60 minutes, c 1, H₂O). Recrystallise *N,N'*-Diacetylchitobiose from aqueous MeOH or aqueous EtOH/1,2-dimethoxyethane. Also, the crystalline solid is suspended in the minimum volume of hot dry MeOH and H₂O is added dropwise, with shaking, until it dissolves, and then set aside at room temperature until crystallisation is complete. The equilibrium rotation, $[\alpha]_D^{25} +18.5$ (c 1, H₂O), is reached after 60 minutes. [Zilliken et al. *J Am Chem Soc* **77** 1296 1955, DOI: 10.1021/ja01610a070; *Beilstein* **18/11** V 147.]

1,3,4,6-Di-*O*-benzylidene-D-mannitol [28224-73-9] $C_{20}H_{22}O_6$, **M 358.4**, **m 192-195°**, **193°**, $[\alpha]_D^{20} -11.9$ (c 0.7, Me₂CO). 1,3,4,6-Di-*O*-benzylidene-D-mannitol recrystallises from Et₂O in long fine needles with λ_{max} at 256nm (ϵ 435) in 95% EtOH, and R_F 0.21 (1:1 CCl₄/EtOAc) on TLC Silica Gel G. [Sinclair *Carbohydr Res* **12** 150 1970, DOI: 10.1016/S0008-6215(00)80238-4; for ORD, CD, NMR, IR and MS see Brecknell et al. *Aust J Chem* **29** 1749 1976, DOI: 10.1071/CH9761749; *Beilstein* **19/11** V 640.]

Digitonin [11024-24-1] $C_{56}H_{92}O_{29}$, **M 1229.3**, **m >270°(dec)**, $[\alpha]_{546}^{20} -63$ (c 3, MeOH). This non-ionic detergent, digitoxin hexa-glycoside, can be recrystallised from aqueous 85% EtOH or MeOH/diethyl ether. It is purified by preparative paper chromatography and developed with the upper phase of a mixture of nBuOH/H₂O/AcOH (4:5:1), and the spot (R_F 0.36) is eluted with 25% CCl₃CO₂H in CHCl₃. It has also been purified by countercurrent distribution. It forms an *ethanolate*, *complexes* with cholesterol and other sterols, and solubilises lipids. It is almost insoluble in H₂O (forming a foam), is more soluble in absolute EtOH (1/75g/100ml), but less in 95% EtOH (0.45g/100ml). Its CMC (critical micellar concentration) is <0.5mM at 20-25°; and micellar average mol wt is 70,000. [Structure: Tschesche & Wulff *Tetrahedron* **19** 621 1963, DOI: 10.1016/S0040-4020(01)98548-5; Ruhenstroth-Bauer & Breitenfeld *Hoppe Seyler's Z Physiol Chem* **302** 111 1955, Grisvold *J Am Pharm Assoc* **23** 664 1934; for papaveraceae alkaloids see Manske *The Alkaloids: Chemistry and Physiology* **10** 467 1968, DOI: 10.1016/S1876-0813(08)60260-2; *Beilstein* **19** IV 1243.]

Digitoxin [71-63-6] $C_{41}H_{64}O_{13}$, **M 764.9**, **m 256-257° (anhydrous)**, $[\alpha]_D^{20} +16.7$ (c 1, CHCl₃), **+4.8** (c 1, dioxane). Digitoxin crystallises from MeOH, aqueous EtOH with 0.5 to 1 H₂O and from H₂O as the *dihydrate*. It also crystallises from CHCl₃/Et₂O as *anhydrous* crystals. The hydrate dehydrates at 120°/vacuum. Its solubility is 2.5% in CHCl₃, 1.7% in EtOH, 0.25% in EtOAc, and 0.001% in H₂O; and has $E_{1\%}^{1cm}$ 202.5 at 219-220nm (50% EtOH). [Stoll et al. *Helv Chim Acta* **37** 1134 1954, DOI: 10.1002/hlca.19540370421; Demoen & Janssen *J Am Pharm Assoc* **42** 635 1953, PMID: 13096411; *Beilstein* **18** IV 1478, **18** V/3 354.]

D(+)-Digitoxose (2,6-dideoxy-D-ribo-hexose) [527-52-6] $C_6H_{12}O_4$, **M 148.2**, **m 110°**, **112°**, $[\alpha]_{546}^{20} +57$ (c 1, H₂O). Crystallise D(+)-digitoxose from MeOH/Et₂O, Et₂O, EtOAc, EtOAc/Et₂O/petroleum ether or Me₂CO/Et₂O and dry it over P₂O₅/vacuum. It has $[\alpha]_D^{20} +45.2$ (6 minutes) mutarotating to +50.2° (16 hours constant) (c 1.65, H₂O). [Gut & Prins *Helv Chim Acta* **30** 1223 1947, DOI: 10.1002/hlca.19470300517; Bolliger et al. *Helv Chim Acta* **35** 93 1952, DOI: 10.1002/hlca.19520350112; for NMR and abs config of D and L isomers see Tsukamoto et al. *JCS Perkin Trans 1* 2621 1988, DOI: 10.1039/P19880002621; *Beilstein* **1** IV 4191.]

Digoxin [5 β ,20(22)-cardenolide-3 β ,12 β ,14 β -triol-3-(*O*-2,6-dideoxy- β -D-ribo-hexopyranosyl-(1 \rightarrow 4)-*O*-2,6-dideoxy- β -D-ribo-hexopyranosyl-(1 \rightarrow 4)-*O*-2,6-dideoxy- β -D-ribo-hexopyranosyl)-oxy-), 12 β -isomer] [20830-75-5] $C_{41}H_{64}O_{14}$, **M 781.0**, **m 249.3°**, **265(dec)**, $[\alpha]_{546}^{20} +14.0$ (c 10, pyridine), $[\alpha]_{546}^{25} +15.6$ (c 0.5, CHCl₃/MeOH 1:1). Crystallise digoxin from aqueous EtOH, aqueous pyridine, EtOH/CHCl₃, and dry it in a vacuum at 100°. The melting point depends on heating rate, but when placed in a bath at 260° and heated slowly it decomposes at 265°. In EtOH it has λ_{max} at 220nm (ϵ 12,800). Its solubility in H₂O is 6.5mg/100ml at 20°. [Smith *J Chem Soc* 508 1930, DOI: 10.1039/JR9300000508; Hollman *Brit Med J* **312** 912 1996, DOI: 10.1136/bmj.312.7035.912; X-ray: Go et al. *Cryst Struct Commun* **8** 149, 1031 1979, *Beilstein* **18/4** V 381.] A cardiac glycoside; **HIGHLY TOXIC**.

(-)-2,3,4,6-Di-O-isopropylidene-2-keto-L-gulonic acid monohydrate (-DAG) [18467-77-1, 68539-16-2] $C_{12}H_{18}O_7 \cdot H_2O$, **M 292.3**, **m 88-90°**, **100-101°**, **103°**, $[\alpha]_D^{25}$ **-21.6** (c **2.3**, MeOH). Dissolve (-)-DAG in Et₂O, filter, dry (MgSO₄) it, filter it again and evaporate to give a yellow oil. Addition of one drop of H₂O induces crystallisation to the *monohydrate*, which also forms rhombic crystals on recrystallisation from 95% EtOH/H₂O at room temperature. [Flatt et al. *Synthesis* 815 1979, DOI: 10.1055/s-1979-28844; Reichstein & Grüssner *Helv Chim Acta* **17** 311 1934, DOI: 10.1002/hlca.19340170136; Takagi & Jeffrey *Acta Crystallogr Sect B* **34** 2932 1978, DOI: 10.1107/S0567740878009693; *reagent* for the *optical resolution of amines* cf. Mohacsi & Leimgruber *Org Synth* **55** 80 1976, DOI: 10.15227/orgsyn.055.0080; *Beilstein* **19/12** V 520.]

1,2:5,6-Di-O-isopropylidene-D-mannitol [1707-77-3] $C_{12}H_{22}O_6$, **M 262.3**, **m 120-122°**, **121-125°**, **122°**, $[\alpha]_D^{25}$ **+1.2** (c **3**, H₂O). Although quite soluble in H₂O, it gives a purer product when crystallised from small volumes of this solvent (preferably by slow evaporation) to form needles [Baer *J Am Chem Soc* **67** 338 1945, DOI: 10.1021/ja01218a502; for NMR see Curtis et al. *JCS Perkin Trans 1* 1756 1977, DOI: 10.1039/P19770001756]. [*Beilstein* **19/11** V 589.]

Dulcitol (galactitol) [608-66-2] $C_6H_{14}O_6$, **M 182.2**, **m 188-189°**, **b 276-280°/1.1mm**, **pK¹⁸ 13.5**. Crystallise dulcitol from water by addition of EtOH. It is optically inactive and is prepared by reduction of D-galactose. Its *hexaacetate* crystallises from EtOH and has **m 168-169°**. [IR: Thompson et al. *Discuss Faraday Soc* **9** 222 1950, DOI: 10.1039/DF9500900222; Wolfrom & Thompson *Methods in Carbohydrate Chemistry* **II** 67 1963, Academic Press, *Beilstein* **1** IV 2844.]

meso-Erythritol [149-32-6] $C_4H_{10}O_4$, **M 122.1**, **m 122°**, **119-124°**, **b 329-331°/atm**, **pK¹⁸ 13.9**. *meso*-Erythritol crystallises from distilled water or absolute EtOH and is dried at 60° in a vacuum oven. It sublimes at 110° in a high vacuum. It is optically inactive. [Jeanes & Hudson *J Org Chem* **20** 1565 1955, DOI: 10.1021/jo01128a015; IR: Kuhn *Anal Chem* **22** 276 1950, DOI: 10.1021/ac60038a015; *Beilstein* **1** IV 2807.]

Erythrityl tetranitrate {Cordite, nitroglyn, [(2R,3R)-1,3,4-trinitrooxybutan-2-yl] nitrate} [7297-25-8] $C_4H_6N_4O_{12}$, **M 302.1**, **m 61°**. Crystallise cordite from EtOH. It explodes on percussion at ~220-460° and is a coronary vasodilator. Absorption through skin causes a '*nitro headache*'. Not to be confused with the C5 pentaerythritol tetranitrate (see below). For structure-activity relationship of organic nitrates for activation of guanylate cyclase see Schroder & Noack [*Arch Int Pharmacodyn Ther* **290**(2) 235 1987, PMID: 2895614]. [*Beilstein* **1** III 2358, **1** IV 2809.]

D(-)-Fructose (D-levulose, fruit sugar) [57-48-7] $C_6H_{12}O_6$, **M 180.2**, **m 103°**, **103-106°**, **119-122°(dec)**, **d²⁵ 1.694g/cm³**, $[\alpha]_{546}^{20}$ **-190** (after **1 hour**, c **10**, H₂O), **pK²⁵ 12.03**. Dissolve D(-)-fructose in an equal weight of water, add charcoal (previously washed with water to remove any soluble material), filter and evaporate under reduced pressure at 45-50° to give a syrup containing 90% of fructose. After cooling to 40°, the syrup is seeded and kept at this temperature for 20-30 hours with occasional stirring. The crystals are removed by centrifugation, washed with a small quantity of water and dried to constant weight under a vacuum over concentrated H₂SO₄. For higher purity, this material is recrystallised from 50% aqueous ethanol [Tsuzuki et al. *J Am Chem Soc* **72** 1071 1950, DOI: 10.1021/ja01159a004]. [For effect on cardiometabolic health see Malik et al. *J Am Coll Cardiol* **66** 1615 2015, DOI: 10.1016/j.jacc.2015.08.025, PMID: 26429086; *Beilstein* **31** H 321, **1** IV 4401.]

D(+)-Fucose (6-deoxy-d-galactose) [3615-37-0] $C_6H_{12}O_5$, **M 164.2**, **m 144°**, **144-145°**, $[\alpha]_{546}^{20}$ **+89** (after **24 hours**, c **10** in H₂O), **α-form mutarotates: $[\alpha]_D^{20}$ +124** (10 minutes) to **+75.6** (24 hours) (c **9**, EtOH). Crystallise D(+)-fucose from EtOH or 95% EtOH. Its *1,2:3,4-diisopropylidene derivative* has **b 83-84°/0.45mm** and crystallises on seeding with **m 37°** and $[\alpha]_D^{19}$ **-52** (melt). [Schmidt *Methods in Carbohydrate Chemistry* **I** 191 1962, Academic Press, Hockett et al. *J Am Chem Soc* **61** 1658 1939, DOI: 10.1021/ja01876a008; *Beilstein* **31** H 76, **1** IV 4265.]

D-Galactal (1.5-anhydro-2-deoxy-D-lyxo-hex-1-enitol) [21193-75-9] $C_6H_{10}O_4$, **M 146.2**, **m 99-103°**, **100°, 100-102°, 104°, 103-106°**, $[\alpha]_D^{20}$ **-21.3 (c 1, MeOH)**. Recrystallise D-galactal from EtOAc, EtOH or EtOAc/MeOH. [Overend et al. *J Chem Soc* 671 1950, DOI: 10.1039/JR9500000671; Wood & Fletcher *J Am Chem Soc* 79 3234 1957, DOI: 10.1021/ja01569a064; Distler & Jourdan *J Biol Chem* 248 6772 1973, PMID: 4270451; Seeberger et al. *Aldrichim Acta* 30 75 1997; *Beilstein* 17 III/IV 2332.]

Galactaric acid (mucic acid) [526-99-6] $C_6H_{10}O_8$, **M 210.1**, **m 212-213°(dec)**, **230°(dec)**, pK_1^{25} **3.09 (3.29)**, pK_2^{25} **3.63 (4.41)**. Dissolve mucic acid (40g) in the minimum (calculated) volume of N aqueous NaOH (~ 335ml) without heating, decolorise this with charcoal, filter and precipitate it by adding 5N HCl (~ 57ml). Cool for 1 hour at 0°, filter off, wash with cold H₂O and dry it *in vacuo*. All temperatures should be kept below 25°. It is optically inactive. [Barker et al. *J Chem Soc* 4128 1958, DOI: 10.1039/JR9580004128; Lewis et al. *Methods in Carbohydrate Chemistry* II 39 1963, Academic Press, *Beilstein* 3 IV 1292.]

D-Galactonic acid [576-36-3] $C_6H_{12}O_7$, **M 196.2**, **m 141° (hydrate)**, **148° (anhydrous)** pK_{Est} **~3.5**. Crystallise D-galactonic acid from EtOH or aqueous EtOH. It cyclises to **D-galactono-1,4-lactone**, **m 134-136°**, and mutarotates in 1 hour to $[\alpha]_{546}^{30}$ **-92 (c 5, H₂O)**. It can also be obtained from the Na salt by adding 10 times its weight of acetic acid, warming till just brown, cooling, filtering off the crystals and drying them. It has **m 145-146°**, $[\alpha]_D^{25}$ **-13.6 (c 1, H₂O, 2 minutes and mutarotates to -57.6)**. [Brackenbury & Upson *J Am Chem Soc* 55 2512 1933, DOI: 10.1021/ja01333a046; *Beilstein* 3 IV 1257, and for the γ -lactone see below.]

D(-)-Galactono-1,4-lactone [2782-07-2] $C_6H_{10}O_6$, **M 178.1**, **m 134-135°, 134-137°, 137°**, $[\alpha]_D^{20}$ **-78 (c 5, H₂O, 1 hour)**. Crystallise the lactone from EtOH, aqueous EtOH, MeOH or EtOAc. It is also purified by passage through a column of Amberlite IR-120 (H⁺ form), and the effluent and washings are then freeze-dried [Wolf from & Thompson *Methods in Carbohydrate Chemistry* I 67 1963, Academic Press]. The **5,6-O-isopropylidene-D-galactono-1,4-lactone** is purified by chromatography (EtOAc) and has **m 165-167°** (from EtOH/hexane), and **167.5-168.5°** (from Me₂CO/*C₆H₆) [for NMR see Copeland & Stick *Aust J Chem* 31 1371 1978, DOI: 10.1071/CH9781371.] [*Beilstein* 18 III/IV 3026, 18/5 V 18.]

D(+)-Galactosamine hydrochloride [1772-03-8] $C_6H_{13}NO_5 \cdot HCl$, **M 215.6**, **m 181-185°**, $[\alpha]_D^{25}$ **+96.4 (after 24 hours, c 3.2 in H₂O)**, pK_{Est} **~7.7 (free base)**. Dissolve the hydrochloride in a small volume of H₂O. Then add three volumes of EtOH, followed by acetone until faintly turbid and keep overnight in a refrigerator. [Roseman & Ludoweig *J Am Chem Soc* 76 301 1954, DOI: 10.1021/ja01630a098; *Beilstein* 4 IV 2024.]

α -D(+)-Galactose [59-23-4, 3646-73-9 pyranose] $C_6H_{12}O_6$, **M 180.2**, **m 167-168°, 168-170°**, $[\alpha]_D^{25}$ **+80.4 (after 24 hours, c 4 in H₂O)**, pK^{25} **12.48**. α -D-Galactose is crystallised twice from aqueous 80% EtOH at -10°, then dried in a vacuum oven at 90° over P₂O₅ for 10 hours. [Link *Biochemical Preparations* 3 75 1953, Hansen et al. *Biochemical Preparations* 4 2 1955.] Also purify it by recrystallising the dried solid (150g) in hot H₂O (150ml), then adding hot MeOH (250ml) and hot EtOH (500ml), stirring to mix, filtering through a bed of charcoal, and the clear filtrate is stirred to initiate crystallisation. After standing overnight at 10°, the crystals of the **α -anomer** are filtered off by suction, washed with MeOH, then EtOH, and dried (yield 130g); and more can be obtained by evaporation of the filtrate and washing as before. [Wolf from & Thompson *Methods in Carbohydrate Chemistry* I 120 1962, Academic Press, *Beilstein* 1 IV 4336.]

β -D-Galactose [7296-64-2 (pyranose)] $C_6H_{12}O_6$, **M 180.2**, **m 167°**, $[\alpha]_D^{20}$ **+52 (initial, c 4 in H₂O)**. α -D-Galactose (40g) is dissolved in hot H₂O to establish the equilibrium of α - and β - anomers; then the solution is cooled to 0° and poured into absolute EtOH (500ml). Stir vigorously and crystallisation occurs within a few minutes, and more rapidly if seeded, filter the crystals immediately (7g, $[\alpha]_D^{20}$ +65 initial, c 4 in H₂O). This mixture of α - and β - anomers is further separated by dissolving in an equal weight of cold H₂O, filtering and adding to ice cold absolute EtOH (250ml) and stirring for 1 minute when crystals separate, then filter them off. After two such crystallisations, the initial $[\alpha]_D^{20}$ is +53. This can be further purified by shaking with 80% EtOH for 2 minutes, filtering, washing with EtOH and Et₂O, and drying in a vacuum desiccator to give **β -D-galactose** (15g) with **m 167°**, $[\alpha]_D^{20}$ **+52 (initial, c 4 in H₂O)** mutarotating to +80.4. Acetylation of D-galactose with hot NaOAc/Ac₂O gives **β -D-galactopyranoside pentaacetate** **m 142°**, $[\alpha]_D^{25}$ **+25 (c 4 in CHCl₃)**. [Wolf from & Thompson *Methods in Carbohydrate Chemistry* I 120 1962, Academic Press, *Beilstein* 1 IV 4336.]

D(+)-Galacturonic acid [685-73-4, 91510-62-2 (H₂O)] C₆H₁₀O₇, M 194.1, m 159-161°, [α]_D²⁰ +36 (c 6, H₂O, 2h), pK_{Est} ~ 4.8. Crystallisation of the acid from 95% EtOH and drying it in a vacuum desiccator (12mm) over P₂O₅ gives the *monohydrate* mixture of α- and β- anomers (**mostly α-**) as white micro needles, which sinter at ~100-111° and melt at 159-160°, [α]_D²⁰ +107 (initial, c 4 in H₂O mutarotating to +51°). [Link & Sell *Biochemical Preparations* 3 74, 78 1953, *Beilstein* 3 IV 2000.] The **β-anomer** is obtained by warming the α-anomer in EtOH, AcOH or EtOAc and has m 160° (165°, sinters at 140°), [α]_D²⁰ +27 (initial, c 2 in H₂O mutarotating to +55.3° in 24 hours). The *sodium salt* [14984-39-5] M 216.1 has [α]_D²⁰ +27 (c 10 in H₂O after 5 hours). The *phenylhydrazone* has m 141° (from MeOH). [Ehrlich & Schubert *Chem Ber* 62 1974, 2014 1929, DOI: 10.1002/cber.19290620818; Anderson & King *J Chem Soc* 5333 1961, DOI: 10.1039/JR9610005333; *Beilstein* 3 IV 2001.]

Genistin (4',5,7-trihydroxyisoflavone-7-D-glucoside) [529-59-9] C₂₁H₂₀O₁₀, M 432.4, m 256°, [α]_D²² -28 (c 0.6, 0.02N NaOH). Genistin is repeatedly crystallised from hot 80% EtOH/water and treated with charcoal (Nuchar) until free from saponin. The presence of saponin is detected by adding crystals to concentrated H₂SO₄ when the citron yellow colour changes to red, then purple. Pure genistin does not change colour. Its UV in 85% EtOH has λ_{max} at 262.5nm. [Walter *J Am Chem Soc* 63 3273 1941, DOI: 10.1021/ja01857a013; *Beilstein* 18 III/IV 2732.]

α-Gentiobiose (amygdalose, 6-O-α-D-glucopyranosyl-D-glucopyranose) [5995-99-5 (*bi-pyranose*)] C₃₆H₈₆O₁₁, M 342.3, m 86° (2MeOH), 189-195°, 195-197° (anhydrous), [α]_D²⁰ +11 (after 24 hours, c 4, H₂O). Crystallise α-gentiobiose from MeOH (retains solvent of crystallisation). It is best purified by conversion to the **α-octaacetate**, m 191-192° (recrystallise from absolute EtOH or CHCl₃ and excess absolute EtOH), [α]_D²⁰ +51.6 (c 4.3, CHCl₃) directly [Goldstein & Whelan *Methods in Carbohydrate Chemistry* I 313 1962, Academic Press, Hudson & Johnson *J Am Chem Soc* 39 1272 1917, DOI: 10.1021/ja02251a016], or from the **β-octaacetate** (see entry below) obtained by acetylation with Ac₂O and ZnCl₂, and hydrolysed (see entry below). It mutarotates from [α]_D²² +16° (3 minutes) to +9.5 (c 2, H₂O). [Reynolds & Evans *J Am Chem Soc* 60 2559 1938, DOI: 10.1021/ja01277a080; NMR analysis: Bradbury & Collins *Carbohydrate Research* 71 15 1979, DOI: 10.1016/S0008-6215(00)86056-5; *Beilstein* 17/7 V 203.]

β-Gentiobiose (6-O-β-D-glucopyranosyl-D-glucopyranose) [5996-00-9 (*bi-pyranose*), 554-91-6 (*one open ring*)] C₃₆H₈₆O₁₁, M 342.3, m 190-195°, [α]_D²⁰ +8 (after 6 hours, c 3, H₂O). β-Gentiobiose is best purified via the **octaacetate** which is recrystallised from MeOH, EtOH or better from methyl cellosolve by heating at 80°. The octaacetate (15g) is hydrolysed by suspending it in dry 0.05N NaOMe in MeOH (180ml) for 1 hour with occasional shaking at room temperature. Dilute this with H₂O to dissolve suspended matter, pass through Amberlite IR-120 and Duolite A-4 columns and the eluate is evaporated under reduced pressure to a syrup. Residual H₂O is removed by repeated distillation with absolute EtOH under reduced pressure. The syrup is dissolved in methyl cellosolve (~40ml), filtered, nucleated and placed in an oven at 80°. The crystals are filtered off, washed with absolute EtOH (yield 6.7g, 89%), dried and have m 187-189°. Further recrystallisation from methyl cellosolve gives m 190°, and mutarotates from [α]_D²⁸ -1.5 (initial) to +10.6 (final, c 4, H₂O). The pure **β-octaacetate** has m 193° (crystallised from 95% EtOH) and has [α]_D²⁰ -5 (c 1.8, CHCl₃). [Goldstein & Whelan *Methods in Carbohydrate Chemistry* I 313 1962, Academic Press, *Beilstein* 17/7 V 203.]

Glucamine (glycamine, 1-amino-1-deoxy-D-glucitol) [488-43-7] C₆H₁₅NO₅, M 181.2, m 127°, [α]_D²⁰ -8 (c 10, H₂O), pK_{Est} ~9.0. Crystallise glucamine from MeOH or aqueous MeOH and store it in a CO₂-free atmosphere. For the *N-methylglucamine* derivative see below. [Holly et al. *J Am Chem Soc* 72 5416 1950, DOI: 10.1021/ja01168a014; Karrer et al. *Helv Chim Acta* 20 83 1937, Kagan et al. *J Am Chem Soc* 79 3541 1957, DOI: 10.1021/ja01570a063; Groggins & Stirton *Ind Eng Chem* 29 1353 1937, DOI: 10.1021/ie50336a009.]

D-Gluconamide [3118-85-2] C₆H₁₃NO₆, M 195.2, m 142-143°, 144°, [α]_D²³ +31 (c 2, H₂O). Crystallise D-gluconamide from EtOH or 1,2-dimethoxyethane. It mutarotates slowly and hydrolytically in aqueous solution from +30.7 to +13.8 in 80 hours to give **ammonium D-gluconate** which recrystallises from 95% EtOH with m 153-154° and [α]_D²⁵ +11.8 (c 3, H₂O). [Wolfrom et al. *J Am Chem Soc* 80 944 1958, DOI: 10.1021/ja01537a049; *Beilstein* 3 VI 1259.]

D-Glucono- δ -lactone [90-80-2] $C_6H_{10}NO_6$, M 178.1, m 152-153°, 160°(dec), $[\alpha]_{546}^{20}$ +76 (c 4, H_2O). Crystallise D-glucono- δ -lactone from ethylene glycol monomethyl ether and dry it for 1 hour at 110°. It can be freed from other sugars *via* a column of Celite and charcoal (750g of each, 90 x 7.5cm) which is washed with 0.01N formic acid until the pH of the wash is equal to that of the entering acid. The lactone is applied in H_2O and eluted with 0.01N formic acid (7L), then eluted with 7.5% EtOH/0.01N formic acid (8L), then 15% EtOH/0.01N formic acid (8L) which removes pentose and isomaltose (the optical rotation of the eluates are used for sugar detection) and finally elution with aqueous formic acid provides glucolactone which is obtained by evaporating or freeze drying. Its solubility in H_2O is 60%, and 1% in EtOH. A solution in H_2O is slightly acidic, and the lactone dissolves in an equivalent of aqueous NaOH to form *sodium D-gluconate* [527-07-1] M 218.1, has m 200-206°(dec), $[\alpha]_D^{25}$ +12 (c 10, H_2O), pK²⁵ 3.6. [Smith & Whelan *Biochemical Preparations* 10 127 1963, *Beilstein* 3 IV 1255.]

α - and β - Glucosamine (2-amino-2-deoxy-D-glucose) [3416-24-8] $C_6H_{13}NO_5$, M 179.2, m 110°(dec), $[\alpha]_D^{20}$ +28 to +48 (c 5, H_2O), pK²⁴ 7.71. Crystallise the amines from MeOH. The free base has been obtained from the *hydrochloride* (21.5g, see entry below) in a mixture of Et₃N (15ml) and EtOH (125ml) by shaking for 2 days at room temperature, and the solid Et₃N.HCl is filtered off and the process repeated with more Et₃N (3-4 times) until the **α -D-glucosamine** (15g) is free from Cl ions. It has m 88°, $[\alpha]_D^{20}$ +100 mutarotating to +47.5 (c 1, H_2O). When Et₂NH is used as base, the α - to β - conversion is complete giving **β -D-glucosamine**. The *pentaacetate* is purified by dissolving in CHCl₃, treating with charcoal, drying (MgSO₄), evaporating the solvent, and adding a little dry Et₂O to induce crystallisation. It has m 124-126, $[\alpha]_D^{20}$ +113 (c 1, CHCl₃) after 16 hours in a desiccator. [Leaback *Biochemical Preparations* 10 118 1963.] The *N-acetyl derivative*, m 203-205° from MeOH/Et₂O (dry in vacuum P₂O₅) has $[\alpha]_D^{20}$ +75 mutarotating to +41 (c 2, H_2O); this derivative can also be purified by dissolving in the minimum volume of H_2O to which is added 7-8 volumes of EtOH followed by Et₂O until turbid and keeping at ~20° to crystallise. Wash the crystals with MeOH then Et₂O and dry *in vacuo* over P₂O₅. [Horton *Biochemical Preparations* 11 1 1966.]

D-Glucosamine hydrochloride [66-84-2] $C_6H_{13}NO_5 \cdot HCl$, M 215.6, m 190-194°(dec), >300°, $[\alpha]_D^{25}$ +71.8 (after 20 hours, c 4, H_2O), pK²⁴ 7.71. Crystallise the hydrochloride from 3M HCl, water, and finally water/EtOH/acetone as for *galactosamine hydrochloride*. [Purchase & Braun *Org Synth* 26 36 1946, DOI: 10.15227/orgsyn.026.0036; Stacey & Webber *Methods in Carbohydrate Chemistry* I 228 1962, Academic Press.] The salt has also been purified by dissolving in the minimum volume of boiling H_2O (charcoal), filtering and adding a large excess of 95% EtOH (~4 volumes) and stirring vigorously for several hours. Collect the crystals after 4-6 hours to give **α - anomer** which mutarotates from $[\alpha]_D^{25}$ +100 to +72 (equilibrium, c 1, H_2O). A large amount of the **β -anomer** stays in solution. This can be precipitated from the filtrate by adding excess Et₂O. The mixture of α - plus β -anomers has $[\alpha]_D^{25.5}$ +68.8 (c 4.75, H_2O , mutarotating to +70.1) [Leaback *Biochemical Preparations* 10 118 1963]. Note that if Et₂NH is used instead of Et₃N, conversion to the **β -anomer** can be complete (see above). [Stacey et al. *Methods in Carbohydrate Chemistry* I 306 1962, Academic Press, *Beilstein* 4 IV 2018.]

α -D-Glucose [492-62-6] $C_6H_{12}O_6$, M 180.2, m 83° (monohydrate), 146° (anhydrous), mutarotates from $[\alpha]_D^{20}$ +112 to +52.5 (after 24 hours, c 4, H_2O), pK²⁵ 12.46. Recrystallise α -D-glucose slowly from aqueous 80% EtOH, then dry it over P₂O₅ *in vacuo*. Alternatively, crystallise it from water at 55°, then dry it for 6 hours in a vacuum oven between 60-70°/2mm. Its solubilities are: H_2O (~50%), EtOH (1.7%). [Hendricks et al. *J Am Chem Soc* 56 99 1934, DOI: 10.1021/ja01316a029; *Beilstein* 1 IV 4302.] [For equilibrium forms see *Adv Carbohydr Chem* 42 15 1984, DOI: 10.1016/S0065-2318(08)60122-5; Angyal & Pickles *Aust J Chem* 25 1711 1972, DOI: 10.1071/CH9721711.]

β -D-Glucose [50-99-7] $C_6H_{12}O_6$, M 180.2, m 148-150°, 150-152°, mutarotates from $[\alpha]_D^{20}$ +18.7 to +52.5 (after 24 hours, c 4, H_2O). Crystallise β -D-glucose from hot glacial acetic acid or pyridine. Traces of solvent are removed by drying in a vacuum oven at 75° for >3 hours. [Dean & Gottfried *Adv Carbohydr Chem* 5 127 1950, DOI: 10.1016/S0096-5332(08)60337-9; Kjaer & Lindberg *Acta Chem Scand* 13 1713 1959, DOI: 10.3891/acta.chem.scand.13-1713; Whistler & Miller *Methods in Carbohydrate Chemistry* I 130 1962, Academic Press, *Beilstein* 1 IV 4306.] [For equilibrium forms see Angyal *Adv Carbohydr Chem* 42 15 1984, DOI: 10.1016/S0065-2318(08)60122-5; Angyal & Pickles *Aust J Chem* 25 1711 1972, see reference above.]

α -D-Glucose pentaacetate [604-68-2] $C_{16}H_{22}O_{11}$, M 390.4, m 110-111°, 112-113°, $[\alpha]_{546}^{20} +119$, $[\alpha]_D^{20} +102$ (c 5, $CHCl_3$). Crystallise it from MeOH, EtOH or three recrystallisations from 95% EtOH. [Wolfrom & Thompson *Methods in Carbohydrate Chemistry* II 212 1963, Academic Press, *Beilstein* 17/7 V 318.]

β -D-Glucose pentaacetate [604-69-3] $C_{16}H_{22}O_{11}$, M 390.4, m 130-132°, 131-132°, $[\alpha]_{546}^{20} +5$ (c 5, $CHCl_3$). Crystallise the pentaacetate from MeOH or EtOH. It is best purified by recrystallising 160g from 1L of 95% EtOH (charcoal) and filtering hot. It is important that as soon as the temperature of the filtrate cools to ~20° the pentaacetate is filtered off. **Note** that some α -D-isomer will crystallise out in a prolonged crystallisation period. Further crystallisation in this manner and drying in a vacuum over $CaCl_2$ will give **pure** β -D-anomer with m 132°, $[\alpha]_D^{20} +4$ (c 5, $CHCl_3$). [Wolfrom & Thompson *Methods in Carbohydrate Chemistry* II 212 1963, Academic Press, Krahle & Cori *Biochemical Preparations* 1 33 1955, *Beilstein* 17/7 V 319.]

D-Glucose phenylhydrazone [3713-25-5] $C_{12}H_{18}N_2O_5$, M 358.4, m three forms. Crystallise the hydrazone from 70% aqueous EtOH or EtOH/Et₂O. Three forms have been described: ' α ' form m 159°, 160° which mutarotates from $[\alpha]_D^{20} -87$ to -50 (H_2O) [Fischer *Chem Ber* 20 821 1887, DOI: 10.1002/cber.188702001187; Behrend *Justus Liebigs Ann Chem* 353 106 1907, DOI: 10.1002/jlac.19073530104], ' β ' form m 140-141°, 142° which mutarotates from $[\alpha]_D^{20} -2$ to -50 (H_2O) [Behrend & Lohr *Justus Liebigs Ann Chem* 362 78 1908, DOI: 10.1002/jlac.19083620107], and Skraup's form m 115-116° which mutarotates from $[\alpha]_D^{20} -70$ to -47 (H_2O) [Skraup *Monatsh Chem* 10 401 1889, DOI: 10.1007/BF01516449; Butler & Crechter *J Am Chem Soc* 51 3161 1929, DOI: 10.1021/ja01385a042]. These mutarotate to the **formazan**. [*Beilstein* 1 IV 4322, Mester & Major *J Am Chem Soc* 77 4297 1955, DOI: 10.1021/ja01621a037; Stanek et al. *The Monosaccharides*, Academic Press 1963, pp 539-541, 543.]

D-Glucuronic acid [6556-12-3] $C_6H_{10}O_7$, M 194.1, m 159-161°, 165°, $[\alpha]_D^{20} +36$ (c 6, H_2O , 2 hours, mutarotating from +11.5°), pK_2^{20} 3.18. Crystallise the acid from EtOH or EtOAc, wash it with MeOH and dry it *in vacuo* to give the ' **β** ' form. Heating converts it to the lactone (see below). The **sodium salt monohydrate** [207300-70-7] M 234.1 has m ~136-138°(dec) $[\alpha]_D^{20} +21$ (c 2, H_2O after 2 hours). [Sutter & Reichstein *Helv Chim Acta* 21 1210 1938, DOI: 10.1002/hlca.193802101150; *Beilstein* 3 H 886, 3 IV 1997.]

D-Glucurono-6,3-(δ)-lactone [32449-92-6] $C_6H_8O_6$, M 176.1, m 175-177°, $[\alpha]_{546}^{20} +22$ (after 24 hours, c 10, H_2O). Dissolve the lactone or mixture of lactone and acid in H_2O and concentrate the solution on a steam bath until crystallisation begins. Cool rapidly to room temperature with stirring. After 2 hours the product is filtered off, washed with cold EtOH and dried to m 174-175° and $[\alpha]_D^{25} +19.8$ (c 5.2, H_2O). The amount of **free acid** can be obtained by titration of an ice-cold aqueous solution with standard alkali. It can be recrystallised from EtOH, EtOH/ H_2O or MeOH, and the highest recorded m is 180°. [Stacey *J Chem Soc* 1529 1939, DOI: 10.1039/JR9390001529; Mehlretter et al. *J Am Chem Soc* 73 2424 1951, DOI: 10.1021/ja01150a005; *Beilstein* 18 III/IV 3055, 18/5 V 33.]

D(+)-Glycogen [9005-79-2] $(C_6H_{10}O_5)_n$, M 25,000-100,000, m 270-280°(dec), $[\alpha]_{546}^{20} +216$ (c 5, H_2O). A 5% aqueous solution (charcoal) of D(+)-glycogen is filtered, and an equal volume of EtOH is added. After standing overnight at 3° the precipitate is collected by centrifugation, washed with absolute EtOH, then EtOH/diethyl ether (1:1) and diethyl ether, and dried. [Sutherland & Wosilait *J Biol Chem* 218 459 1956, <http://www.jbc.org/content/218/1/459>.]

Glycyrrhizic acid ammonium salt (3 H_2O) (Glycyrrhizin) [53956-04-0] $C_{42}H_{62}O_{16}.NH_3$, M 823.0, m 210°(dec), 220°(dec, sintering at 170°), $[\alpha]_{546}^{20} +60$ (c 1, 50% aqueous EtOH), $[\alpha]_D^{20} +46.9$ (c 1.5, 40% EtOH), $pK_{Est} \sim 4.0$. It is the salt of a glucosylglucoside of a pentacyclic triterpene acid present in *glycyrrhiza* root (licorice). Crystallise the ammonium salt from glacial acetic acid, then dissolve it in ethanolic ammonia and evaporate. The **pentahydrate** forms needles from 75% aqueous EtOH, m 212-217°(dec), with UV λ_{max} at 248nm (ϵ 11,400, in 0.1mM EtOH). It has an intensely sweet taste. The **free acid, 18 β -glycyrrhetinic acid (Enoxolone)** [1449-05-4] M 470.7 crystallises from 75%v/v EtOH/ H_2O with m 284-285°, with UV similar to the glucoside salt. [Karrer et al. *Helv Chim Acta* 4 100 1921, DOI: 10.1002/hlca.19210040108; Lythgoe & Trippett *J Chem Soc* 1983, 1987 1950, DOI: 10.1039/JR9500001983; Marsh & Levvy *Biochem J* 63 9 1956, DOI: 10.1042/bj0630009; *Beilstein* 18 IV 5156.]

Heparin (from pig intestinal mucosa) [9005-49-6] ($\text{C}_{12}\text{H}_{19}\text{NO}_{20}\text{S}_3$)_n, $M_r \sim 12,000\text{--}15,000$, amorphous, $[\alpha]_D^{20} \sim +55$ (H_2O). Most likely contaminants are mucopolysaccharides including heparin sulfate and dermatan sulfate. Purify heparin by precipitation with cetylpyridinium chloride from saturated solutions of high ionic strength. [Cifonelli & Roden *Biochemical Preparations* **12** 12 1968, Wolfrom et al. *J Org Chem* **29** 540 1964, DOI: 10.1021/jo01026a005; Foster & Huggard *Adv Carbohydr Chem* **10** 335-368 1955, DOI: 10.1016/S0096-5332(08)60397-5] Useful anticoagulant, with protamine sulfate [9007-31-2] as antidote.

Heparin (sodium salt) [9041-08-1] $M_r \sim 3000$ (low Mol Wt, Bovine), amorphous, $[\alpha]_D^{20} +47$ (c 1.5, H_2O). Dissolve the salt in 0.1M NaCl (1g/100ml) and precipitate it by adding EtOH (150ml). [Wolfrom et al. *J Org Chem* **29** 540 1964, DOI: 10.1021/jo01026a005; Foster & Huggard *Adv Carbohydr Chem* **10** 335-368 1955, DOI: 10.1016/S0096-5332(08)60397-5.]

***n*-Heptyl- β -D-glucopyranoside** [78617-12-6] $\text{C}_{13}\text{H}_{26}\text{O}_6$, M 278.4, m 74-77°, 76-77°, $[\alpha]_D^{20} -34.2$ (c 5, H_2O). Purify this non-ionic detergent by repeated crystallisation from Me_2CO which is a better solvent than EtOAc. Its CMC (critical micellar concentration) in H_2O is 79mM. The *acetate* has m 66-68.5° and $[\alpha]_D^{20} -20.5$ (c 4, CHCl_3) [Pigman & Richtmyer *J Am Chem Soc* **64** 369 1942, DOI: 10.1021/ja01254a041]. [Beilstein **17** IV 2936.]

Heptyl- β -D-1-thioglucopyranoside [85618-20-8] $\text{C}_{13}\text{H}_{26}\text{O}_5\text{S}$, M 294.4, m 96.1-98.3°, 98-99°. The *tetraacetyl* derivative is purified by silica gel column chromatography, eluting with a $^*\text{C}_6\text{H}_6/\text{Me}_2\text{CO}$ gradient (up to 5% of Me_2CO), and recrystallising from *n*-hexane to give colourless needles m 72-74° (Erbing & Lindberg *Acta Chem Scand* **B30** 611 1976, DOI: 10.3891/acta.chem.scand.30b-0611; gave m 69-70°). Hydrolysis with an equivalent of base in methanol gives the *desired glucoside*. It provides a clear yellow solution in MeOH at 0.1g/ml. This is a non-ionic detergent for reconstituting membrane proteins and has a critical micelle concentration of 30 mM. [Shimamoto et al. *J Biochem (Tokyo)* **97** 1807 1985, PMID: 2863264; Saito & Tsuchiya *Chem Pharm Bull Jpn* **33** 503 1985, DOI: org/10.1248/cpb.33.503; for enzymic syntheses see Shinoyama et al. *Agric Biol Chem* **55** 1679 1991, <http://dx.doi.org/10.1080/00021369.1991.10870814>].

D(-)-Isoascorbic acid (araboascorbic acid, 5R-[R-1,2-dihydroxyethyl-3,4-dihydroxy-5H-furan-2-one]) [89-65-6] $\text{C}_6\text{H}_8\text{O}_6$, M 176.1, m 169°, 174°(0.5 H_2O , dec), $[\alpha]_D^{25} -16.8$ (c 2, H_2O), $[\alpha]_D^{20} +77$ (c 2, Me_2CO , acetonilidene derivative), pK^{18} 3.99 (4.23). Crystallise D(-)-isoascorbic acid from H_2O , EtOH or dioxane. Its UV has λ_{max} is at 245nm with ϵ 7,500 (EtOH). [Reichstein et al. *Helv Chim Acta* **17** 510, 516 1934, DOI: 10.1002/hlca.19340170157; Heslop et al. *J Chem Soc* 225 1944, DOI: 10.1039/JR9440000225; Beilstein **18** III/IV 3037, 18/5 V 26.]

2-Keto-L-gulonic acid (xylo-2-hexulonic acid) [526-98-7, 342385-52-8 $x\text{H}_2\text{O}$] $\text{C}_6\text{H}_{10}\text{O}_7$, M 194.1, m 159-162°, 171°, $[\alpha]_D^{20} -34.8$ (c 2, MeOH), $[\alpha]_D^{18} -48$ (c 1, H_2O), $pK_{\text{Est}} \sim 2.2$. Crystallise 2-keto-L-gulonic acid from half its weight of water, wash it with Me_2CO and dry it *in vacuo*. [Reichstein & Grüssner *Helv Chim Acta* **17** 311 1934, DOI: 10.1002/hlca.19340170136; NMR: Crawford et al. *J Am Chem Soc* **102** 2220 1980, DOI: 10.1021/ja00527a012; Beilstein **3** IV 1985.]

Lactobionic acid (4-O- β -D-galactopyranosyl-D-gulonic acid) [96-82-2] $\text{C}_{12}\text{H}_{22}\text{O}_{12}$, M 358.3, m 113-118°, 128-130°, mutarotates from $[\alpha]_D^{20} +53$ to $+22.6$ (c 3, after 4 hours in H_2O), pK^{25} 3.6. Crystallise lactobionic acid from water by addition of EtOH. The *calcium salt monohydrate* [110638-68-1] forms needles which give the *anhydrous* salt on crystallising from small volumes of absolute EtOH with $[\alpha]_D^{20} +23.5$ (c 3, H_2O). [NMR: Taga et al. *Bull Chem Soc Jpn* **51** 2278 1978, DOI: org/10.1246/bcsj.51.2278; Beilstein **17** III/IV 3392, 17/7 V 436.] Heating the acid in dioxane, dehydrates it to *lactobiono- δ -lactone* with m 195-196° and it mutarotates from $[\alpha]_D^{20} +53$ to $+22.6$ during 4 hours. [Fischer & Meyer *Chem Ber* **22** 362 1988, DOI: 10.1002/cber.18890220182; Ruff & Ollendorff *Chem Ber* **33** 1798 1900, DOI: 10.1002/cber.19000330262.]

α -Lactose monohydrate (β -D-galactopyranosyl-(1 \rightarrow 4)-D-glucose, milk sugar) [63-42-3, 5989-81-1]

C₁₂H₂₂O₁₁ · H₂O, M 360.3(hydrate), m 202.8°(dec), 220°(dec), 253-255° (252.4°), [α]_D²⁰ +52.3 (c 4.2, H₂O), (+55.4 also reported), pK²⁵ 12.2 (OH). Obtained from whey, α-lactose crystallises from water below 93.5° as the *hydrate* which can be dried at 80°/14mm. [Horst & Wiersma *Recl Trav Chim Pays-Bas* **72** 878 1953, DOI: 10.1002/recl.19530721009; *Beilstein* **17** III/IV 3066.] It is an analytical standard.

Lactulose (4-O-β-D-galactopyranosyl-D-fructose) [4618-18-2] C₁₂H₂₂O₁₁, M 342.2, m 168.5-169°(dec), [α]_D²⁰ -57 (c 1, H₂O). Crystallise lactulose from MeOH or 50% MeOH. It mutarotates from [α]_D²⁰ -11.9 to -50.7 (c 1, H₂O). [Montgomery & Hudson *J Am Chem Soc* **52** 2101, 2104 1930, DOI: 10.1021/ja01368a060; *Beilstein* **17** III/IV 3094, **17/7** V 214.] NMR in Me₂SO at 24° shows 0% α-pyranose, 27% β-pyranose, 20% α-furanose and 52% β-furanose forms [Angyal *Adv Carbohydr Chem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5].

Lanatoside A (digitoxigenin monoacetyl-tetraglycoside) [17575-20-1] C₄₉H₇₆O₁₉, M 969.1, m 245-248°, [α]_D²⁰ +32 (EtOH). Crystallise lanatoside A from MeOH or 95% aqueous EtOH. Its solubility is 1/16000 (H₂O), 1/225 (CHCl₃), 1/40 (EtOH) and 1/20 (MeOH). [Stoll & Kreis *Helv Chim Acta* **16** 1049 1933, DOI: 10.1002/hlca.193301601136; Kuhn et al. *Helv Chim Acta* **45** 881 1962, DOI: 10.1002/hlca.19620450318; *Beilstein* **18** III/IV 1480.] It is **cardiotonic**.

Lanatoside B (gitoxigenin monoacetyl-tetraglycoside) [17575-21-2] C₄₉H₇₆O₂₀, M 985.1, m 233°(dec), 245-248°(dec), [α]_D²⁰ +35 (MeOH). Crystallise lanatoside B from MeOH. Its solubility is: 1/5600 (CHCl₃), 1/40 (EtOH) and 1/20 (MeOH), but is insoluble in H₂O. [Stoll & Kreis *Helv Chim Acta* **16** 1049 1933, DOI: 10.1002/hlca.193301601136; Okano et al. *Chem Pharm Bull Jpn* **5** 171 1957, DOI: org/10.1248/cpb1953.5.171; *Beilstein* **18** III/IV 2465.] It is **cardiotonic**.

Lanatoside C (digoxigenin monoacetyl-tetraglycoside) [17575-22-3] C₄₉H₇₆O₂₀, M 985.1, m 246-248°, [α]_D²⁰ +33.5 (c 1.85, 95% EtOH). Crystallise lanatoside C from MeOH. Its solubility is: 1/17000 (H₂O), 1/1500-2000 (CHCl₃) and 1/45 (EtOH). [Stoll & Kreis *Helv Chim Acta* **16** 1049 1933, DOI: 10.1002/hlca.193301601136; Stoll & Kreis *Helv Chim Acta* **35** 1318 1952, DOI: 10.1002/hlca.19520350431; Okada et al. *Chem Pharm Bull Jpn* **23** 2039 1975, DOI: org/10.1248/cpb.23.2039; *Beilstein* **18** III/IV 2455.] It is **cardiotonic**.

α-L(+)-Lyxose [87-72-9, 1949-78-6] C₅H₁₀O₅, M 150.1, m 106-108°, [α]_D²⁰ +14 after 1 hour (c 6, H₂O). The **α-anomer** crystallises from propan-1-ol or EtOH, and the **β-anomer** crystallises from propan-2-ol. The **2,4-dinitrophenylhydrazone** has m 171-172° and [α]_D²⁰ -31 (pyridine). In D₂O it has 21% of the **α-pyranose** form. The **2-methyl ether** has m 120-121° and [α]_D²⁰ +6 (c 1, H₂O). [Angyal *Adv Carbohydr Chem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5; Bentley *J Am Chem Soc* **79** 1720 1957, DOI: 10.1021/ja01564a053; *Beilstein* **1** I 439, **1** IV 4232.]

β-D(-)-Lyxose [(2R,3S,4S,5R)-tetrahydro-2H-pyran-2,3,4,5-tetraol] [608-47-9, 107655-34-5, 1114-34-7] C₅H₁₀O₅, M 150.1, m 118-119°, 120-122°, 152°, [α]_D²⁰ -14 (c 4, H₂O), α-anomer has m 105-107° mutarotates from [α]_D²⁰ +5.6 to -18.8 (c 4, H₂O). Crystallise β-D-lyxose from EtOH or aqueous 80% EtOH by slow crystallisation. Dry it under vacuum at 60°, and store it in a vacuum desiccator over P₂O₅ or CaSO₄. [*Beilstein* **1** IV 4230, Overend et al. *J Chem Soc* 3487 1961, DOI: 10.1039/JR9610003487.] The ¹H NMR in D₂O has δ: α-H (pyranose) 5.39 (J 4.0 Hz), β-H (pyranose) 5.19 (J ~0 Hz), α-H (furanose) 5.26 (J 4.0 Hz), β-H (furanose) 5.24 (J 4.5 Hz), and at 31° in D₂O it consists of 70% α-pyranose, 28% β-pyranose, 1.5% α-furanose and 0.5% β-furanose [Angyal & Pickles *Aust J Chem* **25** 1711 1972, DOI: 10.1071/CH9721711].

Maltose monohydrate (4-O-α-D-glucopyranosyl-D-glucose) [6363-53-7] C₁₂H₂₂O₁₁ · H₂O, M 360.3, m 109-110°, 118°, mutarotates from [α]_D²⁰ +111.7 to +130.4 (c 4, H₂O). Purify maltose by chromatography from aqueous solution on a charcoal/Celite (1:1) column, wash it with water to remove glucose and other monosaccharides, then elute it with aqueous 75% EtOH. Crystallise it from water, aqueous EtOH or EtOH containing 1% nitric acid. Dry it as the **monohydrate** at room temperature under vacuum over H₂SO₄ or P₂O₅. Also purify it by dissolving it in MeOH, evaporating to a syrup which on standing for 12 hours in contact with

1/10th its volume of H₂O gives crystals of the *monohydrate*. Its iodine number is 55.5. The **osazone** has **m 200°(dec)** and $[\alpha]_D^{20} +58$ (c 1.4, H₂O). [Haworth et al. *J Chem Soc* 791 1937, DOI: 10.1039/JR9370000791; Haworth & Leitch *J Chem Soc Trans* **115** 809 1919, DOI: 10.1039/CT9191500809; for maltose & melibiose see Haworth et al. *J Chem Soc* 3146 1927, DOI: 10.1039/JR9270003146; *Beilstein* **17** III/IV 3057, **17** V 189.]

D-Mannitol (Mannite) [69-65-8] C₆H₁₄O₆, **M 182.2**, **m 166.1°**, **168-170°**, $[\alpha]_{546}^{20} +29$ (c 10, after 1 hour in 8% borax solution), $[\alpha]_D^{22} -4.0$ (c 0.5, DMF), **pK¹⁸ 13.5**. D-Mannitol is crystallised from EtOH, MeOH or H₂O and dried at 100°. [Thamsen *Acta Chem Scand* **6** 270, 279, 280 1952, DOI: 10.3891/acta.chem.scand.06-0270; *Beilstein* **1** IV 2841.]

Mannitol hexanitrate [15825-70-4] C₆H₈N₆O₁₈, **M 452.2**, **m 112-113°**, $[\alpha]_D^{22} +45.8$ (c 1.4, EtOH). The hexanitrate crystallises from EtOH or dilute EtOH as silky white needles. **EXPLOSIVE (on detonation)**. [Hayward *J Am Chem Soc* **73** 1974 1951, DOI: 10.1021/ja01149a023; Patterson & Todd *J Chem Soc* 2876 1929, DOI: 10.1039/JR9290002876; *Beilstein* **1** IV 2841.]

α-D(+)-Mannose [3458-28-4, 31103-86-3] C₆H₁₂O₆, **M 180.2**, **m 132°**, $[\alpha]_D^{20}$ mutarotates from **+29.9 to +14** (c 4, H₂O). Crystallise α-D(+)-mannose repeatedly from EtOH, aqueous 80% EtOH, AcOH or MeOH/propan-2-ol and then dry it *in vacuo* over P₂O₅ at 60°. [For ¹H NMR and equilibrium forms see Angyal *Adv Carbohydr Chem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5; and Angyal & Pickles *Aust J Chem* **25** 1711 1972, DOI: 10.1071/CH9721711; *Beilstein* **1** IV 4328.]

D(+)-Melezitose hydrate (O-α-D-glucopyranosyl-(1 → 3)-β-fructofuranosyl-(2 → 1)-α-D-glucopyranose [597-12-6, 207511-10-2] C₁₈H₃₂O₁₆ · xH₂O, **M 540.5**, **m 153-154°(dec)**, **2H₂O**, **160°(dec)**, $[\alpha]_D^{20} +88$ (c 2, H₂O for dihydrate) and $[\alpha]_D^{20} +91.7$ (c 2, H₂O for anhydrous). D(+)-Melezitose crystallises from aqueous EtOH as the *monohydrate* and water as the *dihydrate*, and is then dried at 110° (*anhydrous*). It is also purified by dissolving in an equal volume of H₂O, filtering into a crystallising dish and allowing to stand (loosely covered) for several weeks undisturbed at 20°. The crystals of clear prisms are wiped carefully and dried in air. They *effloresce* at once losing 3.35% of their weight, and after 3 days in air the loss is for 1H₂O from the *dihydrate*. Drying at 110° for 6 hours *in vacuo* yields the *anhydrous* form which after 2 days in air absorbs H₂O to give the *monohydrate*. The *monohydrate* is also obtained by dissolving it in an equal weight of H₂O at 60° and adding 4 volumes of 95% EtOH, and drying in air overnight. [Richtmyer & Hudson *J Org Chem* **11** 610 1946, DOI: 10.1021/jo01175a026; *Beilstein* **17** III/IV 3815, **17/8** V 414.]

D(+)-Melibiose (2H₂O) (6-O-α-D-galactopyranosyl-D-glucose) [585-99-9, *monohydrate* 66009-10-7] C₁₂H₂₂O₁₁ · xH₂O, **M 360.3**, **m 84-85°**, **178-181°**, **182°**, **184-185°**, $[\alpha]_D^{20} +135$ (c 5, after 10 hours H₂O). D(+)-Melibiose crystallises as a *hydrate* from water or aqueous EtOH. The **α-anomer** is obtained by recrystallising 1g from a mixture of 0.35ml of H₂O and 0.2ml of EtOH. It crystallises easily with **m 179°** and mutarotates from $[\alpha]_D^{20} +166$ to +142.3 (220 minutes, c 4, H₂O). Crystallisation from MeOH gives the *anhydrous* form which it hydrates to the *monohydrate* in air. The **β-anomer dihydrate** has **m 85-86°** and mutarotates from $[\alpha]_D^{20} +123.5$ to +143.1 (c 4, H₂O concentration for anhydrous). [Haworth et al. *J Chem Soc* 3146 1927, DOI: 10.1039/JR9270003146; Fletcher & Diehl *J Am Chem Soc* **74** 5774 1952, DOI: 10.1021/ja01142a512; *Beilstein* **17** III/IV 3075, **17/7** V 206.]

N-Methyl-D(-)-glucamine (Meglumine) [6284-40-8] C₇H₁₇NO₅, **M 195.2**, **m 128-129°**, **129-131.5°**, $[\alpha]_{546}^{20} -19.5$ (c 2, H₂O), **pK²⁸ 9.62**. Crystallise N-methyl-D(-)-glucamine from MeOH. Its solubility in H₂O is 10%. [Karrer & Herkenrath *Helv Chim Acta* **20** 83 1937, DOI: 10.1002/hlca.19370200109, also for other N-alkyl derivatives, *Beilstein* **4** IV 1914.]

N-Methyl α-L-glucosamine [42852-95-9] C₇H₁₅NO₅, **M 193.2**, **m glass**, $[\alpha]_D^{25} -65$ (c 1, MeOH) **pK_{Est} ~9**. The *hydrochloride* crystallises from EtOH as *hygroscopic* needles with **m 160-163°**, $[\alpha]_D^{25} +103$ mutarotating to -88 after 24 hours (c 0.6, H₂O), and gives the *free base* as a glass. The *pentaacetate* crystallises from CHCl₃/Et₂O with **m 160.5-161.5°**, $[\alpha]_D^{25} -100$ (c 0.7, CHCl₃), and the *N-acetate* crystallises from CHCl₃/MeOH with **m 165-166°**, $[\alpha]_D^{25} -51$ (c 0.4, H₂O). [Kuehl et al. *J Am Chem Soc* **68** 536 1946, DOI: 10.1021/ja01207a520; **69** 3032 1947, DOI: 10.1021/ja01204a029; Lemieux & Wolfrom *Adv Carbohydr Chem* **3**

337 1948, DOI: 10.1016/S0096-5332(08)60034-X; *Beilstein* **4** IV 2032.]

Methyl α -D-glucoside (methyl α -D-glucopyranoside) [97-30-3] $C_7H_{14}O_6$, **M 194.2**, **m 168°**, **166-169°**, $[\alpha]_D^{20}$ **+158.9** (c **10**, H_2O), **pK²⁵ 13.71**. Crystallise methyl α -D-glucoside from MeOH or EtOH. Its solubility in H_2O is 10%. [Ferrier et al. *Carbohydr Research* **27** 55 1973, DOI: 10.1016/S0008-6215(00)82424-6; *Beilstein* **17/7** V 13.]

Methyl β -D-glucoside (methyl β -D-glucopyranoside) [7000-27-3] $C_7H_{14}O_6$, **M 203.2** (**0.5 H_2O**), **m 107-109°**, $[\alpha]_D^{20}$ **-33** (c **10**, H_2O). Crystallise methyl β -D-glucoside from MeOH or EtOH. Its solubility in H_2O is 10%. [Ferrier et al. *Carbohydr Research* **27** 55 1973, DOI: 10.1016/S0008-6215(00)82424-6; *Beilstein* **17/7** V 10.]

4-Methylumbellifer-7-yl- α -D-glucopyranoside [17833-43-1] $C_{16}H_{18}O_8$, **M 338.3**, **m 171-173°(hydrate)**, **209-210°**, **221-222°**, $[\alpha]_D^{20}$ **+162°** (c **0.5**, pyridine). Recrystallise 4-methylumbellifer-7-yl- α -D-glucopyranoside from hot H_2O or EtOH. [Courtin-Duchateau & Veyrières *Carbohydr Research* **65** 23, 29 1978, DOI: 10.1016/S0008-6215(00)84209-3; *Beilstein* **18** IV 443.]

4-Methylumbellifer-7-yl- β -D-glucopyranoside [18997-57-4] $C_{16}H_{18}O_8$, **M 338.3**, **m 211-213°**, **211°**, $[\alpha]_D^{20}$ **-68°** (c **0.5**, pyridine), **-89.5** (c **0.5**, H_2O for half hydrate). 4-Methylumbellifer-7-yl- β -D-glucopyranoside crystallises as the *half hydrate* from hot H_2O . It has $E^{mM} = 13.9$ at 317.5nm (MeOH). [Constantzas & Kocourek *Coll Czech Chem Commun* **24** 1099 1959, DOI: org/10.1135/ccccc19591099; De Re et al. *Ann Chim (Rome)* **49** 2089 1959, Courtin-Duchateau & Veyrières *Carbohydr Research* **65** 23, 29 1978, DOI: 10.1016/S0008-6215(00)84209-3; *Beilstein* **18** III/IV 5152, **8** IV 433, **18/7** V 616.]

Naringin (4',5,7-trihydroxyflavanone 7-rhamnoglucoside) [10236-47-2] $C_{27}H_{32}O_{14}$, **M 580.5**, **m ~83°** (**6 H_2O**), **166°** (also reported), **171°** (**2 H_2O**), $[\alpha]_D^{19}$ **-90** (c **1**, EtOH), $[\alpha]_{546}^{20}$ **-107** (c **1**, EtOH). This bitter principle from grape juice crystallises from water to give the hydrate with 6-8 H_2O which when dried at 110° gives the *dihydrate*. Its solubility in H_2O is 0.1% at 40° and 10% at 75°. The **2,4-dinitrophenylhydrazone** crystallises from aqueous dioxane with **m 246-247°** [Douglass et al. *J Am Chem Soc* **73** 4023 1951, DOI: 10.1021/ja01152a515]. [Pulley & von Loesecke *J Am Chem Soc* **61** 175 1939, DOI: 10.1021/ja01870a056; *Beilstein* **18** III/IV 2637, **18** V 528.]

2-Nitrophenyl- β -D-galactopyranoside [369-07-3] $C_{12}H_{15}NO_8$, **M 301.3**, **m 185-190°**, **193°**, **193-194°**, $[\alpha]_D^{18}$ **-51.9** (c **1**, H_2O). Purify 2-nitrophenyl- β -D-galactopyranoside by recrystallisation from EtOH. [Seidman & Link *J Am Chem Soc* **72** 4324 1950, DOI: 10.1021/ja01165a601; Snyder & Link *J Am Chem Soc* **75** 1758 1953, DOI: 10.1021/ja01103a531]. It is a *chromogenic substrate* for β -galactosidases [Jagota et al. *J Food Sci* **46** 161 1981, DOI: 10.1111/j.1365-2621.1981.tb14554.x]. [*Beilstein* **17/7** V 52.]

4-Nitrophenyl- α -D-galactopyranoside [7493-95-0] $C_{12}H_{15}NO_8$, **M 301.3**, **m 166-169°**, **173°**, $[\alpha]_D^{25}$ **+248** (c **1**, H_2O). Purify 4-nitrophenyl- α -D-galactopyranoside by recrystallisation from H_2O or aqueous EtOH. The *monohydrate* has **m 85°** which resolidifies and melts again at **151-152°** (the *hemihydrate*), then resolidifies again and melts at **173°** to give the *anhydrous* form. Drying the monohydrate at 60° yields the hemihydrate, and drying at 100° gives the anhydrous compound. The *tetraacetate* has **m 147°** after drying at 100°. [Jermyn *Aust J Chem* **15** 569 1962, DOI: 10.1071/CH9620569; Helfrich & Jung *Justus Liebigs Ann Chem* **589** 77 1954, DOI: 10.1002/jlac.19545890108.] It is a *chromogenic substrate* for α -galactosidase [Dangelmaier & Holmsen *Anal Biochem* **104** 182 1980, DOI: 10.1016/0003-2697(80)90296-1]. [*Beilstein* **17/7** V 55.]

4-Nitrophenyl- β -D-galactopyranoside [3150-24-1] $C_{12}H_{15}NO_8$, **M 301.3**, **m 178°**, **178-181°**, **181-182°**, $[\alpha]_D^{20}$ **-83** (c **1**, H_2O). Purify the galactoside by recrystallisation from EtOH. [Hobikoshi *J Biochem (Tokyo)* **35** 39 1942, Goebel & Avery *J Exptl Medicine* **50** 521 1929, DOI: 10.1084/jem.50.4.521; Snyder & Link *J Am Chem Soc* **75** 1758 1953, DOI: 10.1021/ja01103a531.] It is a *chromogenic substrate* for β -galactosidases [Buoncore et al. *J Appl Biochem* **2** 390 1980]. [*Beilstein* **17/7** V 55.]

4-Nitrophenyl- α -D-glucopyranoside [3767-28-0] $C_{12}H_{15}NO_8$, **M 301.3**, **m 206-212°**, **216-217°** (sinters at

210°, $[\alpha]_D^{20}$ **+215 (c 1, H₂O)**. Purify 4-nitrophenyl- α -D-glucopyranoside by recrystallisation from H₂O, MeOH or EtOH. [Jermyn *Aust J Chem* **7** 202 1954, DOI: 10.1071/CH9540202; Montgomery et al. *J Am Chem Soc* **64** 690 1942, DOI: 10.1021/ja01255a060.] It is a **chromogenic substrate** from α -glucosidases [Oliveira et al. *Anal Biochem* **113** 188 1981, DOI: 10.1016/0003-2697(81)90064-6], and is a **substrate** for glucansucrases [Binder & Robyt *Carbohydr Research* **124** 287 1983, DOI: 10.1016/0008-6215(83)88464-X].

4-Nitrophenyl- β -D-glucopyranoside [2492-87-7] **C₁₂H₁₅NO₈, M 301.2, m 164°, 164-165°, 165°, $[\alpha]_D^{20}$ -107 (c 1, H₂O)**. Purify 4-nitrophenyl- β -D-glucopyranoside by recrystallisation from EtOH or H₂O. [Montgomery et al. *J Am Chem Soc* **64** 690 1942, DOI: 10.1021/ja01255a060; Snyder & Link *J Am Chem Soc* **75** 1758 1953, DOI: 10.1021/ja01103a531.] It is a **chromogenic substrate** for β -glucosidases [Weber & Fink *J Biol Chem* **255** 9030 1980, PMID: 6773958]. [Beilstein **17/7** V 53.]

N-Nonanoyl-n-methylglucamine (Mega-9) [85261-19-4] **C₁₆H₃₃NO₆, M 335.4, m 87-89°**. It is a non-ionic detergent which is purified as for n-decanoyl-N-methylglucamine above. [Hildreth *Biochem J* **207** 363 1982, DOI: 10.1042/bj2070363.]

Nonyl- β -D-glucopyranoside [69984-73-2] **C₁₅H₃₀O₆, M 306.4, m 67.5-70°, 70-71°, $[\alpha]_D^{20}$ -34.4 (c 5, H₂O), $[\alpha]_D^{25}$ -28.8 (c 1, MeOH)**. Purify nonyl- β -D-glucopyranoside by recrystallisation from Me₂CO or hexane/Et₂O and store it in well-stoppered containers as it is *hygroscopic*. It is a non-ionic detergent with a CMC (critical micellar concentration) of 6.5mM. [Pigman & Richtmyer *J Am Chem Soc* **64** 369 1942, DOI: 10.1021/ja01254a041.] It is a UV transparent non-ionic detergent for solubilising membrane proteins [Schwendener et al. *Biochem Biophys Res Commun* **100** 1055 1981, DOI: 10.1016/0006-291X(81)91930-6]. [Beilstein **17** III/IV 2937, **17/7** V 39.]

1-O-Octyl- β -D-glucopyranoside [29836-26-8] **C₁₄H₂₈O₆, M 292.4, m 62-65°, 63.8-65°, $[\alpha]_D^{20}$ -34° (c 4, H₂O)**. Purify octyl- β -D-glucopyranoside by recrystallisation from Me₂CO. It is *hygroscopic* and should be stored in a well-stoppered container. [Noller & Rockwell *J Am Chem Soc* **60** 2076 1938, DOI: 10.1021/ja01276a017; Pigman & Richtmyer *J Am Chem Soc* **64** 369 1942, DOI: 10.1021/ja01254a041.] It is a UV transparent non-ionic dialysable detergent for solubilising membrane proteins. Its CMC (critical micellar concentration) is 20-25mM in H₂O (20-25°), with average micellar wt of 25,000 and aggregation number of 84. The **2,3,4,6-tetraacetate** [38954-67-5] **C₂₂H₃₆O₁₀, M 460.5**, has **m 50-52°**, and $[\alpha]_D^{20}$ -16 (c 1.8, CHCl₃). The **α -D-isomer** with $[\alpha]_D^{20}$ +118 (c 1, MeOH) has similar solubilising properties. [Lazo & Quinn *Anal Biochem* **102** 68 1980, DOI: 10.1016/0003-2697(80)90318-8; Stubbs et al. *Biochim Biophys Acta* **426** 46 1976, DOI: 10.1016/0005-2736(76)90428-4; Beilstein **17/7** V 38.]

Pectic acid [9046-40-6] **(C₆H₈O₆)_n, M_r ~500,000, amorphous, $[\alpha]_D$ +250 (c 1, 0.1M NaOH)**. Citrus pectic acid (500g) is refluxed for 18 hours with 1.5L of 70% EtOH, and the suspension is filtered hot. The residue is washed with hot 70% EtOH and finally with ether. It is dried in a current of air, ground and dried for 18 hours at 80° under vacuum. [Morell & Link *J Biol Chem* **100** 385 1933, <http://www.jbc.org/content/100/2/385>.] It can be further purified by dispersing it in water and adding just enough dilute NaOH to dissolve the pectic acid, then passing the solution through columns of cation- and anion-exchange resins [Williams & Johnson *Ind Eng Chem (Anal Ed)* **16** 23 1944, DOI: 10.1021/i560125a007], and precipitating with two volumes of 95% EtOH containing 0.01% HCl. The precipitate is worked with 95% EtOH, then Et₂O, dried and ground. [Rees & Walsh *Angew Chem Int Ed* **16** 214 1977, DOI: 10.1002/anie.197702141; Rees *Adv Carbohydr Chem* **24** 267 1969, DOI: 10.1016/S0065-2318(08)60352-2.]

Pectin (1-4 linked heteropolysaccharide) [9000-69-5] **M_r 30,000-100,000, amorphous**. Dissolve the pectin in hot water to give a 1% solution, then cool, and make it to about 0.05M in HCl by addition of concentrated HCl, and precipitate it by pouring it slowly, with vigorous stirring into two volumes of 95% EtOH. After standing for several hours, the pectin is filtered through a nylon cloth, then re-dispersed in 95% EtOH and set aside overnight. The precipitate is filtered off, washed with EtOH/Et₂O, then Et₂O and dried in air. [Rees & Walsh *Angew Chem Int Ed* **16** 214 1977, DOI: 10.1002/anie.197702141; Rees *Adv Carbohydr Chem* **24** 267

1969, DOI: 10.1016/S0065-2318(08)60352-2.]

Pentaerythritol (2,2-bis[hydroxymethyl]-1,3-propanediol) [115-77-5] $C(CH_2OH)_4$, M 136.2, m 253-258°, 260.5°, 268-269°, b 276°/30mm. Reflux pentaerythritol with an equal volume of MeOH, then cool, and the precipitate is collected and dried at 90°. It can also be recrystallised from dilute aqueous HCl. After sublimation under high vacuum at 200° it has **m 265.5°**. Its solubility in H₂O is 10%. [*Beilstein* 18 III 2361, 1 IV 2812.]

Pentaerythritol tetraacetate [597-71-7] $C_{13}H_{20}O_8$, M 304.3, m 83-84°, 84-86°. Crystallise pentaerythritol tetraacetate from hot water, then leach it with cold water until the odour of acetic acid is no longer detectable. It also crystallises from 95% EtOH after dissolving in CHCl₃, washing with saturated NaHCO₃, then H₂O, drying over anhydrous CaCl₂ and evaporating. It has been prepared by acetolysis of the **tetranitrate** (see next entry) in 95% yield [Wolfrom et al. *J Am Chem Soc* 73 874 1951, DOI: 10.1021/ja01146a534]. [Breusch & Oğuzer *Chem Ber* 88 1511 1955, DOI: 10.1002/cber.19550881007; LeFèvre et al. *J Chem Soc* 16 1958, DOI: 10.1039/JR9580000016; *Beilstein* 1 IV 1812, 2 IV 264.]

Pentaerythritol tetranitrate [78-11-5] $C_5H_8N_4O_{12}$, M 316.2, m 140.1°, 141.3°, decomp >150°. Crystallise pentaerythritol tetranitrate from acetone or acetone/EtOH. When crystallised from H₂O at 0°, it may have **m 26-28°** (hydrate?). It detonates more easily than TNT on percussion. The **O-acetate**, when crystallised from EtOH, has **m 87-88°**. Although it has been distilled at 60°/2mm, distillation should **NOT** be attempted as it is **VERY EXPLOSIVE**. The **autoignition** temperature is **150°**. It is a vasodilator and used in the management of angina. [Marans et al. *J Am Chem Soc* 76 1304 1954, DOI: 10.1021/ja01634a032; Camp et al. *J Am Chem Soc* 77 751 1955, DOI: 10.1021/ja01608a059; *Beilstein* 1 IV 2816, 2 IV 264.]

Pentaerythrityl laurate (pentaerythrityl tetra-*n*-dodecanoate) [13057-50-6] $C_{53}H_{109}O_8$, M 865.3, m 50°, 50.8-51.8°. Crystallise the laurate from Me₂CO, Et₂O or petroleum ether. [Breusch & Oğuzer *Chem Ber* 88 1511 1955, DOI: 10.1002/cber.19550881007, and includes a large number of other esters.]

2-Phenylethyl-β-D-thiogalactoside [63407-54-5] $C_{14}H_{20}O_5S$, M 300.4, m 108°, $[\alpha]_D^{23}$ -32.2 (c 5, MeOH). Recrystallise the thiogalactoside from H₂O and dry in air to give the 1.5H₂O which has **m 80°**. The **anhydrous** surfactant is obtained by drying it at 78° over P₂O₅. [Helferich & Türk *Chem Ber* 89 2215 1956, DOI: 10.1002/cber.19560891002.]

Phenyl-β-D-galactopyranoside [2818-58-8] $C_{12}H_{16}O_6$, M 256.3, m 153-154°, 146-148°, 155-156°(dried at 105°), $[\alpha]_D^{20}$ -42 (c 1, H₂O). Recrystallisation of phenyl-β-D-galactopyranoside from H₂O gives the 0.5H₂O. [Conchie & Hay *Biochem J* 73 327 1959, DOI: 10.1042/bj0730327; IR: Whistler & House *Anal Chem* 25 1463 1953, DOI: 10.1021/ac60082a013.] It is an **acceptor substrate** for fucosyltransferase [Chester et al. *Eur J Biochem* 69 583 1976 DOI: 10.1111/j.1432-1033.1976.tb10944.x]. [*Beilstein* 17/7 V 47.]

Phenyl-β-D-glucopyranoside [1464-44-4] $C_{12}H_{16}O_6$, M 256.3, m 174-175° 174-176°, 176°, 176-178°, $[\alpha]_D^{20}$ -72.2 (c 1 for dihydrate, H₂O). Phenyl-β-D-glucopyranoside crystallises from H₂O with 2H₂O and can be dried *in vacuo* at 100°/P₂O₅. The dry preparation has $[\alpha]_D^{25}$ -70.7 (c 2, H₂O). [Robertson & Waters *J Chem Soc* 2729 1930, IR: Bunton et al. *J Chem Soc* 4419 1955, DOI: 10.1039/JR9550004419; Takahashi *Yakugaku Zasshi (J Pharm Soc Jpn)* 74 706 1954, Whistler & House *Anal Chem* 25 1463 1953, DOI: 10.1021/ac60082a013; UV: Lewis *J Am Chem Soc* 57 898 1935, DOI: 10.1021/ja01308a032.] It is a **substrate** for β-D-glucosidase [deBryne *Eur J Biochem* 102 257 1979, DOI: 10.1111/j.1432-1033.1979.tb06288.x]. [*Beilstein* 17 III/V 2946, 17/7 V 46.]

Phlorizin (2H₂O) (Phloridzin, phloretin 2'-O-β-D-glucoside) [60-81-1] $C_{21}H_{24}O_{10}$, M 436.4, m 106-109°, 110°, $[\alpha]_{546}^{20}$ -62 (c 3.2, EtOH). This flavonoid glucoside crystallises as the **dihydrate** from water and causes glycosuria. [Brazy & Dennis *Am J Physiol* 234 1279 1978, for effect of phloretin and theophylline on 3-O-methylglucose transport by intestinal epithelial cells see Randles & Kimmich *Am J Physiol (cell physiol)* 234 C64 1978, PMID: 629334; Zemplén & Bognár *Chem Ber* 75 1040 1942, DOI: 10.1002/cber.19420750903; *Beilstein* 17/7 V 177.] It is a glucoside in *Malus* (apple) species of trees, and is sweet with a bitter after taste.

D(+)-Raffinose (5H₂O) (Melitose, 6-O- α -D-galactopyranosyl-D-glucopyranosyl- β -D-fructo-furanose [17629-30-0 (5H₂O), 512-69-6 (anhydrous)] C₁₈H₃₂O₁₆, 5H₂O, M 594.5, m 80°, 80-82°, [α]_D²⁰ +124 (c 10, H₂O), [α]_D²⁰ +105 (c 1 for pentahydrate, H₂O), pK₁²⁵ 12.40, pK₂²⁵ 13.44, pK₃²⁵ 13.52. D(+)-Raffinose crystallises from H₂O, 90% aqueous EtOH or MeOH as the *pentahydrate*. The *anhydrous* sugar has m 132-135°. It has R_F 0.8 on TLC (Silica Gel, and 1:3:3 CHCl₃/butanone:/MeOH). The *undecaacetate* has been purified through an alumina column by elution with CHCl₃, and recrystallised from EtOH/MeOH/H₂O (3:2:5), with m 99-100°, and [α]_D²⁰ +92.8 (c 5.14, EtOH). [For pK see Coccioli & Vicedomini *Ann Chim (Rome)* **66** 269, 275 1976, and for ¹H NMR see Suami et al. *Carbohydr Research* **26** 234 1973, DOI: 10.1016/S0008-6215(00)85044-2; *Beilstein* **17** III/IV 3801, **17/8** V 403.]

L(+)- α -Rhamnose (H₂O) (6-deoxy-L-mannose) [10030-85-0 (H₂O), 3615-41-6 (anhydrous)] C₆H₁₂O₅, H₂O, M 182.2, m 90-92°, 101°, 105°, [α]_D¹⁸ -6.8 mutarotating to +9.1 (c 1, H₂O). Crystallise the rhamnose from H₂O or EtOH. It crystallises easily as the *monohydrate* by evaporating a solution in MeOH (90%) and H₂O (10%). It is also purified by dissolving in a small volume of EtOH, adding a few drops of H₂O and cooling. The ¹H NMR in D₂O at 44° shows 60% α -pyranose and 40% β -pyranose forms [Angyal *Adv Carbohydr Chem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5.] [Smith *J Chem Soc* 1035 1940, DOI: 10.1039/JR9400001035; McGeachin & Beevers *Acta Cryst* **10** 227,230 1957, DOI: 10.1107/S0365110X57000687; *Beilstein* **1** IV 4261.]

D(+)-Ribonic acid- γ -lactone [5336-08-3] C₅H₈O₅, M 148.12, m 80°, 84-86°, 85-87°, [α]_D²⁰ +18.3 (c 5, H₂O). Purify D(+)-ribonic acid- γ -lactone by recrystallisation from EtOAc. The *tribenzoate* has m 54-56° (from AcOH), [α]_D²⁵ +27 (c 2.37, Me₂NCHO), and the **3,5-O-benzylidene derivative** has m 230-231.5° (needles from Me₂CO-petroleum ether) and [α]_D²⁵ -177 (CHCl₃). [Chen & Joullié *J Org Chem* **49** 2168 1984, DOI: 10.1021/jo00186a018; Zinner & Voigt *J Carbohydr Research* **7** 38 1968, DOI: 10.1016/S0008-6215(00)81432-9.]

α -D(-)-Ribose [50-69-1] C₅H₁₀O₅, M 150.1, m 88-92°, 90°, [α]_D²⁰ -24 (after 24 hours, c 10, H₂O), pK₁²⁵ 12.22. Crystallise α -D(-)-ribose from aqueous 80% EtOH, dry it under vacuum at 60° over P₂O₅ and store it in a vacuum desiccator. It exhibits complex mutarotation with : [α]_D¹⁰ -23.1 (1.5 minutes), -21.3° (5 minutes), -19.5 (10 minutes), -19.1 (30 minutes), -21.2 (60 minutes), -23.1 (120 minutes), -23.7 (300 minutes), (c 4.5, H₂O) [Phelps et al. *J Am Chem Soc* **56** 747 1934, DOI: 10.1021/ja01318a506]. The ¹H NMR in D₂O at 44° shows 17% α -pyranose, 59% β -pyranose, 9% α -furanose and 15% β -furanose forms with **furanose** α -H at 5.34ppm (*J* = 3.0Hz) and β -H at 5.31 (*J* = 1.7Hz) [Angyal *Adv Carbohydr Chem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5; Angyal & Pickles *Aust J Chem* **25** 1711 1972, DOI: 10.1071/CH9721711]. The *phenylhydrazone* crystallises from aqueous pyridine in yellow needles, m 163-164°, and the *benzylphenylhydrazone* has m 127-128° [Sowden *J Am Chem Soc* **72** 808 1950, DOI: 10.1021/ja01158a041.] [*Beilstein* **1** IV 4211.]

(+)-Rutin (quercetin-3-rubinoside) See Vitamin P in ‘Physiologically Active....’, this chapter.

Saccharides. They are separated by anion-exchange chromatography. [Walborg & Kondo *Anal Biochem* **37** 320 1970, DOI: 10.1016/0003-2697(70)90054-0.]

D(-)-Salicin [2-(hydroxymethyl)phenyl- β -D-glucopyranoside] [138-52-3] C₁₃H₁₈O₇, M 286.3, m 204-208°, [α]_D²⁵ -63.5 (c ca 3, H₂O). Crystallise D(-)-salicin from EtOAc, EtOH or water and sublime it at 190-195°/12mm. [Armour et al. *J Chem Soc* 412 1961, DOI: 10.1039/JR9610000412; for IR see Pearl & Darling *J Org Chem* **24** 731 1959, DOI: 10.1021/jo01088a001; *Beilstein* **17** III/IV 2986, **17/7** V 113.]

Sennoside A (bianthraquinonyl-bis-glucoside *R,R*-enantiomer) [81-27-6] C₄₂H₃₈O₂₀, M 862.7, m 220-240°(dec), [α]_D²⁰ -164 (c 0.1, Me₂CO/H₂O 6:4), [α]_D²⁰ -24 (c 0.2, 70% aqueous dioxane). Sennoside A forms yellow crystals from aqueous acetone, 2-ethoxyethanol or large volumes of H₂O. [Stoll et al. *Helv Chim Acta* **32** 1892 1949, DOI: 10.1002/hlca.19490320613; *Beilstein* **17** III/IV 3403.] It is an analytical chromatography standard, and the laxative principle of *senna*.

Sennoside B (bianthraquinonyl-bis-glucoside *R,S*-enantiomer) [128-57-4] $C_{42}H_{38}O_{20}$, **M 862.7**, **m 209-212°(dec)**, $[\alpha]_D^{20}$ **-100** (c 2, Me_2CO/H_2O 7:1), $[\alpha]_D^{20}$ **-67** (c 0.2, **70% aqueous dioxane**). Sennoside B forms yellow crystals from aqueous acetone or large volumes of H_2O . [Stoll et al. *Helv Chim Acta* **32** 1892 1949, DOI: 10.1002/hlca.19490320613; *Beilstein* **17** III/IV 3402.] It is an analytical chromatography standard.

Sinigrin monohydrate (Myronate **K**, 1-thio- β -D-glucopyranose 1-[N-(sulfooxy)-3-butenimide] monopotassium salt) [64550-88-5] $C_{10}H_{18}NO_{10}S_2.K$, **M 415.5**, **m 125-127°, 128°, 127-129°, 179°(anhydrous)**, $[\alpha]_D^{20}$ **-17** (c 0.2, H_2O), pK_{Est} **<0**. Purify sinigrin by recrystallising it three times from EtOH and once from MeOH. The *tetraacetate* has **m 193-195°**, $[\alpha]_D^{20}$ **-16** (c 0.14, H_2O). [Benn & Ettlinger *JCS Chem Commun* 445 1965, DOI: 10.1039/C19650000445; Kjaer et al. *Acta Chem Scand* **10** 432 1956, DOI: 10.3891/acta.chem.scand.10-0432; Marsh & Waser *Acta Cryst (Sect B)* **26** 1030 1970, DOI: 10.1107/S0567740870003539.] It is a β -D-thioglucopyranoside *substrate* for thiogluconidase, and is present in horseradish. [MacLeod & Rossiter *Phytochem* **25** 1047 1986, DOI: 10.1016/S0031-9422(00)81551-4]. [*Beilstein* **31** H 476, **17** III/IV 3738.]

α -Solanine (solan-5-en-3 β -yl-[O³- β -D-glucopyranosyl-O²- α -L-rhamnopyranosyl- β -D-galactopyranoside], a solanidine triglycoside) [20562-02-1] $C_{45}H_{73}NO_{15}$, **M 868.1**, **m 285°(dec), 286°(dec) (sintering >190°)**, $[\alpha]_D^{20}$ **-58** (c 0.8, pyridine), pK^{15} **6.66**. Recrystallise α -solanine from EtOH, 85% aqueous EtOH, MeOH or aqueous MeOH as *dihydrate* **m 276-278°**. Its solubility in H_2O is 25mg/l, and 5% in pyridine, but it is very soluble in Et_2O and $CHCl_3$. The *hydrochloride* is gummy or amorphous but has been crystallised (**m ~212° dec**). It is present in potato sprouts and has insecticidal properties. [Kuhn et al. *Chem Ber* **88** 1492 1955, DOI: 10.1002/cber.19550881005; *Beilstein* **21** III/IV 1402.] For solanidine (aglycone) [80-78-4] see p 988.

Solasonine (solasodine-3-O-triglycoside) [19121-58-5] $C_{45}H_{73}NO_{16}$, **M 884.1**, **m 301-303° (sinters at ~296°)**, $[\alpha]_D^{20}$ **-75** (c 0.5, MeOH), pK_{Est} **~7.7**. Solasonine crystallises from aqueous 80% dioxane or MeOH in needles. [Bell & Briggs *J Chem Soc* 1 1942, DOI: 10.1039/JR9420000001; Briggs et al. *J Chem Soc* 4645 1961, DOI: 10.1039/JR9610004645; Briggs et al. *J Chem Soc* 2848 1963, DOI: 10.1039/JR9630002848.] The *picate* crystallises from 30% aqueous EtOH with **m 197-198°(dec)** [Briggs & Cambie *J Chem Soc* 1422 1958, DOI: 10.1039/JR9580001422]. [*Beilstein* **27** III/IV 2006.] For solasodine (aglycone) [126-17-0] see p 988.

D(-)-Sorbitol (D-glucitol) [50-70-4] $C_6H_{14}O_6$, **M 182.2**, **m 89-93° (hemihydrate), 98-100°, 110-111° (anhydrous)**, $[\alpha]_{546}^{20}$ **-1.8** (c 10, H_2O), pK^{60} **13.00**. Crystallise D(-)-sorbitol (as *hemihydrate*) several times from EtOH/water (1:1), then dry it by fusing and storing over anhydrous $MgSO_4$. [Koch et al. *J Am Chem Soc* **75** 953 1953, DOI: 10.1021/ja01100a054; *Beilstein* **1** IV 2839.]

Starch [9005-84-9] $(C_6H_{10}O_5)_n$, **M (162.1)_n**. Starch is de-fatted by Soxhlet extraction with Et_2O or 95% EtOH. For fractionation of starch into ‘amylose’ and ‘amylopectin’ fractions, see Lansky et al. [*J Am Chem Soc* **71** 4066 1949, DOI: 10.1021/ja01180a056].

Streptozotocin (N-[methylnitrosocarbamoyl]- α -D-glucosamine, streptozocin) [18883-66-4] $C_8H_{15}N_3O_7$, **M 265.2**, **m 111-114°(dec), 114-115°(dec), 115°(dec with evolution of gas), 121°(dec)**, $[\alpha]_D^{20}$ **~+39** (H_2O , may vary due to mutarotation). Recrystallise streptozotocin from 95% EtOH. It is soluble in H_2O , MeOH and Me_2CO . It has UV with λ_{max} at 228nm (ϵ 6360) in EtOH. The *tetraacetate* has **m 111-114°(dec)**, and $[\alpha]_D^{25}$ **+41** (c 0.78, 95% EtOH) after recrystallisation from EtOAc. [Herr et al. *J Am Chem Soc* **89** 4808 1967, DOI: 10.1021/ja00994a053; NMR: Wiley et al. *J Org Chem* **44** 9 1979, DOI: 10.1021/jo01315a003.] It is a potent methylating agent for DNA [Bennett & Pegg *Cancer Res* **41** 2786 1981, PMID: 6454479].

D(+)-Sucrose (β -D-fructofuranosyl- α -D-glucopyranoside) [57-50-1] $C_{12}H_{22}O_{11}$, **M 342.3**, **m 160-186°, 185-187°, 186-188°**, $[\alpha]_{546}^{20}$ **+78** (c 10, H_2O), $[\alpha]_D^{20}$ **+66** (c 26, H_2O), pK^{25} **12.62**. Crystallise D(+)-sucrose from water (solubility: 1g in 0.5ml H_2O at 20°, 1g in 0.2ml in boiling H_2O). It is soluble in EtOH (0.6%) and MeOH (1%). *Sucrose diacetate hexaisobutyrate* [27216-37-1] $C_{40}H_{62}O_{19}$, **M 846.9**, is purified by melting and, while molten, treated with $NaHCO_3$ and charcoal, then filtered, and has **m -7°, b 288°/atm**, d^{25} **1.146g/ml**, $[\alpha]_D^{20}$ **+49** (c 1, $CHCl_3$). [Hynes & Le Page ‘Sucrose, a convenient test crystal for absolute structures’ *J Appl Cryst* **24** (4) 352 1991, DOI: 10.1107/S0021889891002492; *Beilstein* **17/8** V 399.]

D(+)-Sucrose octaacetate [126-14-7] $C_{28}H_{38}O_{19}$, **M 678.6**, **m 82-85°, 83-85°, b 260°/1mm**, $[\alpha]_{546}^{20}$ **+71 (c 2.5, EtOH)**. After three recrystallisations from EtOH or 95% EtOH (charcoal), the **m** of the *octaacetate* rises to **88-90°**, or Et₂O with **m 89°** and $[\alpha]_D^{25}$ **+58.5° (c 2.6, EtOH)**. It has a bitter taste. [Linstead et al. *J Am Chem Soc* **62** 3260 1940, DOI: 10.1021/ja01868a509; Lemieux & Huber *J Am Chem Soc* **78** 4117 1956, DOI: 10.1021/ja01597a070; *Beilstein* **17/8** V 410.]

D(-)-Tagatose [87-81-0] $C_6H_{12}O_6$, **M 180.2**, **m 131-132°, 134-135°**, $[\alpha]_{546}$ **-6.5 (c 1, H₂O)**. Crystallise D(-)-tagatose from EtOH/H₂O (6:1). It mutarotates from $[\alpha]_D^{22}$ **+2 (2 minutes)** to **-5.0 (30 minutes)** (c 4, H₂O). The *phenylosazone* crystallises from aqueous EtOH with **m 185-187°(dec)**, and $[\alpha]_D^{23}$ **+47 (c 0.82, 2-methoxyethanol)**. [Totton & Lardy *J Am Chem Soc* **71** 3076 1949, DOI: 10.1021/ja01177a037; Gorin et al. *Canad J Chem* **33** 1116 1955, DOI: 10.1139/v55-130; Reichstein & Bosshard *Helv Chem Acta* **17** 753 1934, DOI: 10.1002/hlca.19340170194; Wolfrom & Bennett *J Org Chem* **30** 1284 1965, DOI: 10.1021/jo01015a532; *Beilstein* **1** IV 4414.] In D₂O at 27° the ¹H NMR showed the following ratios: α-pyranose (79), β-pyranose (16), α-furanose (1) and β-furanose (4) [Angyal *Adv Carbohydr Chem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5; Angyal & Pickles *Aust J Chem* **25** 1711 1972, DOI: 10.1071/CH9721711]. *Sugar substitute*, not commonly found in nature, but is produced using L-arabinose isomerase as biocatalyst and D-galactose as substrate [Review: Deok-Kun Oh *Appl Microbiol Biotechnol* **76** 1 2007, DOI: 10.1007/s00253-007-0981-1].

Thevetin A (cardenolide glycoside) [37933-66-7] $C_{42}H_{64}O_{19}$, **M 872.9**, **m softens at 194°, m 208-210°**, $[\alpha]_D^{20}$ **-72 (c 1.48, MeOH)**. This cardiac glycoside of *Thevetia nerifolia* Juss seeds crystallises from H₂O. The *acetyl derivative* crystallises from MeOH/Et₂O at -15° with **m 145-149°**, and $[\alpha]_D^{26}$ **-54.2 (c 1.86, CHCl₃)**. Store at -20°. [Block et al. *Helv Chim Acta* **43** 652 1960, DOI: 10.1002/hlca.19600430306; for ¹³C NMR see Tori et al. *Tetrahedron Lett* **18** 717 1977, DOI: 10.1016/S0040-4039(01)92735-2; *Beilstein* **18** III/IV 2552, **18/4** V 439.] Unlike Thevetin B (below, which occurs with it), it has an aldehyde group at C10 in the steroid moiety.

Thevetin B (cardenolide glycoside) [11018-93-2, 27127-79-3] $C_{42}H_{66}O_{18}$, **M 858.9**, **m 197-201°**, $[\alpha]_D^{24}$ **-61.4 (c 1.5, MeOH)**. This cardiac glycoside of *Thevetia nerifolia* Juss seeds crystallises (as *trihydrate*) from isopropanol. Dry it at 100°/0.01mm to give the *hemihydrate (very hygroscopic)*. [Block et al. *Helv Chim Acta* **43** 652 1960, DOI: 10.1002/hlca.19600430306; ¹³C NMR: Tori et al. *Tetrahedron Lett* **18** 717 1977, DOI: 10.1016/S0040-4039(01)92735-2; for thevetin B evaluation in serum by fluorescence polarisation immunoassay using anti-digitoxin antibodies see Uber-Bucek et al. *J Pharm Biomed Anal* **10**(6) 413 1992, PMID: 1420463; *Beilstein* **18** III/IV 1493.]

α,α'-D(+)-Trehalose dihydrate (α-D-glucopyranosyl-α-D-glucopyranoside) [6138-23-4] $C_{12}H_{22}O_{11} \cdot 2H_2O$, **M 378.3**, **m 96.5-97.5°, 94-100° (dihydrate), 214-216° (anhydrous)**, $[\alpha]_D^{20}$ **+180 (dihydrate, c 4, H₂O)**, $[\alpha]_D^{20}$ **+199 (anhydrous, c 4, H₂O)**. α,α'-D(+)-Trehalose crystallises (as the *dihydrate*) from aqueous EtOH. Dry it at 13°. For the *anhydrous* compound dissolve 10g in pyridine (200ml) and distil off this solvent at atmospheric pressure, and when the temperature rises to 115.3° all the H₂O is removed and 73ml of distillate is collected. Most of the anhydrous material crystallises out at this stage. The crystals are collected (6.8g), washed with Et₂O to give 6.1g of *anhydrous* product. Higher yields are obtained by slightly more prolonged distillation. [Birch *J Chem Soc* 3489 1965, DOI: 10.1039/JR9650003473; X-ray cryst: Brown et al. *Acta Cryst* **28B** 3145 1972, DOI: 10.1107/S0567740872007654; *Beilstein* **17/8** V 3.]

D(+)-Turanose [3-O-α-D-glucopyranosyl-D-fructose] [547-25-1] $C_{12}H_{22}O_{11}$, **M 342.3**, **m 168-170°**, $[\alpha]_D^{20}$ **+88 (c 4, H₂O)**. Crystallise D(+)-turanose from H₂O by addition of EtOH (its solubility is 5.3% in 95% EtOH). Form **m 157°** is obtained by crystallisation from hot MeOH, and mutarotates from $[\alpha]_D^{20}$ **+27.3 to +88 (c 4, H₂O)**. The *phenylosazone* crystallises from 15 parts of 95% EtOH with **m 200-205°**, $[\alpha]_D^{20}$ **24.5** mutarotating to **+33 [in 24 hours, c 0.82, pyridine/EtOH (4:6)]**. [Pascu *Methods in Carbohydrate Chemistry* **I** 353 1962, Academic Press, *Beilstein* **17/7** V 213.] In D₂O at 36° the ¹H NMR showed the following ratios: α-pyranose (<4), β-pyranose (39), α-furanose (20) and β-furanose (41) [Angyal *Adv Carbohydr Chem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5]. Although it is a carbon source for micro-organisms and many bacteria and fungi. It is not metabolised in higher plants but can be taken into plant cells by sucrose transporters where it is involved in sugar signalling.

Ustilagic acid (Ustizeain B, di-D-glucosyldihydroxyhexadecanoic acid) [8002-36-6] $C_{37}H_{62}O_{17}$, M ~780, m 146-147°, $[\alpha]_D^{23}$ +7 (c 1, pyridine), $pK^{25} \sim 4.9$. Ustilagic acid is a mixture of partly acetylated di-D-glucosyldihydroxyhexadecanoic acid which crystallises from Et_2O . It has also been purified from corn smut fungal culture by dissolving it in hot MeOH, filtering and concentrating by blowing a current of air until the solution becomes turbid, then heating to 50° and adding 4 volumes of H_2O (also at 50°), and cooling very slowly. Filter off the white solid and dry it in air. It crystallises from Et_2O , and is soluble in MeOH, butan-1,2-diol, poorly soluble in EtOH, *n*-BuOH, Me_2CO and insoluble in H_2O , EtOAc and $*C_6H_6$. [Lemieux et al. *Can J Chem* **29** 409 1951, DOI: 10.1139/v51-049; for structure and hydrolysis to **ustilic acid A**, m 114-115°, $[\alpha]_D^{20}$ -6.3 (MeOH) which gave a **methyl ester** m 80-81.5°, $[\alpha]_D^{20}$ -0.2 ($CHCl_3$) see Lemieux *Can J Chem* **29** 415 1951, DOI: 10.1139/v51-050; for structure and configurations of **ustilic acids** see Lemieux *Can J Chem* **31** 396 1953, DOI: 10.1139/v53-056; for biosynthesis see Boothroyd et al. *Can J Biochem Physiol* **33** 289 1955; DOI: 10.1139/o55-039.]

Vicine (2,4-diamino-5-β-D-glucopyranosidoxy-6-hydroxypyrimidine) [152-93-2] $C_{10}H_{16}N_4O_7$, M 304.3, m 243-244°(dec), $[\alpha]_D^{20}$ -12.1 (c 4, 0.2N NaOH). Crystallise Vicine from water (1%) or aqueous 85% EtOH, and dry it at 135°, then or over P_2O_5 at ~25°. [For isolation from mixed vetch seeds (*Vicia sativa*) and complete structure determination including UV spectra see Bendich & Clements *Biochim Biophys Acta* **12** 462 1953, DOI: 10.1016/0006-3002(53)90166-8; *Beilstein* **31** H 163, **25** III/IV 4285.]

Xanthorhamnin {xanthene rhamnoside, 3-[(2*S*,3*R*,4*S*,5*R*,6*R*)-6-[(2*R*,3*R*,4*R*,5*S*,6*S*)-3,5-dihydroxy-6-methyl-4-[(2*S*,3*R*,4*R*,5*R*,6*S*)-3,4,5-trihydroxy-6-methyloxan-2-yl]oxyoxan-2-yl]oxymethyl]-3,4,5-trihydroxy-oxan-2-yl]oxy-2-(3,4-dihydroxyphenyl)-5-hydroxy-7-methoxychromen-4-one} [1324-63-6] $C_{34}H_{42}O_{20}$, M 770.7, m 195°, $[\alpha]_D^{20}$ +3.75 (EtOH), pK_1^{24} 8.69, pK_2^{24} 11.28, pK_3^{24} 12.22. Crystallise xanthorhamnin from a mixture of ethyl and isopropyl alcohols, dry it in air, then dry it further for several hours at 110°. The UV (EtOH) has λ_{max} at 258 and 362nm. [Nystrom et al. *J Org Chem* **22** 1272 1957, DOI: 10.1021/jo01361a619; *Beilstein* **18** III/IV 3498.] It has been isolated from buckthorn berries (*Rhamnus catharticus*), and forms a **pentahydrate**, M 860.8, which holds its H_2O tenaciously.

α-D(+)-Xylose (wood sugar) [58-86-6] $C_5H_{10}O_5$, M 150.1, m 146-147°, 153-154°, 154-158°, $[\alpha]_D^{20}$ +92 mutarotating to +18.8 (16 hours, c 10, H_2O), pK^{18} 12.14. α-D(+)-Xylose forms needles or prisms (which have a very sweet taste) by slow crystallisation from aqueous 80% EtOH or absolute EtOH, which are then dried at 60° *in vacuo* over P_2O_5 . Store it in a vacuum desiccator over $CaSO_4$. One gram dissolves in 0.8ml H_2O . [Bragg & Hough *J Chem Soc* 4347 1957, Hudson & Yanovsky *J Am Chem Soc* **39** 1013 1917, DOI: 10.1021/ja02250a019; Monroe *J Am Chem Soc* **41** 1002 1919, DOI: 10.1021/ja02227a010; *Beilstein* **1** IV 4223.] In D_2O at 31°, the 1H NMR showed the following ratios: α-pyranose (36.5), β-pyranose (63), α-furanose + β-furanose (~1) [Angyal *Adv Carbohydr Chem* **42** 15 1984, DOI: 10.1016/S0065-2318(08)60122-5; Angyal & Pickles *Aust J Chem* **25** 1711 1972, DOI: 10.1071/CH9721711; *Beilstein* **1** IV 4223]. The enantiomeric **α-L(-)-xylose** [609-06-3] has identical properties except for the optical rotation which is negative.

CAROTENOIDS

Introduction

Carotenoids are polyene pigments that are mostly naturally occurring in bacteria, plants and animals. They have been isolated from the natural sources and obtained first by extraction with solvents and then purified by column chromatography through Al_2O_3 of various grades, $\text{Ca}(\text{OH})_2$ alone or with CaCO_3 , MgO or Silica Gel and eluted with solvents of various polarities. The progress of separation can be followed visually because the bands of most carotenoids are of various colours. The bands can be collected by elution, or the column can be extruded and the bands cut out and extracted with a polar solvent, e.g. MeOH. This chromatography can be repeated with the separate bands, and finally the carotenoids are recrystallised to analytical purity. The purity can be checked by TLC on Silica Gel or Al_2O_3 plates or paper chromatography and eluted in two dimensions. Gas-liquid or HPLC has been used for preparative work as well as for checking the purity and identifying them using internal standards such as tocopherol acetate (vitamin E acetate) and retinyl acetate.

Carotenoids are generally light sensitive, easily oxidised by air and are affected by traces of acid, e.g. in solvents. These cause the polyenes to bleach or polymerise. The necessary precautions are therefore required to minimize these effects during isolation, purification and storage. They are identified by their UV-VIS spectra, and their molar extinction coefficients at specific wavelengths (λ_{max}) have been used for characterisation and for quantitation. More recently ORD, CD, NMR, IR and mass spectroscopy have been used extensively.

Antheraxanthin [5,6-epoxy-zeaxanthin, (3R,3'S,5'R,6'S)-5',6'-dihydro-5',6'-epoxy- β,β -carotene-3,3'-diol] [9Z- 68831-78-7, all-trans- 640-03-9] $\text{C}_{40}\text{H}_{56}\text{O}_3$, M 584.8, m 205°, λ_{max} 460.5, 490.5nm, in CHCl_3 . This xanthophyll cycle pigment is a bright yellow *accessory* pigment found in many organisms that perform photosynthesis (e.g. algae, euglenoids) and plants. Likely impurities are violaxanthin (see later) and mutatoxanthin (which contains a dihydrofuran ring derived from the epoxy group of antheraxanthin). Purify it by chromatography on columns of $\text{Ca}(\text{OH})_2$ and of ZnCO_3 . It crystallises from $^*\text{C}_6\text{H}_6/\text{MeOH}$ as needles or thin plates. Store it in the dark, under N_2 or argon at -20° . In $\text{CHCl}_3/\text{SbCl}_2$ it gives a blue colour with λ_{max} at 587nm and gives a stable blue colour with conc HCl. [Karrer & Oswald *Helv Chim Acta* **18** 1303 1935, DOI: 10.1002/hlca.193501801182; Karrer & Jucker *Helv Chim Acta* **28** 300 1945, DOI: 10.1002/hlca.660280137.]

β -Apo-4'-carotenal [12676-20-9] $\text{C}_{30}\text{H}_{40}\text{O}$, M 416.7, m 139°, $A_{1\text{cm}}^{1\%}$ 2640 at 461nm. Recrystallise the carotenal from $\text{CHCl}_3/\text{EtOH}$ mixture or *n*-hexane. The λ_{max} of the absorption spectra of the radical anions and cations of such polyenals formed by pulse radiolysis are linearly dependent on the number of conjugated double bonds [Land et al. *JCS Faraday Trans 1* **74** 538 1978, DOI: 10.1039/F19787400538; Bobrowski & Das *J Phys Chem* **91** 1210 1987, DOI: 10.1021/j100289a035; Beilstein **7** IV 1782.]

trans- β -Apo-8'-carotenal [1107-26-2] $\text{C}_{30}\text{H}_{40}\text{O}$, M 416.7, m 136-139°. Recrystallise β -apo-8'-carotenal from $\text{CHCl}_3/\text{EtOH}$ mixture or *n*-hexane. The absorption spectra of the radical anion of this C_{30} aldehyde in CTAB and Triton X100 micelles suggested an alcohol-like nature of the environment probed by the long-chain polyenal. For effect of conjugation on spectra see preceding entry [Bobrowski & Das *J Phys Chem* **91** 1210 1987, DOI: 10.1021/j100289a035; Land et al. *JCS Faraday Trans 1* **74** 538 1978, DOI: 10.1039/F19787400538; Beilstein **7** III 2622, **7** IV 1782.] It has been stated to form red to deep violet crystals, or a crystalline powder with a metallic lustre, and is sensitive to oxygen and light and should be stored in the dark under inert gas. Also described as *Apocarotenal*, which has an orange to orange-red (C.I. Food Orange 6) colour used in foods, pharmaceutical and cosmetic products.

β -Apo-8'-carotenoic acid ethyl ester [1109-11-1] $\text{C}_{32}\text{H}_{44}\text{O}_2$, M 460.7, m 134-138°, $A_{1\text{cm}}^{1\%}$ 2550 at 449nm, CI Food Orange 7; CI (1075) No 40825, INS No 160f. Recrystallise the ester from petroleum ether or petroleum ether/EtOAc. The A_{475}/A_{449} is between 0.82 to 0.86. The colour of a sample in Me_2CO disappears after successive additions of a 5% solution of NaNO_2 and 1N H_2SO_4 . Carr-Price reaction: a solution of a sample in CHCl_3 turns blue on addition of excess of Carr-Price reagent TS. See also colour reaction with $\text{CF}_3\text{CO}_2\text{H}$ [Dugan et al. *Anal Chem* **36** 114 1964, DOI: 10.1021/ac60207a035]. Store it in the dark under N_2 or argon at

-20°. [Beilstein 9 IV 2701 for Me ester] **β -Apo-8'-carotenoic acid** and its esters (ethyl and glyceryl) were obtained from sunflower oil oxidation [Yanishlieva et al. *J Am Oil Chem Soc* **78** 641 2001, DOI: 10.1007/s11746-001-0319-7].

β -Apo-8'-carotenoic acid methyl ester [16266-99-2] $C_{31}H_{42}O_2$, M 512.7, m 136-137°, $A_{1\text{cm}}^{1\%}$ 2575 at 446nm and 2160 at 471nm, in petroleum ether. Recrystallise the methyl ester from petroleum ether or petroleum ether/EtOAc. Store it in the dark in an inert atmosphere at -20°. [See ethyl ester above, Beilstein 9 IV 2701.]

Astacene (β , β -carotene-3,3',4,4'-tetraone, Astacin not to be confused with the metalloproteinases) [514-76-1] $C_{40}H_{48}O_4$, M 592.8, m 228°, 232-233°, 240-243°(evacuated tube), $\epsilon_{1\text{cm}}^{1\%}$ 550,000 at 498nm (pyridine). Astacene is a red pigment found in marine organisms (protozoa, algae, crustaceans, sponges, fish, reptiles) together with astaxanthin (see below) from which it is possibly formed by autoxidation [Kuhn & Lederer *Chem Ber* **66** 488 1933, DOI: 10.1002/cber.19330660410]. A probable impurity is astaxanthin (3,3'-dihydroxy- β , β -carotene-4,4'-dione) which may be its precursor. It has been prepared by adding a solution of **canthaxanthene** (1.0g, see [514-78-3] below) in C_6H_6 (15ml) to *t*-BuOK in *t*-BuOH (1.40N, 130ml) and shaken with O_2 at 20° for 30 hours. Water (200ml) is added, then 0.5N HCl (800ml) and extracted with $CHCl_3$ (300ml). The extract is washed with aqueous $NaHCO_3$ (3 x 300ml), H_2O (2 x 200ml), diluted with C_6H_6 (50ml) and the organic layer is evaporated *in vacuo*. The residue is recrystallised from $CHCl_3$ -EtOH (1:5) to give **astacene** (450mg, ~90%) as deep purple leaflets m 232-233° with λ_{max} (pyridine) 498nm (ϵ 100 x 10³), (CS_2) 513nm, (hexane) 477nm, (EtOH) 483nm, and (KOH-EtOH) 478nm; the IR ($CHCl_3$) has ν_{max} at 3410br (ϵ ~90), 1610 (ϵ 1200), 1550, 1327, 969 (ϵ 895) cm^{-1} ; the 1H NMR (60MHz, $CDCl_3$, TMS) has δ at 1.30 (12H, C-1 Me_2 and C-1' Me_2), 2.10 (6H, C-5 Me and C-5' Me), 2.02 (6H, C-9 Me and C-9' Me), 2.02 (6H, C-13 Me and C-13' Me), and 6.06 (C-2 H and C-2' H) (14 olefinic H-7,7', 8,8', 10,10', 11,11', 12,12', 14,14', and 15,15' are not included); and m/z 592 M^{+} . [Cooper et al. *JCS Perkin Trans I* 2195 1975, DOI: 10.1039/P19750002195.] Alternatively, purify **astacin** by chromatography on alumina/fibrous clay (1:4) or mixed chromatograms on sucrose using C_6H_6 /light petroleum 1:4 as eluant; or by partition between petroleum ether and MeOH (alkaline). It crystallises from pyridine/water in purple leaflets or needles with a metallic lustre. Store it in the dark under N_2 at -20°. It is very soluble in pyridine, dioxane, CS_2 , $CHCl_3$ and dilute aqueous NaOH, slightly soluble in AcOH, EtOAc and C_6H_6 , but almost insoluble in H_2O , Et_2O , petroleum ether and MeOH. Its **diacetate (enol-acetate)**, obtained by Ac_2O /pyridine treatment at 25° for 34 hours has m 232-233° (dec) (from $CHCl_3$ /EtOH, 1:5) or 235° (dec) (from pyridine + H_2O), and forms violet/black needles with λ_{max} (pyridine) at 497nm (ϵ 109 x 10³) and (EtOH) 474nm; the IR ($CHCl_3$) has ν_{max} at 1756 (ϵ 530), 1639 (ϵ 1150), 1555 and 972 cm^{-1} ; the 1H NMR (60MHz, $CDCl_3$, TMS) has δ at 1.35 (12H, C-1 Me_2 and C-1' Me_2), 2.04 (6H, C-5 Me and C-5' Me), 2.04 (6H, C-9 Me and C-9' Me), 2.02 (6H, C-13 Me and C-13' Me), and 2.29 (2 OAc) (14 olefinic H-7,7', 8,8', 10,10', 11,11', 12,12', 14,14', and 15,15' are not included); and m/z 576 M^{+} . The **dipalmitate (enol-ester, Astacein)** has m 121°, and forms red square leaflets from petroleum ether. [Davis & Weedon *Proc Chem Soc* 182 1960, DOI: 10.1039/PS9600000161 for first page of Notes is 161; Widmer et al. *Helv Chim Acta* **65** 671 1982, DOI: 10.1002/hlca.19820650306; Karrer et al. *Helv Chim Acta* **17** 412 1934, DOI: 10.1002/hlca.19340170147; 745 1934, DOI: 10.1002/hlca.19340170192; **18** 96 1935, DOI: 10.1002/hlca.19350180114; **19** 479 1936, DOI: 10.1002/hlca.19360190174; for synthesis see Choi & Koo *J Org Chem* **70** 3328 2005, DOI: 10.1021/jo0501011; for isolation from brown trout see Steven *Nature* **160** 507 1947, PubMed: 20267555; Beilstein H **30** 102, **7** III 4797.]

Astaxanthin (3,3'-dihydroxy- β , β -carotene-4,4'-dione) [472-61-7] $C_{40}H_{52}O_4$, M 596.8, m 182-183° (for \pm , see below). It is a potent antioxidant from marine algae, red yeast and other plant and marine animal sources. It was isolated from lobster eggs, is present in salmon, the red feathers of some birds (*Laniarius* spp, and flamingo), and occurs in flower petals of some **Ranunculaceae**. Racemic astaxanthin has been synthesised from astacene (see above) by reduction with KBH_4 to β , β -carotene-3,4,3',4'-tetra-ol followed by oxidation with 2,3-dichloro-5,6-dicyano-*p*-benzoquinone and purification by preparative TLC on Kieselgel (20% Me_2CO /light petroleum) to give (\pm)-**astaxanthin** m 182-183° as red needles, after crystallisation from Me_2CO /light petroleum. It has λ_{max} (MeOH) 472nm, (hexane) 466-467nm, ($CHCl_3$) 485 and (CS_2) 503nm; the IR ($CHCl_3$) has ν_{max} at 3620, 3510, 1660, 1610 and 975 cm^{-1} ; the 1H NMR (60MHz, $CDCl_3$, TMS) has δ at 1.31 (6H, C-1 Me and C-1' Me), 1.20 (6H, C-1 Me and C-1' Me), 1.94 (6H, C-5 Me and C-5' Me), 1.98 (12H, C-9 Me, C-9' Me, C-13 Me and C-13' Me), (14 olefinic H-7,7', 8,8', 10,10', 11,11', 12,12', 14,14', and 15,15' are not included); and m/z 592.386 M^{+} .

TLC in the above system or Micro-Cel did not separate it from natural (3*S*,3'*S*)-astaxanthin from the common lobster or sea water crayfish. Synthetic (\pm)-**astaxanthin** combined with the appropriate apoprotein to form **α -crustacyanin** with λ_{\max} (MeOH) at 630nm. Its **diacetate** has **m 203-205°** (blue-black needles from pyridine/H₂O, **m 189-191°** also reported), the UV has λ_{\max} (CHCl₃) with 482nm, and is also indistinguishable by mixed TLC on Kieselgel, Micro-Cel or alumina from natural (3*S*,3'*S*)-**astaxanthindiacetate**, **m 187-189°**. All manipulations have to be carried out under argon to avoid their ready oxidation to **astacin** (see above). [Cooper et al *JCS Perkin Trans I* 2195 1975, DOI: 10.1039/P19750002195; Kienzel & Mayer *Helv Chim Acta* **61** 2609 1978, DOI: 10.1002/hlca.19780610732.]

An efficient synthesis of natural (3*S*,3'*S*)-**astaxanthin** from (4*R*,4'*R*)-4-hydroxy-2,2,6-trimethylcyclohexanone in 67% yield and high optical purity has been described and was purified by recrystallisation from CH₂Cl₂/MeOH to give dark violet crystals, **m 216-218°**, with R_F 0.30 on TLC with kieselgel (CH₂Cl₂/Me₂CO/HCOOH, 95:5:3); its **UV** has λ_{\max} nm (ϵ), (CHCl₃) at 492 (1920), 488 (125,000), and (CH₂Cl₂) 324(14,500), 298 (15,500), ~285sh (~14,000), 251 (14,500), ~225 (~15,000); and **IR** (KBr) has ν_{\max} at 3486m (OH), 1664s (C=O, conjugated), 1607m, 1557s (C=O, conjugated), 1399w, 1390m, 1366w (gem Me₂), 1076s, 1039m (OH) and 969s (CH=CH, *trans*) cm⁻¹. [Widmer et al. *Helv Chim Acta* **64** 2405 1981, DOI: 10.1002/hlca.19810640749]. Other high yielding syntheses have been achieved [Becher et al. *Helv Chim Acta* **64** 2419 1981, DOI: 10.1002/hlca.19810640750; Kienzle & Mayer *Helv Chim Acta* **61** 2609 1978, DOI: 10.1002/hlca.19780610732], and the absolute configuration of natural (3*S*,3'*S*)-**astaxanthin**, **m 219-220°** has been determined by X-ray analysis of a crystalline synthetic intermediate [Widmer et al. *Helv Chim Acta* **64** 2405 1981, DOI: 10.1002/hlca.19810640749] and comparison of CD spectra [Englert et al. *Helv Chim Acta* **60** 1209 1977, DOI: 10.1002/hlca.19770600409]. Its **¹H NMR** (270MHz, CDCl₃, TMS) has δ for olefinic protons at 6.21 (H-7 and H-7'), 6.43 (H-8 and H-8'), 6.30 (H-10 and H-10'), 6.66 (H-11 and H-11'), 6.45 (H-12 and H-12'), ~6.30 (H-14 and H-14') and ~6.68 (H-15 and H-15') with $J_{7,8}$ or $J_{7,8'}$ = 16.3Hz, $J_{10,11}$ or $J_{10',11'}$ = 11.5Hz and $J_{11,12}$ or $J_{11',12'}$ = 14.7Hz; δ for Me groups at 1.21 and 1.32 (6H, C-1 Me and C-1' Me), 1.95 (6H, C-5 Me and C-5' Me), 2.00 (6H, C-9 Me and C-9' Me), 1.99 (6H, C-13 Me and C-13' Me); and δ for cyclohexenone protons at 1.82 (2H, axial H-2 and H-2'), 2.16 (2H, equatorial H-2 and H-2'), 4.32 (2H, H-3 and H-3'), and 3.70 (2H, 3-OH and 3'-OH, J = 1.8Hz,) with $J_{2,2'}$ (gem) or $J_{2',2'}$ (gem) = 12.7Hz, $J_{2e,3a}$ (vic) = 5.7Hz and $J_{2a',3a'}$ (vic) = 14.0Hz, $J_{2e',3a'}$ (vic) = 5.7Hz and $J_{2a',3a'}$ (vic) = 14.0Hz; the **¹³C NMR** (23MHz, CDCl₃, TMS) has δ for methyl group carbons at 12.8 (13,13'), 12.6 (9,9'), 13.9 (5,5'), 26.2 (1 or 1'), 30.8 (1' or 1); for end group carbons 36.8 (1,1'), 45.6 (2,2'), 69.2 (3,3'), 200.3 (4,4'), 126.9 (5,5'), 162.4 (6,6'); and for olefinic carbons at 123.2 (7,7'), 142.2 (8,8'), 134.7 (9,9'), 135.1 (10,10'), 124.6 (11,11'), 139.7 (12,12'), 136.7 (13,13'), 133.8 (14,14') and 130.7 (15,15'); and the **CD** spectrum in CH₂Cl₂ has λ_{\max} ($\Delta\epsilon$) at 224 (+12.8), 249 (-14.4), 280 (+ 12.5), 323 (-23.1), 384 (+6.7) and 521 (-3.2, very broad) [Englert et al. *Helv Chim Acta* **60** 1209 1977, DOI: 10.1002/hlca.19770600409].

It is the prosthetic group in many carotenoproteins, mainly in invertebrates, which vary from colourless to bright blue. It reduces blood glucose and improves some parameters in diabetic metabolism. It ameliorates blood flow and vascular tone in hypertension models and regulates **connexin 43** in *in vitro* studies, and may be a cancer chemopreventive drug. It is obtained on a large scale from the heterobasidiomycetous yeast *Phaffia rhodozyma* and plays an important role in carotene biosynthesis [Schroeder & Johnson *J Biol Chem* **270** 18374 1995, DOI: 10.1074/jbc.270.31.18374]. For an efficient synthesis see Choi & Koo *J Org Chem* **70** 3328 2005, DOI: 10.1021/jo0501011.

Bixin (6,6'-diapo- ψ,ψ -carotenedioic acid monomethyl ester) [6983-79-5 for *cis*- or **α -bixin**] C₂₅H₃₀O₄, **M 394.5**, **m 198°**, 217°(dec), λ_{\max} (CHCl₃) **209, 475 and 443nm**, **pK_{Est} ~4.3**. The natural half ester, **α -bixin Annatto** (a pigment from *Bixa orellana* shrub native of S. America, India, E. Africa, Caribbean and Phillippines) has a *cis*-double bond conjugated with the ester group and is unstable when isolated. It readily isomerises to the stable *all-trans* halfester **α -bixin** (*trans*-bixin, also present in nature) [39937-23-0] in CHCl₃ solution containing I₂. Crystallise bixin from Me₂CO (violet prisms). Synthetic ***cis*-methylbixin** crystallised from EtOAc with **m 163°**, whereas the synthetic ***all-trans*-methylbixin**, which also crystallises from EtOAc has **m 204-205°**, distinguished by ¹H NMR spectroscopy. [For synthesis of methylbixins see Pattenden et al. *J Chem Soc (C)* 235 1970, DOI: 10.1039/J39700000235]. [For the synthesis, UV and IR of Me and Et diesters see Isler et al. *Helv Chim Acta* **40** 1242 1957, DOI: 10.1002/hlca.19570400515; Buchta & Andree *Chem Ber* **92** 3111 1959, DOI: 10.1002/cber.19590921214]. [Beilstein **2** III 2020, **2** IV 2455, H **30** 110.] The **free acid**, **Norbixin**, a dicarboxylic acid, and the esters have been used to colour margarine and impart a red colour to cheddar cheese.

Canthaxanthin (*trans*) (β , β -carotene-4,4'-dione) [514-78-3] $C_{40}H_{52}O_2$, M 564.9, m 211-212°, 213°, ϵ_{mol} 124,100 (at 466nm, petroleum ether), 118,000 (at 484nm * C_6H_6). Purify canthaxanthin by chromatography on a column of deactivated alumina or magnesium oxide, or on a thin layer plate of silica gel G (Merck), using dichloromethane/diethyl ether (9:1) to develop the chromatogram. Crystallise it from CH_2Cl_2 or $CH_2Cl_2/MeOH$ (violet crystals). Store it in the dark and in an inert atmosphere at -20°. [For an efficient synthesis see Choi & Koo *J Org Chem* **70** 3328 2005, DOI: 10.1021/jo0501011; for 'Isolation and light-stimulated expression of canthaxanthin and spirilloxanthin biosynthesis genes from the photosynthetic bacterium *Bradyrhizobium* sp. strain ORS278', see Giraud & Verméglio *Methods in molecular biology* (Clifton NJ) **892** 173 2012, PMID: 22623302.] [Beilstein **7** III 4364, **7** IV 2680.] It has been found in birds and fish.

Capsanthin [(3*R*,3'*S*,5'*R*)-3,3'-dihydroxy- β , κ -carotene-6-one] [465-42-9] $C_{40}H_{56}O_3$, M 584.9, m 170°, 177-178°, λ_{max} 484 (ϵ_{mol} 121,000), inflexion at 515 nm, in hexane. It has been isolated from paprika extract (main carotenoid), and possible impurities are zeaxanthin and capsorubin (below). Purify capsanthin by chromatography on a column of alumina (grade IV) and develop with 0.2% EtOH in * C_6H_6 . It crystallises from petroleum ether in red needles, IR has ν_{max} at 3600, 1664 cm^{-1} , and $\epsilon_{1033}/\epsilon_{1664} \sim 4.1$. [Barber et al. *J Chem Soc* 4019 1961, DOI: 10.1039/JR9610004019; for ORD & absolute configuration see Bartlett et al. *J Chem Soc (C)* 2527 1969, DOI: 10.1039/J39690002527.] The *di-O-acetate* is purified on an alumina (grade II) column in * C_6H_6 and it is recrystallised from MeOH with m 160-162°. [Faigle & Karrer *Helv Chim Acta* **44** 1257 1961, DOI: 10.1002/hlca.19610440510; Beilstein **8** II 415, **8** III 3047, **8** IV 2657.] It has anti-neoplastic properties.

Capsorubin (3,3'-dihydroxy- κ , κ -carotene-6,6'dione) [470-38-2] $C_{40}H_{56}O_4$, M 600.9, m 218°, λ_{max} 443, 468, 503 nm, in hexane. Possible impurities are zeaxanthin and capsanthin. Purify capsorubin by chromatography on a column of $CaCO_3$ or MgO. Crystallise it from *benzene/petroleum ether or CS_2 . [Beilstein **1** III 3327, **8** III 3873, **8** IV 3304.] It is a red pigment in paprika which contains capsanthin-capsorubin synthase whose gene has been cloned [Bouvier et al. *Plant Mol. Biol.* **36**: 785 1998, DOI: 10.1023/A:1005966313415].

α -Carotene (6'*R*- α -carotene) [7488-99-5] $C_{40}H_{56}$, M 536.9, m 184-188°, 187.5° (evacuated tube), $[\alpha]_{D}^{20} +385^\circ$ (c 0.08, * C_6H_6), $[\alpha]_{D}^{25} +538^\circ$, λ_{max} 422, 446, 474 nm, in hexane, $A_{1cm}^{1\%}$ 2725 (at 446nm), 2490 (at 474nm), ϵ_{mol} 145,300 (at 455nm, hexane), and 456 (at 485nm, * C_6H_6). Purify α -carotene by chromatography on columns of calcium hydroxide, alumina or magnesia. Crystallise it from $CS_2/MeOH$, toluene/MeOH, diethyl ether/petroleum ether, or acetone/petroleum ether. Store it in the dark, under N_2 or Ar at -20°. It gives a blue colour with λ_{max} at 542nm when mixed with $SbCl_3$ in $CHCl_3$. [Karrer & Walker *Helv Chim Acta* **16** 641 1933, DOI: 10.1002/hlca.19330160193; Eugster et al. *Helv Chim Acta* **52** 1729 1969, DOI: 10.1002/hlca.19690520627; Eugster & Karrer *Helv Chim Acta* **38** 610 1955, DOI: 10.1002/hlca.19550380307; Strain *J Biol Chem* **105** 523 1934, <http://www.jbc.org/content/105/3/523>; Beilstein **5** III 2457, **5** IV 2620.]

all-trans- β -Carotene [7235-40-7] $C_{40}H_{56}$, M 536.9, m 178-179°, 179-180°, 180°, 181°, 183° (evacuated capillary), ϵ_{mol} 138,900 (at 450 nm, petroleum ether), 124,300 (at 462nm, * C_6H_6), CI Food Orange **5**, INS No. 160a(i); CI (1975) No. 40800. It forms purple prisms when crystallised from * $C_6H_6/MeOH$ and red rhombs from petroleum ether. Its solubility in hexane is 0.1% at 0°. It is **oxygen sensitive** and should be stored under N_2 at -20° in the dark. It gives a deep blue colour with λ_{max} at 590nm when mixed with $SbCl_3$ in $CHCl_3$. Its UV (* C_6H_6) has λ_{max} at 429nm, 454 and 484nm. The principal peak at λ_{max} 454nm has $A_{1cm}^{1\%}$ 2000. [Synthesis: Surmatis & Ofner *J Org Chem* **26** 1171 1961, DOI: 10.1021/jo01063a048; Milas et al. *J Am Chem Soc* **72** 4844 1950, DOI: 10.1021/ja01166a541.] β -Carotene is also purified by column chromatography (Al_2O_3 activity I-II). It is dissolved in petroleum ether/* C_6H_6 (10:1), applied to the column and eluted with petroleum ether/EtOH; the desired fraction is evaporated and the residue is recrystallised from * $C_6H_6/MeOH$ (violet-red plates). [For UV see Inhoffen et al. *Justus Liebigs Ann Chem* **570** 54, 68 1950, DOI: 10.1002/jlac.19505700105; Review: Fleming *Selected Organic Synthesis* (J Wiley, Lond) pp. 70-74 1973.] Alternatively, it can be purified by chromatography on a magnesia column, thin layer of Kieselguhr or magnesia. Crystallise it from $CS_2/MeOH$, Et_2O /petroleum ether, acetone/petroleum ether or toluene/MeOH. Store it in the dark, under an inert atmosphere, at -20°. Recrystallise it also from 1:1 EtOH/ $CHCl_3$. [Bobrowski & Das *J Phys Chem* **89** 5079 1985, DOI: 10.1021/j100269a038; Johnston & Scaiano *J Am Chem Soc* **108** 2349 1986, DOI: 10.1021/ja00269a035; Strain *J Biol Chem* **105** 523 1934, <http://www.jbc.org/content/105/3/523>; **4** 1

1957, *Beilstein* **5** II 638, **5** III 2453, **5** IV 2617.]

δ-Carotene (6R-ε,ψ-carotene) [472-92-4] $C_{40}H_{56}$, **M 536.9**, **m 140.4°**, **142°** (evacuated capillary) $[\alpha]_D^{25} +352$ (c **16** hexane), $[\alpha]_{Cd} +317$ (CS₂), λ_{max} **430, 456, 486nm** (hexane). δ-Carotene crystallises from CS₂/hexane/EtOH as red needles. The *racemic* carotene is purified through an alumina (Grade II) column by elution with 15% *C₆H₆/petroleum ether (b 60-80°), and the main band eluent is evaporated and the residue is crystallised from MeOH/petroleum ether (b 60-80°) to give **δ-carotene** with **m 150-151°**. Store at -20°. [Porter & Anderson *Arch Biochem Biophys* **32** 21 1951, DOI: 10.1016/0003-9861(51)90233-0; Synthesis: Manchand et al. *J Chem Soc* 2019 1965, DOI: 10.1039/JR9650002019; for Absolute Configuration see Buchecker & Eugster *Helv Chim Acta* **54** 327 1971, DOI: 10.1002/hlca.19710540131; *Beilstein* **5** III 2453, **5** IV 2617.]

γ-Carotene (β,ψ-carotene) [472-93-5, 10593-83-6] $C_{40}H_{56}$, **M 536.9**, **m 152-153.5°** (synthetic), **177.5°** (polymorph), $A_{lm}^{1\%}(\lambda_{max})$ **2055** (at **437nm**), **3100** (at **462nm**), **2720** (at **494nm**) in hexane. Purify γ-carotene by chromatography on alumina [Grade II in petroleum ether (b 60-80°) and elute with *C₆H₆], or magnesia columns. When crystallised from *C₆H₆/MeOH (2:1), it had **m 177.5°**. Store it in the dark, under an inert atmosphere at 0°, or in an evacuated tube at -20°. The purity is verified by TLC on Ca(OH)₂/Kieselgel (8:2) using petroleum ether (b 60-80°) as eluant. [Manchand et al. *J Chem Soc* 2019 1965, DOI: 10.1039/JR9650002019; Ruegg et al. *Helv Chim Acta* **44** 985 1961, DOI: 10.1002/hlca.19610440414; *Beilstein* **5** III 2453, **5** IV 2617.]

ξ-Carotene [38894-81-4] $C_{40}H_{56}$, **M 536.9**, **m 38-42°**, λ_{max} **378, 400, 425nm**, $A_{lm}^{1\%}(\lambda_{max})$ **2270** (**400nm**), in petroleum ether. Purify ξ-carotene by chromatography on 50% magnesia-HyfloSupercel, developing with hexane and eluting with 10% EtOH in hexane. It crystallises from toluene/MeOH. [Gorman et al. *J Am Chem Soc* **107** 4404 1985, DOI: 10.1021/ja00301a006.] Store it in the dark under N₂ or argon at -20°. Also purify it like γ-Carotene. [*Beilstein* **5** IV 2623.]

Citranaxanthin [5,9,14,18-tetramethyl-20-(2,6,6-trimethyl-1-cyclohex-1-enyl)- 3,5,7,9, 11,13,15,17,19-eicosanonen-2-one] [3604-90-8] $C_{33}H_{44}O$, **M 456.7**, **m 155-156°**, $A_{lm}^{1\%}(\lambda_{max})$ **410** (**349nm**), **275** (**466nm**) in hexane (food additive **E161i**). Purify it by chromatography on a column of 1:1 MgO and HyfloSupercel (diatomaceous filter aid). Crystallise it from petroleum ether (deep violet crystals). Store it in the dark under N₂ or argon at 0°.

Crocetin diethyl ester (8,8'-diapo-ψ,ψ-carotenedioic acid diethyl ester) [5056-14-4] $C_{24}H_{32}O_4$, **M 384.5**, **m 216-218°**, **218-219°**, **222.5°**, $A_{lm}^{1\%}(\lambda_{max})$ **2340** (at **400nm**), **3820** (at **422nm**), **3850** (at **450nm**) in petroleum ether. Purify the diethyl ester by chromatography on a column of silica gel G. Recrystallise it from *benzene. Store it in the dark, under N₂ or argon, at 0°. The *dimethyl ester*, purified in the same way, has **m 225-227°** and similar spectra. [For synthesis, UV and IR see Isler et al. *Helv Chim Acta* **40** 1242 1957, DOI: 10.1002/hlca.19570400515; *Beilstein* **2** III 2018, H **30** 106.] The *free acid*, **Crocetin (all-trans tetramethylhexadeca-2,4,6,8,10,14-heptaenedioic acid)** [27876-94-4, di-Na salt 64603-92-5] $C_{20}H_{24}O_4$, **M 328.4**, forms brick red crystals from acetic anhydride with **m 286°** and **pK₁²⁵ 4.39**. [See details in Gainer & Grabiak Patent, US 7759506 B2, publication date July 20, 2010; Buchta & Andree *Chem Ber* **93** 1349 1960, DOI: 10.1002/cber.19600930617.]

α-Cryptoxanthin (all-trans-3R,6R'-β, ε-carotene-3-ol, zeinoxanthin, physoxanthin) [24480-38-4] $C_{40}H_{56}O$, **M 552.9**, **m 175-176°** (racemate **157.5-158.5°**), $[\alpha]_D^{25}$ **-508** (c **0.4** Me₂CO), **CD_{max} Δε -3.6** (**218nm**), **+4.4** (**245nm**) and **-9.0** (**283nm**), λ_{max} (hexane) **421, 446, 474nm**, ϵ_{mol} **145,500** (at **446nm**, hexane), **130,000** (at **459nm**, *C₆H₆). It crystallises from *C₆H₆/MeOH in red needles. The *racemate* is purified through an Al₂O₃ column, eluting with *C₆H₆ and recrystallising from MeOH/Et₂O. It has λ_{max} (ϵ 10⁻³) (hexane) at 421, 444, 471nm (90, 133, 122). [Loeber et al. *J Chem Soc (C)* 404 1971, DOI: 10.1039/J39710000404; Goodfellow et al. *J Chem Soc (D)* 1578 1970, DOI: 10.1039/C29700001578; *Beilstein* **6** IV 5113.]

β-Cryptoxanthin (all-trans-3R-β,β-carotene-3-ol, caricaxanthin) [472-70-8] $C_{40}H_{56}O$, **M 552.9**, **m 169°** (natural), (racemate **m 172-173°**), ϵ_{mol} **131,900** (at **449nm**, petroleum ether). Purify it by chromatography on MgO, CaCO₃ or deactivated alumina, using EtOH or diethyl ether to develop the column. Crystallise it from

*C₆H₆/EtOH (metallic prisms), or needles from *C₆H₆. Store it in the dark under N₂ or argon at -20°. The **acetate** has **m 117.5°**. The **racemate** is purified through a column of alumina (grade IV), eluted with *C₆H₆ then EtOAc/*C₆H₆ (1:9) and recrystallised from petroleum ether (b 60–80°) with **m 172–173°**. [Loeber et al. *J Chem Soc (C)* 404 1971, DOI: 10.1039/J39710000404; Goodfellow et al. *JCS Chem Commun* 1578 1970, DOI: 10.1039/C29700001578; Isler et al. *Helv Chim Acta* **40** 456 1957, DOI: 10.1002/hlca.19570400225; *Beilstein* **6** III 3772, **6** IV 5111.]

3,30-Diketospirilloxanthin (all-*trans*-2,31-dimethoxy-2,6,10,14,19,23,27,31-octamethyl-4,6,8,10,12,-14,16,18,20,22,24,26,28-dotriacontatridecene-3,30-dione, 2,2'-diketospirilloxanthin) [24009-17-4] C₄₂H₅₆O₄, **M 624.9**, **m 225–227°**, $\epsilon_{1\text{m}}^{1\%}(\lambda_{\text{max}})$ **550 (at 349nm), 820 (at 422nm), 2125 (at 488nm), 2725(516nm), 2130(at 551nm) in hexane**. Purify it by chromatography on a column of partially deactivated alumina. Recrystallise it from acetone/petroleum ether. Store it in the dark, in an inert atmosphere at 0°. [cf. *Beilstein* **1** III 2297, **1** IV 2750.]

Dolichols [11029-02-0; 2067-66-5 for C₁₀₀H₁₆₄O, **M 1382.4**] **C₈₀–C₁₀₅ polypren-mono-ols**. Crystallise each dolichol 6 times from petroleum ether/EtOH at -20°. The pure individual **prenol** should run as an entity on a chromatogram on paraffin impregnated paper, with acetone as the mobile phase. [For isolation, UV, IR, ¹H NMR and characterisation of a C₁₀₀ dolichol from pig liver see Burgos et al. *Biochem J* **88** 470 1963, DOI: 10.1042/bj0880470] Dolichols are a group of long-chain polyunsaturated alcohols made up of isoprene units (head to tail) where the terminal isoprene unit is saturated and the terminal carbon atom possesses a hydroxyl group. Their biosynthesis proceeds *via* the HMG-CoA reductase (mevalonate) pathway, which is **inhibited** by **Statins**, where at first a *cis* (Z) prenyltransferase catalyses the condensation of farnesyl pyrophosphate (FPP) with varying numbers of isopentenyl pyrophosphates (depending on the transferase). This produces polyprenyl pyrophosphates, known as **dehydrodolichyl diphosphates**, which lose the diphosphate group to form **dehydrodolichols**, and finally the terminal isoprene residue is reduced to produce the α-saturated polyprenyl alcohols, the **dolichols**. The lengths of these straight-chain isoprene residues can run up to more than C₁₀₀. They are widely distributed in the plant and animal kingdoms, and play an important role in the mammalian nervous systems including brain tissue. [For occurrence in mammalian nervous system see Fedorow et al. *J Neurochemistry* **92** 990 2005, DOI: 10.1111/j.1471-4159.2004.02975.x; and Ward & Guan et al. 'Identification and quantification of dolichols and dolichoic acids in neuromelanin from **substantia nigra** of the human brain' *J Lipid Res* **48**(7) 1457–1462 2007, DOI: 10.1194/jlr.C700008-JLR200; for biological role see Chojnacki & Dallner *Biochem J* **251** 1 1988, DOI: 10.1042/bj2510001, PMID: 3291859; for metabolic disorders of dolichol biosynthesis defects see Cantagrel & Lefeber *J Inherit Metab Dis* **34** 859 2011, DOI: 10.1007/s10545-011-9301-0]. Dolichols tend to be formed and found with varying poly-isoprene lengths together and named according to the number of isoprene residues, e.g. Dol-14 to Dol-18 [for recent results of structural studies see Skorupinska-Tudek et al. *Chem Rec* **8**(1) 33–54 2008, DOI: 10.1002/tcr.20137].

Echinenone (β,β-caroten-4-one) [432-68-8] C₄₀H₅₄O, **M 550.8**, **m 178–179°**, $A_{1\text{m}}^{1\%}(\lambda_{\text{max}})$ **2160 (at 458nm) in petroleum ether**. Purify this xanthophyll (found in some cyanobacteria; and isolated from sea urchins) by chromatography on partially deactivated alumina or magnesia, or by using a thin layer plate of silica gel G with 4:1 cyclohexane/diethyl ether as the developing solvent. Recrystallise it from *C₆H₆/MeOH. Store it in the dark at -20°. The **oxime** crystallises from *C₆H₆ with **m 208°**. [For isolation from sea urchins see Lederer & Moore 'Echinenone as a Provitamin A', *Nature* **137**(3476): 996–996 1936, DOI: 10.1038/137996b0; for isolation from *Micrococcus roseus* see Schwartzel & Cooney *J bacteriol* **104** 272–274 1970, PMID: 5473895; *Beilstein* **7** III 2858, **7** IV 1881.]

Ethyl bixin (6,6'-diapo-Ψ,Ψ-carotenedioic acid monomethyl ester monoethyl ester) [6895-43-8] **M 436.6**, **m 138°**. Crystallise the ester from EtOH. **Diethyl-norbixin**, C₂₈H₃₆O₄, **M 436.6**, prepared by synthesis, purified through Al₂O₃/CH₂Cl₂ and eluted with EtOH, then recrystallised from *C₆H₆ has **m 199°**, and UV(petroleum ether) with λ_{max} at 431, 455 and 488nm ($E_{1\text{cm}}^{1\%}$ 2340, 3580 and 3590) [synthesis, UV and IR: Isler et al. *Helv Chim Acta* **40** 1242 1957, DOI: 10.1002/hlca.19570400515]. [cf. Bixin above, *Beilstein* **2** III 2020, **2** IV 2356.]

Lutein See xanthophyll.

Lycopene (all-trans- Ψ,Ψ -carotene) [502-65-8] $C_{40}H_{56}$, M 536.9, m 172-173°, 174°, ϵ_{mol} 184,900 (470nm petroleum ether), 180,600 (at 487nm $*C_6H_6$). This bright red carotenoid pigment which is found in tomato species crystallises from CS_2 /MeOH, diethyl ether/petroleum ether, or acetone/petroleum ether. It can also be purified by column chromatography on deactivated alumina, $CaCO_3$, calcium hydroxide or magnesia. Its solubility in hexane is 1g/L at $\sim 15^\circ$. The deep red solid is oxygen sensitive and should be stored in the dark, in an inert atmosphere. Also purified like γ -Carotene. [Beilstein 1 III 1076, 1 IV 1165.]

Lycoxanthin (alltrans- Ψ,Ψ -carotene-16,16'-ol) [19891-74-8] $C_{40}H_{56}O$, M 552.9, m 168°, 173-174°, ϵ $\frac{1\%}{lcm}$ 3360 (472.5nm), also λ_{max} 444 and 503nm in petroleum ether. Crystallise lycoxanthin from diethyl ether/light petroleum, $*C_6H_6$ /petroleum ether (red-brown round crystal aggregates) or CS_2 (purple needles). Purify it also by chromatography on columns of $CaCO_3$, $Ca(OH)_2$ or deactivated alumina, and washing with $*benzene$ and eluting with 3:1 $*C_6H_6$ /MeOH. Its UV(Me₂CO) has λ_{max} at 448, 474 ($E \frac{1\%}{lcm}$ 3080) and 505nm; also λ_{max} ($*C_6H_6$) at 458, 487 and 521nm ($10^{-3}\epsilon$ 141.8, 162.7 and 108.8). Store it in the dark, in an inert atmosphere, at -20° . The *monoacetate* forms deep purple needles from $*C_6H_6$ /MeOH m 137°. The *dipalmitate ester* crystallises from $*C_6H_6$ /MeOH and has m 76°. [For revised structure of lycoxanthin and lycophyll see Chohnoky, Szabolcs & Waight *Tetrahedron Lett* 9 1931 1968, DOI: 10.1016/S0040-4039(01)99058-6; for stereochemistry see Kelly et al. *Acta Chem Scand* 25 1607 1971, DOI: 10.3891/acta.chem.scand.25-1607; for total synthesis see Kjosén & Liaaen-Jensen *Acta Chem Scand* 25 1500 1971, DOI: 10.3891/acta.chem.scand.25-1500; Beilstein 1 III 2051, 1 IV 2368.]

Methylbixin (6,6'-diapo- Ψ,Ψ -carotenedioic acid dimethyl ester) [26585-94-4] $C_{26}H_{32}O_4$, M 408.5, m stable *trans* form 203°, 205-206° (corr), unstable *cis* form 164°, λ_{max} 405, 425, 450, 484nm in hexane. Crystallise the dimethyl ester from EtOH/ $CHCl_3$, or $*benzene$ (blue-purple needles with m 203°). Also purify it by chromatography on alumina (Grade III) and eluting with 9:1 $*C_6H_6$ /petroleum ether (b 60-80°), and recrystallising from EtOAc. [Pattenden et al. *J Chem Soc (C)* 235 1970, DOI: 10.1039/J39700000235; Beilstein 2 III 2020, 2 IV 2356.] Its UV(petroleum ether) has λ_{max} 432, 456 and 490nm ($E \frac{1\%}{lcm}$ 2650, 4050 and 4050) [for synthesis, UV and IR see Isler et al. *Helv Chim Acta* 40 1242 1957, DOI: 10.1002/hlca.19570400515; Buchta & Andree *Chem Ber* 92 3111 1959, DOI: 10.1002/cber.19590921214]. See Bixin above.

Physalien (all-trans β -carotene-3,3'-(*R,R*)-diol dipalmitate) [144-67-2] $C_{72}H_{116}O_4$, M 1045.7, m 98.5-99.5°, $A \frac{1\%}{lm} (\lambda_{max})$ 1410 (at 449nm), 1255 (at 478nm) in hexane. Purify it by chromatography on water-deactivated alumina, using hexane/diethyl ether (19:1) to develop the column. It crystallises from $*benzene$ /EtOH in red needles. Store in the dark, in an inert atmosphere, at 0° . [See Zeaxanthin dipalmitate.] [Beilstein 6 III 5970.]

Phytoene (7,7',8,8',11,11',12,12'-octahydro- Ψ,Ψ -carotene) [540-04-5; 15-cis- 13920-14-4] $C_{40}H_{64}$, M 544.9, $A \frac{1\%}{lm} (\lambda_{max})$ 850 (at 287nm) in hexane, λ_{max} 275, 287 and 297nm. Purify phytoene by chromatography on columns of magnesium oxide-Supercel (a diatomaceous filter aid) or alumina [For total synthesis see Davis et al. *J Chem Soc (C)* 2154 1966, DOI: 10.1039/J39660002154; for isolation and properties see Rabourn et al. *Arch Biochem Biophys* 48 267 1954, DOI: 10.1016/0003-9861(54)90341-0]. Store it as a solution in petroleum ether under nitrogen at -20° . [Beilstein 1 IV 1155.]

Phytofluene (all-trans- 5,6,7,8,9,10,10',9',8',7',6',5'-dodecahydrolycopene) [540-05-6; 15-cis- 27664-65-9] $C_{40}H_{62}$, M 542.9, b 140-185°(bath temperature)/0.0001mm, $A \frac{1\%}{lm} (\lambda_{max})$ 1350 (at 348nm) in petroleum ether, λ_{max} 331, 348, 267. Purify the viscous orange oil by chromatography on partially deactivated alumina [Goodwin *Biochem J* 53 538 1953, Kushwaha et al. *J Biol Chem* 245 4708 1970, PMID: 4393960; Davis et al. *J Chem Soc (C)* 2154 1966, DOI: 10.1039/J39660002154]. Store it as a solution in petroleum ether under N_2 at -20° . [Beilstein 1 III 1072, 1 IV 1159.]

Prolycopene (all-cis [*Z*]- Ψ,Ψ -carotene or lycopene) [2361-24-2] $C_{40}H_{56}$, M 536.5, m 111°, λ_{max} 443.5, 470

nm in petroleum ether. Purify polycopene by chromatography on deactivated alumina [Kushwaha et al. *J Biol Chem* **245** 4708 1970, PMID: 4393960]. Crystallise it from petroleum ether. Store it in the dark, under N₂ or argon at -20°. [Beilstein **1** III 1079, **1** IV 1167.] It is a naturally occurring stereoisomer of lycopene [Zechmeister et al. *Proc Natl Acad Sci U S A*. 1941 Oct 15; **27**(10) 468–474 1941, PMCID: PMC1078363]. For the stepwise isomerisation of polycopene, a polycis-carotenoid, to *all-trans*-lycopene, and characterisation see Magoon & Zechmeister [*Arch Biochem Biophys* **69** 535-47 1957, DOI: 10.1016/0003-9861(57)90518-0]

Proneurosporene (3,4,7',8'-tetrahydrolycopene, 7,9,9-tricis-neurosporene) [10467-46-6] C₄₀H₅₈, M 538.5, λ_{max} 408, 432, 461 nm, ε_{1cm}^{1%} 2040 (at 432nm) in hexane. Purify proneurosporene by chromatography on deactivated alumina [Kushwaha et al. *J Biol Chem* **245** 4708 1970, PMID: 4393960;]. Store it in the dark, in an inert atmosphere at 0°. [Beilstein **1** IV 1156.]

Retinol (vitamin A alcohol), retinoic acid (vitamin A acid), retinyl acetate (vitamin A acetate), retinal (vitamin A aldehyde), retinyl palmitate (vitamin A palmitate) See in 'Physiologically Active..', this chapter.

Spirilloxanthin [rhodoviolascein, 2,31-dimethoxy-2,6,10,14,19,23,27,31-octamethyl-dotriaconta-4,6,8,10,12,14,16,18,20,22,24,26,28-tridecaene, (3*E*,3'*E*)-1,1'-Dimethoxy-3,3',4,4'-tetrahydro-1,1',2,2'-tetrahydro-ψ,ψ-carotene] [928-13-2, 22076-36-4, 26645-74-9, 34255-08-8] C₄₂H₆₀O₂, M 596.9, m 216-218°, 218°, λ_{max} 463, 493, 528 nm, ε_{1cm}^{1%} 2680 (at 493nm) in petroleum ether (b 40-70°). Spirilloxanthin crystallises from CHCl₃/petroleum ether, acetone/petroleum ether, *C₆H₆/petroleum ether or *C₆H₆ as deep red spindle-like crystals. Purify it also by chromatography on a column of CaCO₃/Ca(OH)₂ mixture, Ca(OH)₂ or deactivated alumina and elute it with a *C₆H₆/MeOH mixture. It gives a blue colour with SbCl₃ in CHCl₃ with λ_{max} at 642nm. [Polgar et al. *Arch Biochem Biophys* **5** 243 1944; for synthesis see Surmatis & Ofner *J Org Chem* **28** 2735 1963, DOI: 10.1021/jo01045a057; Karrer & Koenig *Helv Chim Acta* **23** 460 1940, DOI: 10.1002/hlca.19400230160; for 'Isolation and light-stimulated expression of canthaxanthin and spirilloxanthin biosynthesis genes from the photosynthetic bacterium *Bradyrhizobium* sp. strain ORS278', see Giraud & Verméglio *Methods in molecular biology (Clifton NJ)* **892** 173 2012, PMID: 22623302.] Store it in the dark in an inert atmosphere at -20°. [Beilstein **1** III 2297, **1** IV 2750.]

Squalane (Cosbiol, 2,6,10,15,19,23-hexamethyltetracosane, perhydrosqualene) [111-01-3] C₃₀H₆₂, M 422.8, m -38°, b 176°/0.05mm, 210-215°/1mm, 274°/10mm, ~350°/760mm, d₄²⁰ 0.80785, n_D²⁰ 1.416. Purify squalane by fractional distillation *in vacuo* or evaporative distillation. It is soluble in petroleum ether, *C₆H₆, Et₂O and CHCl₃, slightly soluble in alcohols, Me₂CO and AcOH but insoluble in H₂O. Small quantities can be purified by TLC as for squalene below. It is used as a marker in GLC and HPLC. [Staudinger & Leupold *Helv Chim Acta* **15** 221 1932, DOI: 10.1002/hlca.19320150120; Sax & Stross *Anal Chem* **29** 1700 1957, DOI: 10.1021/ac60131a044; Mandai et al. *Tetrahedron Lett* **22** 763 1981, DOI: 10.1016/0040-4039(81)80145-1; Beilstein **1** IV 593.]

Squalene (all-trans-2,6,10,15,19,23-hexamethyl-2,6,10,14,18,22-tetracosahexaene) [111-02-4] C₃₀H₅₀, M 410.7, m -5°, ~75°, b 203°/0.1mm, 213°/1mm, 285°/25mm, d₂₅²⁵ 0.8670, n_D²⁰ 1.4905. Crystallise squalene repeatedly from Me₂CO (1.4ml/g) using a Dry-ice bath, wash the crystals with cold acetone, then freeze the squalene under vacuum. Squalene is further purified by passage through a column (or TLC plate) of silica gel or chromatographed on activated alumina, using petroleum ether as eluent and stored in a vacuum in the dark. Dauben et al. [*J Am Chem Soc* **74** 4321 1952, DOI: 10.1021/ja01137a022] purified squalene *via* its **hexachloride** which is anti-bactericidal. Squalene is a key intermediate in sterol biosynthesis. [Capstack et al. *J Biol Chem* **240** 3258 1965, PMID: 14321361; Krishna et al. *Arch Biochem Biophys* **114** 200 1966, DOI: 10.1016/0003-9861(66)90322-5; Heilbron & Thompson *J Chem Soc* 883 1929, DOI: 10.1039/JR9290000883; Karrer et al. *Helv Chim Acta* **13** 1084 1930, DOI: 10.1002/hlca.19300130532; UV: Farmer et al. *J Chem Soc* 5411943, DOI: 10.1039/JR9430000541; Beilstein **1** III 1071.] Squalene is abundant in nature and particularly in shark liver oil. In animals it is the main natural precursor of sterols (*via* lanosterol), and in plants also it is the main precursor of sterols (*via* cycloartenol).

Violaxanthin (5S,6S,5'S,6'S-diepoxyzeaxanthin) [126-29-4] $C_{40}H_{56}O_4$, **M 600.9**, **m 200°, 207-208°, $[\alpha]_{D}^{18} +35$ (c 0.1, $CHCl_3$)**. Crystallise violaxanthin from MeOH (orange prisms) by adding a little Et_2O in MeOH, or CS_2 (red brown crystals). Also purify it by column chromatography, and the purity is checked by TLC (see δ -carotene). It has λ_{max} at 415, 440 and 469nm. [Kuhn & Winterstein *Ber* **64** 326 1931, Karrer & Morf *Helv Chim Acta* **14** 1044 1931, DOI: 10.1002/hlca.19310140513; for absolute configuration see Bartlett et al. *J Chem Soc (C)* 2527 1969, DOI: 10.1039/J39690002527; *Beilstein* **19** III/IV 1139.] It is a food additive and is biosynthesised from zeaxanthin (below) by epoxidation.

Xanthophyll (Lutein, 3R,3'R- α -carotene-3,3'-diol) [127-40-2] $C_{40}H_{56}O_2$, **M 568.9**, **m 183°, 193°, 196°, $[\alpha]_{D}^{18} +165$ (c 0.7, $*C_6H_6$)**[NB: Cd I and II have λ 533 & 537 nm], $\epsilon_{1\%}^{1cm}(\lambda_{max})$ **1750 (423nm), 2560 (446nm), 2340 (477.5nm) and ϵ_{mol} 144,800 (at 445nm) in EtOH; λ_{max} in CS_2 446, 479 and 511nm, and ϵ_{mol} 127,000 (at 458nm, $*C_6H_6$)**. Crystallise lutein from MeOH (1g/700ml; copper-coloured prisms with a metallic lustre) or from diethyl ether by adding MeOH. Also purify it by chromatography on columns of magnesia or calcium hydroxide, and recrystallise it from $CS_2/EtOH$. The UV has λ_{max} (ϵ) (dioxane) at 268nm (35,000), 333nm (15,500), 429nm (100,000), 453nm (152,000) and 481nm (142,000). It may be purified via the **dipalmitate ester** which crystallises from EtOH (red needles) with **m 92°**. Store it in the dark, in an inert atmosphere, e.g. argon [Zechmeister *Chem Rev* **34** 267 1944, DOI: 10.1021/cr60108a004; Buchecker et al. *Helv Chim Acta* **57** 631 1974, DOI: 10.1002/hlca.19740570317; Karrer *Helv Chim Acta* **34** 2160 1951, DOI: 10.1002/hlca.19510340708; for absolute configuration see Goodfellow et al. *JCS Chem Commun* 1578 1970, DOI: 10.1039/C29700001578; *Beilstein* **6** II 1026, **6** III 5871, **6** IV 7017.] The yellow colour of chicken egg yolks, fat and skin comes from ingested xanthophyll and zeaxanthin.

Zeaxanthin [alltrans- β -carotene-3,3'(R,R')-diol] [144-68-3] $C_{40}H_{56}O_2$, **M 568.9**, **m 205-206° (natural), 207°, 215.5°, $[\alpha]_{D}^{18} -44°$ (c 0.7, $CHCl_3$)**, [NB: Cd I and II have λ 533 & 537 nm], λ_{max} **275nm (log ϵ 4.34), 450nm (log ϵ 5.14), 480nm (log ϵ 5.07) in EtOH, ϵ_{mol} 132,900 (at 452nm, Me_2CO)**. Zeaxanthin, one of the most common carotenoid alcohols in nature, forms yellow plates (with a blue lustre) from MeOH, EtOH and with 0.5 MeOH from $*C_6H_6/MeOH$. The **diacetate** (from natural) has **m 154-155°** (from $CH_2Cl_2/MeOH$), the UV has λ_{max} ($\epsilon_{1\%}^{1cm}$) at 452, 480nm (2210, 1945). The **dipalmitate (Physalein)** (from natural source) has **m 95-96°** (from $CH_2Cl_2/MeOH$), the UV has λ_{max} ($\epsilon_{1\%}^{1cm}$) at 452, 480nm (1335, 1190). [Isler et al. *Helv Chim Acta* **39** 2041 1956, DOI: 10.1002/hlca.19560390717; Hlubucek et al. *JCS Perkin Trans 1* 848 1974, DOI: 10.1039/P19740000848.] The **racemate** is purified by chromatography on an alumina (Grade IV) column by elution with a gradient of petroleum ether (b 60-80°), $*C_6H_6$ and $EtOAc$. The red band gives a solid which recrystallises from MeOH with **m 211°**, and the UV has λ_{max} at 439infl, 462, 589nm (ϵ 10^{-3} : 90, 126, 110 respectively). [Loeber et al. *J Chem Soc (C)* 404 1971, DOI: 10.1039/J39710000404, *Beilstein* **6** II 1026, **6** III 5865, **6** IV 7017.]

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STEROIDS

This section also includes steroidal hormones, steroidal alkaloids and cardenolides.

(-)-3- β -Acetoxy-5-etiolic acid [3- β -acetoxy-5-etiocholenic acid, androst-5-ene-17- β -carboxylic acid] [7150-18-7, 51424-66-9] $C_{22}H_{32}O_4$, **M 306.5**, **m 238-240°**, **241-242°**, **243-245°**, **246-247°**, $[\alpha]_D^{20}$ **-19.9** (**c 1**, **Me₂CO**), **-36** (**c 1**, **dioxane**), **-33.5°** (**CHCl₃**), **pK_{Est} ~ 4.7**. The acid is purified by recrystallisation from Me₂CO, Et₂O/pentane, or AcOH, dried in a vacuum oven (105°/20mm) and sublimed at high vacuum. [Staunton & Eisenbraun *Org Synth* **42** 4 1962, DOI: 10.15227/orgsyn.042.0004; Steiger & Reichstein *Helv Chim Acta* **20** 1040 1937, DOI: 10.1002/hlca.193702001150.] A useful acid for the *optical resolutions* of alcohols [J. Jacques et al. *Enantiomers, Racemates, and Resolutions* Wiley-Interscience, New York, p 333 1981].

21-Acetoxypregnenolone (3 β -hydroxy-21-acetoxypregn-5-en-20-one) [566-78-9] $C_{23}H_{34}O_4$, **M 374.3**, **m 184-185°**. Crystallise 21-acetoxypregnenolone from Me₂CO by allowing the solvent to evaporate in a vacuum to a small volume, collect the colourless needles and wash them with Et₂O, then dry it *in vacuo*. The crystals become opaque on standing. [Steiger & Reichstein *Helv Chim Acta* **20** 1164 1937, DOI: 10.1002/hlca.193702001158.]

Adrenosterone (Reichstein's G, androst-4-ene-3,11,17-trione) [382-45-6] $C_{19}H_{24}O_3$, **M 300.4**, **m 214-217°**, **219-222°**, **220-224°**, **224-226°**, $[\alpha]_{546}^{20}$ **+364** (**c 0.18**, **EtOH**), $[\alpha]_D^{20}$ **+300** (**c 1**, **CHCl₃**). Dissolve adrenosterone in Me₂CO, decolorise it with charcoal, filter, add H₂O, Me₂CO evaporate and the solid is recrystallised from aqueous EtOH. Also recrystallise it from Et₂O or Et₂O/pentane and dry it at 110°/0.1mm for 2 hours. It can be sublimed under high vacuum. [Reichstein *Helv Chim Acta* **20** 953, 1937 DOI: 10.1002/hlca.193702001138; 979 1937, DOI: 10.1002/hlca.193702001141, Mason et al. *J Biol Chem* **116** 267 1936, <http://www.jbc.org/content/116/1/267>; *Beilstein* **7** III 4601.]

Aldosterone (18-aldocorticosterone, Reichstein's X) [52-39-1] $C_{21}H_{28}O_5$, **M 360.5**, **m 108-112°(hydrate)**, **164°(anhydrous)**, $[\alpha]_D^{25}$ **+161** (**c 1**, **CHCl₃**). Crystallise aldosterone from aqueous acetone. It exists in solution as an equilibrium mixture of free aldehyde and its cyclic hemiacetal, favouring the hemiacetal. The *d*-**isomer** is considered as the biologically active one. The **21-acetate** crystallises from Me₂CO/Et₂O or CH₂Cl₂/EtOAc and has **m 198-199°**, and $[\alpha]_D^{25}$ **+121.7** (**c 0.7**, **CHCl₃**). [Barton et al. *JCS Perkin Trans 1* 2243 1975, DOI: 10.1039/P19750002243; *Beilstein* **8** IV 3491.]

5 α -Androstane (etioallocholane) [5 α - 438-22-2, 24887-75-0] $C_{19}H_{32}$, **M 260.5**, **m 50-50.5°**, $[\alpha]_D$ **+2** (**c 0.12**, **CHCl₃**). Purify 5 α -androstane by chromatography through Al₂O₃ (grade III) and elute with petroleum ether then recrystallise it from MeOH (2x) and Me₂CO/MeOH. It sublimes at 60°/high vacuum. Also dissolve it in petroleum ether, filter it through silica gel, evaporate and recrystallise it. [Steiger & Reichstein *Helv Chim Acta* **20** 817 1937, DOI: 10.1002/hlca.193702001117; Prelog et al. *Helv Chim Acta* **27** 66 1944, DOI: 10.1002/hlca.19440270109; IR, NMR: Halkes & Havinga *Recl Trav Chim Pays-Bas* **84** 889 1965, DOI: 10.1002/recl.19650840710; Allinger et al. *Tetrahedron* **27** 5093 1971, DOI: 10.1016/S0040-4020(01)90764-1; *Beilstein* **5** III 1110, **5** IV 1211.]

5 β -Androstane (etiocholane, testane) [438-23-3] $C_{19}H_{32}$, **M 260.5**, **m 78-79°**, **78-80°**, $[\alpha]_D^{20}$ **+2.03**, $[\alpha]_{546}^{18}$ **+5** (**c 1**, **CHCl₃**). Crystallise etiocholane from acetone. The method of purification for 5 α -androstane (above) could be used here. [Shoppee *Helv Chim Acta* **27** 246, 260 1944, DOI: 10.1002/hlca.19440270128; Butenandt & Dannerbaum *Hoppe Seyler's Z Physiol Chem* **229** 192 1934, *Beilstein* **5** III 1110, **5** IV 1211.]

4-Androstene-3,17-dione (androstenedione) [63-05-8] $C_{19}H_{26}O_2$, **M 286.4**, **m 170-171°**, **173-174°**, $[\alpha]_D^{20}$ **+196** (**c 0.13**, **EtOH**). Crystallise the dione from hexane. It is soluble in *C₆H₆, CHCl₃ and EtOH. It has λ_{max} at 239nm. It is a precursor of estrone or testosterone and has androgenic activity. [Ruzicka & Wettstein *Helv Chim Acta* **18** 986 1935, DOI: 10.1002/hlca.193501801132; *Beilstein* **7** III 3636, **7** IV 2381.]

Androsterone (Cis) (3 α ,5 α -3-hydroxyandrostane-17-one) [53-41-8] C₁₉H₃₀O₂, M 290.4, m 181-184°, 185-185.5°, [α]_D²⁰ +118 (c 1, EtOH), [α]_D²⁰ +96 (c 0.7, EtOH). Crystallise androsterone from Me₂CO/Et₂O or Me₂CO and sublime it at high vacuum. The *acetate* [1164-95-0] crystallises from Et₂O, Me₂CO/Et₂O or aqueous EtOH, and sublimates at high vacuum with m 165-166°, and [α]_D²⁵ +87 (c 2, EtOH). [Ruzicka et al. *Helv Chim Acta* **17** 1389 1934, DOI: 10.1002/hlca.193401701169; Marker *J Am Chem Soc* **57** 1755 1935, DOI: 10.1021/ja01312a505; Göndös & Orr *JCS Chem Commun* 1239 1982, DOI: 10.1039/C39820001239; Beilstein **8** IV 642.]

epi-Androsterone (Trans, 3 β -hydroxy-5 α -androstane-17-one) [481-29-8] C₁₉H₃₀O₂, M 290.4, m 172-173°, (161-162° *dl*-), [α]_D²⁰ +115 (c 1, MeOH), [α]_D²⁰ +44.4 (c 0.27, CHCl₃). Purify *epi*-androsterone via the acetate, hydrolyse this and recrystallise it from CHCl₃/hexane or aqueous EtOH. The *acetate* [1239-31-2] is purified by chromatography and when crystallised from petroleum ether has m 103-104°, and [α]_D +68.5 (c 1, CHCl₃). The *oxime* has m 194-196° (from MeOH), and [α]_D +17.5 (c 6.2, CHCl₃). The *racemic ketone* is sublimed at 130°/high vacuum and after two crystallisations from methylcyclohexane it gives prisms with m 161-162° (which changed crystal form at 140-145°). [Ruzicka & Wettstein *Helv Chim Acta* **18** 1264 1935, DOI: 10.1002/hlca.193501801176; Johnson et al. *J Am Chem Soc* **75** 2275 1953, DOI: 10.1021/ja01105a530; **78** 6331 1956, DOI: 10.1021/ja01605a020; Cardwell et al. *J Chem. Soc* 361 1953, DOI: 10.1039/JR9530000361; Beilstein **8** IV 462.] It is a steroid hormone with weak androgenic activity.

Betamethasone (9 α -fluoro-11 β ,17 α ,21-trihydroxy-16 β -methylpregna-1,4-diene-3,20-dione) [378-44-9] C₂₂H₂₉FO₅, M 392.5, m 231-136°(dec), 235-237°(dec), [α]_D²⁰ +108 (c 1, Me₂CO). Betamethasone crystallises from ethyl acetate, and has λ_{\max} at 238nm (log ϵ 4.18) in MeOH. The *21-acetate* [287-24-6] crystallises from Me₂CO/Et₂O (charcoal) with m 196-201° (205-208°) and has [α]_D²⁰ +140 (CHCl₃). [Taub et al. *J Am Chem Soc* **82** 4012 1960, DOI: 10.1021/ja01500a053; Oliveto et al. *J Am Chem Soc* **80** 6687 1958, DOI: 10.1021/ja01557a058; Beilstein **8** IV 3501.]

Bisnorcholanic acid (pregnane-20-carboxylic acid) [28393-20-6] C₂₂H₃₆O₂, M 332.5, m 214° (α -form), 242° (β -form), 210-211° (γ -form), 184° (δ -form), 181° (ϵ -form), pK_{Est} ~5.0. Bisnorcholanic acid crystallises from EtOH (α -form), or acetic acid (*all forms*). The α -form has [α]_D²⁰ -7.5 (c 0.8, EtOH), and its *ethyl ester* has m 106-107°; the β -form has [α]_D²⁰ +23.3 (c 0.86, EtOH), and its *ethyl ester* is an oil; the γ -form has [α]_D²⁰ 0, and its *ethyl ester* has m 82-83°; the δ -form has [α]_D²⁰ +31.7 (c 0.76, EtOH), and its *methyl ester* has m 99°; the ϵ -form has [α]_D²⁰ +14.3 (c 0.8, EtOH), and its *methyl ester* has m 117°. [For synthesis from cholanolic acid see Hollander & Gallagher *J Biol Chem* **162** 549 1946, <http://www.jbc.org/content/162/3/549>; Wieland et al. *Hoppe Seyler's Z Physiol Chem* **161** 80 1926, Beilstein **9** III 2650.]

Brassicasterol [24 β (R)-methylcholesta-5,22E-dien-3 β -ol, 5,22-cholestadien-24 β (R)-methyl-3 β -ol, 24R-Ergosta-5,22E-dien-3 β -ol] [474-67-9] C₂₈H₄₆O, M 398.7, m 148°, 149-150°, 157-158°, [α]_D²⁰ -60, -66, (c 1.2, CHCl₃). It is purified from the 7,22E-isomer, which is formed during synthesis, by TLC on Silica Gel G-Celite (1:1) with 30% EtOAc/hexane as eluant and recrystallised from MeOH or EtOH [Anastasia et al. *JCS Perkin I* 379 1983, DOI: 10.1039/P19830000379]. In a different synthesis from the *enantiomeric* 23,24-cyclopropane precursor 3 β -acetates, and 22E,24 β (R)-3 β -brassicasteryl acetate, m 152-154°, was obtained and purified by HPLC [using two Altex Ultrasphere™ ODS columns 5 μ m (10mm id x 25cm) and MeOH mobile phase], has t_R = 118 minutes] then crystallised from MeOH. The *acetate* has the ¹H NMR (360MHz, CDCl₃, TMS) with δ at 5.37 (m, 1H, H-C6), 5.18 (m, 2H, H-C22 and H-C23), 4.60 (m, 1H, H-C3), 2.03 (s, 3H, OCOCH₃), 1.02 (s, 3H, H₃C19), 1.01 (d, 3H, J = 6.49Hz, H₃C21), 0.91 (d, 3H, J = 6.80Hz, H₃C28), 0.83 (d, 3H, J = 6.5Hz) and 0.83 (d, 3H, J = 6.5Hz, H₃C26), 0.81 (d, 3H, J = 6.5Hz, H₃C27), 0.69 (s, 3H, H₃C18); and HRMS m/z 380.3452 (100, M-HOAc), C₂₈H₄₄⁺ requires 380.3443. [Lang & Djerassi *Helv Chim Acta* **65** 407 1982, DOI: 10.1002/hlca.19820650144; Rubinstein et al. *Phytochemistry* **15** 195 1976, DOI: 10.1016/S0031-9422(00)89083-4]. The *acetate* (5mg) in dry Et₂O is de-acetylated with LiAlH₄ (4 fold) at ~25°/30 minutes, excess LiAlH₄ is decomposed with saturated aqueous Na₂SO₄, H₂O is added and the free sterol is extracted into Et₂O, dried (MgSO₄), evaporated and the residue is recrystallised from MeOH to give pure (22,24R)-brassicasterol m 156-158°, with one peak in HPLC (as above but with an ODS-2 column) at t_R = 55 minutes. Its ¹H NMR (360MHz, CDCl₃, TMS) has δ at 5.35 (m, 1H, H-C6), 5.181 (m, 2H, H-C22 and H-C23), 3.53 (m,

1H_{ax}, H-C3), 1.011 (d, 3H, $J = 6.34\text{Hz}$, H₃C21), 1.010 (s, 3H, H₃C19), 0.910 (d, 3H, $J = 6.84\text{Hz}$, H₃C28), 0.834 (d, 3H, $J = 6.10\text{Hz}$) and 0.817 (d, 3H, $J = 6.22\text{Hz}$, H₃C26), 0.815 (d, 3H, $J = 6.22\text{Hz}$, H₃C27), 0.693 (s, 3H, H₃C18); and the HRMS has m/z at 398.3545 (M^+), C₂₈H₄₆O requires 398.3549. [Lang & Djerassi *Helv Chim Acta* **65** 407 1982, DOI: 10.1002/hlca.19820650144; Rubinstein et al. *Phytochemistry* **15** 195 1976, DOI: 10.1016/S0031-9422(00)89083-4; Sheikh & Djerassi *Steroids* **26** 129 1975, DOI: 10.1016/0039-128X(75)90010-0]. It is a plant sterol (phytosterol) present in canola oils and rape seed, but is also found in some marine algae, diatoms and shellfish (oysters). It *inhibits* the enzyme Δ^{24} -sterol reductase that is involved in mammalian cholesterol biosynthesis [Fernandez et al. *Biochem J* **366** 109 2002, DOI: 10.1042/bj20011777]. **Brassino steroids** have plant growth regulating activities [Anastasia et al. *JCS Perkin I* 379 1983, DOI: 10.1039/P198300003793].

Crinosterol [24 α (S)-methylcholesta-5,22E-dien-3 β -ol, 5,22-cholestadien-24 α (S)-methyl-3 β -ol, 24S-Campesta-5,22E-dien-3 β -ol] [17472-78-5] C₂₈H₄₆O, M 398.7, m 147-148°, 152-154° [α]_D²⁶ -47.2 (c 1.2, CHCl₃). Crinosterol occurs in sponges, oysters and diatoms. It is the **24- α -methyl epimer of Brassicasterol** (see [474-67-9] above). They sometimes occur together in Nature and sometimes are formed together in synthesis; purification of one invariably requires the removal of the other. **Crinosteryl acetate** m 148-150°, when isolated from a synthesis was separated from the **24- β -methyl epimer** by HPLC (see brassicasterol above), had $t_R = 112$ minutes, and was recrystallised from MeOH, or *alternatively*, it is prepared by acetylation of crinosterol, then purified by TLC [10% w/w AgNO₃/silica gel and developed with pure CHCl₃] to give homogeneous (by GLC) **acetate** with m 157-158° (compare with above). Crinosteryl acetate has IR (Nujol) with ν_{\max} at 1720, 970, 955 and 795 cm⁻¹, and the ¹H NMR (360MHz, CDCl₃, TMS) with δ at 5.375 (m, 1H, H-C6), 5.160 (m, 2H, H-C22 and H-C23), 4.601 (m, 1H_{ax}, H-C3), 2.033 (s, 3H, OCOCH₃), 1.019 (s, 3H, H₃C19), 1.002 (d, 3H, $J = 6.68\text{Hz}$, H₃C21), 0.909 (d, 3H, $J = 6.78\text{Hz}$, H₃C28), 0.835 (d, 3H, $J = 6.52\text{Hz}$) and 0.817 (d, 3H, $J = 6.63\text{Hz}$, H₃C26), ~0.815 (d, 3H, $J = 6.6\text{Hz}$, H₃C27), 0.690 (s, 3H, H₃C18); and HRMS m/z 380.3420 (100, M-HOAc), C₂₈H₄₄ requires 380.3443. [Lang & Djerassi *Helv Chim Acta* **65** 407 1982, DOI: 10.1002/hlca.19820650144; Anastasia et al. *JCS Perkin I* 379 1983, DOI: 10.1039/P19830000379; for a stereoselective synthesis see Anastasia et al. *JCS Perkin I* 2365 1983, DOI: 10.1039/P19830002365; Rubinstein et al. *Phytochemistry* **15** 195 1976, DOI: 10.1016/S0031-9422(00)89083-4;]. Deacetylation with LiAlH₄ (see brassicasterol above) gives **crinosterol** which is purified by TLC [silica gel developed with 2% v/v EtOH/CHCl₃] has m 152-154° (reported twice, and 147-148° reported twice, both recrystallised from MeOH or Me₂CO), and has ¹H NMR (360MHz, CDCl₃, TMS) has δ at 5.349 (m, 1H, H-C6), 5.162 (m, 2H, H-C22 and H-C23), 3.53 (m, 1H_{ax}, H-C3), 1.010 (s, 3H, H₃C19), 1.001 (d, 3H, $J = 6.21\text{Hz}$, H₃C21), 0.910 (d, 3H, $J = 6.86\text{Hz}$, H₃C28), 0.836 (d, 3H, $J = 6.45\text{Hz}$) and 0.817 (d, 3H, $J = 6.62\text{Hz}$, H₃C26), 0.815 (d, 3H, $J = 6.62\text{Hz}$, H₃C27), 0.693 (s, 3H, H₃C18); and HRMS m/z 398.3551 (M^+), C₂₈H₄₆O requires 398.3549. [Lang & Djerassi *Helv Chim Acta* **65** 407 1982, DOI: 10.1002/hlca.19820650144; Anastasia et al. *JCS Perkin I* 2365 1983, DOI: 10.1039/P19830002365; Rubinstein & Goad *Phytochemistry* **13** 485 1974, DOI: 10.1016/S0031-9422(00)91239-1; see Rubinstein et al. *Phytochemistry* **15** above]. **Brassino steroids** have plant growth regulating activities [Anastasia et al. *JCS Perkin I* 379 1983, DOI: 10.1039/P19830000379].

Campesterol (24R-24-methylcholest-5-en-3 β -ol) [474-62-4] C₂₈H₄₈O, M 400.7, m 156-159°, 157-158°, [α]_D²⁴ -35.1 (c 1.2, CHCl₃). This plant sterol (phytosterol) is recrystallised twice from hexane and once from Me₂CO. The **benzoyl derivative** has m 158-160° and [α]_D²³ -8.6 (CHCl₃); and the **acetyl derivative** has m 137-138° (EtOH) and [α]_D²³ -35.1 (c 2.9, CHCl₃) [Fernholz & MacPhillamy *J Am Chem Soc* **63** 1155 1941, DOI: 10.1021/ja01849a079]. [Beilstein **6** III 2680.] Its structure is similar to that of cholesterol, except that it has a 24(R)-Me group in the side-chain, and competes with it in mammalian organs.

CHAPS (3-[(3-cholamidopropyl)dimethylammonio] -1-propanesulfonate betaine) [75621-03-3 anhydrous], [313223-04-0 hydrate] C₃₂H₅₈N₂O₇S, xH₂O, M 614.88, m 157°(dec), pK_{Est(1)} ~2 (acidic), pK_{Est(2)} ~10 (basic). Triturate this zwitterionic detergent (~50g) with MeOH (200ml), then triturate it with Me₂CO (500ml), collect it by vacuum filtration and dry it thoroughly at ~20°. Recrystallise it at 0° from absolute MeOH followed by drying *in vacuo* to constant weight. It should give one spot with R_F 0.32 by TLC on silica gel G in 95% MeOH/5% NH₄OH and visualised with iodine vapour; the tertiary amine precursor has R_F 0.40. It has low UV absorption (1% in H₂O: A_{280nm} is 0.029 and A_{260nm} is 0.035), and its CMC (critical micellar concentration)

is 8mM. It is soluble in H₂O, MeOH, Me₂SO but insoluble in Me₂CO, MeCN, hexane, toluene and xylene. [Hjelmeland et al. *Anal Biochem* **130** 72 1983, DOI: 10.1016/0003-2697(83)90651-6; Matuo et al. *Methods Enzymol* **198** 511 1991, DOI: 10.1016/0076-6879(91)98050-G.]

CHAPSO (3-[(3-cholamidopropyl)dimethylammonio]-2-hydroxy-1-propanesulfonate betaine) [82473-24-3] C₃₂H₅₈N₂O₈S, xH₂O, M 630.89, m 184-196°, [α]_D²⁰ +19 (c 2, H₂O), pK_{Est(1)} ~<2 (acidic), pK_{Est(2)} ~9 (basic). Dissolve this zwitterionic detergent (~60g) in 40% aqueous MeOH and stir it with the mixed-bed ion-exchange resin RG-501 x 8 (100g, Bio-Rad) until the pH of the supernatant is ~7. Filter off the resin and evaporate the filtrate to dryness under reduced pressure. Check for homogeneity by TLC on silica gel G in 95% MeOH/5% NH₄OH and visualize the spot with iodine vapour (see CHAPS above). [Hjelmeland et al. *Anal Biochem* **130** 72 1983, DOI: 10.1016/0003-2697(83)90651-6; Matuo et al. *Methods Enzymol* **198** 511 1991, DOI: 10.1016/0076-6879(91)98050-G.]

Chenodeoxycholic acid (chenodiol, 3α,7α-dihydroxycholan-24-oic acid) [474-25-9] C₂₄H₄₀O₄, M 392.6, m (119°), 143°, 165-167°, [α]_D²⁰ +14 (c 2, EtOH), pK_{Est} ~4.9. This major bile acid in vertebrates (~80mg) is chromatographed on silica gel (5g) and eluted with CHCl₃/EtOAc (3:2) and crystallised from EtOAc/hexane. It has IR with ν_{max} at 1705 cm⁻¹(CHCl₃). It also crystallises from EtOAc, EtOAc/heptane after purifying via the poorly soluble Na and K salt if necessary. [Kametani et al. *J Org Chem* **47** 2331 1982, DOI: 10.1021/jo00133a020; Beilstein **10** IV 1604.]

5α-Cholanic acid (allocholanic acid) [468-98-4, 546-98-4] C₂₄H₄₀O₂, M 360.6, m 170, [α]_D²⁰ +22 (CHCl₃), pK_{Est} ~4.9. Purify 5α-cholanic acid by chromatography on silica gel and eluting with MeOH/EtOAc/hexane (1:5:10) then recrystallising from Et₂O/hexane, MeOH or AcOH. The *methyl ester* has m 91° (from MeOH) and ¹H NMR (CDCl₃) has δ for C18 & C19 at 0.66 and 0.78 [see 5β-cholanic acid below, Mandava et al. *Steroids* **23** 357 1974, DOI: 10.1016/0039-128X(74)90087-7]. [Demir et al. *Org Prepn Proced Int* **19** 197 1987, DOI:10.1080/00304948709356187; Stoll et al. *Helv Chim Acta* **18** 644 1935, DOI: 10.1002/hlca.19350180184; Beilstein **9** III 2656, **9** IV 1992.]

5β-Cholanic acid (ursocholanic acid) [546-18-9, 25312-65-6] C₂₄H₄₀O₂, M 360.6, m 164-165°, 164-167°, [α]_D²⁰ +21.7 (CHCl₃), pK_{Est} ~4.9. Crystallise 5β-cholanic acid from EtOH, Me₂CO or AcOH. The *ethyl ester* has m 93-94° (from 80% EtOH), b 273°/12mm, [α]_D²⁰ +21 (CHCl₃), and the *methyl ester* has m 86-87° (from MeOH) after purification through Al₂O₃ and eluting with *C₆H₆; and has [α]_D²⁰ +21.2 (CHCl₃). The ¹H NMR (CDCl₃) for the C18 & C19 protons has δ at 0.65 and 0.92 [see 5α-cholanic acid above, Mandava et al. *Steroids* **23** 357 1974, DOI: 10.1016/0039-128X(74)90087-7]. [Stoll et al. *Helv Chim Acta* **18** 644 1935, DOI: 10.1002/hlca.19350180184; Huang-Minlon *J Am Chem Soc* **71** 3301 1949, DOI: 10.1021/ja01178a008; Beilstein **9** III 2656, **9** IV 1992.]

Cholesta-3,5-diene [747-90-0] C₂₇H₄₄, M 368.6, m 77-78°, 78°, 78-79°, 79-80°, 79.5-80.5°, 80-81°, [α]_D¹⁶ -116 (c 1.1, CHCl₃). This diene has been prepared in many ways, but several of these are from solvolysis of epicholesteryl toluene-*p*-sulfonate using KOAc/Ac₂O (38 hours at 75°), then methanolic KOAc (reflux at 3.5 hours), followed by reduction with LiAlH₄/Et₂O (20 hours at 20°) in ca 60% yield [Schmid & Kägi *Helv Chim Acta* **35** 2194 1952, DOI: 10.1002/hlca.19520350704], or AgOAc/AcOH, KOAc/MeOH or KOAc/AcOH (reflux ~3 hours 92-99% yield), and with neutralised Al₂O₃ at 20° [Evans & Shoppee *J Chem Soc* 540 1953, DOI: 10.1039/JR9530000540; Shoppee et al. *J Chem Soc* 2876 1955, DOI: 10.1039/JR9550002876], KOAc/aqueous Me₂CO [Becker & Wallis *J Org Chem* **20** 353 1955, DOI: 10.1021/jo01121a013], and diphenylguanidinium salts accelerate solvolysis [Shoppee & Williams *J Chem Soc* 686 1955, DOI: 10.1039/JR9550000686]. After the usual workup involving washing with NaHCO₃ and H₂O, the diene is separated from other steroids and impurities by chromatography through Al₂O₃ (various grades, including activity II-III, neutralised, basic or activated at 250-310°) and usually eluted with pentane, pentane/*C₆H₆ or Et₂O/MeOH. It is then recrystallised from Me₂CO, Et₂O/MeOH or Et₂O/EtOH, dried *in vacuo* and stored at -20°. The UV has λ_{max} (log ε) at 228.5 (4.308), 235 (4.344) and 243 (4.163) nm.

Cholesta-5,7-dien-3β-ol (7-dehydrocholesterol, 3β-hydroxy-5,7-cholestadiene, Provitamin D₃) [434-16-2] C₂₇H₄₄O, M 384.6, m 150-151° (anhydrous), 148-152°, [α]_D²⁰ -113.6 (c 1, CHCl₃), [α]_D²⁰ -127 (*C₆H₆). It

occurs in humans and higher animals, and has been isolated from pig skin or the horned snail (*Buccinum undatum*). It is purified by recrystallisation from MeOH, Et₂O/MeOH or aqueous Et₂O (plates) and holds to H₂O tenaciously. It is insoluble in H₂O but is soluble in most organic solvents. Store it at -20° in the dark and absence of air which oxidises it slowly. Irradiation with UV light produces vitamin D₃ and like all provitamin D have λ_{\max} at 260, 270, 281 and 293.5 nm. The *acetate* [1059-86-5] has **m 129-130°**, and $[\alpha]_{\text{D}}^{20}$ **-85.3 (c 1.2, *C₆H₆)**, crystallises from MeOH; and the *benzoate* has **m 139-140°**, and $[\alpha]_{\text{D}}^{20}$ **-53.2 (c 1.2, CHCl₃)**, and crystallises from CHCl₃/Me₂CO in plates. [Bernstein et al. *J Org Chem* **14** 433 1949, DOI: 10.1021/jo01155a015; *Beilstein* **6** IV 4153]. Not to be confused with *dihydrocholesterol*.

Cholesta-5,7-dien-3 α -ol [57651-84-0] **C₂₇H₄₄O**, **M 384.6**, **m 125-126°**, $[\alpha]_{\text{D}}^{20}$ **-70.5 (CHCl₃)**, crystallises from MeOH in needles, and its *acetate*, has **m 114.5°**, and $[\alpha]_{\text{D}}^{20}$ **-35 (CHCl₃)**. It also crystallises from MeOH in needles. [Aberhart et al. *J Org Chem* **41** 1067 1976, DOI: 10.1021/jo00868a031]. [for NMR of steroids see Kirk et al. *JCS Perkin Trans 2* 1567 1990, DOI: 10.1039/P29900001567; and Anastasia et al. *Steroids* **49** 543 1987, DOI: 10.1016/0039-128X(87)90094-8; and for ORD see Weiss et al. *Tetrahedron* **21** 3105 1965, DOI: 10.1016/S0040-4020(01)90764-1].

Cholesta-5,24-dien-3 β -ol (Desmosterol, 24-dehydrocholesterol, 3 β -hydroxy-5,24-cholestadiene, natural 3 β ,20*R*-form) [313-04-2] **C₂₇H₄₄O**, **M 384.6**, **m 117-118°, 119-119.5°, 120.5-121°, 121.5°, 121.5-122.5°**, $[\alpha]_{\text{D}}^{20}$ **-38.2 (c 1.14, CHCl₃)**. It occurs in chick embryos, rat skins, red algae, barnacles and *Funtmia latifolia* seeds. This is a useful intermediate for the synthesis of vitamin D metabolites [Koreeda et al. *J Org Chem* **45** 1172 1980, DOI: 10.1021/jo01294a056]. The sterol in hexane can be purified by chromatography on a AgNO₃/silicic acid/Super-Cel column in *C₆H₆, then re-chromatographed on Alumina F20, eluting with *C₆H₆ followed by *C₆H₆/EtOAc (1:1 v/v) and recrystallised from distilled MeOH or from Me₂CO/MeOH. It has IR (KBr) with ν_{\max} at 1373, 1056, 1022, 959, 950, 835 and 800 cm⁻¹; and HRMS has *m/z* 384.3421 (M⁺), C₂₇H₄₄O requires 384.3389 [Apfel *J Org Chem* **43** 2284 1978, DOI: 10.1021/jo00405a044]. The *acetate* [2665-04-5] **m 99°**, $[\alpha]_{\text{D}}^{28}$ **-44.8 (c 1, CHCl₃)**, can be obtained by refluxing 3-chlorocholesta-5,24-diene with AcOH/AcOK for 2.5 hours, cooling, adding H₂O, extracting with Et₂O, drying (MgSO₄), evaporating the extract, followed by chromatography through silica gel, eluting with *C₆H₆, evaporating and the white crystalline residue is recrystallised from Me₂CO/MeOH. The *acetate* has ¹H NMR (60MHz, CCl₄, TMS) with δ at 0.68 (s, 3H, H₃C18), 0.98 (s, 3H, H₃C19), 1.00 (s, 3H, H₃C21), 1.58 (s, 3H, H₃C26 or H₃C27), 1.65 (s, 3H, H₃C27 or H₃C26), 1.94 (s, 3H, 3-OCOCH₃), 4.50 (m, 1H_{ax}, H-C3), 5.00 (m, H, H-C24) and 5.35 (m, 1H, H-C5). Hydrolysis of the acetate with MeOH/KOH gave *desmosterol* with **m 119-120°**, and $[\alpha]_{\text{D}}^{28}$ **-41 (CHCl₃)**. [Ochi et al. *Steroids* **30** 795 1977, DOI: 10.1016/0039-128X(87)90094-8]. *Desmosterol* has also been obtained from the readily available *3-hydroxy-bisnorcholenic acid* in several synthetic steps and finally via the 3-tetrahydropyranyl ether which was hydrolysed in THF containing a few drops of concentrated HCl at 40°/5 minutes to give the *sterol m 120-122°* in 70% yield after recrystallisation from MeOH. It has ¹H NMR (60MHz, CDCl₃, TMS) with δ at 0.41 (s, H₃C18), 0.51 (d, *J* = 6Hz, H₃C21), 0.61 (s, H₃C19), 1.97 and 1.02 (2s, H₃C26, H₃C27), 2.10 (1H_{ax}, H-C3), 3.03 (24-H) and 3.20 (6-H). [Dasgupta et al. *J Org Chem* **39** 1658 1974, DOI: 10.1021/jo00925a013.] Store it at -20°.

Cholesta-5,24-dien-3 α -ol [67392-80-7] **M 384.6**, has **m 137-139°** (from MeOH), $[\alpha]_{\text{D}}^{28}$ **-43.5 (CHCl₃)**, and its *acetate* [67383-62-4] **M 426.6**, has **m 112-115°** (from MeOH), $[\alpha]_{\text{D}}^{28}$ **-47.5 (CHCl₃)**.

The **3 β ,20*S*-form** [59532-46-6] **M 384.6**, has **m 125-127°** (from MeOH), and is obtained by dehydration of 20*S*-3 β ,25-dihydroxycholest-5-ene with dioxane/H₂SO₄ (50% yield). It has IR (KBr) with ν_{\max} at 3000 (OH) cm⁻¹, and ¹H NMR (60MHz, CDCl₃, TMS) with δ at 0.70 (s, H₃C18), 0.84 (d, H₃C21), 1.02 (s, H₃C19), 1.62 and 1.70 (2s, H₃C26, H₃C27). It is an *inhibitor* of the *cholesterol side-chain cleavage enzyme system* in bovine adrenocortical preparations [Burstein et al. *Steroids* **27** 361 1976, DOI: 10.1016/0039-128X(76)90057-X]; and is the *immediate precursor* of cholesterol in the *Bloch biosynthetic pathway*.

5 α -Cholestane [481-21-0] **C₂₇H₄₈**, **M 372.7**, **m 80°, 80-80.5°**, $[\alpha]_{\text{D}}^{20}$ **+29.5**, $[\alpha]_{\text{D}}^{20}$ **+24**, $[\alpha]_{\text{D}}^{18}$ **+30.2**, (c 2, CHCl₃). 5 α -Cholestane is prepared by catalytic reduction of cholestene with EtOAc/PtO₂/H₂, and recrystallised from EtOAc or Et₂O/EtOH. Store it at room temperature. The ¹H NMR (100MHz, CDCl₃, TMS) signals of the methyl groups have δ at 0.645 (s, 13-Me), 0.778 (s, 10-Me), 0.898 (, *J* = 6.0Hz, 20-Me) and 0.861 (d, *J* = 6.3Hz, 25-Me₂) [Wittstruck et al. *JCS Perkin Trans I* 1403 1977, DOI: 10.1039/P19770001403], and its CD spectrum in hexane has a $\Delta\epsilon$ value of +0.5 at the longest wavelength of 183nm [Kirk et al. *Tetrahedron Lett* 1355 1973,

DOI: 10.1016/S0040-4039(01)95941-6]. **Note** that catalytic hydrogenation of cholest-4-ene in acidic medium provides **5- α -cholestane** whereas hydrogenation in neutral media results in **5- β -cholestane** [Windaus *Chem Ber* **52** 170 1919, DOI: 10.1002/cber.19190520122]. [Ruzicka et al. *Helv Chim Acta* **16** 327, 334 1933, DOI: 10.1002/hlca.19330160146; *Beilstein* **5** III 1132, **5** IV 1227.]

5 β -Cholestane (coprostan, ψ -cholestane, pseudocholestane) [481-20-9, 25269-18-5 + 21 other numbers + 105308-41-6] **C₂₇H₄₈, M 372.7, m 70°, 71-72°, [α]_D²⁰ +25 (c 2, CHCl₃)**. Catalytic hydrogenation (Pt₂O/Et₂O) of 3-chlorocholest-4-ene (see above) followed by crystallisation from Me₂CO [Young et al. *J Am Chem Soc* **81** 1452 1959, DOI: 10.1021/ja01515a042], or catalytic reduction (Pt₂O/Et₂O) followed by chromatography through an Al₂O₃ column and eluting with petroleum ether followed by recrystallisation of the residual oil from Et₂O/EtOH give high yields of coprostan [Nickon & Bagli *J Am Chem Soc* **83** 1498 1961, DOI: 10.1021/ja01467a049] (see previous entry). It also crystallises in needles from EtOH. Store it at room temperature. Its CD spectrum in hexane has a $\Delta\epsilon$ value of +1.1 at the longest wavelength of 184nm (see previous entry) [Kirk et al. *Tetrahedron Lett* 1355 1973, DOI: 10.1016/S0040-4039(01)95941-6].

5 α -Cholestan-3 β -ol (β -cholestanol, dihydrocholesterol) [80-97-7] **C₂₇H₄₈O, M 388.7, m 142-143°(monohydrate), [α]_D²⁰ +28 (c 1, CHCl₃), [α]_D²⁰ +27.4 (in CHCl₃)**. Purify 5- α -cholestan-3 β -ol *via* acetylation, crystallisation and de-acetylation, then recrystallisation from EtOH or slightly aqueous EtOH, or MeOH. Its solubility is: 0.5% (MeOH) and 1% (EtOH) at 25°. [Mizutani & Whitten *J Am Chem Soc* **107** 3621 1985, DOI: 10.1021/ja00298a036.] The **acetate** has **m 114-115°** from EtOAc/MeOH and [α]_D²⁰ +13 (c 2, CHCl₃). [Bruce & Ralls *Org Synth Coll Vol* **2** 191 1943, DOI: 10.15227/orgsyn.017.0045; *Beilstein* **6** IV 3577.]

5 β -Cholestan-3 α -ol (Epicoprostanol, epi-5 β -cholestanol) [516-88-7, 516-92-7] **C₂₇H₄₈O, M 388.7, m 113-114°, 115-116°, 116-117°, 118°, [α]_D²⁰ +32 (c 1.8, CHCl₃)**. It is the main product from the acetolysis of 3 β -chlorocoprostan (KOAc/AcOH, reflux for 6 hours) involving a **Walden inversion**, and purified as in the following entry [Bridgewater & Shoppee *J Chem Soc* 1709 1953, DOI: 10.1039/JR9530001709]. It has also been obtained by reduction of 5 β -coprostan-3-one with LiAlH₄ to give a 94% yield of epicoprostanol **m 115-116°**, which did not form a precipitate with digitonin, and the small amount of digitonide formed gave a small quantity of coprostanol **m 100°** (~4% yield and crystallised from MeOH, see below) [Shoppee & Summers *J Chem Soc* 687 1950, DOI: 10.1039/JR9500000687]. *Alternatively*, the residue is purified as described in the succeeding entry or recrystallised from Me₂CO or Et₂O/*C₆H₆ to give the desired **3 α -ol**, after drying *in vacuo* at 80°. **Note** that crystals from EtOH have **m 115-116°**, and from Me₂CO have **m 117-118°**. Store it at room temperature. The **3 α -yl-acetate** (obtained by catalytic reduction of coprostan-3-one, Et₂O/MeOH/PtO₂/H₂, followed by chromatographic separation as below and acetylation) has **m 87-89°, 89-91°, and [α]_D¹⁸ +42 (c 1.42, CHCl₃)**. [Shoppee & Summers *J Chem Soc* 687 1950, DOI: 10.1039/JR9500000687; Ruzicka et al. *Helv Chim Acta* **17** 1407 1934, DOI: 10.1002/hlca.193401701171.]

5 β -Cholestan-3 β -ol (coprostan-3 β -ol) [360-68-9] **C₂₇H₄₈O, M 388.7, m 100-101°, 101°, 102°, [α]_D²⁰ +28 (c 1.8, CHCl₃)**. It is the main product from the acetolysis of 3 α -chlorocoprostan (KOAc/AcOH, reflux for 6 hours) involving a **Walden inversion**. Unlike epicoprostanol (above), coprostanol is precipitated by a 1.33% solution of digitonin in 95% EtOH, the digitonide can be collected, decomposed by dissolving in pyridine, precipitating the digitonin with excess of Et₂O, filtering off digitonin and the coprostanol is obtained by evaporation of the organic solvent. A possible impurity is the **3 α -ol** from which it can be separated by chromatography on Al₂O₃ and eluting with Et₂O/*C₆H₆ (1:24) whereby the **3 α -ol** runs first (**m 116-117°**, see preceding entry) and further elution gives the pure **3 β -coprosterol (m 99-101°)**. Crystallise it from MeOH (needles) or aqueous EtOH. Store it at room temperature. The **3 β -yl-acetate** from acetylation with Ac₂O (also obtained by catalytic reduction of coprostan-3-one, AcOH/PtO₂/H₂, followed by chromatographic separation as above and acetylation) has **m 91°, 90- 91°, and [α]_D¹⁸ +14.5 (c 1.8, CHCl₃)**. Its solubility is 0.79% in H₂O at 25°. [Bridgewater & Shoppee *J Chem Soc* 1709 1953, DOI: 10.1039/JR9530001709; Shoppee & Summers *J Chem Soc* 687 1950, DOI: 10.1039/JR9500000687; see Ruzicka et al. *Helv Chim Acta* **17** 1407 1934, DOI: 10.1002/hlca.193401701171.]

Cholest-2-ene [15910-23-3] **C₂₇H₄₆, M 370.6, m 75-76°, [α]_D²⁰ +66 (c 1.65, CHCl₃)**. Purify cholest-2-ene by chromatography on Al₂O₃ (Spence H) and elute with petroleum ether (b 40-60°), then recrystallise the residue

from EtOAc/MeOH. Also recrystallise it from MeOH or diethyl ether/acetone. [Alt & Barton *J Chem Soc* 4284 1954, DOI: 10.1039/JR9540004284; Bergbreiter & Chandran *J Am Chem Soc* **109** 174 1987, DOI: 10.1021/ja00235a027; *Beilstein* **5** III 1320, **5** IV 1507.]

Cholesterol (cholest-5-en-3 β -ol) [57-88-5] $C_{27}H_{46}O$, **M 386.7**, **m 147-149°**, **148.9-149.4°**, **b 360°/atm**, $[\alpha]_D^{25}$ **-35** (hexane), $[\alpha]_D^{20}$ **-39** (c **2**, $CHCl_3$). Crystallise cholesterol from ethyl acetate, EtOH or isopropyl ether/MeOH. [Hiromitsu & Kevan *J Am Chem Soc* **109** 4501 1987, DOI: 10.1021/ja00249a012.] For extensive details of purification through the dibromide, see Fieser [*J Am Chem Soc* **75** 5421 1953, DOI: 10.1021/ja01117a508] and Schwenk and Werthessen [*Arch Biochem Biophys* **40** 334 1952, DOI: 10.1016/0003-9861(52)90119-7], and by repeated crystallisation from acetic acid; see Fieser [*J Am Chem Soc* **75** 4395 1953, DOI: 10.1021/ja01114a003]. Like many sterols, cholesterol gives colour reactions with concentrated H_2SO_4 . When cholesterol is dissolved in a small volume of $CHCl_3$ and mixed with concentrated H_2SO_4 , the colour of the organic layer becomes crimson, then changes to purple, and on further standing in air it turns to blue, then green and finally yellow. The H_2SO_4 layer develops a green fluorescence. [*Beilstein* **6** III 2607, **6** IV 4000; Fieser **1** 141.]

Cholesteryl acetate [604-35-3] $C_{29}H_{48}O_2$, **M 428.7**, **m 112-114°**, **113-115°**, **115-116°**, $[\alpha]_{546}^{20}$ **-51** (c **5**, $CHCl_3$), $[\alpha]_D^{20}$ **-39** (c **2**, $CHCl_3$). Crystallise the acetate from *n*-pentanol or Me_2CO . Also purify it by chromatography through silica gel and eluting with MeOH. [*Beilstein* **6** III 2607, **6** IV 4004.]

Cholesteryl myristate [1989-52-2] $C_{41}H_{72}O_2$, **M 597.0**, **m 69-71°**, **84°**, $[\alpha]_D^{20}$ **-25.4** (c **1**, $CHCl_3$). Crystallise the myristate ester from Me_2CO , EtOH/EtOAc or EtOH/Et₂O or *n*-pentanol. Purify it also by column chromatography on silica gel and eluting with MeOH then evaporating to dryness. Recrystallise it and finally, dry it *in vacuo* over P_2O_5 and store it at -20°. [Labarrere et al. *Analyt Chem* **30** 1466 1958, DOI: 10.1021/ac60141a008; Malanik & Malát *Anal Chim Acta* **76** 464 1975, DOI: 10.1016/S0003-2670(01)85421-0; *Beilstein* **6** III 2638].

Cholesteryl oleate [303-43-5] $C_{45}H_{78}O_2$, **M 651.1**, **m 44-47°**, **48.8-49.4°**, $[\alpha]_D^{20}$ **-24** (c **1**, $CHCl_3$). Purify the oleate ester by chromatography on silica gel and eluting with MeOH. [*Beilstein* **6** III 2642.]

3 α -5 β -7 α -12 α -Cholic acid (3 α -7 α -12 α -trihydroxy-5 β -cholan-24-oic acid, Cholalin) [81-25-4] $C_{24}H_{40}O_5$, **M 408.6** (anhydrous), **426.6** (hydrate) **m 196-198°**, **198-200°**, **200-201°**, $[\alpha]_D^{20}$ **+37**, $[\alpha]_{546}^{20}$ **+41** (c **0.6**, EtOH), **pK_a²⁰ 4.98**. This bile acid crystallises from H_2O , aqueous AcOH (as *hydrate*), wet Et₂O (as *hydrate*) or EtOH (as *alcoholate*). Dry it under vacuum at 94° to give the *anhydrous acid m 198°*. Its solubility (w/v) at 15° is 0.028% in H_2O , 3.06% in EtOH, 0.12% in Et₂O, 0.51% in $CHCl_3$, 0.036% in $*C_6H_6$, 2.82% in Me_2CO and 15.2% in AcOH. When an alcoholic solution of cholic acid + I_2 is added to aqueous KI, it forms a molecular compound $(C_{24}H_{40}O_5I)_4 \cdot KI \cdot H_2O$. The *methyl ester* is *dimorphic* with **m 155-156°** and **162°** (from 95% aqueous EtOH), and $[\alpha]_D^{20}$ **+25** (EtOH); and the *ethyl ester* has **m 155° (162-163°** after recrystallisation from EtOAc/petroleum ether (b 30-60°) 2:8. [Cortese *J Am Chem Soc* **59** 2532 1937, DOI: 10.1021/ja01291a014]. [Anderson et al. *Biochem J* **67** 323 1957, DOI: 10.1042/bj0670323; **85** 236 1962, DOI: 10.1042/bj0850236; Ekwall et al. *Acta Chem Scand* **11** 590 1957, DOI: 10.3891/acta.chem.scand.11-0590.] The acid has been used as a *non-denaturing ionic detergent*, and for *solubilising membrane-bound proteins* in their native state [Hooper *Biochem Soc Trans* **14** 586 1986, DOI: 10.1042/bst0140586]. It has also been used in *liposome preparations* [Yang & Lundahl *Anal Biochem* **218** 210 1994, DOI: 10.1006/abio.1994.1162.] [*Beilstein* **10** III 2162, **10** IV 2072.]

4,5-Coprosten-3-ol (cholest-4-ene-3 β -ol, allocholesterol) [517-10-2] $C_{27}H_{46}O$, **M 386.7**, **m 132°**, $[\alpha]_D^{24}$ **+43.7**, (c **1**, $*C_6H_6$). Purify 4,5-coprosten-3-ol by dissolving it in Et₂O, adding an equal volume of MeOH and removing the Et₂O with a stream of CO_2 until crystallisation begins. The sterol crystallises in needles when cooled in an ice-salt bath. Dry it *in vacuo*. It is precipitated by digitonin, and gives a red colour with 90% trichloroacetic acid (*Rosenheim reaction*). The *acetate* crystallises from aqueous MeOH **m 85°** (long white needles). [Schoenheimer & Evans *J Biol Chem* **114** 567 1936, <http://www.jbc.org/content/114/3/567>; for the separation from cholesterol see Stoll *Hoppe Seyler's Z Physiol Chem* **246** 10 1937, *Beilstein* **6** IV 3577.]

Corticosterone (11 β , 21-dihydroxypregn-4-en-3,20-dione, Reichstein's substance H) [50-22-6] $C_{21}H_{30}O_4$, M 346.5, m 179-183°, 180-181°, 180-182°, 181-184°, $[\alpha]_D^{15} +223$ (c 1.1, EtOH), $[\alpha]_D^{23-25} +194$ (c 0.1, dioxane). Purify corticosterone by recrystallisation from Me_2CO (trigonal plates), EtOH or isoPrOH. It has UV with λ_{max} at 240nm, and gives an orange-yellow solution with strong fluorescence on treatment with concentrated H_2SO_4 . It is insoluble in H_2O but soluble in organic solvents. [Reichstein & Euw *Helv Chim Acta* **21** 1197 1938, DOI: 10.1002/hlca.193802101149; **27** 1287 1944, DOI: 10.1002/hlca.194402701164; Mason et al. *J Biol Chem* **114** 613 1936, <http://www.jbc.org/content/114/3/613>; ORD: Foltz et al. *J Am Chem Soc* **77** 4359 1955, DOI: 10.1021/ja01621a052; NMR: Shoolery & Rogers *J Am Chem Soc* **80** 5121 1958, DOI: 10.1021/ja01552a031.] The **21-O-acetyl derivative** [1173-26-8] crystallises from Me_2CO/Et_2O with m 152.5-153°, and $[\alpha]_D^{20} +195$ (c 0.6, Me_2CO); and the **21-O-benzoyl derivative** crystallises from AcOH/ Et_2O with m 201-202° [Reichstein *Helv Chim Acta* **20** 953 1937, DOI: 10.1002/hlca.193702001138]. [Beilstein **8** IV 2907.] This drug activates both **mineralocorticoid** [Krozowski & Funder et al. *Proc Natl Acad Sci USA (PNAS)* **80** 6056 1983, PubMedID 6310613] and **glucocorticoid** [Rousseau et al. *J Mol Biol* **67** 99 1972, DOI:10.1016/0022-2836(72)90389-0] **receptors**.

Cortisol See hydrocortisone [50-23-7] below.

Cortisone (17 α ,21-dehydrocorticosterone, 17 α ,21-dihydroxy-4-pregnene-3,11,20-trione, Reichstein's substance Fa) [53-06-5] $C_{21}H_{28}O_5$, M 360.5, m 223-228°, 230-231°, $[\alpha]_D^{20} +225$ (c 1, EtOH). Crystallise cortisone from 95% EtOH or acetone. The UV has ϵ 14,000 $M^{-1}cm^{-1}$ at 237nm (EtOH). [Hems *J Pharm Pharmacol* **5** 408 1953, PMID: 13070171; Beilstein **8** IV 3480.] It is a corticosterone analogue with *ca* twice the anti-inflammatory potency, but with much lower Na ion retention potency.

Cortisone-21-acetate [50-04-4] $C_{23}H_{30}O_6$, M 402.5, m 237-240°, 242-243°, $[\alpha]_D^{20} +227$ (c 1, $CHCl_3$). Crystallise cortisone-21-acetate from acetone or $CHCl_3$. The UV has ϵ 15,800 $M^{-1}cm^{-1}$ at 238nm in dioxane. [Sarett *J Biol Chem* **162** 601 1946, PMID: 21018769; Beilstein **8** III 4058, **5** IV 3481.]

Deoxycholic acid [83-44-3] $C_{23}H_{38}O_6$, M 392.6, m 171-174°, 176°, 176-178°, $[\alpha]_D^{20} +64$ (c 1, EtOH), $[\alpha]_D^{20} +55$ (c 2.5, EtOH), pK²⁵ 6.58. Reflux the bile acid with CCl_4 (50ml/g), filter, evaporate under vacuum at 25°, recrystallise the residue from acetone and dry it under vacuum at 155° [Trenner et al. *J Am Chem Soc* **76** 1196 1954, DOI: 10.1021/ja01633a094]. A solution of (cholic acid-free) material (100ml) in 500ml of hot EtOH is filtered, evaporate it to less than 500ml on a hot plate, and pour it into 1500ml of cold diethyl ether. The precipitate, filtered off by suction, is crystallised twice from 1-2 parts of absolute EtOH, to give an **alcoholate**, m 118-120°, which is dissolved in EtOH (100ml for 60g) and poured into boiling water. After boiling until free of the EtOH, the precipitate is filtered off, dried, ground and dried to constant weight *in vacuo* [Sobotka & Goldberg *Biochem J* **26** 555 1932, DOI: 10.1042/bj0260555]. Deoxycholic acid is also freed from fatty acids and cholic acid by silica gel chromatography and elution with 0.5% acetic acid in ethyl acetate [Tang et al. *J Am Chem Soc* **107** 4058 1985, DOI: 10.1021/ja00299a048]. It can also be recrystallised from butanone. Its solubility in H_2O at 15° is 0.24g/L, but in EtOH it is 22.07g/L. [Beilstein **10** IV 1608.]

11-Deoxycorticosterone (21-hydroxyprogesterone) [64-85-7] $C_{21}H_{30}O_3$, M 330.5, m 141-142°, $[\alpha]_D^{20} +178$ and $[\alpha]_{546} +223$ (c 1, EtOH). Crystallise 11-deoxycorticosterone from diethyl ether. [Schindler et al. *Helv Chim Acta* **24** 360 1941, DOI: 10.1002/hlca.19410240152; Steiger & Reichstein *Helv Chim Acta* **20** 1164 1937, DOI: 10.1002/hlca.193702001158.]

11-Deoxycorticosterone acetate (21-acetoxy-4-pregnen-3,20-dione) [56-47-3] $C_{23}H_{32}O_4$, M 372.5, m 154-159°, 154-160°, 155-157°, 155-161°, $[\alpha]_D^{20} +174$ (c 1, dioxane), $[\alpha]_D^{22-24} +196$ (c 1, $CHCl_3$). 11-Deoxycorticosterone acetate recrystallises from EtOH as needles or Me_2CO /hexane, and sublimes at high vacuum. It is partly soluble in MeOH, Me_2CO , Et_2O and dioxane but insoluble in H_2O . [Reichstein & Euw *Helv Chim Acta* **23** 136 1940, DOI: 10.1002/hlca.19400230116; Romo et al. *J Am Chem Soc* **79** 5034 1957, DOI: 10.1021/ja01575a055; NMR: Shoolery & Rogers *J Am Chem Soc* **80** 5121 1959, DOI: 10.1021/ja01552a031; Beilstein **8** IV 2195.]

Dexamethasone (9- α -fluoro-16- α -methylprednisolone, prednisolone F) [50-02-2] $C_{22}H_{29}FO_5$, M 392.5, m 262-264°, 268-271°, $[\alpha]_D^{25} +77.5$ (c 1, dioxane). Dexamethasone has been recrystallised from Et₂O or small volumes of EtOAc. Its solubility in H₂O is 10 mg/100ml at 25°; and is freely soluble in Me₂CO, EtOH and CHCl₃. Store it below 8°. A stock solution of 20µg/ml is generally made by adding 1ml of absolute EtOH to 100µg of steroid, stir gently to dissolve then add 49ml of sterile medium while mixing. [Arth et al. *J Am Chem Soc* **80** 3161 1958, DOI: 10.1021/ja01545a061; for the β -methyl isomer see Taub et al. *J Am Chem Soc* **82** 4012 1960, DOI: 10.1021/ja01500a053; see *Beilstein* **8** IV 3501.] This is a glucocorticoid steroidal anti-inflammatory drug.

Dexamethasone 21-acetate (9- α -fluoro-16- α -methylprednisolone-21-acetate, prednisolone F acetate) [1177-87-3] $C_{24}H_{31}FO_6$, M 434.5, m 215-225°, 229-231°, $[\alpha]_D^{25} +77.6$ (c 1, dioxane), +73 (c 1, CHCl₃). Dexamethasone 21-acetate is purified on neutral Al₂O₃ using CHCl₃ as eluent, the fractions are evaporated, and the residue is recrystallised from CHCl₃. Store it below 8°. It has λ_{max} at 239nm (ϵ 14,900) in EtOH. It is a glucocorticoid steroidal anti-inflammatory agent. [Oliveto et al. *J Am Chem Soc* **80** 4431 1958, DOI: 10.1021/ja01549a090]. [*Beilstein* **8** IV 3501.]

Dexamethasone 21-phosphate disodium salt (Baldex, Colvasone, Dexabene, Desone among many trade names) [2392-39-4] $C_{22}H_{28}FNa_2O_8P$, M 516.4, m 233-255°, $[\alpha]_D^{25} +74$ (for anhydrous salt, c 1, H₂O). Recrystallise it from EtOH and dry it *in vacuo*. It has λ_{max} at 238-239nm (ϵ 14,000) in EtOH. Store it below 8°, but ethanolic solutions should be stored at ~0°. Like the preceding two drugs it is also an *anti-inflammatory* agent, but is more soluble in H₂O. It is a *pro-drug* and is converted to dexamethasone *in vivo*; it also *stimulates glutamine uptake* in the *cerebral cortex*.

Digitoxigenin (3 β ,14 β -dihydroxy-20(22)-norcholenic acid lactone) [143-62-4] $C_{23}H_{34}O_4$, M 374.5, m 253°, 249-255°, $[\alpha]_{546}^{20} +17$ (c 1, MeOH). Dissolve digitoxigenin in EtOAc, wash it with H₂O, dry it (Na₂SO₄), evaporate and recrystallise it from 40% aqueous EtOH or aqueous MeOH then EtOAc/Et₂O. The UV has λ_{max} at 218nm (ϵ 14,500 M⁻¹cm⁻¹). The *3-acetate* crystallises from Me₂CO/Et₂O with m 222-227°, and $[\alpha]_D^{20} +21$ (c 1, CHCl₃). [Wyss et al. *Helv Chim Acta* **43** 664 1960, DOI: 10.1002/hlca.19600430308; Cruz et al. *J Org Chem* **42** 3580 1977, DOI: 10.1021/jo00442a029; *Beilstein* **18** IV 1468.]

Dihydrotachysterol [67-96-9] $C_{28}H_{46}O$, M 398.7, m 125-127°, $[\alpha]_D^{20} +97$ (CHCl₃). It was prepared by reduction of *tachysterol* (see below) with Na/isoPrOH. Crystallise the sterol from 90% MeOH, and UV with λ_{max} at 242, 251 and 261nm ($E_{1cm}^{1\%}$ 760, 1010 and 650) in EtOH. The *acetate* has m 108-110° and $[\alpha]_D^{20} +32.8^\circ$ (CHCl₃); and UV with λ_{max} at 242, 251 and 261nm ($E_{1cm}^{1\%}$ 780, 910 and 600) in EtOH. The *propionate* has m 97-98° and $[\alpha]_D^{20} +37$ (CHCl₃); and UV with λ_{max} 242, 251 and 261nm ($E_{1cm}^{1\%}$ 750, 860 and 570) in EtOH. The *2,4-dinitrobenzoate* has m 164-165° (yellow needles from Me₂CO/MeOH) and $[\alpha]_D^{18} +138$ (c 0.58, Me₂CO)[Werder *Hoppe Seyler's Z Physiol Chem* **260** 119 1939, Werder USPat 2228491 (to Winthrop 1941); Windaus et al. *Justus Liebigs Ann Chem* **499** 188 1932, DOI: 10.1002/jlac.19324990116; *Beilstein* **6** III 2833, **6** IV 3994, 4161.] This is a synthetic *vitamin D analogue* which is activated in the liver and has a rapid onset of action, albeit a shorter half-life.

Diosgenin (25R-spirostaen-3 β -ol) [512-04-9] $C_{27}H_{42}O_3$, M 414.6, m 204-207°, $[\alpha]_D^{25} -129$ (in Me₂CO). Crystallise the saponin (present in wild yams) from acetone, and chromatograph it on Al₂O₃ and elute with *C₆H₆/Et₂O (9:1), then recrystallise it from MeOH. Its solubility is ~4% in H₂O and 5% in CHCl₃. The *acetate* crystallises from AcOH with m 198°; and has $[\alpha]_D^{20} -119$ (pyridine). [Marker et al. *J Am Chem Soc* **65** 1199 1943, DOI: 10.1021/ja01246a051; Mazur et al. *J Am Chem Soc* **82** 5889 1960, DOI: 10.1021/ja01507a028; *Beilstein* **19** IV 862.]

α -Ecdyson (2 β ,3 β ,14 α ,22(R),25-pentahydroxy-7-cholesten-6-one) [3604-87-3] $C_{27}H_{44}O_6$, M 464.7, m 239-242°, 242°, $[\alpha]_D^{20} +72$ (c 1, EtOH). Recrystallise α -ecdyson from tetrahydrofuran/petroleum ether and from H₂O as a *hydrate*. It has been purified by chromatography on Al₂O₃ and elution with EtOAc/MeOH. It has λ_{max} at 242nm (ϵ 12,400). Its *acetate* has m 214-216° from EtOAc/petroleum ether, and the *2,4-dinitrophenyl-*

hydrazone has **m 170-175°**(dec) from EtOAc. [Karlson et al. *Justus Liebigs Ann Chem* **662** 1 1963, DOI: 10.1002/jlac.19636620102; Karlson *Pure Appl Chem* **14** 75 1967, DOI: org/10.1351/pac196714010075; *Beilstein* **8** IV 3613.]

β-Ecdyson (β-echdysterone, crustecdysol, 20-hydroxyecdysol) [5289-74-7] **C₂₇H₄₄O₇, M 480.7, m 240-242°, 245-247°, [α]_D²⁰ +66 (c 1, MeOH)**. Crystallise β-ecdysol from water, tetrahydrofuran/petroleum ether, MeOH and EtOAc after chromatographic purification. It has λ_{max} (EtOH) at 240nm (ε 12,670 M⁻¹cm⁻¹). [Also IR & NMR: Hueppi & Siddall *J Am Chem Soc* **89** 6790 1967, DOI: 10.1021/ja01001a087; Kametani et al. *Tetrahedron Lett* **21** 4855 1980, DOI: 10.1016/0040-4039(80)80158-4; *Beilstein* **8** IV 343.]

(+)-Equilenin (3-hydroxyestra-1,3,5,7,9-pentaen-17-one) [517-09-9] **C₁₈H₁₈O₂, M 266.3, m 250-252°, 258-259° (vac), [α]_D¹⁶ +87 (c 0.1, dioxane)**. Crystallise (+)-equilenin from EtOH (solubility is 0.63% at 18°, 2.5% at 78°), aqueous EtOH or *C₆H₆ (Norite) and dry it in a vacuum. It sublimes on melting and at 170-180°/0.01mm. The **acetate** crystallises from MeOH with **m 165-167°** and [α]_D²⁰ +71 (c 1, CHCl₃). [Bachmann et al. *J Am Chem Soc* **62** 824 1940, DOI: 10.1021/ja01861a036; *Beilstein* **8** III 1522, 1523, 1525, **8** IV 1420.]

Ergosterol (Provitamin D₂) [57-87-4] **C₂₈H₄₄O, M 396.7, m 160°, 165-166°, [α]_D²⁰ -171 (c 1, CHCl₃), [α]_D²⁰ -135 (c 1, CHCl₃)**. Crystallise ergosterol from EtOAc, then from ethylene dichloride or EtOH/*C₆H₆ (3:1). It has been purified by conversion to the **isobutyl ester** which crystallises in massive snow-white crystals from Et₂O/Me₂CO (1:3) which have **m**: turbid at **148°**, melts to a turbid liquid at **159°** and becomes clear at **162°**, which had [α]_D²⁰ -84 (c 1, CHCl₃), followed by hydrolysis. [When crystallised from EtOH, it forms the **1,5-hydrate m 168°**. The water is difficult to remove giving an **amorphous** solid **m 166-183°, b 250°/high vacuum**. It is light sensitive. The **benzoate** has **m 169-171°**, after crystallisation from Me₂CO/*C₆H₆ (4:1) [whose crystals becomes highly charged after prolonged standing at 0°] and has [α]_D²⁰ -177 (c 1, CHCl₃). [For highly purified material and its esters see Bills & Honeywell *J Biol Chem* **80** 15 1928, <http://www.jbc.org/content/80/1/15>; for UV of sterols see Hogness et al. *J Biol Chem* **120** 239 1937, <http://www.jbc.org/content/120/1/239>. *Beilstein* **6** IV 4407.]

17α-Estradiol [1,3,5(10)-estratriene-3,17α-diol] [57-91-0] **C₁₈H₂₄O₂, M 272.4, m 223, [α]_D²⁰ +57 (c 1, EtOH)**. 17α-Estradiol recrystallises from aqueous EtOH (80%) as the **hemihydrate** and differs from the **β-anomer** (below) by not precipitating with digitonin in 80% aqueous EtOH. The **diacetate** [1474-52-8] crystallises from aqueous EtOH in needles with **m 139-140°**. The **3-benzoate** crystallises in three forms **m 158°, 153° and 63°**. [See references for 17β-Estradiol below.]

17β-Estradiol (1,3,5-estratrien-3,17β-diol, 17-β-estradiol was incorrectly called α-estradiol) [50-28-2] **C₁₈H₂₄O₂, M 272.4, m 173-179°, 176-178°, [α]_D²⁰ +76 to +83 (c 1, dioxane)**. 17β-Estradiol (**previously known as α-estradiol**) is purified by chromatography on SiO₂ (toluene/EtOAc 4:1) and recrystallised from CHCl₃/hexane or 80% EtOH. It is stable in air, is insoluble in H₂O, and is precipitated by digitonin. The UV has λ_{max} at 225 and 280 nm. The **diacetate** [3434-88-6] has **m 97-98°** and forms leaflets from aqueous EtOH. The **3-benzoate** [50-50-0] crystallises from aqueous MeOH with **m 193°** and [α]_D²⁵ +58 to 63 (c 1, dioxane). [Miescher & Scholz *Helv Chim Acta* **20** 263 1937, DOI: 10.1002/hlca.19370200137; 1237 1937, DOI: 10.1002/hlca.193702001168; *Biochem J* **32** 1273 1938, DOI: 10.1042/bj0321273b; Oppolzer & Roberts *Helv Chim Acta* **63** 1703 1980, DOI: 10.1002/hlca.19800630638; Inhoffen & Zühlsdorff *Chem Ber* **74** 1911 1941, DOI: 10.1002/cber.19410741215; *Beilstein* **6** IV 6611.] It is a primary female sex hormone.

β-Estradiol-6-one (1,3,5-estratriene-3,17β-diol-6-one) [571-92-6] **C₁₈H₂₂O₃, M 286.4, m 278-280°, 281-283°, [α]_D²⁰ +4.2 (c 0.7, EtOH)**. β-Estradiol-6-one forms plates from EtOH. The **3,17-diacetate** has **m 173-175°** after recrystallisation from aqueous EtOH. [Longwell & Wintersteiner *J Biol Chem* **133** 219 1940, <http://www.jbc.org/content/133/1/219>.] The UV has λ_{max} at 255 and 326nm in EtOH [Slaunwhite et al. *J Biol Chem* **191** 627 1951, PMID: 14861208]. [*Beilstein* **8** IV 2398.]

Estriol (1,3,5-estratrien-3β,16α,17β-triol) [50-27-1] **C₁₈H₂₄O₃, M 288.4, m 278.5-284°, 283°, [α]_D²⁰ +66 (c 1, dioxane)**. Crystallise estriol from EtOH/ethyl acetate. Also purify it by countercurrent distribution with

cyclohexane/EtOAc (1:1) and EtOH/H₂O (1:1). The UV (EtOH) has λ_{max} at 280nm (ϵ 2,090 M⁻¹cm⁻¹). [Huffmann & Lott *J Am Chem Soc* **71** 719 1949, DOI: 10.1021/ja01170a096; Leeds et al. *J Am Chem Soc* **76** 2943 1954, DOI: 10.1021/ja01640a026; *Beilstein* **6** IV 7550.] Estrogenic.

Estrone (1,3,5-Estratrien-3-ol-17-one, Folliculin) [53-16-7] C₁₈H₂₂O₂, M 270.4, m 260-261°, polymorphs have m 254° and 256°, [α]_D²⁰ +198 (c 1, dioxane), pK²⁵ 0.91. Purify estrone by chromatography on silica gel, eluting with 2:1 hexane/EtOAc and recrystallising from EtOH or Et₂O/EtOH. [Danishefsky & Cain *J Am Chem Soc* **98** 4975 1976, DOI: 10.1021/ja00432a044.] The *acetate* [901-93-9] crystallises from EtOH with m 125-127°. [*Beilstein* **8** III 1171.] Estrogenic.

Estrone 3-O-sulfamate [1,3,5(10)-estratrien-17-one 3-sulfamate] [148672-09-7] C₁₈H₂₃NO₄S, M 349.5. Estrone 3-O-sulfamate is purified by silica gel flash chromatography to give a product with one spot on TLC. It is an *active site-directed inhibitor of estrone sulfatase*, i.e. a time-dependent enzyme inactivator. [Woo et al. *J Med Chem* **39** 1349 1996, DOI: 10.1021/jm950931z; Purohit et al. *Biochemistry* **34** 11508 1995, DOI: 10.1021/bi00036a025.]

17- α -Ethinylestradiol [57-63-6] C₂₀H₂₄O₂, M 296.4, m 141-146°, 145-146°, [α]_D²⁰ +4 (c 1, CHCl₃). 17- α -Ethinylestradiol forms a *hemihydrate* on recrystallising from MeOH/H₂O. It dehydrates on melting and remelts on further heating at m 182-184°. The UV has λ_{max} at 281nm (ϵ 2040) in EtOH. Its solubility is 17% in EtOH, 25% in Et₂O, 20% in Me₂CO, 25% in dioxane and 5% in CHCl₃. [Petit & Muller *Bull Soc Chim Fr* 121 1951.] The *diacetyl* derivative has m 143-144° (from MeOH) and [α]_D²⁰ +1 (c 1, CHCl₃) [Mills et al. *J Am Chem Soc* **80** 6118 1958, DOI: 10.1021/ja01555a055]. [*Beilstein* **6** IV 6877.] Estrogenic.

Etiocholanolic acid (5 β ,17 β -androstane-17-carboxylic acid) [438-08-4] C₂₀H₃₂O₂, M 304.5, m 228-229°, pK_{Est} ~4.7. Crystallise etiocholanolic acid from glacial acetic acid (needles) and sublime it at 160°/0.002mm. The *ethyl ester* has m 77.5-78.5° (from EtOH) [α]_D²⁰ +52.5 (c 0.61, pyridine). [Jacobs & Elderfield *J Biol Chem* **108** 497 1935, <http://www.jbc.org/content/108/2/497>; Weiland et al. *Hoppe Seyler's Z Physiol Chem* **161** 80 1926, *Beilstein* **9** III 2644.]

Finasteride [(5 α ,17 β)-N-(1,1-dimethylethyl)-3-oxo-4-azaandrost-1-ene-17-carboxamide] [98319-26-7] C₂₃H₃₆N₂O₂, M 372.5, m 252-254°, ~257°, [α]_D²⁵ -59 (c 1, MeOH). Several 17-carbamoyl-4-azaandrost-1-en-3-ones have been synthesised from androsthenones essentially by cleaving the Steroid A ring, replacing C-4 by a nitrogen atom and altering the C-17 substituent while keeping most of the rest of the steroid intact. The study was devised to find an active *testosterone 5 α -reductase inhibitor* as is the case with *Finasteride*. The purification of the crude products was performed by chromatography on silica gel (e.g. one or two PrePaks, 325g each) in CH₂Cl₂ and eluting the azasteroidcarboxamide with a solvent mixture, e.g. CH₂Cl₂-Me₂CO (or cyclohexane-EtOAc 3:1) and finally EtOAc, checking the fractions by analytical HPLC using a μ -Porasil column or TLC run on silica gel coated plates (e.g. 250 μ m analytical, 1000 or 2000 μ m thicknesses) and visualised in the usual way (e.g. spraying with H₂SO₄ and heating in a hot oven until spots show colour). The desired fractions are collected, evaporated and the solid recrystallised from hexane, Et₂O, EtOAc or mixtures of these or aqueous EtOH. *Finasteride* is soluble in most organic solvents but is a very weak base (both nitrogen atoms are linked in amide bonds) and it is poorly soluble in aqueous 0.1N NaOH and 0.1N HCl. [Rasmusson et al. 'Azasteroids: structure-activity relationships for *inhibition of 5 α -reductase and of androgen receptor binding*' *J Med Chem* **29** 2298 1986, DOI: 10.1021/jm00161a028; Rasmusson & Reynolds US Patent 4760071 A to Merck & Co, Juky 26 1988; Bhattacharya et al. *J Am Chem Soc*, **110** 3318 1988, DOI: 10.1021/ja00218a062.]

Finasteride is a *competitive inhibitor* of steroid 5 α -reductase, i.e. it suppresses the conversion of testosterone to dihydrotestosterone, and reduces prostatic dihydrotestosterone levels and prostate size *in vivo*. It is 100-fold more potent as a competitive inhibitor of the rat NADPH: Δ 4-3-oxosteroid-5 α -oxidoreductase (steroid 5 α -reductase) type 1 enzyme (K_i = 3-5 nM), and of the human type 1 enzyme (K_i greater than or equal to 300 nM). It is orally active and has been used to restore baldness. [Thigpen & Russell 'Four-amino acid segment in steroid 5 α -reductase 1 confers sensitivity to *Finasteride*, a competitive inhibitor' *J Biol Chem* **267** 8577 1992, PMID: 1314830.]

Gitoxigenin (3 β ,14,16 β ,21-tetrahydroxy-20(22)-norcholenic acid lactone) [545-26-6] C₂₃H₃₄O₅, M 390.5, m 223-226°, 234°, 239-240° (anhydrous by drying at 60°), [α]_D²⁰ +30 (c 1, MeOH). Recrystallisation of gitoxigenin from aqueous EtOH produces plates of the *sesquihydrate* which dehydrate on drying at 100° *in vacuo*. It also crystallises from Me₂CO/MeOH and from EtOAc (the crystals contain 1 mol of EtOAc) with [α]_D²¹ +24.8 (c 1, dioxane). It has UV with λ_{max} at 310, 485 and 520nm in 96% H₂SO₄. On heating with ethanolic HCl it yields *digitaligenin* with loss of H₂O. [Smith *J Chem Soc* 23 1931, DOI: 10.1039/JR9310000023; Beilstein 8 IV 2456.]

Glycocholic acid (N-cholyglycine) [475-31-0] C₂₆H₄₃NO₆, M 465.6, m 130° (hydrate) 154-155°, 165-168°(anhydrous), [α]_D²⁰ +37 (c 1, EtOH), pK_a²⁵ 4.4. Glycocholic acid crystallises from hot water as the *sesquihydrate*. Dry it at 110° *in vacuo*. An analytical sample is prepared by suspending the acid (4g) in H₂O (400ml) at ~20°, heating to boiling with slow stirring, filtering hot and allowing to cool to ~20°. The acid is filtered off, washed with H₂O, dried in air, recrystallised from 5% aqueous EtOH, washed well and dried over P₂O₅ in a moderate vacuum to constant weight. Recrystallisation from EtOH/EtOAc, and drying, gave the *anhydrous* acid. [Cortese & Bauman *J Am Chem Soc* 57 1393 1935, DOI: 10.1021/ja01311a002; Bergstrom & Norman *Acta Chem Scand* 7 1126 1953, DOI: 10.3891/acta.chem.scand.07-1126; Beilstein 10 IV 2077.] It emulsifies oils and fats.

Glycodeoxycholic acid monohydrate [N-(3 α -12 α -dihydroxy-5 β -cholan-24-oyl)glycine] [360-65-6] C₂₆H₄₃NO₅, M 467.6, m 186-177°(dec), 187-188°, [α]_D²³ +45.9 (c 1, EtOH), pK_{Est} ~ 4.4. Glycodeoxycholic acid recrystallises from H₂O or aqueous EtOH with 1 mol of H₂O and is dried at 100° *in vacuo*. Its solubility in EtOH is ~5%. [UV: Lindstedt & Sjövall *Acta Chem Scand* 11 421 1957, DOI: 10.3891/acta.chem.scand.11-0421.] The *Na salt* recrystallises from EtOH/Et₂O with m 245-250° and [α]_D²³ +41.2 (c 1, H₂O) [Wiand Hoppe Seyler's *Z Physiol Chem* 106 181 1919, Cortese *J Am Chem Soc* 59 2532 1937, DOI: 10.1021/ja01291a014]. [Beilstein 10 IV 1611.] It is an emulsifying bile acid.

Hecogenin (25R-5 α -spirostan-3 β -ol-12-one) [467-55-0] C₂₇H₄₂O₄, M 430.6, m 245-250°, 253°, 264-266°, 268°, [α]_D²³ +8 (c 1, CHCl₃). The steroidal saponin (~35 mg) in EtOAc is chromatographed on Al₂O₃ and eluted with *C₆H₆/Et₂O, and the residue on evaporation is recrystallised from Me₂CO. [Mazur et al. *J Am Chem Soc* 82 5889 1960, DOI: 10.1021/ja01507a028; Marker et al. *J Am Chem Soc* 69 2167 1947, DOI: 10.1021/ja01201a032; Beilstein 19 III/IV 2581.]

Hecogenin acetate [915-35-5] C₂₉H₄₄O₅, M 472.7, m 265-268°(dec), [α]_D²⁰ -6.0 (c 1, CHCl₃). Crystallise the *acetate* from MeOH. [Beilstein 19 IV 2583.]

Hydrocortisone (Cortisol, 11 β ,17 α ,21-trihydroxypregn-4-ene-3,20-dione) [50-23-7] C₂₁H₃₀O₅, M 362.5, m 212-213°, 214-217°, 218-221°, 220-222°, [α]_D²² +167 (c 1, EtOH). Recrystallise hydrocortisone from EtOH or isoPrOH. It is bitter tasting and has UV with λ_{max} at 242 nm (log ϵ 4.20). Its solubility at 25° is: H₂O (0.28%), EtOH (1.5%), MeOH (0.62%), Me₂CO (0.93%), CHCl₃ (0.16%), propylene glycol (1.3%) and Et₂O (0.35%). It gives an intense green colour with concentrated H₂SO₄. [Wendler et al. *J Am Chem Soc* 72 5793 1950, DOI: 10.1021/ja01168a551; Beilstein 8 IV 3422.] It is a glucocorticoid hormone released in response to stress and low blood glucose. Topically used to reduce swelling and rashes from sting bites and allergies.

Hydrocortisone acetate (21-acetoxy-11 β ,17 α -trihydroxypregn-4-ene-3,20-dione) [50-03-3] C₂₃H₃₂O₆, M 404.5, m 218-221.5°, 221-223°, 222-225°, [α]_D²⁵ +166 (c 0.4, dioxane), +150.7 (c 0.5, Me₂CO). The acetate recrystallises from Me₂CO/Et₂O or aqueous Me₂CO as *hygroscopic* monoclinic crystals. Its UV has λ_{max} at 242 nm (A_{1cm}^{1%} 390) in MeOH. Its solubility at 25° is: H₂O (0.001%), EtOH (0.45%), MeOH (0.04%), Me₂CO (1.1%), CHCl₃ (0.5%), Et₂O (0.15%), and it is very soluble in Me₂NCHO. [Wendler et al. *J Am Chem Soc* 74 3630 1952, DOI: 10.1021/ja01134a049; Antonucci et al. *J Org Chem* 18 70 1953, DOI: 10.1021/jo01129a013; Beilstein 8 IV 3424.]

19-Hydroxy-4-androsten-3,17-dione [510-64-5] C₁₉H₂₆O₃, M 302.4, m 167-169°, 168-170°, 169-170°, 172-173°, [α]_D²⁰ +190 (c 1, CHCl₃). Recrystallise 19-hydroxy-4-androsten-3,17-dione from Me₂CO/hexane or Et₂O/

hexane. It has UV with λ_{\max} at 242nm in EtOH or MeOH. The *19-acetoxy* derivative has $[\alpha]_{\text{D}}^{25} +185$ (CHCl₃) and λ_{\max} with 237.5nm in EtOH. [Ehrenstein & Dünnenberger *J Org Chem* **21** 774 1956, DOI: 10.1021/jo01113a015; *Beilstein* **8** IV 2162.]

25-Hydroxycholesterol (cholest-5-en-3 β ,25-diol) [2140-46-7] C₂₇H₄₆O₂, M 402.7, m 177-179°, 178-180°, 181.5-182.5°, $[\alpha]_{\text{D}}^{25} -39$ (c 1.05, CHCl₃). 25-Hydroxycholesterol forms colourless needles from MeOH [Riediker & Schwartz *Tetrahedron Lett* **22** 4655 1981, DOI: 10.1016/S0040-4039(01)83005-7]. The *3 β -acetoxy derivative* has m 142-142.8° (from Me₂CO) and $[\alpha]_{\text{D}}^{25} -40.4$ (c 2, CHCl₃). The *3 β ,25-diacetoxy derivative* has m 119-120.5° (from MeOH) and $[\alpha]_{\text{D}}^{25} -35.5$ (CHCl₃). [Dauben & Bradlow *J Am Chem Soc* **72** 4248 1950, DOI: 10.1021/ja01165a117; Ryer et al. *J Am Chem Soc* **72** 4247 1950, DOI: 10.1021/ja01165a116; *Beilstein* **6** IV 6437.] It suppresses interleukin-1-driven inflammation downstream of type 1 interferon [Reboldi et al. *Science* **345** 679 2014, DOI: 10.1126/science.1254790].

18-Hydroxy-11-deoxycorticosterone (18,21-dihydroxypregn-4-en-3,20-dione tautomeric with 18,20-epoxy-20,21-dihydroxypregn-4-en-3-one) [379-68-0] C₂₁H₃₀O₄, M 346.5, m 168-170°, 171-173°, 191-195°, 200-205°, $[\alpha]_{\text{D}}^{20} +151$ (c 1, CHCl₃). Recrystallise 18-hydroxy-11-deoxycorticosterone from Et₂O/Me₂CO to give crystals m 200-205°. When it is recrystallised from Me₂CO, it has m 191-195°. It has UV with λ_{\max} at 240nm. The *21-O-acetoxy-18-hydroxy derivative* has m 158-159° (from Et₂O/*C₆H₆), and the *21-O-acetoxy-18,20-epoxy derivative* has m 149-154° (from Et₂O). [Kahnt et al. *Helv Chim Acta* **38** 1237 1955, DOI: 10.1002/hlca.19550380519; Pappo *J Am Chem Soc* **81** 1010 1959, DOI: 10.1021/ja01513a068.]

17 β -Hydroxy-17 α -methyl-3-androsterone (Mestanolone, Ermalone) [521-11-9] C₂₀H₃₂O₂, M 304.5, m 192-193°. Dissolve mestanolone in Et₂O, wash it with N NaOH, H₂O, dry it (Na₂SO₄), evaporate and recrystallise it from EtOAc. The *semicarbazone* has m 235-236° (from EtOH). [Ruzicka et al. *Helv Chim Acta* **18** 1487 1935, DOI: 10.1002/hlca.193501801203.] It is an oral androgenic (anabolic) steroid.

17 α -Hydroxy-6 α -methylprogesterone (Medroxyprogesterone) [520-85-4] C₂₂H₃₂O₃, M 344.5, m 220°, $[\alpha]_{\text{D}}^{25} +75$ (CHCl₃). If it contains the *epi-isomer* (TLC), then dissolve it in CHCl₃, bubble dry HCl gas to epimerise it, evaporate and recrystallise it from chloroform. The UV has λ_{\max} at 241nm (ϵ 16,150) in EtOH. The *17-acetate* [71-58-9] crystallises from MeOH with m 207-208° and $[\alpha]_{\text{D}}^{25} +61$ (CHCl₃). Its UV has λ_{\max} at 240nm (ϵ 15,950) in EtOH. [Babcock et al. *J Am Chem Soc* **80** 2904 1958, DOI: 10.1021/ja01544a079; *Beilstein* **8** IV 2212.] It is a steroidal progestin drug.

α -Hyodeoxycholic acid [83-49-8] C₂₄H₄₀O₄, M 392.6, m 196-197°, 200-201°, $[\alpha]_{\text{D}}^{20} +8$ (c 2, EtOH), pK_{Est} ~4.9. Crystallise α -hyodeoxycholic acid from EtOAc or Me₂CO. The K salt separates in needles from an alcoholic solution of the acid when an equivalent of KOMe is added (see lithocholic acid [434-13-9]). [Weiland & Gumlish *Hoppe Seyler's Z Physiol Chem* **215** 18 1933, Windaus & Bohne *Justus Liebigs Ann Chem* **433** 278 1923, DOI: 10.1002/jlac.19234330126; *Beilstein* **10** III 1631.]

Lanosterol [79-63-0] C₃₀H₅₀O, M 426.7, m 138-140°, $[\alpha]_{\text{D}}^{20} +62.0$ (c 1, CHCl₃). If very impure, then it should be acetylated, converted to the *dibromide acetate* [crystallised from EtOAc with slow cooling, m 168-170°, $[\alpha]_{\text{D}}^{20} +214^\circ$ (CHCl₃)], de-brominated with Zn dust to give the acetate (below) which is recrystallised from 3-4 parts of Me₂CO/MeOH (4:1) and hydrolysed as for stigmasterol (below). Recrystallise it from anhydrous MeOH. Dry it *in vacuo* over P₂O₅ for 3 hours at 90°. The purity is checked by proton magnetic resonance. The *acetate* crystallises from MeOH with m 131-133°, $[\alpha]_{\text{D}}^{25} +62$ (c 1, CHCl₃). [Block & Urech *Biochemical Preparations* **6** 32 1958, Van Tamelen et al. *J Am Chem Soc* **104** 6479, 6480 1982, DOI: 10.1021/ja00387a069; *Beilstein* **6** III 2880, **6** IV 4188.] It is the precursor of animal and fungus steroids.

Lithocholic acid (3 α -hydroxy-5 β -cholan-24-oic acid) [434-13-9] C₂₄H₄₀O₃, M 376.6, m 183-188°, 184-186°, $[\alpha]_{\text{D}}^{23} +35$ (c 1, EtOH), pK_{Est} ~4.8. Lithocholic acid can be purified by conversion to the rather insoluble Na or K salt by addition of the equivalent amount of aqueous NaOH or KOH, filtering off the alkali salt, washing it with ice cold H₂O, dissolving it in the least volume of boiling H₂O, acidifying with dilute HCl (slight excess), filtering off the acid, washing with cold H₂O and drying it thoroughly in a vacuum. Recrystallise it

from Me₂CO, EtOH or acetic acid. The **methyl ester** crystallises from MeOH, with 0.5 mol of MeOH, and has **m 92-93°**, $[\alpha]_{\text{D}}^{25} +34$ (MeOH). It has also been purified by recrystallisation from petroleum ether (b 40-60°) and, after chromatography on Al₂O₃ in petroleum ether, gave a **labile form m 92-93°** which is transformed to the **stable form m 125-126°** after standing for 2 days in a vacuum desiccator. [Hoehn & Mason *J Am Chem Soc* **62** 569 1940, DOI: 10.1021/ja00387a069; Sarel & Yanuka *J Org Chem* **24** 2018 1959, DOI: 10.1021/jo01094a602; *Beilstein* **10** IV 785.] It has been used in a study to investigate the regulation of hepatic phospholipid and bile acid homeostasis through SMAD3 activation by TGFβ [Matsubara et al. *J Lipid Res* **53** 2698 2012, PMID: 23034213].

6-α-Methylprednisolone (Medrol, 11β,17-21-trihydroxy-6α-methylpregna-1,4-dien-3,20-dione) [83-43-2] C₂₂H₃₀O₅, **M 347.5**, **m 226-237°, 228-237°, 240-242°**, $[\alpha]_{\text{D}}^{24} +91$ (c 0.5, dioxane). Recrystallise medrol from EtOAc. The UV has λ_{max} at in 95% EtOH 243nm (ε 14,875). The **21-acetoxy derivative** has **m 205-208°** (from EtOAc), and $[\alpha]_{\text{D}}^{24} +95$ (c 1, CHCl₃). [Spero et al. *J Am Chem Soc* **78** 6213 1956, DOI: 10.1021/ja01604a078; Fried et al. *J Am Chem Soc* **81** 1235 1959, DOI: 10.1021/ja01514a055; for ¹H NMR see Slomp & McGarvey *J Am Chem Soc* **81** 2200 1959, DOI: 10.1021/ja01518a047; *Beilstein* **8** IV 3498.] Useful for the treatment of patients with cardiac arrest [Mentzelopoulos et al. *Arch Intern Med* **169** 15 2009, DOI: 10.1001/archinternmed.2008.509].

17α-Methyltestosterone (Android, Mesterone) [58-18-4] C₂₀H₃₀O₂, **M 302.5**, **m 163.5°, 164-165°, 167°, 162-168°**, $[\alpha]_{\text{D}}^{20} +87$ (c 1, dioxane), $[\alpha]_{\text{D}}^{20} +82.3$ (c 1, EtOH) $A_{1\text{cm}}^{1\%}$ **241nm is 495-530 (EtOH)**. This **anabolic steroid** is crystallised from hexane or hexane/*benzene. It has $E_{1\text{cm}}^{1\%}$ 495-530 at 241nm (EtOH). The colour reaction with 2,4-dinitrophenyl hydrazine is used for assaying it. [Gornall & Macdonald *J Biol Chem* **201** 279 1953, PMID: 13044797.] In another **colour reaction** the sterone (1mg) in acetic acid (0.2ml) + 88% H₃PO₄ (2ml) is allowed to stand for 1 hour when it becomes fluorescent. After 1 hour it is diluted with acetic acid (~3ml) and provides a strong yellow fluorescence with the intensity of 50-100 times that of estrone. [Stuart & Stuckey *J Pharm Pharmacol* **1** 130 1949, DOI: 10.1111/j.2042-7158.1949.tb12388.x; Oppenauer *Recl Trav Chim Pays-Bas* **56** 137 1937, DOI: 10.1002/recl.19370560206; *Beilstein* **8** IV 1010.]

Norcholanic acid (5β-24-norcholan-23-oic acid) [511-18-2] C₂₃H₃₈O₂, **M 346.5**, **m 175.5-176.5°, 177°, 186°**, $[\alpha]_{\text{D}}^{20} +32$ (EtOH), **pK_{Est} ~4.8**. Recrystallise the acid from AcOH. The **methyl ester** has **m 74°** (needles from MeOH), and the **ethyl ester** has **m 65-66°**. [For **synthesis** from 12-ketonorcholanic acid, via the semicarbazone then reduction with NaOEt see Cook & Haselwood *J Chem Soc* 428 1934, DOI: 10.1039/JR9340000428; Yanuka et al. *Tetrahedron Lett* 1725 1968, DOI: 10.1016/S0040-4039(01)99037-9; *Beilstein* **9** III 2652, **10** IV 2083.]

α-Oestradiol See estradiol above [57-91-0], and **β-Oestradiol-3-benzoate** see estradiol 3-benzoate above [50-50-0].

Ouabain [3-[(6-deoxy-α-L-mannopyranosyl)oxy]-1,5,11a,14,19-pentahydroxycard-20(22)-enolide. **G-Strophanthin, Acocantherine**] [630-60-4, 312619-45-7 (hydrate), 11018-89-6 (octahydrate)] C₂₉H₄₄O₁₂, **M 728.8** (8 H₂O), **m 190°(dec), 200-202°(dec)**, $[\alpha]_{\text{D}}^{20} -30$ (c 1, H₂O). It crystallises from water as the **octahydrate**. Dry it at 130°. It decomposes at 190° when dry. Store it in the dark as it is light sensitive, but it is stable in air. Its solubility (g/100ml) in H₂O is 1.3 (~25°), 20 (~100°), and in EtOH it is 1.0 (~25°) and 12.5 (~78°). It is highly **TOXIC** cardiac glycoside as it is a **strong inhibitor of cation transport** and of Na⁺ and K⁺ **ATPase**. At lower doses it is used to treat hypotension and some arrhythmia. [Fürstenwerth 'Ouabain - The Insulin of the Heart'. *Int J Clin Pract* **64** 1591 2010, DOI: 10.1111/j.1742-1241.2010.02395.x, PMID: 20946265.] [*Beilstein* **18/5** V 625.]

Pancuronium bromide (2β,16β-dipiperidino-5α-androstan-3α,17β-diol diacetate dimetho-bromide)

[15500-66-0] $C_{35}H_{60}N_2O_4 \cdot Br_2$, **M 732.7**, **m 212-215°, 215°**. The bromide forms odourless crystals with a bitter taste which are purified through acid-washed Al_2O_3 and eluted with isoPrOH/EtOAc (3:1) to remove impurities (e.g. the monomethobromide) and eluted with isoPrOH to give the pure dibromide which is recrystallised from CH_2Cl_2/Me_2CO or isoPrOH/ Me_2CO . It is soluble in H_2O (10%) and $CHCl_3$ (3.3%) at 20°. It is a non-depolarising muscle relaxant, and used in *euthenasia*. [Buckett et al. *J Med Chem* **16** 1116 1973, DOI: 10.1021/jm00268a011.]

Prednisolone acetate (21-acetoxypregna-1,4-diene-11 β -17 α -diol-3,20-dione) [52-21-1] $C_{23}H_{30}O_6$, **M 402.5**, **m 237-239°, 240-242°, 240-243°, 244°**, $[\alpha]_D^{20}$ **+116** (c 1, dioxane). Recrystallise prednisolone acetate from EtOH, Me_2CO , Me_2CO /hexane, and it has UV with λ_{max} at 243nm in EtOH. [Joly et al. *Bull Soc Chim Fr* 366 1958; Herzog et al. *J Am Chem Soc* **77** 4781 1955, DOI: 10.1021/ja01623a027; *Beilstein* **8** IV 3468.]

Prednisone (17,21-dihydroxypregna-1,4-diene-3,11,20-trione, deltacortisone, dehydrocortisone) [53-03-2] $C_{21}H_{26}O_5$, **M 358.5**, **m 238°** (dec), $[\alpha]_D^{20}$ **+172** (c 0.5, dioxane), λ_{max} **238nm** (log ϵ **4.18**) in MeOH. Crystallise prednisone from acetone/hexane, then recrystallise it from Me_2CO . The *monoacetate* crystallises from Me_2CO /hexane with **m 227-233°(dec)**, and $[\alpha]_D^{25}$ **+186** (c 1, dioxane); and the *diacetate* crystallises from Me_2CO /hexane with **m 219-221°(dec)**, and $[\alpha]_D^{25}$ **+125** (c 1, $CHCl_3$). [Herzog et al. *Tetrahedron* **18** 581 1962, DOI: 10.1016/S0040-4020(01)92709-7; *Beilstein* **8** IV 3531.]

5 α -Pregnane (allopregnane) [641-85-0] $C_{21}H_{36}$, **M 288.5**, **m 84.5-85°**, $[\alpha]_D^{20}$ **+21.7** (c 1.3, $CHCl_3$). Recrystallise 5 α -pregnane several times from Me_2CO . The melting point is lowered (e.g. to 50-71°) if it is contaminated with the **5 β -isomer** (see below). [Butenandt et al. *Chem Ber* **64** 2529 1931, DOI: 10.1002/cber.19310640933; Steiger & Reichstein *Helv Chim Acta* **21** 161 1938, DOI: 10.1002/hlca.19380210128; *Beilstein* **5** III 1120, 1121, 1125.]

5 β -Pregnane (17 β -ethyletiocholane) [481-26-5] $C_{21}H_{36}$, **M 288.5**, **m 82-83°, 83.5°**, $[\alpha]_D^{20}$ **+21.2** (c 0.75, $CHCl_3$). Crystallise 5 β -pregnane from MeOH or Me_2CO . The mixed melting point with allopregnane (above) is ~50-71°. [Butenandt et al. *Chem Ber* **64** 2529 1931, DOI: 10.1002/cber.19310640933; Steiger & Reichstein *Helv Chim Acta* **21** 161 1938i, DOI: 10.1002/hlca.19380210128.] 5 β -Pregnane is the parent compound of progesterones.

5 α -Pregnane-3 α ,20 β R-diol [566-57-5] $C_{21}H_{36}O_2$, **M 320.5**, **m 207-209°**, $[\alpha]_D^{26}$ **+12** (c 1.1, $CHCl_3$). Crystallise 5 α -pregnane-3 α ,20 α -diol from Me_2CO . Its *diacetate* [6170-22-5] **M 404.5**, has two melting points **134-140°** and **152.4°**, and has $[\alpha]_D^{26}$ **+40.5** (c 0.8, $CHCl_3$). It is a *progesterone metabolite* in urine during pregnancy. [See references below.]

5 α -Pregnane-3 α ,20 α S-diol [566-58-5] $C_{21}H_{36}O_2$, **M 320.5**, **m 248°**, $[\alpha]_D^{20}$ **+17** (c 0.15, EtOH). Crystallise 5 α -pregnane-3 α ,20 α -diol from EtOH. Its *diacetate* [6003-18-5] has **m 142°** and $[\alpha]_D^{20}$ **+18** (c 0.4, $*C_6H_6$). It is a *progesterone metabolite* in urine during pregnancy. [See references below.]

5 β -Pregnane-3 α ,20 β R-diol [80-91-1] $C_{21}H_{36}O_2$, **M 320.5**, **m 239°**, $[\alpha]_D^{20}$ **+10** (c 1, EtOH). Crystallise pregnane-diol (which is abundant in the urine of pregnant women) from EtOH or Me_2CO and dry it *in vacuo*. It can be oxidised to progesterone (see below) and it is not precipitated by digitonin. Its *diacetate* [6100-28-3] has **m 112-113°** and $[\alpha]_D^{26}$ **+60** (c 1, $CHCl_3$). It is a *progesterone metabolite* in urine during pregnancy. [Marrian *Biochem J* **23** 1090 1929, DOI: 10.1042/bj0231090; Johnson et al. *J Chem Soc* 1302 1954, DOI: 10.1039/JR9540001302; Mattox & Vrieze *J Org Chem* **32** 708 1967 DOI: 10.1021/jo01278a045; *Beilstein* **6** III 4778, **6** IV 6111.]

5 β -Pregnane-3 α ,20 α S-diol [80-92-2] $C_{21}H_{36}O_2$, **M 320.5**, **m 243-244°**, $[\alpha]_{546}^{20}$ **+31** (c 1, EtOH). Crystallise the diol from acetone. The *diacetate* [1174-69-2] crystallises from petroleum ether with **m 180°** (also **182-183°**) and $[\alpha]_D^{20}$ **+35** (c 1.1, $CHCl_3$). [Marrian *Biochem J* **23** 1090 1929, DOI: 10.1042/bj0231090; Fish et al. *J Biol Chem* **143** 715 1942, <http://www.jbc.org/content/143/3/715>; Hirschmann *J Biol Chem* **140** 797 1941, <http://www.jbc.org/content/140/3/797>; Johnson et al. *J Chem Soc* 1302 1954, DOI: 10.1039/JR9540001302; Glick & Hirschmann *J Org Chem* **27** 3212 1962, DOI: 10.1021/jo01056a054; *Beilstein* **6** III 4778, **6** IV 6111.]

Progesterone [57-83-0] $C_{21}H_{30}O_2$, M 314.5, m 126°, 128.5°, $[\alpha]_{D}^{20}$ +230 (c 1, EtOH), +214.7 (c 1.3, Me₂CO). The α -form crystallises from EtOH with m 127-131°. The β -form crystallises from petroleum ether or aqueous petroleum ether/aqueous Et₂O with m 119-120° or 121°. It also crystallises from Et₂O, Me₂CO/EtOAc, MeOH, aqueous Et₂O, aqueous MeOH, wet petroleum ether, Et₂O/petroleum ether, petroleum ether/*C₆H₆, Et₂O/pentane and isopropyl ether. The UV has λ_{max} at 240nm with log ϵ 4.25 (EtOH). [Wintersteiner & Allen *J Biol Chem* **107** 321 1934, <http://www.jbc.org/content/107/1/321>; *Beilstein* **7** III 3648, **7** IV 2395.] It is produced in the *corpus luteum* and by the placenta as *antagonist* of estrogens.

Rubijervine (slanid-5-ene-3 β -12 α -diol) [79-58-3] $C_{27}H_{43}NO_2$, M 413.6, m 242-244°, 240-246°, $[\alpha]_{D}^{20}$ +19 (EtOH), $[\alpha]_{D}^{27.5}$ +20 (c 1.04, MeOH), pK_{Est} ~7.0. Rubijervine (from *Liliaceae*) crystallises from 95% EtOH as colourless rods. It has solvent of crystallisation and is dried at 120°/2mm. It is precipitated by digitonin. The *hydrobromide* crystallises from MeOH/Me₂CO (needles) with m 265-270°(dec). The *diacetate* crystallises from MeOH with m 160-163°. The *3-benzoate* gives colourless prisms from *C₆H₆ with m 156-159° and $[\alpha]_{D}^{27.5}$ +22 (c 1.6, CHCl₃). [Pelletier & Locke *J Am Chem Soc* **79** 4531 1957, DOI: 10.1021/ja01573a077; Jacobs & Craig *J Biol Chem* **148** 41 1943, <http://www.jbc.org/content/148/1/41>; for stereochemistry from X-ray studies see Hohne et al. *Tetrahedron* **22** 673 1966, DOI: 10.1016/0040-4020(66)80036-4; *Beilstein* **21** III/IV 2310.]

β -Sitosterol (stigmast-5-ene-3 β -ol) [83-46-5] $C_{29}H_{50}O$, M 414.7, m 136.5-137.5°, 140°, $[\alpha]_{D}^{20}$ -42 (c 2, CHCl₃). Crystallise this phytosterol from EtOH, MeOH, Me₂CO, Me₂CO/EtOH or Me₂CO/MeOH. It has also been purified by zone melting. The *acetate* crystallises from MeOH or EtOH as plates with m 127-128° and $[\alpha]_{D}^{20}$ -41 (c 2, CHCl₃). The *benzoate* crystallises from EtOH as needles with m 146-147° and $[\alpha]_{D}^{20}$ -13.8 (c 2, CHCl₃). [Fujimoto & Jacobson *J Org Chem* **29** 3377, 3381 1964, DOI: 10.1021/jo01034a060; Shoppee *J Chem Soc* 1032 1948, DOI: 10.1039/JR9480001032; Shoppee *J Chem Soc* 1043 1948, DOI: 10.1039/JR9480001043; Heilbron et al. *J Chem Soc* 344, 347 1941, DOI: 10.1039/JR9410000344; *Beilstein* **6** III 2696.]

Smilagenin (25R-spirostan-3 β -ol, isosarsasapogenin) [126-18-1] $C_{27}H_{44}O_3$, M 416.6, m 185°, $[\alpha]_{D}^{25}$ -69, $[\alpha]_{D}^{25}$ -80 (c 0.3, CHCl₃). Chromatograph smilagenin on active Al₂O₃ and elute with *C₆H₆, then recrystallise it from Me₂CO, aqueous EtOH (m 187-188°) or MeOH. The *acetate* crystallises from MeOH with m 152° and $[\alpha]_{D}^{25}$ -59.6, $[\alpha]_{D}^{25}$ -68.9 (c 0.25, CHCl₃). [Askew *J Chem Soc* 1399 1936, DOI: 10.1039/JR9360001399 and 1402 1936, DOI: 10.1039/JR9480001043; Scheer et al. *J Am Chem Soc* **77** 641 1955, DOI: 10.1021/ja01608a033; *Beilstein* **19** III/IV 826.]

Solanidine (solanid-5-en-3 β -ol) [80-78-4] $C_{27}H_{43}NO$, M 397.6, m 218-219°(sublimes), $[\alpha]_{D}^{20}$ -29 (c 0.5, CHCl₃), pK¹⁵ 6.66. Solanidine crystallises from CHCl₃/MeOH, aqueous EtOH or aqueous MeOH as needles. TLC on Al₂O₃ plates using CH₂Cl₂/MeOH (98:2) gives a spot at R_F 0.47. The *hydrochloride* crystallises from aqueous EtOH with m 345°(dec). The *acetate* crystallises from EtOH with m 208°. [Schreiber & Rönsch *Tetrahedron* **21** 645 1965, DOI: 10.1016/S0040-4020(01)82235-3; Kessar et al. *Tetrahedron* **27** 2153 1971, DOI: 10.1016/S0040-4020(01)91614-X; Reichstein & Reich *Ann Rev Biochem* **15** 155 1946, DOI: 10.1146/annurev.bi.15.070146.001103; *Beilstein* **21** III/IV 1398, **27** III/IV 2000.] This is a poisonous steroidal alkaloid from *Solanaceaea* family.

α -Solanine (solan-5-en-3 β -yl-[O³- β -D-glucopyranosyl-O²- α -L-rhamnopyranosyl- β -D-galactopyrano-side]) See 'Carbohydrates' in this Chapter.

Solasodine (22 α ,25R-spirosol-5-en-3 β -ol) [126-17-0] $C_{27}H_{43}NO_2$, M 413.6, m 202°, $[\alpha]_{D}^{25}$ -100 (c 2, MeOH), pK²⁵ 7.7. Solasodine crystallises (as *monohydrate*) from MeOH as lustrous plates (on heating, the plates change to needles as they melt and resolidify in needles), or aqueous 80% EtOH, and sublimes at high vacuum. After recrystallisation from H₂O, m 198-199°, then from Me₂CO/H₂O, m 199-201°, it has $[\alpha]_{D}^{22}$ -109.3 (c 0.581, CHCl₃). On TLC with silica gel G (*C₆H₆/absolute EtOH, 8:2) it has R_F 0.45. The IR (KBr) has ν_{max} at 10.3, 10.4, 11.2, 11.5 μ (azaooxaspirane bands). [Schreiber et al. *Tetrahedron* **20** 1939 1964, DOI: 10.1016/S0040-4020(01)98462-5; Uhle *J Org Chem* **27** 656 1962, DOI: 10.1021/jo01049a525; Kessar et al.

Tetrahedron **27** 2153 1971, DOI: 10.1016/S0040-4020(01)91614-X; *Beilstein* **27** III/IV 2000.] This is a poisonous steroidal alkaloid from *Solanaceae* plants and is teratogenic.

Stigmasterol (3 β -hydroxy-24-ethylcholesta-5,22-diene) [83-48-7] C₂₉H₄₈O, M 412.7, m 160-164°, 170°, [α]_D²² -51 (CHCl₃), [α]_D²⁰₅₄₆ -59 (c 2, CHCl₃). Stigmasterol is best purified *via* the **tetrabromide-acetate**. The impure sterol (3g) is acetylated with Ac₂O (60ml) by refluxing for 1.5 hour. The mixture is cooled at 20° for 1 hour, and the crude acetate is collected. The acetate (3g) in Et₂O (30ml) is then treated with Br₂/AcOH (38ml, from 5g Br₂ in 100ml AcOH), and after cooling at 6° overnight, the tetrabromoacetate is filtered off and washed with Et₂O. After six recrystallisations from CHCl₃/MeOH the **tetrabromoacetate** has m 194-196°. This product (1g) in AcOH (12ml) and Zn dust (1g) is refluxed for 1.5 hours, filtered hot, diluted with H₂O (30m L) and extracted with Et₂O. The extract is washed with dilute aqueous sodium sulfite, then H₂O, the extract is dried (Na₂SO₄) and the stigmasterol acetate (~550mg) is recrystallised (4x) from EtOH and twice from MeOH/CHCl₃ (2:1) to give the **acetate** with m 139-148°. This acetate (400mg) is hydrolysed in boiling 10% alcoholic KOH (1ml) for 1 hour. Then H₂O (30ml) is added and the mixture is extracted with Et₂O. The extract is washed with aqueous Na₂CO₃, then H₂O, the solvent is distilled off and the residue is recrystallised (3x) from 95% EtOH to give ~110mg of pure stigmasterol. It is dried in a vacuum over P₂O₅ for 3 hours at 90°. The purity is checked by NMR. The **acetate** crystallises from MeOH with m 145°, [α]_D²⁵ -56 (c 2, CHCl₃). [Byerrum & Ball *Biochemical Preparations* **7** 86 1959, Thornton et al. *J Am Chem Soc* **62** 2006 1940, DOI: 10.1021/ja01865a028; Colin et al. *Anal Chem* **51** 1661 1979, DOI: 10.1021/ac50047a018; *Beilstein* **6** IV 4170.] This is a common phytosterols in plant fats and like phytosterols is generally broken down in the bile.

Tachysterol [3- β , 6E,22,E)-9,19-secoergosta-5(10),6,8,22-teraene-3-ol] [115-61-7] C₂₈H₄₄O, M 396.7, oil, [α]_D¹⁸ -70 (c 0.12, Et₂O), [α]_D¹⁸ -86.3 (petroleum ether). Windaus first obtained tachysterol by irradiation of 7-dehydrocholesterol where he readily obtained adducts with citraconic anhydride, or maleic acid [Windaus et al. *Justus Liebigs Ann Chem* **499** 188 1932, DOI: 10.1002/jlac.19324990116; Windaus et al. *Justus Liebigs Ann Chem* **492** 226 1932, DOI: 10.1002/jlac.19324920111]. This sterol is among the products which lead to the antirachitic agent *ergocalciferol* (*calciferol* Vitamin D₂). Its name is derived from the great ease with which it forms these adducts (Gr. *Tachys*, swift). Steroids which possess the 5,6 and 7,8 conjugated double bonds in ring B undergo this photolytic reaction with UVB light which first causes epimerisation of the angular 10-methyl group followed by scission of the C9—C10 bond. These include *ergosterol*, *lumisterol* and *precalciferol*. The citraconic anhydride adduct of the latter is obtained in ~30% yield by irradiation of *precalciferol* in Et₂O with UVB light at ~0°/~7.5hrs under N₂, followed by treatment with the anhydride. Recrystallisation from AcOH gave **tachysterol-citraconic anhydride acetate** m 161-162°, [α]_D +75 (c 1, CHCl₃). This adduct gave **calciferol 3,5-dinitrobenzoate** with m 158-159°, [α]_D +57 (c 1, *C₆H₆) on treatment with 3,5-dinitrobenzoyl chloride/pyridine in *C₆H₆ [Velluz et al. US 2847426A (to Roussell UCLAF 1956)]. The sterol is an oil with UV λ_{\max} (10⁻³ε) at 264nm (18), 280nm (23), and 294nm (in C₄H₆), which is characterised by its adducts, the **4-methyl-3,5-dinitrobenzoate** m 155° (yellow crystals from AcOH) and by reduction to *dihydotachysterol* [67-96-9] above. It is a precursor of the active vitamin D₂ and the interconversions of provitamin D, lumisterol, previtamin D and tachysterol in the photolytic process have been revisited by Havinga et al. [*Tetrahedron* **11** 276 1960, DOI:10.1016/S0040-4020(01)93178-3]. Tachysterol is insoluble in H₂O and MeOH but soluble in most organic solvents and is **not** precipitated by *digonin*. It should be stored at low temperature in the dark under N₂ because it is very sensitive to oxidation by air, to heat and light transforms it. [For structure see Grundmann *Z Physiol* **252** 151 1938; and for stereochemistry see Inhoffen et al. *Chem Ber* **88** 1424 1955, DOI: 10.1002/cber.19550880913; Verloop et al. *Recl Trav Chim Pays-Bas* **76** 689 1957, DOI: 10.1002/recl.19570760903; Braude & Wheeler *J Chem Soc* 329 1955, DOI: 10.1039/JR9550000329; Dimroth *Chem Ber* **70** 1631/1937, DOI: 10.1002/cber.19370700731.]

Taurocholic acid (3 α , 7 α ,12 α -trihydroxy-5 β -cholestan-24-oic acid *N*-(2-sulfoethyl) amide, *N*-coloyotaurine) [81-24-3] C₂₆H₄₅NO₇S, M 515.6 (free acid), m ~125°(dec), [α]_D +38.8 (c 2, EtOH), pK²⁵ 1.4. The acid is present in bile and is isolated as an amorphous pale yellow powder. Crystallise it from EtOH/EtOAc/Et₂O (amorphous m 125°) or EtOH/Et₂O [Josephson *Biochem J* **29** 1484 1935, DOI: 10.1042/bj0291484]. The anhydrous acid is **hygroscopic**, freely soluble in H₂O and EtOH but insoluble in Et₂O and EtOAc. It is hydrolysed by acids or alkalis to cholic acid and taurine. The **sodium salt (hydrate)** [312693-83-7,

345909-26-4 (xH_2O)] crystallises from aqueous EtOH/Et₂O with **m 231-232°**, and $[\alpha]_D^{23} +23.6$ (c 2.5, H₂O); and it has UV λ_{\max} (H₂SO₄) at 303, 389 and 480nm. [cf Cortese *J Am Chem Soc* **59** 2532 1937, DOI: 10.1021/ja01291a014; *Beilstein* **10** III 2177, **10** IV 2078.] Its sodium salt is present in the bile of mammals.

Taurodeoxycholic acid sodium salt monohydrate (3a,12a-dihydroxy-5b-cholestan-24-oic acid N-(2-sulfoethyl) amide Na salt monohydrate, N-[desoxycholy]taurine Na salt H₂O) [1180-95-6, 207737-97-1 ($x H_2O$)] C₂₆H₄₄NO₆SNa, **M 539.7**(anhydrous), **m 168°(dec)**, **171-175°**, $[\alpha]_D^{23} +37$ (c 1, H₂O), **pK²⁵ 1.4 (free acid)**. The Na salt is dissolved in the smallest volume of H₂O, a saturated solution of aqueous NaCl/Et₂O is added and the mixture is stored at 0° for 24 hours. Then shake the mixture well, keep it in the cold for another day and filter the crystals with gentle shaking, wash them with ice-cold saturated aqueous NaCl saturated with Et₂O, dry them over CaCl₂ and extract them with absolute EtOH. Add ~10ml of H₂O to the solid followed by enough Et₂O to incipient cloudiness. Store it overnight at 0°. Add ice-cold Et₂O to make 250ml, collect the crystals, wash them with Et₂O then petroleum ether and dry them in air. The purification can be repeated with NaCl and Et₂O with ~85% recovery. **Note** that precipitation will not occur unless enough H₂O is present. Its solubility in H₂O is 10%. [Cortese *J Am Chem Soc* **59** 2532 1937, DOI: 10.1021/ja01291a014.] It forms mixed micelles with micellar average Mol Wt of 3100, aggregation number 6, and CMC (critical micellar concentration) of 1-4mM at 20-25°. It is a useful **anionic detergent** for solubilising membrane proteins [Hajjar et al. *J Biol Chem* **258** 192 1983, PMID: 6848493]. [*Beilstein* **10** IV 1611.] The **free bile acid** [(3 α ,5 β ,12 α ,20R,24Z)-3,12-dihydroxy-N-(2-sulfoethyl)cholan-24-imidic acid] [516-50-7] C₂₆H₄₅NO₆S, **M 499.7** has **m 204-208°** (**141-144°** also reported), pK_a -0.94, and a solubility in H₂O of 41mg/ml at 25° [for preparation of the **acid dihydrate** see Parenti, *Eur Pat Appl* (1990), 3 pages. CODEN: EPXXDW EP 400695 A219901205 CAN 114:82269 AN 1991:82269; and Bloch & Watkins for the 'Determination of conjugated bile acids in human bile and duodenal fluid by reverse-phase HPLC' *J Lipid Res* **19**(4) 510 1978, PMID: 659989]. The deoxycholytaurine rescues human colon cancer cells from apoptosis by signaling activated by EGFR-dependent PI3K/Akt. Bile acids are **steroidal physiological amphipathic detergents** that facilitate digestion, excretion, absorption, and transport of fats and sterols in the intestine and liver.

Testosterone (17- β -hydroxyandrost-4-ene-3-one) [58-22-0] C₁₉H₂₈O₂, **M 288.4**, **m 155°**, $[\alpha]_{546}^{20} +130$ (c 1, dioxane). Crystallise testosterone from aqueous acetone, hexane or isoPrOH. The long needles that separated from EtOH/AcOH were used for X-ray crystallography [Roberts et al. *JCS Perkin Trans 2* 1978 1973, DOI: 10.1039/P29730001978.] The **acetate** [1045-69-8] crystallises from MeOH or aqueous Me₂CO, with **m 140-141°** and $[\alpha]_D^{20} +87.8$ (c 1, EtOH). [Ruzicka et al. *Helv Chim Acta* **18** 1478 1935, DOI: 10.1002/hlca.193501801201; and **19** 99 1936 DOI: 10.1002/hlca.19360190115; 842 1936, DOI: 10.1002/hlca.19360190115; *Beilstein* **8** IV 974.] It is secreted in male testes and to a lesser extent in female ovaries of mammals.

Testosterone propionate [57-85-2] C₂₂H₃₂O₃, **M 344.5**, **m 120°**, **118-122°**, $[\alpha]_{546}^{20} +102$ (c 2, dioxane). Crystallise the propionate from aqueous EtOH, or Et₂O/petroleum ether (**m 121°**), and its UV has λ_{\max} at 240nm (EtOH), and $[\alpha]_{546}^{20} +114$ (c 1, CHCl₃). Also purify it by HPLC. Its solubility in H₂O very low at 1.48mg/L at 25°. [Ercoli & de Ruggieri *J Am Chem Soc* **75** 650, 652 1953, DOI: 10.1021/ja01099a040; polymorphism: Brandstätter-Kuhnert & Kofler *Microchim Acta* 847, 850 1959, DOI: 10.1007/BF01216751; *Beilstein* **8** IV 977.]

Ursodiol (ursodeoxycholic acid, 3 α ,7 β -dihydroxy-5 β -cholan-24-oic acid, 7 β -hydroxylithocholic acid, ursodeoxycholic acid) [128-13-2] C₂₄H₄₀O₄, **M 392.5**, **m 203°**, $[\alpha]_D^{20} +60$ (c 0.2, EtOH), **pK_{Est} ~4.8**. Recrystallise ursodiol from wet Et₂O, EtOH or EtOH/MeOH. It is almost insoluble in H₂O, sparingly soluble in Et₂O, very slightly soluble in CHCl₃ but freely soluble in AcOH. The **diformate** has **m 170°**, and the **diacetate** has **m 100-102°**. It is an **anticholelithogenic** drug. [Iwasaki Hoppe Seyler's *Z Physiol Chem* **244** 181, 183 1936, Ward et al. *Drugs* **27** 95 1984, *Beilstein* **10** III 1635, **10** IV 1604.] It is present in bear bile.

PHYSIOLOGICALLY ACTIVE COMPOUNDS (including miscellaneous low-molecular-weight bioactive substances, drugs, antibiotics, coenzymes, vitamins, lipids, phospholipids, nucleosides, nucleotides polynucleotides and useful reagents)

NOTE: Several physiologically active compounds are also present in previous chapters, and their bio-activities are briefly mentioned in their respective entries. They should be located from the CAS Registry Numbers Index.

Acetoacetyl coenzyme A trisodium salt trihydrate [102029-52-7, anhydrous free acid 1420-36-6] $C_{25}H_{37}N_7Na_3O_{18}P_3S \cdot 3H_2O$, M 971.6, 851.6(anhydrous free acid), pK_1 4.0 (NH_2), pK_2 6.4 (PO_4^-). Purification can be carried out by passage through a DEAE-cellulose formate column, then through a Dowex 50 (H^+) column to remove Na^+ ions, concentrated by lyophilisation and redissolved in H_2O . It is commercially available as a solution of 0.05g/ml of H_2O . The concentration of acetoacetylcoenzyme A is determined by the method of Stern et al. [*J Biol Chem* **221** 15 1956, PMID: 13345795]. It is stable at pH 7-7.5 for several hours at 0° (half-life *ca* 1-2 hours). At room temperature it is hydrolysed in *ca* 1-2 hours at pH 7-7.5. At pH 1.0/ 20° it is more stable than at neutrality. A solution of the trisodium salt (0.05g/ml H_2O) adjusted to pH 5 with 2N NaOH can be stored frozen for several weeks. It is stable at pH 2-3/- 17° for at least 6 months. Store at -20° . [Hersch & Jencks *J Biol Chem* **242** 3468 1967, <http://www.jbc.org/content/242/15/3468>; Clikenbeard et al. *J Biol Chem* **250** 3108 1975, PMID: 164460; Simon & Shemin *J Am Chem Soc* **75** 2520 1953, DOI: 10.1021/ja01106a522; Moffatt & Khorana *J Am Chem Soc* **81** 1265 1959, DOI: 10.1021/ja01514a073; Salam et al. *Biochem J* **258** 563 1989 DOI: 10.1042/bj2580563; *Beilstein* **26** III/IV 3668.]

Acetylcholine bromide [66-23-9, anion-free 51-84-3] $C_7H_{16}NO_2 \cdot Br$, M 226.1, m 140-143°, 143°, 146°. The bromide is a *hygroscopic* solid, but less so than the hydrochloride salt. It crystallises from EtOH as prisms or needles. Some hydrolysis occurs in boiling EtOH, particularly if it contains some H_2O . It can also be recrystallised from EtOH or MeOH by adding dry Et_2O . [Siegal & Sapru *Essential Neuroscience* Revised 1st ed, Philadelphia: Lippincott, Williams & Wilkins **Ch 15** pp 255–267. 2006; *Beilstein* **4** IV 1446.] It is a neuro-modulator and transmitter.

Acetylcholine chloride [60-31-1] $C_7H_{16}NO_2 \cdot Cl$, M 181.7, has m 148-150°, 151°. It is very soluble in H_2O (>10%), and is very *hygroscopic*. If pasty, dry it in a vacuum desiccator over H_2SO_4 until a solid residue is obtained. Dissolve this in absolute EtOH, filter it and add dry Et_2O , when the hydrochloride separates. Collect by filtration and store it under very dry conditions. [Jones & Major *J Am Chem Soc* **52** 307 1930, DOI: 10.1021/ja01364a043.] The *chloroplatinate* crystallises from hot H_2O in yellow needles and can be recrystallised from 50% EtOH, m 242-244° [Dudley *Biochem J* **23** 1064 1929, DOI: 10.1042/bj0231064]; other m given is 256-257°. The *perchlorate* crystallises from EtOH as prisms m 116-117°. [Bell & Carr *J Am Pharm Assoc* **36** 272 1947, DOI: 10.1002/jps.3030360907; for heat of hydrolysis see Annis & Eley *Biochem J* **53** 34 1953, DOI: 10.1042/bj0530034; *Beilstein* **4** IV 1446.] It is a neuro-modulator and transmitter [Yu & Dayan 'Uncertainty, neuromodulation, and attention' *Neuron* **46**(4) 681 2005, DOI: 10.1016/j.neuron.2005.04.026, PMID: 15944135].

***N*⁴-Acetylcytosine** [14631-20-0] $C_6H_7N_3O_2$, M 153.1, m >300°, 326-328°, $pK_{Est(1)}$ ~1.7, $pK_{Est(2)}$ ~10.0. If TLC or paper chromatography shows that it contains unacetylated cytosine, then reflux it in Ac_2O for 4 hours, cool at 3-4° for a few days, collect the crystals, wash them with cold H_2O , then EtOH and dry at 100° . It is insoluble in EtOH and dissolves in H_2O with difficulty, but crystallises in prisms from hot H_2O . It is hydrolysed by 80% aqueous AcOH at $100^\circ/1$ hour. [UV: Brown et al. *J Chem Soc* 2384 1956, DOI: 10.1039/JR9560002384; Codington et al. *J Am Chem Soc* **80** 5164 1958, DOI: 10.1021/ja01552a038.] It forms an *Hg salt* [Fox et al. *J Am Chem Soc* **79** 5060 1957, DOI: 10.1021/ja01575a063; for the synthesis of cytidine see Schwartz & Lerner *J Org Chem* **40** 24 1975, DOI: 10.1021/jo00889a005]. [*Beilstein* **25** III/IV 3657.]

***N*-Acetylhistamine** [*N*-(2-1{3}*H*-imidazol-4-yl)ethylacetamide] [673-49-4] $C_7H_{11}N_3O$, M 153.2, m 147-148°, 148-149°, pK^{25} 6.99. It is purified by recrystallisation from Et_2O /EtOH or $EtOAc$ /EtOH (needles), and dried *in vacuo*. It sublimes at $148^\circ/0.05$ mm. It is slightly soluble in H_2O and EtOH, soluble in Me_2CO and very soluble in Et_2O . The *nitrate* has m 170°, the *picrate* is *dimorphic* with m 169-171° (from EtOH/ Et_2O) and m 181-183° (from EtOH). [Tabor & Mosettig *J Biol Chem* **80** 703 1949, PMID: 18135805; Nagarajan et al.

Indian J Chem, Sect B **16** 629, 633 1977, *Beilstein* **25** II 304, **25** III/IV 2053, **25/9** V 523.]

O-Acetyl- β -methylcholine chloride [Methacholine chloride, Amechol, Provocholine, 2-acetoxypyrpyl-ammonium chloride] [62-51-1] $C_8H_{18}NO_2 \cdot Cl$, **M 195.7**, **m 171-173°**, **172°**, **172-173°**. It forms white *hygroscopic* needles from Et_2O and is soluble in H_2O , $EtOH$ and $CHCl_3$. It decomposes readily in alkaline solutions and slowly in H_2O . It should be handled and stored in a dry atmosphere at 2-8°. The *bromide* is less hygroscopic, and the *picrate* has **m 129.5-131°** (from $EtOH$). [For racemate, heat of hydrolysis see Annis & Eley *Biochem J* **53** 34 1953, DOI: 10.1042/bj0530034; for *acetyl β -methylthiocholine bromide* see Hansen *Acta Chem Scand* **13** 159 1959, DOI: 10.3891/acta.chem.scand.13-0159; *Beilstein* **4** IV 1670.] A cholinergic agent, used mainly to diagnose bronchial hyper-reactivity [see Birnbaum & Barreiro 'Methacholine challenge testing: identifying its diagnostic role, testing, coding, and reimbursement' *Chest* **131**(6) 1932 2007, DOI: 10.1378/chest.06-1385, PMID: 17565027]. It is an anti-hypertensive and anti-anginal agent.

N-Acetyl neuraminic acid (NANA, O-Sialic acid, 5-acetamido-3,5-dideoxy-D-glycero-D-galacto-2-nonulosonic acid, lactaminic acid) [131-48-6] $C_{11}H_{19}NO_9$, **M 309.3**, **m 159°(dec)**, **181-183°(dec)**, **186°(dec)**, **185-187°(dec)**, $[\alpha]_D^{25}$ -33 (c 2, H_2O), **pK²⁵ 2.6**. A Dowex-1x8 (200-400 mesh) in the formate form is used, and is prepared by washing with 0.1M NaOH, then 2N sodium formate; excess formate is removed by washing with H_2O . N-Acetyl neuraminic acid in H_2O is applied to this column, washed with H_2O , then eluted with 2N formic acid at a flow rate of 1ml/minute. Fractions (20ml) are collected and tested (Bial's orcinol reagent, cf. *Biochemical Preparations* **7** 1 1960). NANA elutes at a formic acid molarity of 0.38, and the Bial positive fractions are collected and lyophilised. The residue is recrystallised from aqueous AcOH: suspend 1.35g of residue in AcOH, heat rapidly to boiling, add H_2O dropwise until the suspension dissolves (do not add excess H_2O), filter hot and then keep at +5° for several hours until crystallisation is complete. Collect NANA and dry it in a vacuum over P_2O_5 . Alternatively, dissolve 1.35g of NANA in 14ml of H_2O , filter, add 160ml of MeOH followed by 360ml of Et_2O . Then add petroleum ether (b 40-60°) until heavy turbidity. Cool at 20° overnight. The yield of NANA is ca 1.3g. Dry it over P_2O_5 at 100°/1mm to constant weight. It *mutarotates* in Me_2SO : $[\alpha]_D^{20}$ -115° (after 7 minutes) to -32° (after 24 hours). It is commercially available as an aqueous solution (0.01g/ml). [For IR and synthesis see Cornforth et al. *Biochem J* **68** 57 1958, DOI: 10.1042/bj0680057; Zillikin & O'Brien *Biochemical Preparations* **7** 1 1960; for ^{13}C NMR and $1-^{13}C$ synthesis see Benzing-Nguyen & Perry *J Org Chem* **43** 551 1978, DOI: 10.1021/jo00398a005; Danishefsky & DeNinno *J Org Chem* **51** 2615 1986, DOI: 10.1021/jo00363a047; Gottschalk, *The Chemistry and Biology of Sialic Acids and Related Substances*, Cambridge University Press, London, ISBN 10: 0521051231 ISBN 13: 9780521051231; 1960, *Beilstein* **4** IV 3288.]

N-Acetyl penicillamine (N-acetyl-3-mercapto-D-valine) [D- 15537-71-0, DL -59-53-0] $C_7H_{13}NO_3S$, **M 191.3**, **m 183°**, **186-187° (DL-form)**, **189-190° (D-form)**, **D-form** $[\alpha]_D^{25}$ +18 (c 1, 50% $EtOH$), **pK_{Est(1)} ~3.0 (CO₂H)**, **pK_{Est(2)} ~8.0 (SH)**. Both forms are recrystallised from hot H_2O . A pure sample of the **D-form** is obtained after five recrystallisations. [Crooks in *The Chemistry of Penicillin* Clarke, Johnson and Robinson eds, Princeton University Press, 470 1949, ISBN: 9780691653471, ISBN: 9781400874910; Review: Chain et al. (*Antibiotics* Oxford University Press) **2** 1949, *Beilstein* **4** III 1662.] **S-Nitroso-N-acetylpenicillamine** [67776-06-1] $C_7H_{12}N_2O_4S$, **M 220.3**, is an S-nitrosothiol used as a model in nitric oxide studies of biological systems, particularly related to vasodilation [Zhang & Hogg 'S-Nitrosothiols: Cellular Formation and Transport' *Free Radical Biology and Medicine* **38** 831 2005, DOI: 10.1016/j.freeradbiomed.2004.12.016, PMID: 15749378].

p-Acetylphenyl sulfate potassium salt [38533-41-4] $C_8H_7O_5S \cdot K$, **M 254.3**, **m dec on heating**, **pK_{Est} ~2.1**. Purify the salt by dissolving it in the minimum volume of hot water (60°) and adding $EtOH$, with stirring, then leave at 0° for 1hour. The crystals are filtered off and recrystallised from H_2O until free of Cl^- and SO_4^{2-} ions. Dry it in a vacuum over P_2O_5 at room temperature. It is a *specific substrate for arylsulfatases* which hydrolyse it to p-acetylphenol [UV has λ_{max} at 327nm (ϵ 21700 $M^{-1}cm^{-1}$)] [Milsom et al. *Biochem J* **128** 331 1972, DOI: 10.1042/bj1280331].

S-Acetylthiocholine bromide [25025-59-6] $C_7H_{16}NOS \cdot Br$, **M 242.2**, **m 217-223°(dec)**. It is a *hygroscopic* solid which can be recrystallised from ligroin/ $EtOH$ (1:1), dried and kept in a vacuum desiccator. Crystallisation from $*C_6H_6/EtOH$ gives **m 227°** or from propan-1-ol the **m** is 213°. [Hansen *Acta Chem Scand* **11** 537 1957,

DOI: 10.3891/acta.chem.scand.11-0537; for hydrolysis studies see Heilbronn *Acta Chem Scand* **12** 1481 1958, DOI: 10.3891/acta.chem.scand.12-1481; Heilbronn *Acta Chem Scand* **12** 1492 1958, DOI: 10.3891/acta.chem.scand.12-1492; *Beilstein* **4** IV 1585.] It is a skin and eye irritant.

S-Acetylthiocholine chloride [6050-81-3] $C_7H_{16}NOS$. **Cl**, **M 197.7**, has **m 172-173°**. The chloride can be purified in the same way as the bromide, and it can be prepared from the iodide. A few milligrams dissolved in H_2O can be purified by applying onto a Dowex-1 Cl^- resin column (prepared by washing with $N HCl$ followed by CO_3^{2-} -free H_2O until the pH is 5.8). After equilibration for 10 minutes, elution is started with CO_3^{2-} -free distilled H_2O , and 3ml fractions are collected and their OD values at λ 229nm are measured. The fractions with appreciable absorption are pooled and lyophilised at 0-5°. **Note** that at higher temperatures decomposition of the ester is appreciable; hydrolysis is appreciable at pH >10.5/20°. The residue is dried *in vacuo* over P_2O_5 , checked for traces of iodine (add concentrated H_2SO_4 and heat, violet vapours are released), and recrystallised from propan-1-ol. [For hydrolysis studies see Heilbronn in previous entry; and for electrical conductivity studies see El-Hammamy et al. *Adv Appl Sci Res* **2**(1) 90-94 2011, pelagiaresearchlibrary.com/advances.../AASR-2011-2-1-90-94.pdf; Gal & Roth *Clin Chim Acta* **2** 316 1957, DOI: 10.1016/0009-8981(57)90009-8; *Beilstein* **4** IV 1585.]

S-Acetylthiocholine iodide [1866-15-5] $C_7H_{16}NOS$. **I**, **M 289.2**, has **m 203-204°, 204°, 204-205°**. Recrystallise the iodide from propan-1-ol (or *iso*-PrOH, or EtOH/Et₂O) until almost colourless and dry it in a vacuum desiccator over P_2O_5 . Its solubility in H_2O is 10% w/v at ~25°. A 0.075M (21.7mg/ml) solution in 0.1M phosphate buffer pH 8.0 is stable for 10-15 days if kept refrigerated. Store it away from light. It is commercially available as a 1% solution in H_2O . [Ellman et al. *Biochemical Pharmacology* **7**, 88 1961, DOI: 10.1016/0006-2952(61)90145-9; Hansen *Acta Chem Scand* **13** 151 1959, **11** 537 1957, DOI: 10.3891/acta.chem.scand.11-0537; Gal & Roth *Clin Chim Acta* **2** 316 1957, DOI: 10.1016/0009-8981(57)90009-8; for hydrolysis studies see Heilbronn in previous entry; *Beilstein* **4** III 726, **4** IV 1585.]

Actinomycin C (Cactinomycin) [8052-16-2] $C_{63}H_{88}N_{12}O_{16}$, **M 1268.6**. (A commercial mixture of Actinomycin C_1 ~5%, C_2 ~30% and C_3 ~65% is available). **Actinomycin C_1 (native)** crystallises from EtOAc as red crystals, is soluble in $CHCl_3$, $*C_6H_6$ and Me_2CO , and has **m 246-247°(dec)**, with $[\alpha]_D^{20}$ -328 (0.22, MeOH) and λ_{max} 443nm (ϵ 25,000) and 240nm (ϵ 34,000). **Actinomycin C_2 (native)** crystallises as red needles from EtOAc and has **m 244-246°(dec)**, with $[\alpha]_D^{20}$ -325 (c 0.2, MeOH), λ_{max} 443nm (ϵ 25,300) and 240nm (ϵ 33,400). **Actinomycin C_3 (native)** recrystallises from cyclohexane, or $*C_6H_6/MeOH/cyclohexane$ as red needles with **m 238-241° (dec)**, $[\alpha]_D^{20}$ -321 (c 0.2, MeOH), and λ_{max} 443nm (ϵ 25,000) and 240nm (ϵ 33,300). [Brockman & Lackner, *Chem Ber* **101** 1312 1968, DOI: 10.1002/cber.19681010425.] It is **light sensitive**. [*Beilstein* **27** III/IV 9642.] **TOXIC**.

Actinomycin D (Dactinomycin) [50-76-0] $C_{62}H_{86}N_{12}O_{16}$, **M 1255.4**, has **m 241-243°(dec), 251-253°(dec)**, $[\alpha]_D^{22}$ -296 (c 0.22, MeOH). It crystallises as bright red (red-orange) rhombic crystals from absolute EtOH or from MeOH/EtOH (1:3). It will also crystallise from EtOAc/cyclohexane (**m 246-247° dec**), $CHCl_3$ /petroleum ether (**m 245-246° dec**), and EtOAc/MeOH/ $*C_6H_6$ (**m 241-243° dec**). Its solubility in MeCN is 1mg/ml. $[\alpha]_D^{20}$ varies from -296° to -327° (c 0.2, MeOH). λ_{max} (MeOH) 445, 240nm (log ϵ 4.43, 4.49), λ_{max} (MeOH, 10N HCl, 1:1) 477nm (log ϵ 4.21) and λ_{max} (MeOH, 0.1N NaOH) 458, 344, 285 (log ϵ 3.05, 4.28, 4.13). It is **HIGHLY TOXIC**, light sensitive and anti-neoplastic. [Bullock & Johnson, *J Chem Soc* 3280 1957, DOI: 10.1039/JR9570003280; *Beilstein* **27** III/IV 9642.]

Adenosine-5'-diphosphate [adenosine-5'-pyrophosphate, ADP] [58-64-0] $C_{10}H_{15}N_5O_{10}P_2$, **M 427.2**, $[\alpha]_D^{25}$ -25.7 (c 2, H_2O), pK_1^{25} <2 (PO_4H), pK_2^{25} <2 (PO_4H), pK_3^{25} 3.95 (NH_2), pK_4^{25} 6.26 (PO_4H). It is characterised by conversion to the **acridine salt** by addition of alcoholic acridine (1.1g in 50ml), filtering off the yellow salt and recrystallising from H_2O . Store at -20°. The **salt** has **m 215°(dec)**, and UV with λ_{max} at 259nm (ϵ 15,400) in H_2O . [Baddiley & Todd *J Chem Soc* 648 1947, DOI: 10.1039/JR9470000648; 582 1949, DOI: 10.1039/JR9490000582; cf. LePage *Biochemical Preparations* **1** 1 1949, Martell & Schwarzenbach *Helv Chim Acta* **39** 653 1956, DOI: 10.1002/hlca.19560390302]. [*Beilstein* **26** III/IV 2369.]

Adenosine-3'-monophosphoric acid hydrate [3'-adenylic acid, 3'-AMP] [84-21-9] $C_{10}H_{16}N_5O_8P$. xH_2O , **M 347.3(anhydrous)**, **m 197°(dec, as $2H_2O$), 210°(dec)**, **m 210°(dec)**, $[\alpha]_{546}$ -50 (c 0.5, 0.5M Na_2HPO_4), pK_1^{25} 3.65, pK_2^{25} 6.05. It crystallises from large volumes of H_2O in needles as the **monohydrate**, but is not very soluble in boiling H_2O . Under acidic conditions it forms an equilibrium mixture of 2' and 3' adenylic acids *via*

the 2',3'-cyclic phosphate. When heated with 20% HCl, it gives a quantitative yield of furfural after 3 hours, unlike 5'-adenylic acid which only gives traces of furfural. The yellow **monoacridine salt** has **m 175°(dec)**, and the **diacridine salt** has **m 177° (225°)(dec)**. [Brown & Todd *J Chem Soc* 44 1952, DOI: 10.1039/JR9520000044; Ueda & Takaku *Chem Pharm Bull Jpn* 32 1650 1984, DOI: org/10.1248/cpb.32.1650; for phosphorylation of an OH group with *N*-phosphoryl-*N'*-methylimidazole see Takaku et al. *Chem Pharm Bull Jpn* 21 2068 1973, DOI: org/10.1248/cpb.21.444; NMR: Ts'o et al. *Biochemistry* 8 997 1969, DOI: 10.1021/bi00831a033; *Beilstein* 26 III/IV 3607.]

Adenosine-5'-monophosphoric acid monohydrate (5'-adenylic acid, 5'-AMP) [18422-05-4] C₁₀H₁₆N₅O₈P·H₂O, M 365.2, m 178°, 183-188°(dec), 196-200°, 200° (sintering at 181°), [α]_D²⁰ -47.5, [α]₅₄₆ -56 (c 2, in 2% NaOH), -26.0 (c 2, 10% HCl), -38 (c 1, 0.5M Na₂HPO₄), pK₁²⁵ 3.89, pK₂²⁵ 6.14, pK₃²⁵ 13.1. The acid has been recrystallised from H₂O (fine needles) and is freely soluble in boiling H₂O. It crystallises also from H₂O on addition of acetone. Alternatively, purify it by chromatography on Dowex 1 (in formate form), and eluting with 0.25M formic acid. It is then adsorbed onto charcoal (which had been boiled for 15 minutes with M HCl, washed free of chloride ions and dried at 100°) and recovered by stirring three times with isoamyl alcohol/H₂O (1:9 v/v). The aqueous layer from the combined extracts is evaporated to dryness under reduced pressure, and the product is crystallised twice from hot H₂O. [Morrison & Doherty *Biochem J* 79 433 1961, DOI: 10.1042/bj0790433d]. It has UV with λ_{max} at 259nm (ε 15,400) in H₂O at pH 7.0. [Alberty et al. *J Biol Chem* 193 425 1951, PMID: 14907730; Martell & Schwarzenbach *Helv Chim Acta* 39 653 1956, DOI: 10.1002/hlca.19560390302]. The **acridinium salt** has **m 208°** [Baddiley & Todd *J Chem Soc* 648 1947, DOI: 10.1039/JR9470000648; Pettit *Synthetic Nucleotides*, van Nostrand-Reinhold, NY, Vol 1 252 1972, NMR: Sarma et al. *J Am Chem Soc* 96 7337 1974, DOI: 10.1021/ja00830a028; Norton & Allerhand *J Am Chem Soc* 98 1007 1976, DOI: 10.1021/ja00420a025; for IR of **diNa salt** see Miles *Biochem Biophys Acta* 27 46 1958, DOI: 10.1016/0006-3002(58)90291-9]. [*Beilstein* 26 III/IV 3615.]

Adenosine 5'-[β-thio]diphosphate tri-lithium salt [73536-95-5] C₁₀H₁₂N₅O₉P₂S·3Li, M 461.1. Purify it by ion-exchange chromatography on DEAE-Sephadex A-25 using gradient elution with 0.1-0.5M triethylammonium bicarbonate. It has UV with λ_{max} at (H₂O) 259nm (ε 15,000). Store at -20°. [Goody & Echstein *J Am Chem Soc* 93 6252 1971, DOI: 10.1021/ja00752a042; Goody et al. *Biochem Biophys Acta* 276 155 1972, DOI: 10.1016/0005-2744(72)90016-2.]

Adenosine 5'-[α-thio]monophosphate di-lithium salt [19341-57-2, 93839-85-1] C₁₀H₁₂N₅O₆PS·2Li, M 375.2. Purify it as for the diNa salt [Murray & Atkinson *Biochemistry* 7 4023 1968, DOI: 10.1021/bi00851a032, PMID: 4301880]. Dissolve 0.3g in dry MeOH (7ml) and 1M LiI (6ml) in dry Me₂CO containing 1% of mercaptoethanol, and the Li salt is precipitated by adding Me₂CO (75ml). The residue is washed with Me₂CO (4 x 30ml) and dried at 55°/25mm. Store at -20°. It has UV with λ_{max} in HCl, pH 1.2 at 257nm (ε 14,800); (0.015M NaOAc, pH 4.8) 259nm (ε 14,800); and in 0.015M NH₄OH, pH 10.1 at 259nm (ε 15,300). It is an AMP-dependent enzyme inhibitor.

Adenosine-5'-triphosphate (ATP) [56-65-5] C₁₀H₁₆N₅O₁₃P₃, M 507.2, m 187°, [α]₅₄₆ -35.5 (c 1, 0.5 M Na₂HPO₄), pK₁²⁵ 4.00, pK₂²⁵ 6.48. ATP is purified by precipitating it as the **barium salt** on adding excess barium acetate solution to a 5% solution of ATP in water. The precipitate is filtered off, washed with distilled water, dissolved in 0.2M HNO₃ and again precipitated with barium acetate. The precipitate, after several washings with distilled water, is redissolved in 0.2M HNO₃, and slightly more than an equivalent of 0.2M H₂SO₄ is added to precipitate all the barium as BaSO₄ which is filtered off. The ATP is then precipitated by addition of a large excess of 95% ethanol. It is filtered off, washed several times with 100% EtOH and finally with dry diethyl ether. It is dried *in vacuo*. [Kashiwagi & Rabinovitch *J Phys Chem* 59 498 1955, DOI: 10.1021/j150528a006; *Beilstein* 26 III/IV 3654.]

S-(5'-Adenosyl)-L-homocysteine (SAH) [979-92-0] C₁₄H₂₀N₆O₅S, M 384.4, m 202°(dec), 204°(dec), 205-207°(dec), [α]_D²⁵ +93 (c 1, 0.2N HCl), [α]_D²³ +44 (c 0.1, 0.05N HCl), (pK see SAM hydrochloride below). It has been recrystallised several times from aqueous EtOH or H₂O to give small prisms, and the UV has λ_{max} at 260nm in H₂O. The solubility in 1M HCl is 19.6-20.4 mg/ml at ~25° (faintly yellow solution). Store at -20°. The **picrate** has **m 170°(dec)** from H₂O. [Baddiley & Jamieson *J Chem Soc* 1085 1955, DOI: 10.1039/

JR9550001085; de la Haba & Cantoni *J Biol Chem* **234** 603 1959, PMID: 13641268; Borchardt et al. *J Org Chem* **41** 565 1976, DOI: 10.1021/jo00865a038; NMR: Follmann & Gremels *Eur J Biochem* **47** 187 1974, DOI: 10.1111/j.1432-1033.1974.tb03682.x; *Beilstein* **26** III/IV 3676.]

(+)-S-Adenosyl-L-methionine chloride (SAM hydrochloride) [24346-00-7; zwitterion-(-) 29908-03-0] $C_{15}H_{22}N_6O_5S \cdot HCl$, **M 434.9**, $[\alpha]_D^{25} +32$ (c 3.3, H_2O) $pK_{Est(1)} \sim 2.13$, $pK_{Est(2)} \sim 4.12$, $pK_{Est(3)} \sim 9.28$. Purify it by ion exchange on Amberlite IRC-150 and eluting with 0.1-4M HCl. [Stolowitz & Minch *J Am Chem Soc* **103** 6015 1981, DOI: 10.1021/ja00410a004.] It has been isolated as the **tri-reineckate salt** by adding 2 volumes of 1% solution of ammonium reineckate in 2% perchloric acid. The reineckate salt separates at once but is kept at 2° overnight. The salt is collected on a sintered glass funnel, washed with 0.5% of ammonium reineckate, dried (all operations at 2°) and stored at 2°. To obtain adenosylmethionine, the reineckate is dissolved in a small volume of methyl ethyl ketone (MEK) and centrifuged at room temperature to remove a small amount of solid. The clear dark red supernatant is extracted (in a separating funnel) with a slight excess of 0.1N H_2SO_4 . The aqueous phase is re-extracted with fresh MEK until it is colourless. [Note that reineckates have UV absorption at 305nm (ϵ 15,000), and the optical density at 305nm is used to detect and estimate reineckate ions.] MEK is removed from the aqueous layer containing adenosylmethionine sulfate; the pH is adjusted to 5.6-6.0 and extracted with two volumes of Et_2O . The **sulfate** is obtained by evaporating the aqueous layer in *vacuo*. The **hydrochloride** can be obtained in the same way but using HCl instead of H_2SO_4 . SAM-HCl has a solubility of 10 w/v% in H_2O . The salts are stable in the cold at pH 4-6 but decompose in alkaline media. Store at -70°. [Cantoni *Biochemical Preparations* **5** 58 1957.] The purity of SAM can be determined by paper chromatography [Cantoni *J Biol Chem* **204** 403 1953, www.jbc.org/content/204/1/403, PMID: 13084611; *Methods Enzymol* **3** 600 1957, DOI: 10.1016/S0076-6879(57)03427-8; for electrophoretic methods or enzymic analysis see Cantoni & Vignos *J Biol Chem* **209** 647 1954, PMID: 13192118]. [*Beilstein* **26** III/IV 3676.]

L-Adrenaline [*R*-(*-*)-epinephrine, **L**-(*-*)-(3,4-dihydroxyphenyl)-2-methylaminoethanol] [51-43-4] $C_9H_{13}NO_3$, **M 183.2**, **m** 210°(dec), 211°(dec), 211-212°(dec), 215°(dec), $[\alpha]_{546}^{20} -61$ (c 5, 0.5M HCl), $[\alpha]_D^{20} -52$ (c 2, 5% HCl), $pK_1^{25} 8.88$ (8.75), $pK_2^{25} 9.90$ (9.89), $pK_3^{25} 12.0$ (~13). L-Adrenaline has been recrystallised from EtOH/AcOH/ NH_3 [Jensen *J Am Chem Soc* **57** 1765 1935, DOI: 10.1021/ja01313a002]. It is sparingly soluble in H_2O , readily soluble in acidic or basic solutions but insoluble in aqueous NH_3 , alkali carbonate solutions, EtOH, $CHCl_3$, Et_2O or Me_2CO . It has also been purified by dissolving in dilute aqueous acid, then precipitating it by adding dilute aqueous ammonia or alkali carbonates. It is readily oxidised in air and turns brown on exposure to light and air. (Epinephrine readily oxidises in neutral alkaline solution. This can be diminished if a little sulfite is added). Store it in the dark under N_2 . [Lewis *Br J Pharmacol Chemother* **9** 488 1954, DOI: 10.1111/j.1476-5381.1954.tb00866.x]. The **hydrogen oxalate salt** has **m** 191-192°(dec, evacuated capillary) after recrystallisation from H_2O or EtOH [Petrov et al. *J Chem Soc* 927 1945, DOI: 10.1039/JR9450000927]. [*Beilstein* **13** III/IV 2927.] It is a cardiac stimulant and adrenoreceptor agonist.

Adrenalone hydrochloride (3',4'-dihydroxy-2-methylaminoacetophenone hydrochloride) [62-13-5] $C_9H_{11}NO_3 \cdot HCl$, **M 217.7**, **m** 243°(dec), 244-249°(dec), 248°(dec), 256°(dec), $pK^{25} 5.5$. The salt is synthesised from α -(*p*-toluenesulfonylmethylamino)-3,4-dimethoxyacetophenone by heating with HCl under pressure [DE Patent 277540 to Bayer, 1914]. Alternatively, catechol (*o*-hydroxyphenol) is melted with chloroacetyl chloride and $POCl_3$ whereby catechol monchloroacetate is formed as intermediate which rearranges to 4-chloroacetylcatechol. Treatment of the latter (100g) in EtOH (50ml) with 40% $MeNH_2$ (200ml) with cooling and shaking for 1 hour then allowing to stand overnight at ~25°, gives a salt that is filtered off, washed with a little EtOH, dissolved in dilute HCl. The **free base** is precipitated with ammonia, filtered off washed with a little H_2O , EtOH and Et_2O to give 60g of base which is best converted to the **hydrochloride** for storage. The **hydrochloride** is purified by recrystallisation from EtOH or aqueous EtOH (prisms, decomposing >240°). The solubility at ~25° in H_2O is ~12.5w/v% and in 94%EtOH it is 2.2w/v%. For X-ray crystallography, well developed, water clear crystals were obtained by slow evaporation of an aqueous solution of pure **hydrochloride** [for the crystal structure of the **hydrochloride monohydrate** see Bergin *Acta Crystallogr Section B* **27** 2139 1971, DOI: 10.1107/S0567740871005478]. The **free base** [99-45-6] is obtained by basifying an aqueous solution of the hydrochloride from which it precipitates out, and is recrystallised from H_2O , washed with H_2O , EtOH and Et_2O in which it is poorly soluble, to give microscopic needles **m** 235-236°(dec) (darkening at ~200° and decomposing >230°) [Stolz *Chem Ber* **37** 4149 1904, DOI: not available; DE 152814 to Hoechst.] The **tris-**

benzenesulfonyl derivative has **m 106-107°(dec)** from aqueous EtOH. Similarly, Stolz prepared **2-amino-1-(3,4-dihydroxyphenyl)ethan-1-one** (**3',4'-dihydroxy-2-aminoacetophenone, noradrenalone**) [499-61-6] **C₈H₉NO₃, M 167.2, m** darkens at **200°** and decomposes at **300°**) as pale yellow microscopic crystals, by using NH₃ instead of MeNH₂, and its *hydrochloride* had **m 260°(dec)** (colourless plates from aqueous EtOH). Stolz also prepared **2-ethylamino-1-(3,4-dihydroxyphenyl)ethan-1-one** [pale yellow microscopic crystals from H₂O, **m 185°(dec)**] by using EtNH₂ instead of NH₃, and its *hydrochloride* crystallises as needles **m 260°(dec)** from hot H₂O, but is poorly soluble in EtOH.

It is an andrenergic agonist, is a topical vasoconstrictor, is hemostatic and its activity is similar to that of adrenaline. It is used as a topical anaesthetic. [Gero *J Org Chem* **16** 1222 1951, DOI: 10.1021/jo50002a006; Kindler & Peschke *Arch Pharm* **269** 581, 603 1931, DOI: 10.1002/ardp.19312693504; Beilstein **14** IV 832.]

Allopurinol [Allosig, Progot, **1,5-dihydro-4H-pyrazolo(3,4-d)pyrimidin-4-one**] [315-30-0] **C₅H₄N₄O, M 136.1, m >350°, pK²⁵ 10.2**. It is obtained from 3-amino-4-pyrazolecarboxamide sulfate (75g) in formamide (200ml) at 180-190°/45min, cooled, diluted with cold H₂O (1L) and the *purinol* (48g) is filtered off. It is purified by recrystallisation from H₂O. It has also been purified by dissolving in aqueous 2N NaOH, shake with Norit (charcoal), filter, acidify with 2N HCl to pH 3-4, collect the purinol and recrystallise it from hot H₂O. It does not melt when heated to 350°. Its solubility (w/v%) in H₂O is 0.78 (25°) and 190 (100°), in DMSO it is 4.6 (25°), in CHCl₃ it is 0.6 (25°), in EtOH it is 0.3 (25°), and in *n*-octanol it is <0.01 (25°). It has UV with λ_{max} at 250nm (ε 7600) in 0.1N HCl; 257nm (ε 7200), in 0.1N NaOH, and 252nm (ε 7600) IN MeOH. [Robins *J Am Chem Soc* **78** 784 1956, DOI: 10.1021/ja01585a023; Schmidt & Druey *Helv Chim Acta* **39** 986 1956, DOI: 10.1002/hlca.19560390345].

Allopurinol is a *xanthine oxidase* inhibitor and interferes with uric acid biosynthesis. It is a commonly used drug for the treatment of gout, and renal calculi. [For 'Allopurinol prevents formation of renal calculi in many cases' see *InPharma* **106** 9 1977, DOI: 10.1007/BF03284284; and for 'Allopurinol treatment and its effect on renal function in gout: a controlled study' see Gibson *Ann Rheum Dis* **41** 59 1982, PMCID: PMC1000865.]

Amethopterin (Methotrexate, **4-amino-4-deoxy-N¹⁰-methylpteroyl-L-glutamic acid**) [59-05-2] **C₂₀H₂₂N₈O₅, M 454.4, m 185-204°(dec), [α]_D²⁰ +19 (c 2, 0.1N aqueous NaOH), pK₁ <0.5 (pyrimidine²⁺), pK₂ 2.5 (N⁵-Me⁺), pK₃ 3.49 (α-CO₂H), pK₄ 4.99 (γ-CO₂H), pK₅ 5.50 (pyrimidine⁺)**. Most common impurities are 10-methylpteroyl-glutamic acid, aminopterin and pteroylglutamic acid. Purify the yellow solid by chromatography on Dowex-1 acetate, followed by filtration through a mixture of cellulose and charcoal. It has been recrystallised from aqueous HCl or by dissolution in the minimum volume of N NaOH and acidified until precipitation is complete, filter or *better* collect by centrifugation, wash with H₂O (also by centrifugation) and dry at 100°/3mm. It has UV with λ_{max} at 244 and 307nm (ε 17300 and 19700) in H₂O at pH 1; 257, 302 and 370nm (ε 23000, 22000 and 7100) in H₂O at pH 13. [Momle *Biochemical Preparations* **8** 20 1961, Seeger et al. *J Am Chem Soc* **71** 1753 1949, DOI: 10.1021/ja01173a061.] It is a *potent inhibitor* of dihydrofolate reductase and is used in cancer chemotherapy. [Blakley *The Biochemistry of Folic Acid and Related Pteridines*, North-Holland Publ Co., Amsterdam, NY, pp157-163 1969, ISSN 0071-965X; Beilstein **26** IV 3833.] It is **CARCINOGENIC; HANDLE WITH EXTREME CARE**. Antifolate used extensively in cancer chemotherapy.

Aminopterin (**4-amino-4-deoxypteroyl-L-glutamic acid**) [54-62-6] **C₁₉H₂₂N₈O₅, M 440.4, m 231-235°(dec), [α]_D²⁰ +18 (c 2, 0.1N aqueous NaOH), pK₁ <0.5 (pyrimidine²⁺), pK₂ 2.5 (N⁵-Me⁺), pK₃ 3.49 (α-CO₂H), pK₄ 4.65 (γ-CO₂H), pK₅ 5.50 (pyrimidine⁺)**. Purify aminopterin by recrystallisation from H₂O. It has properties similar to those of methotrexate (above). It has UV at λ_{max} 244, 290 and 355nm (ε 18600, 21300 and 12000) in H₂O at pH 1; 260, 284 and 370nm (ε 28500, 26400 and 8600) in H₂O at pH 13. [Seeger et al. *J Am Chem Soc* **71** 1753 1949, DOI: 10.1021/ja01173a061; Angier & Curran *J Am Chem Soc* **81** 2814 1959, DOI: 10.1021/ja01520a049; Blakley *The Biochemistry of Folic Acid and Related Pteridines*, North-Holland Publ Co., Amsterdam, NY, pp 157-163 1969, ISSN 0071-965X.] For small quantities, chromatograph it on DEAE cellulose with a linear gradient of ammonium bicarbonate pH 8 and increase the molarity from 0.1 to 0.4. Monitoring is by following the UV absorption of the fractions. For larger quantities, a near boiling solution of aminopterin (5g) in H₂O (400ml) is slowly treated with small portions of MgO powder (~0.7g, calcined magnesias) with vigorous stirring until a small amount of MgO remained undissolved and the pH rises from 3-4 to 7-8. Charcoal (1g) is added to the hot solution and filtered immediately through a large sintered glass funnel

of medium porosity and lined with a hot wet pad of Celite (~2-3 mm thick). The filtrate is cooled in ice, and the crystals of the **Mg salt** are collected by filtration and recrystallised from boiling H₂O (200ml). The crystals are washed with EtOH and dried *in vacuo*. The Mg salt is redissolved in boiling H₂O (200ml) and carefully acidified with vigorous agitation with AcOH (2ml). Pure aminopterin (3g) separates in fine yellow needles (**dihydrate**) which are easily filtered. The solid is washed with cold H₂O, then Me₂CO and dried *in vacuo*. If a trace of impurity is still present as shown by DEAE cellulose chromatography or TLC, repetition of the process will remove it; see UV above. [Loo *J Med Chem* **8** 139 1965, DOI: 10.1021/jm00325a036; *Beilstein* **26** IV 3831.] **CARCINOGENIC**. It strongly inhibits **dihydrofolate reductase** from producing tetrahydrofolate then 5,10-methylenetetrahydrofolate consequently arresting the formation of thymidine, a DNA building block.

3-Aminopyridine adenine dinucleotide [21106-96-7] **M 635.4** (see **NAD for pK**) Purify it by ion-exchange chromatography as described [for interaction with dehydrogenases see Fisher et al. *J Biol Chem* **248** 4293 1973, PMID: 4351221; Anderson & Fisher *Methods Enzymol* **66** 81 1980, DOI: 10.1016/0076-6879(80)66441-6].

Amiodarone {2-butyl-3-[3,5-diiodo-4-(β-diethylaminoethoxy)benzoyl]benzofuran, (2-butyl-1-benzofuran-3-yl)-[4-[2-(diethylamino)ethoxy]-3,5-diiodophenyl]methanone; **phen = ar**, hence Amiodarone} [1951-25-3] **C₂₅H₂₉I₂NO₃**, **M 645.3**, **pK²⁵ 6.56 ± 0.06**. The simplest synthesis starts from 2-phenoxy-*n*-hexaldehyde (e.g. 2mmol) which is cyclised to 2-*n*-butylbenzofuran in toluene (15ml) by boiling under reflux in the presence of Amberlyst-15 resin (0.75g) for 20min, filtered to remove the resin, washed with aqueous NaHCO₃, H₂O and dried (MgSO₄), filtered and evaporated to dryness. The residue is distilled (b 80-82°/3mm, 114-116°/8mm) to give the **furan** in 94% yield. A Friedel-Crafts reaction of the furan with *p*-methoxybenzoyl chloride/AlCl₃ gave 2-*n*-butyl-3-(*p*-hydroxybenzoyl)benzofuran in 84% yield, and is followed by treatment with KI and I₂ (to diiodinate the phenolic group) then NaOMe (to form the Na-phenolate) and *N*-(β-chloroethyl)diethylamine to provide **Amiodarone** in 43% yield. The **free base** is a strong enough base to form a stable **hydrochloride**. [Prepn: **FR 1339389**; Tondeur & Binon, **US 3248401** (1963, 1966 to Soc. Belge l'Azote Prod. Chim. Marly); for the synthesis of 2-*n*-butylbenzofuran see Witczak & Kwiecień *Synth Commun* **35** 2223 2005, DOI: 10.1080/00397910500182747; Wang for the 'Synthetic Process for 2-Butyl-3-(hydroxy-3,5-diiodobenzoyl)-benzofuran' Chin. Patent 1,858,042, Nov 8, 2006; Ha et al. *Eur J Clin Pharmacol* **55** 807 2000, DOI: 10.1007/s002280050701.]

Amiodarone hydrochloride (Amiodor) [19774-82-4] **C₂₅H₂₉I₂NO₃ · HCl**, **M 681.8**, has **m 156°**, **m 159.2°**. It is purified by recrystallisation from acetone. Its solubilities at 25° in g/100ml are: CHCl₃(44.5), CH₂Cl₂(19.2), MeOH(9.98), EtOH(1.28), *C₆H₆(0.65), THF(0.60), MeCN(0.32), 1-octanol(0.30), Et₂O(0.17), *n*-PrOH(0.13), H₂O at pH 1.5-7.5(~0.07), hexane(0.03), petroleum ether(0.001) and *iso*-PrOH(<0.001). Its UV (MeOH) has λ_{max} nm (ε) at 208 (4,700), 241 (4,400), 270sh (1,800) and 280sh (1,600), and the IR (KBr) has ν_{max} at 3000-3070(Ar C-H), 2860-2960(aliph C-H), 2200-2900(*tert*-NH⁺), 1636(diAr C=O), 1560-1015(benzofuran C=C), 1580(aliph CH₃), 1285(ketonic C-C), 1250 and 1075(ether C-O-C) and 1225-1025(*tert*-amine C-N) cm⁻¹. [For physicochemical properties see Bonati et al. *J Pharm Sci* **73** 829 1984, DOI: 10.1002/jps.2600730632]

Amiodarone hydrochloride is an **anti-arrhythmic (Class III)** and **anti-angina** drug used in cardiac surgery, cardiac resuscitation, after out-of-hospital cardiac arrest and atrial fibrillation [Kudenchuk et al. *N Engl J Med* **341** 871 1999, PMID: 10486418; Doy et al. *N Engl J Med* **342** 913 2000, DOI: 10.1056/NEJM200003303421302, PMID: 10738049].

Anion exchange resins. These should be conditioned before use by successive washing with water, EtOH and water, and taken through two OH⁻—H⁺—OH⁻ cycles by successive treatment with N NaOH, water, N HCl, water and N NaOH, then washing with water until neutral to give the **OH⁻ form**. (See commercial catalogues on ion-exchange resins.)

R(-)-Apocodeine {(6aR)-5,6,6a,7-tetrahydro-10-methoxy-6-methyl-4H-dibenzo[de,g]quinolin-11-ol} [641-36-1] **C₁₈H₁₉NO₂**, **M 281.3**, **m 124°**, [α]_D²⁵ -27 (0.45, MeOH), **pK_{Est(1)} ~ 7.0**, **pK_{Est(2)} ~ 8.2**. Recrystallise apocodeine from absolute EtOH by boiling and allowing to stand at 0° to give the **alcoholate**, **m 104.5-106.5°**; EtOH is lost slowly at 25°/2mm but readily at 78°/2mm, and the **anhydrous base** has **m 122.5-124.5°**. It is soluble in Et₂O. [Folkers *J Am Chem Soc* **58** 1814 1936, DOI: 10.1021/ja01300a502.] It has also been crystallised from MeOH (small prisms) or MeOH with a little CH₂Cl₂ as a waxy solid and dried at 80°/2mm. It softens above 100° before melting and is sensitive to air and light. The **hydrochloride** [641-36-1] **C₁₈H₁₉NO₂ ·**

HCl, M 317.7, which is formed from the base in EtOH containing an equivalent amount of HCl followed by addition of Et₂O, is recrystallised from 95% EtOH and adding Et₂O until crystals separate. It melts at 260-263° (dec, with softening at 140°) and has $[\alpha]_D^{20}$ -41.3 to -43.3 (c 0.81, H₂O). It is an emetic. [for configuration see Corrodi & Hardegger *Helv Chim Acta* **38** 2038 1955, DOI: 10.1002/hlca.19550380746; for synthesis and pharmacology see Neumeyer et al. *J Med Chem* **16** 1223 1973, DOI: 10.1021/jm00269a601.]

R-Apomorphine [58-00-4, 41372-20-7 (0.5H₂O)] C₁₇H₁₇NO₂, **M 267.3**, **m 195°(dec)**, **pK₁¹⁵ 7.20 (NH₂)**, **pK₂¹⁵ 8.91 (phenolic OH)**. Crystallise R-apomorphine from CHCl₃ and a little petroleum ether, also from Et₂O with 1 mol of Et₂O which it loses at 100°. It sublimes in a high vacuum. It is white but turns green in moist air or in alkaline solution. Its UV has λ_{\max} at 336, 399 (98% EtOH). The *di-O-methylether* is an oil **b 175°/high vacuum**, whose *picrate* crystallises from MeOH and has **m 140° (dec)**. The *di-O-acetate* crystallises from EtOAc/petroleum ether with **m 127-128°**, $[\alpha]_D^{25}$ -88 (c 1, 0.1 N HCl). The *di-O-benzoyl derivative* has **m 156-158°** (from EtOH) and $[\alpha]_D^{18}$ +43.44 (c 3.3, CHCl₃). [Pschorr et al. *Chem Ber* **35** 4377 1902, DOI: 10.1002/cber.19020350496; Beilstein **21** H 246. **NARCOTIC**. It is a dopamine (D1 and D2) receptor agonist, an emetic and an anti-parkinsonian. **R-Apomorphine hydrochloride** [41372-20-7] C₁₇H₁₇NO₂·HCl, **M 312.8**, has **m 285-287°(dec)**, $[\alpha]_D^{20}$ -48 (c 1, H₂O). Crystallise the salt from H₂O (*hemihydrate*) and from EtOH. Crystals turn green on exposure to light. (see previous entry). **NARCOTIC**.

Atenolol {4-[(2-hydroxy-3-*iso*-propylamino)propoxy]phenylacetamide, Tensig} [RS- 29122-68-7, S- 93379-54-5] C₁₄H₂₂N₂O₃, **M 266.3**, **m 146-147°, 150-151° (sealed capillary)**, **pK₂₅ 9.6**. The physiologically active S-enantiomer was synthesised in seven steps starting with *p*-hydroxyacetophenone and involving a *hydrolytic kinetic resolution* (HKR after Jacobsen et al.) of a terminal epoxide using (*R,R*) salen CoIII acetate (see Jacobsen salen catalysts in Catalysts-Part 1, Chapter 5) and *iso*-propylamine. The final step involved converting the methyl 4-[(2*S*-2-hydroxy-3-*iso*-propylamino)propoxy]phenylacetate formed into the amide (*S*-Atenolol) by reaction in MeOH with NH₄OH for 2 hours, evaporating and purifying the drug (72% yield of last step) by recrystallisation from EtOAc. It has **m 148-150°**, $[\alpha]_D^{21}$ -17.0 (c 1, 1 N HCl), IR (KBr) has ν_{\max} at 3325, 3170, 2950, 2839, 1687, 1596, 1506, 1329, 1263, 1139, 1054, 968 and 835 cm⁻¹; the ¹H NMR [Gemini 200 spectrometer, DMSO *d*₆, TMS internal standard] has δ at 1.05 (d, 6H, *J* = 7.0 Hz), 2.70-2.90 (m, 1H), 3.25 (d, 2H, *J* = 6.0 Hz), 3.42 (s, 2H), 3.90-4.10 (m, 2H), 6.18 (br s, 2H), 6.85 (d, 2H, *J* = 7.2 Hz), 7.15 (d, 2H, *J* = 7.2 Hz), 7.55 (br s, 1H); EIMS *m/z* (%): 266(M⁺, 45), 248(21), 208(11), 190(10), 150(22), **132(100)**, 104(31), 78(16), 52(31). The HKR was carried out at room temperature with excellent *enantio* selectivity. The method can be applied to large-scale preparations of S-atenolol without any problems. [Bose & Narsaiah *Bioorg Med Chem* **13** 627 2005, PMID: 15653330.]

It is a **β -blocker** that decreases the activity of the heart by stopping nerve messages to it. It is a prescription drug for the treatment of hypertension, supraventricular tachycardia, atrial and ventricular fibrillation, atrial flutter, myocardial infarction and *angina pectoris*. Stoschitzky et al. [*Chirality* **5** 15 1993, DOI: 10.1002/chir.530050104; Lindner US Patent No 4,652,672 1987] conclude that in man it is the *S*-enantiomer that has the **β -blocking effect**. [Kähönen et al. *Br J Pharmacol* **112** 925 1994, PMCID: PMC1910195; Amery et al. *New Engl J Med* **290** 284 1974, DOI: 10.1056/NEJM197401312900515, PMID: 4148766.]

Atorvastatin {Lipitor, Atorva, 7-[(4-fluorophenyl)- 3-phenyl-4-(phenylaminocarbonyl)-5-propan-2-ylpyrrol-1-yl]- β R, δ R)-3(*R*),5(*R*)-dihydroxyheptan-1-oic acid} [134523-00-5] C₃₃H₃₅FN₂O₅, **M 558.6**, **m 159.9°**. Its solubility in H₂O is <1mg/ml, and 100mg/ml in DMSO. In the final stage of the synthesis of the *Na salt* of the *cis*-drug described by Cho et al. (Patent US 8124790B2 2012 to Medichem Korea Co Ltd) the *tert*-butyl ester in MeOH is cooled to 0° and the pH carefully adjusted to 12 using aqueous 3N NaOH with stirring at ~25 for 2 hours while maintaining the pH constant. After the starting material is consumed, checked by TLC and HPLC, stirring is continued for 2 hours at 0°; the white *Na salt* is filtered off, dried at 35-40° *in vacuo* to give the sodium salt of atorvastatin (6.4g, 74.4%) as a white powder. It has ¹H NMR (DMSO *d*₆) signals at 9.84(s, 1H), 7.4-7.9(m, 2H), 6.9-7.9(m, 12H), 3.93(br, 1H), 3.74(br, 2H), 3.0-3.2(m, 1H), 2.0-1.93 (m, 4H), 1.2-1.36(m, 9H). The lactone, ***R*-(+)-trans-2-(4-fluorophenyl)-5-(1-methylethyl)-N,3-diphenyl-1-[(tetrahydro-4-hydroxy-2-oxo-2H-pyran-6-yl)ethyl]-1H-pyrrole-4-carboxamide**, was prepared by Roth et al. [*J Med Chem* **34** 357 1991, DOI: 10.1021/jm00105a056] from the respective *t*-butyl β -ketoheptanoate with Et₃B/NaBH₄ followed by refluxing in toluene (Dean-Stark trap, to cyclise to the lactone intramolecularly), then the crude colourless foam in CHCl₃ was chromatographed through a silica gel column and eluted with EtOAc/heptane (1:1 v/v) followed by further

chromatography (silica gel; CHCl_3 /propanol 98.5:1.5 v/v) and finally gave pure ***R*-(+)-enantiomer lactone** as a colourless foamy solid mixture (9:1 trans/cis **m 90-97°**). Recrystallisation from toluene/EtOAc gave pure ***R*-form m 148-149°**, with $[\alpha]_D^{23} +24.53$ (c .053 CHCl_3). Hydrolysis of this isomer (10g, 18.5mmol) with NaOH (0.74g, 18.5mmol) in a mixture of tetrahydrofuran/ H_2O (1:2, 90ml) at 0° by warming slowly to 25°, then concentrated and the solid was collected and dried *in vacuo*. Its IR (KBr) has ν_{max} at 3400, 1651, 1598, 1565, 1511, 1438, 1412, 1316, 1224, 1139, 1159, 844, 754 and 702 cm^{-1} ; the ^1H NMR (90MHz, in DMSO d_6) has the following signals downfield from TMS in ppm: 1.34 (d, $J = 7\text{Hz}$, 6H); 1.5 (m, 4H); 1.80 (d of d, $J = 15, 8\text{Hz}$, 1H); 1.99 (d of d, $J = 15, 4\text{Hz}$, 1H); 3-4 (m, 8H); 6.9-7.3 (m, 12H); 7.50 (d, $J = 8\text{Hz}$, 2H) and 9.85 (s, 1H) {compare with above}. The ***S*-(-)-enantiomer lactone** was similarly prepared and found to have a relative potency as a **HMG-CoA reductase inhibitor** of ~13.9 compared to ~500 for the ***R*-(+)-enantiomer**. An elegant synthesis has been accomplished starting from the freely available and cheap **isoascorbic acid** [see 89-65-6] and provides the correct (+)-3*R*, 5*R*- configuration in the heptanoic (or lactone) chain for the most active enantiomer of the drug, without requiring optical resolution steps in which half of the molecule is not needed [Slettinger et al. 'A diastereospecific, non-racemic synthesis of a novel β -hydroxy- δ -lactone HMG-CoA reductase inhibitor' *Tetrahedron Lett* **26** 2951 1985, DOI: 10.1016/S0040-4039(00)98589-7; Bock et al. *Acta Chem Scand* **37B** 341 1983, DOI: 10.3891/acta.chem.scand.37b-0341; Isbell & Frush *Carbohydr Res* **72** 301 1979, DOI: 10.1016/S0008-6215(00)83954-3; review: Roth *Progr Med Chem* **40** 1 2002, DOI: 10.1016/S0079-6468(08)70080-8, PMID: 12516521.] In the form of the free acid, the Na or Ca salts or the lactone these **statins** are lipid lowering agents and **strong inhibitors of 3-hydroxy-3-methylglutamylCoA reductase** which is the first enzyme of the *de novo* biosynthesis of cholesterol (e.g. in mammalian liver) and have the effect of increasing LDL (low density lipoproteins) uptake by hepatocytes, decreasing HDL-C (high density lipoprotein cholesterol) and lowering triglyceride levels. [Roth US patent 4681893, to Warner-Lambert Co, issued 1987-07-21.]

Aureomycin (7-chlorotetracycline) [57-62-5] $\text{C}_{22}\text{H}_{23}\text{ClN}_2\text{O}_8$, **M 478.8**, **m 172-174°(dec)**, $[\alpha]_D^{23} -275$ (MeOH), **pK₁ 3.3, pK₂ 7.44, pK₃ 9.27**. Aureomycin is dehydrated by azeotropic distillation of its solution with toluene. On cooling, the anhydrous material crystallises out and is recrystallised from $^*\text{C}_6\text{H}_6$, then dried under vacuum at 100° over paraffin wax. (If it is crystallised from MeOH, it contains **MeOH** in the crystals which is not removed on drying.) [Stephens et al. *J Am Chem Soc* **76** 3568 1954, DOI: 10.1021/ja01642a064; Laskin & Chan *Biochem Biophys Res Commun* **14** 137 1964, DOI: 10.1016/0006-291X(64)90243-8]. [Beilstein **14** IV 2631.] **Aureomycin hydrochloride (7-chlorotetracycline hydrochloride)** [64-72-2] $\text{C}_{22}\text{H}_{23}\text{ClN}_2\text{O}_8 \cdot \text{HCl}$, **M 514.0**, has **m 210-215°(dec), 234-236°(dec)**, $[\alpha]_D^{25} -23.5$ (H_2O). Purify the salt by dissolving 1g in 20ml of hot water, cooling rapidly to 40°, treating with 0.1ml of 2M HCl, and chilling in an ice-bath. The process is repeated twice. It is also recrystallised from $\text{Me}_2\text{NCHO}/\text{Me}_2\text{CO}$. Store at 2-8°. [Stephens et al. *J Am Chem Soc* **76** 3568 1954, DOI: 10.1021/ja01642a064; UV: McCormick et al. *J Am Chem Soc* **79** 2849 1957, DOI: 10.1021/ja01568a050; Beilstein **14** IV 2631.]

Bacitracin (Altracin, Topitracin, a cyclic polypeptide with a peptide chain tail) [1405-87-4] $\text{C}_{66}\text{H}_{103}\text{N}_{17}\text{O}_{16}\text{S}$, **M 1422.7**, $[\alpha]_D^{23} +5$ (H_2O). Bacitracin has been purified by carrier displacement using *n*-heptanol, *n*-octanol and *n*-nonanol as **carriers** and 50% EtOH in 0.1 N HCl. The pure material gives one spot with $R_F \sim 0.5$ on paper chromatography using $\text{AcOH}:\text{n-BuOH}:\text{H}_2\text{O}$ (4:1:5). [Porath *Acta Chem Scand* **6** 1237 1952, DOI: 10.3891/acta.chem.scand.06-1237.] It has also been purified by ion-exchange chromatography. It is a white powder soluble in H_2O and EtOH but insoluble in Et_2O , CHCl_3 and Me_2CO . It is stable in acidic solution but unstable in base. It is a strong topical antibacterial. [Newton & Abraham *Biochem J* **47** 257 1950, DOI: 10.1042/bj0470257; Synthesis: Munekata et al. *Bull Chem Soc Jpn* **46** 3187 1973, DOI: 10.1246/bcsj.46.3187; 3835 1973, DOI: 10.1246/bcsj.46.3835; Beilstein **27** III/IV 5746.]

***N*⁶-Benzyladenosine** [4294-16-0] $\text{C}_{17}\text{H}_{19}\text{N}_5\text{O}_4$, **M 357.4**, **m 177-179°, 185-187°**, $[\alpha]_D^{25} -68.6$ (c 0.6, EtOH)(see pK of adenosine). Purify it by recrystallisation from EtOH. It has UV with λ_{max} at 266nm (aqueous EtOH/HCl) and 269 nm (aqueous EtOH/NaOH). [Kissman & Weiss *J Org Chem* **21** 1053 1956, DOI: 10.1021/jo01115a621; Beilstein **26** III/IV 3682.] It has plant growth activity [Strnad *J Plant Growth Reg* **15**(4) 179 1996, DOI: 10.1007/BF00190582].

***N*-Benzylpenicillin sodium salt (penicillin G Na salt)** [69-57-8] $\text{C}_{16}\text{H}_{17}\text{N}_2\text{O}_4\text{S} \cdot \text{Na}$, **M 356.4**, **m 215°** (char-

ring and dec), 225° (dec), $[\alpha]_D^{20}$ +269 (c 0.7, MeOH), $[\alpha]_D^{25}$ +305 (c 1, H₂O), pK_a²⁵ 2.76 (4.84 in 80% aqueous EtOH)(for free acid). Purify the salt by dissolving it in a small volume of MeOH (in which it is more soluble than EtOH) and treating gradually with ~5 volumes of EtOAc. This gives an almost colourless crystalline solid (rosettes of clear-cut needles) and recrystallising twice more if slightly yellow in colour. The salt has also been conveniently recrystallised from the minimum volume of 90% Me₂CO and adding an excess of absolute Me₂CO. A similar procedure can be used with wet *n*-BuOH. If yellow in colour, then dissolve (~3.8g) in the minimum volume of H₂O (3ml), add *n*-BuOH and filter through a bed of charcoal. The salt forms long white needles on standing in a refrigerator overnight. More crystals can be obtained on concentrating the mother liquors *in vacuo* at 40°. A further recrystallisation (without charcoal) yields practically pure salt. A good preparation has ~600 Units/mg. The presence of H₂O in the solvents increases the solubility considerably. The solubility in mg/100ml at 0° is 6.0 (Me₂CO), 15.0 (Me₂CO/0.5% H₂O), 31.0 (Me₂CO/1.0% H₂O), 2.4 (methyl ethyl ketone), 81.0 (*n*-butanol) and 15.0 (dioxane at 14°). Alternatively, it is dissolved in H₂O (solubility is ~10%), filtered if necessary and precipitated by addition of EtOH and dried in a vacuum over P₂O₅. A sample can be kept for 24 hours at 100° without loss of physiological activity. It also crystallises from MeOH/EtOAc. [IR: Barnes et al. *Anal Chem* **19** 620 1947, DOI: 10.1021/ac60009a002; *The Chemistry of Penicillin* (Clarke, Johnson and Robinson eds.) Princeton University Press, Princeton NJ, Chapter V 85 1949, Open library OL16386917M; *Beilstein* **27** III/IV 5861.]

Other salts, e.g. the **potassium salt** (M 372.5 [113-58-4]) can be prepared from the Na salt by dissolving it (147mg) ice-cold in H₂O acidified to pH 2, extracting with Et₂O (~50ml), washing once with H₂O, and extracting with 2ml portions of 0.3% KHCO₃ until the pH of the extract rises to ~6.5 (~7 extractions). The combined aqueous extracts are lyophilised, and the white residue is dissolved in *n*-BuOH (1ml, absolute) with the addition of enough H₂O to effect solution. Remove insoluble material by centrifugation and add absolute *n*-BuOH to the supernatant. Crystals should separate on scratching, and after 2.5 hours in a refrigerator they are collected, washed with absolute *n*-BuOH and EtOAc and dried (yield 51.4mg). It also crystallises from aqueous Me₂CO. The **potassium salt** has m 214-217° (dec) (block preincubated at 200°; heating rate of 3°/min) and $[\alpha]_D^{22}$ +285 (c 0.748, H₂O). [*Beilstein* **27** III/IV 5861.]

The **free acid (penicillin G)** (M 334.5, [61-33-6]) has m 186-187° (MeOH/Me₂CO), and m 190-191° (H₂O) $[\alpha]_D^{25}$ +282 (EtOH). [Review: Sheehan & Henery-Logan *J Am Chem Soc* **84** 2983 1962, DOI: 10.1021/ja00874a029; Chain et al. *Antibiotics* (Oxford University Press) **2** 1949, and Cook *Quarterly Reviews (Chemical Society)* **2** 203 1948, DOI: 10.1039/QR9480200203.]

Bergapten (5-methoxypsoralen, 4-methoxy-7H-furo[3,2-g]chromen-7-one) [484-20-8] C₁₂H₈O₄, M 216.2, m 190-193°, 191-193°. Crystallise it from EtOH or aqueous MeOH, and it sublimes *in vacuo*. Its properties are similar to those of its 9-methoxy isomer (xanthotoxin, see below). It is slightly soluble in *C₆H₆, CHCl₃ and AcOH but insoluble in H₂O. Its fluorescence has λ_{ex} at 352nm with λ_{em} at 480nm. It is a DNA intercalator and **possible carcinogen**. [Howell & Robertson *J Chem Soc* 293 1937, DOI: 10.1039/JR9370000293; Boyer et al. *Biochemistry* **27** 3011 1988, DOI: 10.1021/bi00408a052; *Beilstein* **19/6** V 4.] It is present in Bergamot oil [Sakamaki et al. *J Food Hygienic Soc Jpn* **49**(4) 326 2008, DOI: 10.3358/shokueishi.49.326, PMID: 18787320].

(+)-Bicuculine [R-6(5,6,7,8-tetrahydro-6-methyl-1,3-dioxolo[4,5-g]isoquinolin-5-yl)-furo-[3,4-c]-1,3-benzodioxolo-8(6H)-one] [485-49-4] C₂₀H₁₇NO₆, M 367.4, m 177°, 193-195°, 193-197°, 215°, $[\alpha]_D^{20}$ +126 (c 1, CHCl₃), $[\alpha]_D^{20}$ +159 (c 1, CHCl₃), pK_a²⁵ 4.84. It crystallises from CHCl₃/MeOH as plates. The crystals melt at 177°, then solidify and re-melt at 193-195° [Manske *Canad J Research* **21B** 13 1943, DOI: 10.1139/cjr43b-002]. It is soluble in CHCl₃, *C₆H₆, EtOAc but sparingly soluble in EtOH, MeOH and Et₂O. [For Stereochemistry see Bláha et al. *Coll Czech Chem Commun* **29** 2328 1964, DOI: org/10.1135/ccccc19642328; Snatzke et al. *Tetrahedron* **25** 5059 1969, DOI: 10.1016/S0040-4020(01)83253-1; Pharmacol: Curtis et al. *Nature* **226** 1222 1970, DOI: 10.1038/2261222a0; *Beilstein* **27** III/IV 1900]. It is a light sensitive strong GABAA receptor antagonist. [Khawaled et al. *Pflügers Archiv: Eur J Physiol* **438**(3) 314 1999, DOI: 10.1007/s004240050915, ISSN 0031-6768. PMID: 10398861.]

L-erythro-Biopterin (2-amino-4-hydroxy-6-[(1R,2S)-1,2-dihydroxypropyl]pteridine) [22150-76-1] C₉H₁₁N₅O₃, M 237.2, m >300°(dec), $[\alpha]_D^{20}$ -80, $[\alpha]_D^{20}$ -65 (c 2.0, M HCl), pK_a²⁵ 2.23(2.45), pK_a²⁵ 7.89(8.05). Purify L-erythro-biopterin by chromatography on Florisil, which was washed thoroughly with 2M HCl, and eluted with 2M HCl. The fractions with UV-fluorescence are pooled, evaporated *in vacuo* and the

residue is recrystallised. Biopterin is best recrystallised (90% recovery) by dissolving in 1% aqueous NH_3 (ca 100 parts), and adding this solution dropwise to an equal volume of M aqueous formic acid at 100° and allowing to cool at 4° overnight. It is dried at 20° to $50^\circ/0.1\text{mm}$ in the presence of P_2O_5 . [Schircks et al. *Helv Chim Acta* **60** 211 1977, DOI: 10.1002/hlca.19770600125; Armarego et al. *Aust J Chem* **35** 785 1982, DOI: 10.1071/CH9820785.] It also crystallises from ca 50 parts of water or 100 parts of hot 3M aqueous HCl by adding hot 3M aqueous NH_3 and cooling. It has UV with λ_{max} at 212, 248 and 321nm (log ϵ 4.21, 4.09 and 3.94) in H_2O at pH 0.0; 223nm, 235.5, 274.5 and 345nm (log ϵ 4.07, 4.10, 4.18 and 3.82) in H_2O at pH 5.0; 221.5, 254.5 and 364nm (log ϵ 3.92, 4.38 and 3.84) in H_2O at pH 10.0. [Sugimoto & Matsuura *Bull Chem Soc Jpn* **48** 3767 1975, DOI: 10.1246/bcsj.48.1679 *Beilstein* **26** III/IV 4032.] Several urinary pteridine levels are used as **potential biomarkers** for non-invasive diagnosis of cancer [Gamagedara et al. *Clinica Chimica Acta* **1** 14 2011; PMID: 20869359].

D-(+)-Biotin (vitamin H, hexahydro-2-oxo-1H-thieno[3,4-d]imidazole-4-pentanoic acid) [58-85-5] $\text{C}_{10}\text{H}_{16}\text{N}_2\text{O}_3\text{S}$, M **244.3**, m **229-231 $^\circ$** , **230.2 $^\circ$ (dec)**, **230-231 $^\circ$** , **231-233 $^\circ$** , **232-234 $^\circ$ (dec)**, $[\alpha]_{\text{D}}^{20} +108$, $[\alpha]_{\text{D}}^{20} +91.3$ (c 1, 0.1N NaOH), $\text{pK}_{\text{Est}} \sim 4.8$. D-(+)-Biotin crystallises from hot water in fine long needles with a solubility of 22 mg/100ml at 25° . Its solubility in 95% EtOH is 80 mg/100 ml at 25° . Its isoelectric point is at pH 3.5. Store the solid and solutions under sterile conditions because it is susceptible to mould growth. [Confalone *J Am Chem Soc* **97** 5936 1975, DOI: 10.1021/ja00853a061; Wolf et al. *J Am Chem Soc* **67** 2100 1945, DOI: 10.1021/ja01228a013; Ohrui & Emoto *Tetrahedron Lett* 2765 1975, DOI: 10.1016/S0040-4039(00)75234-8; Harris et al. *J Am Chem Soc* **66** 1756 1944, DOI: 10.1021/ja01238a041.] The **(+)-methyl ester** has m **166-167 $^\circ$** (from MeOH/Et₂O), $[\alpha]_{\text{D}}^{22} +57$ (c 1, CHCl_3) [du Vigneaud et al. *J Biol Chem* **140** 643 1941, <http://www.jbc.org/content/140/2/643>; 763 1941]; the **(+)-S-oxide** has m **200-203 $^\circ$** , and $[\alpha]_{\text{D}}^{20} +130$ (c 1.2, 0.1N NaOH) [Melville *J Biol Chem* **208** 495 1954, PMID: 13174559]; the **SS-dioxide** has m **274-275 $^\circ$ (dec)**, **268-270 $^\circ$** , and the **SS-dioxide methyl ester** has m **239-241 $^\circ$** (from MeOH/Et₂O) [Hofmann et al. *J Biol Chem* **141** 207 1941, <http://www.jbc.org/content/141/1/207>]. [*Beilstein* **27** III/IV 7979.] It is a **coenzyme** for lipid and protein metabolism, converting food into glucose. Also vital for maintaining skin, hair and mucous membranes. **D-(+)-Biotin hydrazide** [66640-86-6] $\text{C}_{10}\text{H}_{18}\text{N}_4\text{O}_2\text{S}$, M **258.3**, has m **238-240 $^\circ$** , **245-247 $^\circ$** , $[\alpha]_{\text{D}}^{20} +66$ (c 1, Me_2NCHO). Wash the hydrazide with H_2O , dry it, wash it with MeOH then Et₂O, and dry. Recrystallise it from hot H_2O (clusters of prisms) [Hofmann et al. *J Biol Chem* **144** 513 1942, <http://www.jbc.org/content/144/2/513>]. [*Beilstein* **27** III/IV 7980.]

D-(+)-Biotin N-hydroxysuccinimide ester (+-biotin N-succinimidyl ester) [35013-72-0] $\text{C}_{14}\text{H}_{19}\text{N}_3\text{O}_5\text{S}$, M **341.4**, has m **210 $^\circ$** , **212-214 $^\circ$** , $[\alpha]_{\text{D}}^{20} +53$ (c 1, Me_2NCHO). Recrystallise the ester from refluxing isoPrOH and dry it in a vacuum over P_2O_5 + KOH. Its solubility in dry DMF is ~50mg/ml, and it is stable for at least 30 days at -20° . Store it at -20° . [Jasiewicz et al. *Exp Cell Biol* **100** 213 1976, DOI: 10.1016/0014-4827(76)90344-X.]

D-(+)-Biotin 4-nitrophenyl ester [33755-53-2] $\text{C}_{16}\text{H}_{19}\text{N}_3\text{O}_5\text{S}$, M **365.4**, has m **160-163 $^\circ$** , **163-165 $^\circ$** , $[\alpha]_{\text{D}}^{25} +47$ (c 2, Me_2NCHO containing 1% AcOH), $[\alpha]_{\text{D}}^{25} +51$ also reported. The ester has been recrystallised by dissolving 2g in 95% EtOH (30ml), heated to dissolve, then cooled in an ice-water bath. The crystals are collected, washed with ice-cold 95% EtOH (5ml) and dried over P_2O_5 . Its solubility in MeOH is 25mg/ml at -25° (colourless to green-yellow solution). The R_F on silica plates ($\text{CHCl}_3/\text{MeOH}$ 19:1) is 0.19 [Bodanszky & Fagan *J Am Chem Soc* **99** 235 1977, DOI: 10.1021/ja00443a040].

N-(+)-Biotinyl-4-aminobenzoic acid [6929-40-4] $\text{C}_{17}\text{H}_{21}\text{N}_3\text{O}_4\text{S}$, M **363.4**, has m **295-297 $^\circ$** , **295-300 $^\circ$** , $[\alpha]_{\text{D}}^{23} +56.55$ (c 0.5, 0.1N NaOH), $\text{pK}_{\text{Est}} \sim 4.0$. Dissolve the acid in NaHCO_3 solution, cool and precipitate it by adding N HCl. Collect the solid, dry it at 100° and recrystallise it from MeOH. **Note** that it is hydrolysed by aqueous 3M, 1M and 0.2M HCl at 120° , but can be stored in 5% aqueous NaHCO_3 at -20° without appreciable hydrolysis [Knappe et al. *Biochem Zeitschrift* **338** 599 1963, PMID: 14087327; Wolf et al. *J Am Chem Soc* **73** 4142 1951, DOI: 10.1021/ja01153a027; Bayer & Wilchek *Methods Enzymol* **26** 1 1980]. [*Beilstein* **27** III/IV 7984.] It is a substrate used in biotinidase assays [Craft et al. *Biochemistry* **24** 2471 1985, DOI: 10.1021/bi00331a012; Hayakawa & Oizumi *J Chromatogr* **383**(1) 148 1986, PMID: 3493251].

N-Biotinyl-6-aminocaproic N-succinimidyl ester [72040-63-2] $\text{C}_{20}\text{H}_{30}\text{N}_4\text{O}_6\text{S}$, M **454.5**, has m **149-152 $^\circ$** , **169-171 $^\circ$** . Dissolve ~400mg of the ester in dry propan-2-ol (~25ml) with gentle heating. Reduce the volume to ~10ml by gentle boiling and allow the solution to cool. Decant the supernatant carefully from the white crystals, dry the crystals in a vacuum over P_2O_5 at 60° overnight. This material gives one spot on TLC. [Costello et al.

Clin Chem **25** 1572 1979, PMID: 572747; Kincaid et al. *Methods Enzymol* **159** 605 1988, DOI: 10.1016/0076-6879(88)59058-4; PMID: 2842624.]

N-(+)-Biotinyl-6-aminocaproyl hydrazide (biotin-6-aminohexanoic hydrazide) [109276-34-8] $C_{16}H_{29}N_5O_3S$, **M 371.5**, has **m 189-191°**, **195-196°**, **210°**, $[\alpha]_D^{20} +23$ (c 1, Me₂NCHO). Suspend the hydrazide in ice-water (100mg/ml), stand overnight at 4°, filter and dry the solid in a vacuum. Recrystallise it from isoPrOH. Its solubility in DMSO is ~25mg/ml at ~25°. R_F is 0.26 on SiO₂ plate using CHCl₃/MeOH (7:3) as eluent. [O'Shannessy et al. *Anal Biochem* **163** 204 1987, DOI: 10.1016/0003-2697(87)90114-X.]

N-(+)-Biotinyl-L-lysine (Biocytin) [576-19-2] $C_{16}H_{28}N_4O_4S$, **M 372.5**, has **m 228.5°**, **228-230° (dec)**, **241-243°**, **245-252° (dec, sintering at 227°)**, $[\alpha]_D^{25} +53$ (c 1.05, 0.1 N NaOH). Recrystallise biocytin rapidly from dilute MeOH or Me₂CO. It can also be recrystallised from H₂O by slow evaporation or by dissolving in the minimum volume of H₂O and adding Me₂CO until a solid separates. It is freely soluble in H₂O and AcOH but insoluble in Me₂CO. [Wolf et al. *J Am Chem Soc* **74** 2002 1952, DOI: 10.1021/ja01128a039; **72** 1048 1950, DOI: 10.1021/ja01158a534.] It has been purified by chromatography on superfiltrol-Celite, Al₂O₃ and by countercurrent distribution and then recrystallised [IR: Peck et al. *J Am Chem Soc* **74** 1999 1952, DOI: 10.1021/ja01128a038]. The **hydrochloride** crystallises from aqueous Me₂CO/HCl and has **m 227°(dec)**. [Beilstein **27** III/IV 7984.] Intermediate in biotin metabolism found in serum and urine. For tracing with neuroimaging agents see Mishra et al. [ACS Chemical Neuroscience **1**(2) 129 2010, DOI: 10.1021/cn900010d].

Brefeldin A [1-*R*-2*c*,15*c*-dihydroxy-7*t*-methyl-(1*r*,13*t*)-6-oxa-bicyclo[11.3.0]hexadeca-3*t*,11*t*-dien-5-one, Decumbin] [20350-15-6] $C_{16}H_{24}O_4$, **M 280.4**, **m 200-202°**, **204°**, **204-205°**, $[\alpha]_D^{22} +95$ (c 0.81, MeOH). Brefeldin A was isolated from *Penicillium brefeldianum* and recrystallised from aqueous MeOH/EtOAc or MeOH. Its solubility in H₂O is 0.6mg/ml, 10mg/ml in MeOH and 24.9mg/ml in EtOH. The ***O*-acetate** recrystallises from Et₂O/pentane has **m 130-131°**, and $[\alpha]_D^{22} +17$ (c 0.95, MeOH). [Sigg *Helv Chim Acta* **47** 1401 1964, DOI: 10.1002/hlca.19640470603; UV and IR: Härrä et al. *Helv Chim Acta* **46** 1235 1963, DOI: 10.1002/hlca.19630460419; total synthesis: Kitahara et al. *Tetrahedron Lett* 3021 1979, DOI: 10.1016/S0040-4039(00)71000-8; for X-ray structure see Weber et al. *Helv Chim Acta* **54** 2763 1971, DOI: 10.1002/hlca.19710540839; Beilstein **18** III/IV 1220.] For 'Brefeldin A: insights into the control of membrane traffic and organelle structure' see Klausner et al. [Review *J. Cell Biol.* **116**(5) 1071 1992, DOI: 10.1083/jcb.116.5.1071, PMID: 1740466].

5-Bromo-2'-deoxyuridine [59-14-3] $C_9H_{11}BrN_2O_5$, **M 307.1**, **m 193-197°(dec)**, **217-218°**, $[\alpha]_D^{25} -4.1$ (c 0.1, H₂O), **pK²⁵ ~ 8.1**. Recrystallise the uridine from EtOH or 96% EtOH. It has UV with λ_{max} at 279 nm at pH 7.0, and 279 nm (log ϵ 3.95) at pH 1.9. Its R_F values are 0.49, 0.46 and 0.53 in *n*-BuOH/AcOH/H₂O (4:1:1), *n*-BuOH/EtOH/H₂O (40:11:19) and *i*-PrOH-25% aqueous NH₃-H₂O (7:1:1), respectively. [Stone *Nature* **204** 190 1964, DOI: 10.1038/204190a0; Prystaš & Šorm *Coll Czech Chem Comm* **29** 2956 1964, DOI: org/10.1135/cccc19642956; Beilstein **24** III/IV 1234.] Analogue of thymidine which incorporates into DNA of replicating cells [Lehner et al. *Cell and Tissue Research* **345**(3) 313 2011, DOI: 10.1007/s00441-011-1213-7; PMID: 21837406]. 5-Bromodeoxycytidine deaminated to **5-bromodeoxyuridine** [Russo et al. *Cancer Res* **44**(4) 1702 1984, PMID: 6704976].

5-Bromouridine [957-75-5] $C_9H_{11}BrN_2O_6$, **M 323.1**, **m 180-182°**, **217-218°**, $[\alpha]_D^{22} -11$ (c 2, H₂O), **pK²⁵ 8.1**. Recrystallise it from 96% EtOH. It has UV with λ_{max} at 279nm (log ϵ 3.95) in H₂O pH 1.9. R_F in *n*-BuOH/AcOH/H₂O (4:4:1) is 0.49; in *n*-BuOH/EtOH/H₂O (40:11:9) it is 0.46 and in isoPrOH/25%NH₃/H₂O (7:1:2) it is 0.53 using Whatman No 1 paper. [Prystaš & Šorm *Coll Czech Chem Commun* **29** 2956 1964, DOI: org/10.1135/cccc19642956; Beilstein **31** H 24.] It is a uridine derivative that can be incorporated into RNA and damage DNA. For *in vitro* splicing of pre-mRNA containing bromouridine see Wansink et al. [Molecular biology reports **19**(2) 109 1994, DOI: 10.1007/BF00997156, PMID: 8072491].

Brucine [357-57-3 (anhydrous), 5892-11-5 (4H₂O)] $C_{23}H_{26}N_2O_4$, **M 394.5 (anhydrous)**, **430.5 (2H₂O)**, **466.5 (4H₂O)**, **m 178-179°**, $[\alpha]_{546}^{20} -149.9$ (anhydrous; c 1, in CHCl₃), $[\alpha]_D^{20} -119$ (anhydrous; c 2, in CHCl₃), **pK₁¹⁵ 2.50**, **pK₂¹⁵ 8.16 (pK₂²⁵ 8.28)**. Crystallise brucine once from water or aqueous Me₂CO (as the **tetrahydrate**), then suspend it in CHCl₃ and shake with anhydrous Na₂SO₄ (to dehydrate the brucine, which then dissolves). Precipitate it by pouring the solution into a large bulk of dry petroleum ether (b 40-60°), filter and heat to 120° in a high vacuum [Turner *J Chem Soc* 837 842 1951, DOI: 10.1039/JR9510000837]. The **tetrahydrate** crystallises

from a mixture of EtOH and H₂O as colourless elongated needles [Eeles *Acta Cryst* **6** 809 1953, DOI: 10.1107/S0365110X53002337; *Beilstein* **27** III/IV 7875.] A very useful base for the *optical resolution* of chiral acids. It paralyzes inhibitory neurons, an antagonist at glycine receptors. **BITTER and VERY POISONOUS.**

Brucine sulfate [4845-99-2, 60583-39-3 (7H₂O)] (C₂₃H₂₆N₂O₄)₂ · H₂SO₄ · 7H₂O, **M 887.0 (anhydr)**, has **m ~180°(dec)**, [α]_D²⁰ **-36 (dry; c 1, H₂O)**, [α]_D²⁰ **-24 (c 1, H₂O)**. The *heptahydrate* crystallises from water as colourless laths. [Eeles *Acta Cryst* **6** 809 1953, DOI: 10.1107/S0365110X53002337; *Beilstein* **27** III/IV 7875.]

Butyryl choline iodide [(2-butyryloxyethyl)trimethyl ammonium iodide] [2494-56-6] C₉H₂₀NO₂ · I, **M 301.7, m 85-89°, 87°, 93-94°**. Recrystallise the iodide from isoPrOH or Et₂O. [Tammelin *Acta Chem Scand* **10** 145 1956, DOI: 10.3891/acta.chem.scand.10-0145.] The *perchlorate* has **m 72°** (from isoPrOH). [Aldridge *Biochem J* **53** 62 1953, DOI: 10.1042/bj0530062; *Beilstein* **4** IV 1448.] A substrate for cholinesterase [O'Brien et al. *Biochim Biophys Acta* **526**(1), 129 1978, DOI: 10.1016/0005-2744(78)90297-8], and acetylcholinesterase [EC 3.1.1.7] [Nachmansohn & Wilson *Meth Enzymol* **1** 642 1955, DOI: 10.1016/0076-6879(55)01112-9].

S-Butyryl thiocholine iodide [(2-butyrylmercaptoethyl)trimethyl ammonium iodide] [1866-16-6] C₉H₂₀NOS · I, **M 317.2, m 171-174°, 173-176°**. Recrystallise S-butyryl thiocholine iodide from propan-1-ol and dry it *in vacuo*; store it in the dark under N₂. The *bromide* has **m 150°** (from Me₂CO) or **m 140-143°** (from butan-1-ol). [Gillis *Chem Ind (London)* 111 1957, Hansen *Acta Chem Scand* **11** 537 1957, DOI: 10.3891/acta.chem.scand.11-0537; *Beilstein* **4** IV 1586.]

Carbamazepine (Tegretol, 5H-dibenzo[*b,f*]azepine-5-carboxamide) [298-46-4] C₁₅H₁₂N₂O, **M 236.3, m 189-192°, 190-193°, 191-192°, 204-206°**. Recrystallise from EtOH/*C₆H₆, and dry *in vacuo*, or recrystallise it from EtOH then from *C₆H₆. It is soluble in Me₂CO EtOH and propylene glycol, but not in H₂O. Schindler [US **2948718**, 1960 to Geigy] prepared it in an efficient fume cupboard from 5H-dibenzo[*b,f*]azepine (19.3 g, [256-96-2]) and phosgene (suffocating and poisonous) in toluene to give 5H-dibenzo[*b,f*]azepine-5-carbonyl chloride, **m 168-169°**. Also in an sufficient fume cupboard, the acid chloride in absolute EtOH and NH₃ gas provides the desired *azepine*, **m 204-206°**, after washing with H₂O and recrystallising from absolute EtOH then from *C₆H₆. Its UV has λ_{max} (MeOH) at 237 and 285 nm; the IR has ν_{max} (KBr) at 1678, 1594, 1298, 800, 787, 769 cm⁻¹. It is an *anticonvulsant* and *Na⁺ channel inhibitor*. [Stenger & Roulet *Med Exp Int J Exp Med* **11** 191 1964, ISSN/ISBN: 0258-2589.] It is used in *epilepsy* [Livingston et al. *JAMA* **200** 204 1967, DOI: 10.1001/jama.1967.03120160070009] and pain associated with neuralgia [Scott in 'Carbamazepine' in The History of Epileptic Therapy, History of Medicine Series. CRC Press, 1993, ISBN 9781850703914; Sidebottom & Maxwell *J Clin Pharm Ther* **20** 31 1995, PMID: 7775611.]

Cation exchange resins. These should be conditioned before use by successive washing with water, EtOH and water, and taken through two H⁺-OH⁻-H⁺ cycles by successive M HCl, water, M NaOH, water and M HCl treatment, then washed with water to give the *H⁺ form*. (See commercial catalogues on ion exchange resins).

Cephalosporin C potassium salt [28240-09-7] C₁₆H₂₂N₃O₂S · H₂O, **M 453.5 (nhydrous)**, [α]_D²⁰ **+103 (H₂O)**, **pK₁ <2.6, pK₂ 3.1, pK₃ 9.8**. Purify the salt by dissolving it in the minimum volume of H₂O (filter) and adding EtOH until separation of solid is complete. A solution is stable in the pH range 2.5-8. It has UV with λ_{max} at 260 nm (log ε 3.95) in H₂O. The *Ba salt* has [α]_D²⁰ **+80 (c 0.57, H₂O)** [Woodward et al. *J Am Chem Soc* **88** 852 1966, DOI: 10.1021/ja00956a051; Abraham & Newton *Biochem J* **79** 377 1961, DOI: 10.1042/bj0790377; Hodgkin & Maslen *Biochem J* **79** 393 1961, DOI: 10.1042/bj0790393; see also Abraham *Quarterly Reviews London* **21** 231 1967, DOI: 10.1039/QR9672100231]. [*Beilstein* **27** III/IV 5902.] Antibiotic similar to penicillin.

Chlorambucil [Leukeran, 4-{bis(2-chloroethyl)amino}-phenylbutyric acid] [305-03-3] C₁₄H₁₉Cl₂NO₂, **M 304.2, m 64-66°, pK₁ 5.8 (6.0 at 66°, 50% aqueous Me₂CO), pK₂ 8.0**. Chlorambucil is recrystallised from petroleum ether (flat needles) and has a solubility at 20° of 66% in EtOH, 40% in CHCl₃, 50% in Me₂CO but is insoluble in H₂O [Everett et al. *J Chem Soc* 2386 1953, DOI: 10.1039/JR9530002386]. [As a substrate for the human glutathione transferase P1-1 see Parker et al. *J Mol Biol* **380**(1) 131 2008, DOI: 10.1016/j.jmb.2008.04.066; *Beilstein* **14** IV 1715.] It is a *chemotherapeutic* drug for treating chronic lymphocytic leukemia, low grade Hodgkin and non-Hodgkin lymphoma. **CARCINOGEN.**

Chloramphenicol [Amphicol, 1*R*,2*R*-(-)-2-{2,2-dichloroacetyl-amino}-1-(4-nitrophenyl)-propan-1,3-diol] [56-75-7] $C_{11}H_{12}Cl_2N_2O_5$, **M 323.1**, **m 149-151°**, **150-151°**, **151-152°**, $[\alpha]_D^{20} +20.5$ (c 3, EtOH), $[\alpha]_D^{25} -25.5$ (EtOAc). Purify this broad-spectrum antibiotic by recrystallisation from H₂O (solubility is 2.5mg/ml at 25°) or ethylene dichloride as needles or long plates, and by sublimation at high vacuum. It has $A_{1\text{cm}}^{1\%}$ 298 at λ_{max} 278nm, and it is slightly soluble in H₂O (0.25%) and propylene glycol (1.50%) at 25° but is freely soluble in MeOH, EtOH, BuOH, EtOAc and Me₂CO. [Rebstock et al. *J Am Chem Soc* **71** 2458 1949, DOI: 10.1021/ja01175a065; Controulis et al. *J Am Chem Soc* **71** 2463 1949, DOI: 10.1021/ja01175a066; Long & Troutman *J Am Chem Soc* **71** 2469, 1949, DOI: 10.1021/ja01175a067; 2473 1949, DOI: 10.1021/ja01175a068; Ehrhart et al. *Chem Ber* **90** 2088 1957, DOI: 10.1002/cber.19570900957; *Beilstein* **13** IV 2742.] It inhibits bacterial protein synthesis. **Chloramphenicol palmitate (Vetranal)** [530-43-8] $C_{27}H_{42}Cl_2N_2O_6$, **M 561.5**, has **m 90°**, $[\alpha]_D^{26} +24.6$ (c 5, EtOH). The palmitate crystallises from *benzene or xylene with **m 105-106°** and $[\alpha]_D^{21} -39.5$ (c 2, Et₂O), λ_{max} 267.3nm. [Edgerton et al. *J Am Chem Soc* **77** 27 1955, DOI: 10.1021/ja01606a007; *Beilstein* **13** IV 2753.]

2-Chloroadenosine [146-77-0] $C_{10}H_{12}ClN_5O_4$, **M 301.7**, **m 145-146°(dec)**, **147-149°(dec)**, **pK_{Est(1)} ~ 0.5**, **pK_{Est(2)} ~ 7.6**. Purify 2-chloroadenosine by recrystallisation from H₂O (~1% in cold), and it has UV with λ_{max} at 264 nm (pH 1 and 7) and 265 nm (pH 13) in H₂O. [Brown & Weliky *J Org Chem* **23** 125 1958, DOI: 10.1021/jo01095a626; Schaeffer & Thomas *J Am Chem Soc* **80** 3738 1958, DOI: 10.1021/ja01547a068; IR: Davoll & Lowy *J Am Chem Soc* **74** 1563 1952, DOI: 10.1021/ja01126a066; *Beilstein* **26** III/IV 3725.] It is a selective adenosine A₁ receptor agonist.

Chlorophyll a [479-61-8] $C_{55}H_{72}MgN_4O_5$, **M 983.5**, **m 117-120°**, **150-153°**, **~152.3°(dec)**, **178-180° (sinters at ~150°)**, $[\alpha]_D^{20} -262$ (Me₂CO). It forms green crystals from Me₂CO, Et₂O/H₂O, Et₂O/hexane/H₂O or Et₂O/pentane/H₂O. It is sparingly soluble in MeOH and insoluble in petroleum ether. In alkaline solution it gives a blue-green colour with deep red fluorescence. A very crude chlorophyll mixture has been purified by chromatography on low melting polyethylene (MI 0.044; 'Dow' melting index MI <2) and developed with 70% aqueous Me₂CO. The order of effluent from the bottom of the column is: xanthophylls, chlorophyll *b*, chlorophyll *a*, phaeophytins and carotenes. A mixture of chlorophylls *a* and *b* is best separated by chromatography on sugar, and the order is chlorophyll *b* elutes first followed by chlorophyll *a*. To an Me₂CO/H₂O solution of chlorophylls 200ml of iso-octane are added, and the mixture shaken in a separating funnel and the H₂O is carefully removed. The iso-octane layer is dried (Na₂SO₄) and applied onto a glass column (5cm diameter) dry packed with 1000ml of powdered sucrose which has been washed with 250ml of iso-octane. Elution with 0.5% of isopropanol in iso-octane gives chlorophyll *a*. Keeping the eluate overnight at 0° yields micro crystals which are collected by filtration or centrifugation (Yield 40mg). It is insoluble in H₂O but soluble in many organic solvents. The UV_{EtOH} has λ_{max} **660**, 613, 577, 531, 498, **429** and 409 nm. **Note** that **anhydrous** chlorophylls can be easily enolised, epimerised or allomerised in dry polar organic solvents. [Anderson & Calvin *Nature* **194** 285 1962, DOI: 10.1038/194285a0; Stoll & Weidemann *Helv Chim Acta* **16** 739 757 1933, DOI: 10.1002/hlca.193301601100; NMR: Katz et al. *J Am Chem Soc* **90** 6841 1968, DOI: 10.1021/ja01026a050; **85** 3809 1963, DOI: 10.1021/ja00906a020; for *a* and *b* ORD see Wolf et al. *Justus Liebigs Ann Chem* **704** 208 1967, DOI: 10.1002/jlac.19677040123; Willstätter & Isler *Justus Liebigs Ann Chem* **390** 269 1912, DOI: 10.1002/jlac.19123900302; *Beilstein* **26** III/IV 3243.] **Chlorophyll b** [519-62-0] $C_{55}H_{70}MgN_4O_6$, **M 907.52**, **sinters at 86-92°**, **sinters at 170°**, **dec at 160-170°**, has **m 183-185°**, **190-195°**, $[\alpha]_D^{20} -267$ (Me₂CO/MeOH), $[\alpha]_D^{25} -133$ (MeOH/Pyridine **95:5**). See purification of chlorophyll *a*, and is separated from '*a*' by chromatography on sucrose [UV, IR: Stoll & Weidemann *Helv Chim Acta* **42** 679, 681 1959, DOI: 10.1002/hlca.19590420307]. It forms red-black hexagonal bipyramids or four-sided plates from dilute EtOH and has been recrystallised from CHCl₃/MeOH. It is insoluble in H₂O, but soluble in MeOH, EtOH, EtOAc and insoluble in petroleum ether. [Dougherty et al. *J Am Chem Soc* **88** 5037 1966, DOI: 10.1021/ja00973a056; *Beilstein* **26** III/IV 3787.] These are plant pigments in the light-energy harvesting system.

6-Chloropurine riboside (6-chloro-9-β-D-ribofuranosyl-9H-purine) [2004-06-0, 5399-87-1] $C_{10}H_{11}ClN_4O_4$, **M 286.7**, **m 158-162°(dec)**, **165-166°(sintering at 155°)**, **168-170°(dec)**, $[\alpha]_D^{26} -45$ (c 0.8, H₂O). Purify the riboside by suspending the dry solid (~12 g) in hot MeOH (130 ml) and then adding enough hot H₂O (~560ml) to cause solution, filter and set aside at 5° overnight. The colourless crystals of the riboside are filtered off, washed with Me₂CO, Et₂O and dried at 60°/0.1mm. More material can be obtained by evaporat-

ing the filtrate to dryness and recrystallisation of the residue from MeOH/H₂O (2:1) (15ml/g). It has UV with λ_{max} at 264nm (ϵ 9140) in H₂O. [Robins *Biochemical Preparations* **10** 145 1963, Baker et al. *J Org Chem* **22** 954 1957, DOI: 10.1021/jo01359a027.] It is a substrate for adenosine deaminase.

Chromomycin A₃ [7059-24-7] C₅₇H₈₂O₂₆, M 1183.3, m 185°dec, $[\alpha]_{\text{D}}^{23}$ -57 (c 1, EtOH). Dissolve the anthraquinone glycoside antibiotic (10g) in EtOAc and add to a column of Silica Gel (Merck 0.05-0.2microns, 4x70cm) in EtOAc containing 1% oxalic acid. Elute with EtOAc+1% oxalic acid and check fractions by TLC. Pool the fractions, wash with H₂O thoroughly, dry and evaporate. Recrystallise the residue from EtOAc. The **heptaacetate** has m 214°, $[\alpha]_{\text{D}}^{23}$ -20 (c 1, EtOH). [Miyamoto et al. *Tetrahedron* **23** 421 1967, DOI: 10.1016/S0040-4020(01)83328-7; Harada et al. *J Am Chem Soc* **91** 5896 1969, DOI: 10.1021/ja01049a047; *Beilstein* **17/5** V 673.] It is a fluorescent DNA-binding dye specific for GC residues.

8S,9R-(-)-Cinchonidine [485-71-2] C₁₉H₂₂N₂O, M 294.4, m 204-205°, 210.5°, $[\alpha]_{\text{D}}^{20}$ -127.5 (c 0.5, EtOH), $[\alpha]_{\text{D}}^{20}$ -109.2 (EtOH), pK₁¹⁵ 4.17, pK₂¹⁵ 8.4, pK₁²⁰ 5.80, pK₂²⁰ 10.03. Crystallise cinchonidine from aqueous EtOH (prisms or plates). It is a strong base and readily forms a *mono-* and *di-hydrochloride* and other stable salts. Slightly soluble in Et₂O, but very soluble in EtOH and CHCl₃. For *N-benzylcinchonidinium chloride* see [69257-04-1]. [*Beilstein* **23** III/IV 2824, **23/12** IV 406.] **8R,9S-(+)-Cinchonine** [118-10-5] C₁₉H₂₂N₂O, M 294.4 has m 260-264°, 265°, $[\alpha]_{\text{D}}^{20}$ +268 (c 0.5, EtOH), pK₁¹⁵ 4.28, pK₂¹⁵ 8.35, pK₁²⁰ 5.85, pK₂²⁰ 9.96, and crystallises from EtOH or Et₂O (needles). Its solubility (w/v) is 1.7% in EtOH, 4% in boiling EtOH, 1% in CHCl₃ and is insoluble in H₂O. It is a strong base and readily forms a *mono-* and *di-hydrochloride* and other stable salts. For *N-benzylcinchoninium chloride* see [69221-14-3]. [Rabe *Justus Liebigs Ann Chem* **365** 366, 371 1909, DOI: 10.1002/jlac.19093650214; *Beilstein* **23** III/IV 2819, 2832.] Both bases are used in asymmetric synthesis, and the *optical resolutions* of chiral acids.

Clofazimine [2-(4-chloroanilino)-3-isopropylimino-5-(4-chlorophenyl)-3,4-dihydrophenazine] [2030-63-9] M 473.5, m 210-212°, pK²⁰ 8.37 (8.51). It recrystallises from Me₂CO (dark red crystals). Its solubility in CHCl₃ and EtOH is 7% and 0.1%, at ~25°. Insol. in H₂O. It is antibacterial. [Barry et al. *J Chem Soc* 859 1958, DOI: 10.1039/JR9580000859; *Beilstein* **25** III/IV 3033.] Antibacterial (tuberculostatic and leprostatic)

Clonidine hydrochloride [Catapres, 2-(2,6-dichloroanilino)-2-imidazoline hydrochloride] [4205-91-8] C₉H₉Cl₂N₃. HCl, M 266.6, m 305°, pK²⁵ 5.88 (free base). This antihypertensive is recrystallised from EtOH/Et₂O and dried in a vacuum (solubility in H₂O is 5%). The *free base* [4205-90-7] M 230.1, has m 124-125° and is recrystallised from hexane. [Jen et al. *J Med Chem* **18** 90 1975, DOI: 10.1021/jm00235a020; NMR: Jackman & Jen *J Am Chem Soc* **97** 2811 1975, DOI: 10.1021/ja00843a033.] It is a centrally acting α_2 -adrenergic agonist drug for *controlling hypertension*, treatment of *ADHD*, drugs and alcohol *addiction* and *withdrawal*.

Cloxacillin sodium salt (sodium 3-*o*-chlorophenyl-5-methyl-4-isoxazolyl penicillin monohydrate) [642-78-4, 7081-44-9 (H₂O)] C₁₉H₁₇ClN₃O₅S. Na, M 457.9, m 170°, $[\alpha]_{\text{D}}^{20}$ +163 (H₂O pH 6.0-7.5), pK_{Est} ~ 2.8 (COOH). Purify cloxacillin sodium salt by dissolving it in isoPrOH containing 20% of H₂O, and diluting with isoPrOH to a water content of 5% and chilling. Recrystallise it again in this manner. The sodium salt is collected and dried at 40° in air to give the colourless *monohydrate*. It is soluble in H₂O (5%), MeOH, EtOH, pyridine and ethylene glycol. [Doyle et al. *J Chem Soc* 5838 1963, DOI: 10.1039/JR9630005838; Nayler et al. *Nature* **195** 1264 1962, DOI: 10.1038/1951264a0.] A penicillin-type antibiotic with AmpC β -lactamase activity.

(-)-Cocaine {ecogonine methyl ester benzoate, 2 β -carbomethoxy-3- β -benzoxypitropane, methyl 1*R*-(*exo,exo*)-3-(benzoyloxy)-2-methyl-8-azabicyclo[3.2.1]octane-2-carboxylate} [50-36-2] C₁₇H₂₁NO₄, M 303.4, m 98°, b 187-188°/0.1mm, $[\alpha]_{\text{D}}^{20}$ -15.8 (c 4, CHCl₃), $[\alpha]_{\text{D}}^{20}$ -35 (50% EtOH), pK²⁵ 5.59 and 8.61 (8.39). Recrystallise the addictive drug from EtOH; it sublimes < 90° *in vacuo* in an amorphous form. The *hydrochloride* has m 195° (from MeOH/Et₂O) and $[\alpha]_{\text{D}}^{20}$ -72 (c 2 in H₂O, pH 4.5), -78.5 (50% aqueous EtOH). [Carr & Reynolds *J Chem Soc* **97** 1328 1910, DOI: 10.1039/CT9109701328; Tufariello et al. *J Am Chem Soc* **101** 2435 1979, DOI: 10.1021/ja00503a033.] α -Cocaine is the (+) *enantiomer*. [*Beilstein* **22** I 547, **22** II 150.]

Coccarboxylase tetrahydrate (aneurine pyrophosphoric acid tetrahydrate, thiamine pyrophosphoric acid tetrahydrate) [136-09-4, 68684-55-9] C₁₂H₁₈N₄O₇P₂S. 4H₂O, M 496.4, m 220-222°(sinters at 130-140°),

213-214°, **240-244°(dec)**, **pK_{Est(1)}~2**, **pK_{Est(2)}~6**, **pK_{Est(3)}~9**. Cocarboxylase tetrahydrate crystallises from aqueous Me₂CO. [Wenz et al. *Justus Liebigs Ann Chem* **618** 210 1958, DOI: 10.1002/jlac.19586180124; UV: Melnick *J Biol Chem* **131** 615 1939, <http://www.jbc.org/content/131/2/615>; for X-ray structure see Carlisle & Cook *Acta Cryst (B)* **25** 1359 1969, DOI: 10.1107/S0567740869004043.] The **hydrochloride salt** has **m 242-244°(dec)**, **241-243°(dec)** or **239-240°(dec)** and crystallises from aqueous HCl/EtOH, EtOH containing HCl or HCl/Me₂CO. [Weijlard *J Am Chem Soc* **63** 1160 1941, DOI: 10.1021/ja01849a081; for synthesis see Weijlard & Tauber *J Am Chem Soc* **60** 2263 1938, DOI: 10.1021/ja01276a064; Karrer & Viscontini *Helv Chim Acta* **29** 1901711 1946, DOI: 10.1002/hlca.19460290321; Viscontini et al. *Helv Chim Acta* **32** 1478 1949, DOI: 10.1002/hlca.19490320515; fluorimetric estimation: Burch *Meth Enzymol* **3** 946 1957, DOI: 10.1016/S0076-6879(57)03483-7; Steyn-Parvé *Biochim Biophys Acta* **64** 13 1962, DOI: 10.1016/0006-3002(62)90755-2; for several reviews see Wuest et al. *Ann NY Acad Sci* **98** 383–614 1962, DOI: 10.1111/j.1749-6632.1962.tb30561.x to DOI: 10.1111/j.1749-6632.1962.tb30583.x; *Beilstein* **27** III/IV 1777.]

Coenzyme A trihydrate [85-61-0] **C₂₁H₃₆N₇O₁₆P₃S. 3H₂O**, **M 821.6**, **pK₁ 4.0** (adenine NH₂), **pK₂ 6.5** (PO₄H), **pK₃ 9.6** (SH). The white powder is best stored in an inert atmosphere in the dark in sealed ampoules after drying *in vacuo* over P₂O₅ at 34°. It has UV with λ_{max} at 259 nm (ε 16,800) in H₂O. [Buyske et al. *J Am Chem Soc* **76** 3575 1954, DOI: 10.1021/ja01642a065.] It is soluble in H₂O but insoluble in EtOH, Et₂O and M₂CO. It is readily oxidised in air and is best kept as the more stable **trilithium salt** [Moffatt & Khorana *J Am Chem Soc* **83** 663 1961, DOI: 10.1021/ja01464a036; see also Beinert et al. *J Biol Chem* **200** 385 1953, PMID: 13034796; De Vries et al. *J Am Chem Soc* **72** 4838 1950, DOI: 10.1021/ja01166a532; Gregory et al. *J Am Chem Soc* **74** 854 1952, DOI: 10.1021/ja01123a530; and Baddiley *Adv Enzymol* **16** 1 1955, DOI: 10.1002/9780470122617.ch1]. [*Beilstein* **26** III/IV 3663.] It is an enzyme cofactor.

Coenzyme Q₀ (Ubiquinone, **2,3-dimethoxy-5-methyl-1,4-benzoquinone**, **3,4-dimethoxy-2,5-toluquinone**, **fumigatin methyl ether**) [605-94-7] **C₉H₁₀O₄**, **M 182.2**, **m 56-58°, 58-60°, 59°, 59-60°**. It crystallises in red needles from petroleum ether (b 40-60°) and sublimes at high vacuum at a bath temperature of 46-48° [Anslow et al. *J Chem Soc* 439 1938, DOI: 10.1039/JR9380000439; UV in EtOH: Vischer *J Chem Soc* 815 1953, DOI: 10.1039/JR9530000815; for UV in cyclohexane see Morton et al. *Helv Chim Acta* **41** 2343 1958, DOI: 10.1002/hlca.19580410745; Aghoramurthy et al. *Chem Ind (London)* 1327 1954]. [*Beilstein* **8** IV 2721.]

Coenzyme Q₄ (Ubiquinone-4, **2,3-dimethoxy-5-methyl-6-[3,7,11,15-tetramethyl-hexadeca-2*t*,6*t*,10*t*,14-tetraenyl]-1,4-benzoquinone**) [4370-62-1] **C₂₉H₄₂O₄**, **M 454.7**, **m 30°, 33-45°**. It is a red oil which can be purified by chromatography on SiO₂ plates and eluted with Et₂O/hexane. The purity is checked by HPLC (silica column using 7% Et₂O/hexane). It has UV with λ_{max} at 270 nm (ε 14,800) in petroleum ether. [NMR and MS: Naruta *J Org Chem* **45** 4097 1980, DOI: 10.1021/jo01309a006; cf. Morton *Biochemical Spectroscopy* (Adam Hilger, London, 1975) p 491]. It has also been dissolved in MeOH/EtOH (1:1 v/v) and kept at 5° until crystals appear [Lester et al. *Biochim Biophys Acta* **32** 492 1958, DOI: 10.1016/0006-3002(59)90624-9; Belogrudov et al. *Arch Biochem Biophys* **392**(1) 48 2001, DOI: 10.1006/abbi.2001.2448, PMID: 11469793]. It is an essential component of cellular electron transport.

Coenzyme Q₉ (Ubiquinone-9, **2,3-dimethoxy-5-methyl-6-[3,7,11,15,19,23,27,31,35-nonamethylhexatriaconta-2*t*,6*t*,10*t*,14*t*,18*t*,22*t*,26*t*,30*t*,34-nonaenyl]-1,4-benzoquinone**) [303-97-9] **C₅₄H₈₂O₄**, **M 795.3**, **m 40.5-42.5°, 44-45°, 45°**. The yellow crystals are purified by recrystallisation from petroleum ether and by chromatography on SiO₂ plates and eluted with Et₂O/hexane. The purity can be checked by HPLC (silica column using 7% Et₂O/hexane). It has UV with λ_{max} at 270nm (ε 14,850) in petroleum ether. [NMR and MS: Naruta *J Org Chem* **45** 4097 1980, DOI: 10.1021/jo01309a006; Lester et al. *Biochem Biophys Acta* **32** 492 1958, DOI: 10.1016/0006-3002(59)90624-9; cf. Morton *Biochemical Spectroscopy* (Adam Hilger, London, 1975) p 491, IR: Lester et al. *Biochim Biophys Acta* **33** 169 1959, DOI: 10.1016/0006-3002(59)90511-6; UV: Rüegg et al. *Helv Chim Acta* **42** 2616 1959, DOI: 10.1002/hlca.19590420733; Shunk *J Am Chem Soc* **81** 5000 1959, DOI: 10.1021/ja01527a062; *Beilstein* **8** IV 3313.] A membrane component of proton/electron transport.

Coenzyme Q₁₀ **C₅₉H₉₀O₄**, **M 863.4**, (Ubiquinone-10, **2,3-dimethoxy-5-methyl-6-[3,7,11,15,19,23,27,31,35,-39-decamethyltetra-conta-2*t*,6*t*,10*t*,14*t*,18*t*,22*t*,26*t*,30*t*,34*t*,38-decaenyl]-1,4-benzoquinone**) [303-98-0] **M 795.3**, **m 48-49°, 49°, 49.5-50.5°, 50°, 48-52°**. Purify this yellow-orange solid by recrystallisation from EtOH,

EtOH/Me₂CO or Et₂O/EtOH and by chromatography on silica gel using isoPrOH/Et₂O (3:1) to give orange crystals. It has UV with λ_{\max} at 270nm (ϵ 15,170) in petroleum ether. [Terao et al. *J Org Chem* **44** 868 1979, DOI: 10.1021/jo01319a051; for NMR and MS see Naruta et al. *J Org Chem* **45** 4097 1980, DOI: 10.1021/jo01309a006; for properties of the coenzyme Q compounds see Lester et al. *Biochem Biophys Acta* **33** 169 1959, DOI: 10.1016/0006-3002(59)90511-6; NMR: Planta et al. *Helv Chim Acta* **42** 1278 1959, DOI: 10.1002/hlca.19590420423; Morton *Biochemical Spectroscopy* (Adam Hilger, London, 1975) p 491].

Colcemid {**Demecolcine**, (7S)-6,7-dihydro-1,2,3,10-tetramethoxy-7-(methylamino)-benzo[a]heptalen-9(5H)-one} [477-30-5] C₂₁H₂₅NO₅, M 371.4, m 182-185°, 183-185°, 186°, [α]_D²⁰ -129 (c 1, CHCl₃). Colcemid is purified by chromatography on silica and eluting with CHCl₃/MeOH (9:1), and by recrystallisation from EtOAc/Et₂O to form yellow prisms. Store it at -20°. Its UV in EtOH has λ_{\max} at 243nm (ϵ 30,200) and 350nm (ϵ 16,3000). [Synthesis, IR, NMR, MS: Capraro & Brossi *Helv Chim Acta* **62** 965 1979, DOI: 10.1002/hlca.19790620406; *Beilstein* **8** IV 3319.] It is less toxic than colchicine (see below), and an antineoplastic agent affecting the mitotic cycle: Rieder & Palazzo *J Cell Sci* **102** 387 1992, PMID: 1506421.

Colchicine {N-[(7S)-1,2,3,10-tetramethoxy-5,6,7,9-tetrahydro-9-oxo-benzo[a]heptalen-7-yl]-acetamide} [64-86-8] C₂₂H₂₅NO₆, M 399.5, m 150-160° (dec), 155-157° (dec), 156-157°, [α]_D²⁰ -570 (c 1, H₂O), [α]_D²⁰ -443 (c 1.7, H₂O), [α]_D²⁰ -120 (c 1, CHCl₃), pK_a²⁰ 1.85 and 12.4. Commercial material contains up to 4% desmethylcolchicine. Purify colchicine by chromatography on alumina and eluting with CHCl₃ [Ashley et al. *J Chem Soc* 677 1944, DOI: 10.1039/JR9440000677]. Alternatively, an acetone solution on alkali-free alumina has been used, and eluting with acetone [Nicholls & Tarbell *J Am Chem Soc* **75** 1104 1953, DOI: 10.1021/ja01101a028]. It crystallises as yellow needles from EtOAc or CHCl₃ with solvent of crystallisation which can be removed at ~70°. It is soluble in Et₂O (0.5%), *C₆H₆ (1%) and H₂O (4%). It is sold as ‘**Colgout**’ for the treatment of gout and **binds to tubulin**. [Schreiber et al. *Helv Chim Acta* **44** 540 1961, DOI: 10.1002/hlca.19610440225; Scott et al. *Tetrahedron* **21** 3605 1965, DOI: 10.1016/S0040-4020(01)96977-7; van Tamelen et al. *Tetrahedron* **14** 8 1961, DOI: 10.1016/0040-4020(61)80083-5; *Beilstein* **14** IV 946.]

Compactin [Mevastatin, 1S,7R,8S,8aR)-8-{2-[(2R,4R)-4-Hydroxy-6-oxotetrahydro-2H-pyran-2-yl]ethyl}-7-methyl-1,2,3,7,8,8a-hexahydronaphthalen-1-yl (2S)-2-methylbutanoate} [73573-88-3] C₂₃H₃₄O₅, M 390.5, m 151-153°, 152°, [α]_D²² +283 (c 0.48, acetone). Purify compactin by recrystallisation from aqueous EtOH. Its UV (EtOH) has λ_{\max} at 230, 237 and 246nm (log ϵ 4.28, 4.30 and 4.11); and its IR (KBr) has ν_{\max} 3520, 1750 (lactone CO) and 1710 (CO ester) cm⁻¹. [Clive et al. *J Am Chem Soc* **110** 6914 1988 DOI: 10.1021/ja00228a067; Review: Rosen & Heathcock *Tetrahedron* **42** 4909 1986, DOI: 10.1016/S0040-4020(01)88045-5; IR, NMR, MS: Brown et al. *JCS Perkin Trans 1* 1165 1976, DOI: 10.1039/P19760001165.] It is a **statin** and a **potent inhibitor of 3-hydroxy-3-methylglutaryl-CoA reductase** (HMG-CoA reductase) which inhibits cholesterol biosynthesis, and lowers cholesterol levels [Brown et al. *J Biol Chem* **253** 1121 1978, PMID: 624722; Nakamura & Abeles *Biochemistry* **24** 1364 1985, DOI: 10.1021/bi00327a014; *Beilstein* **18/3** V 145].

Creatinine [2-amino-1-methylimidazolidin-4-one, 2-imino-1-methyl-4(3H)-oxoimidazolidine] [60-27-5] C₄H₇N₃O, M 113.1, m 295° (dec), ~305° (dec), pK₁²⁵ 4.80, pK₂²⁵ 9.2. Likely impurities are creatine and ammonium chloride. Dissolve it in dilute HCl, then neutralise with ammonia. Recrystallise it from H₂O by adding excess of Me₂CO. The **picrate** crystallises from 23 volumes of boiling H₂O and has m 220-221° (dec). [King *J Chem Soc* 2377 1930, DOI: 10.1039/JR9300002374; *Beilstein* **25** III/IV 3543.] Serum creatinine is an important indicator of renal function. The ratio of creatinine in blood to that in urine allows calculation of the **GFR** (glomerular filtration rate), an indication of renal health. [Taylor *Clinical Chemistry* John Wiley & Sons, NY. pp. 4, 58–62 1989; Shemesh et al. *Kidney Int.* **28**(5) 830 1985, DOI: 10.1038/ki.1985.205. PMID: 2418254

Crotaline (monocrotaline, 12,13-dihydroxy-(13 β -14 β H)-14,19-dihydro-20-norcrotalanan-11,15-dione) [315-22-0] C₁₆H₂₃NO₆, M 325.4, m 196-197° (dec), 197-198° (dec), 203° (dec), 204° (dec), [α]_D²⁰ -55 (c 1, EtOH). Crotaline (a pyrrolizidine alkaloid) forms prisms from absolute EtOH and crystallises also from CHCl₃. Its UV(96% EtOH) has λ_{\max} at 217nm (log ϵ 3.32). [Adams et al. *J Am Chem Soc* **74** 5612 1952, DOI: 10.1021/ja01142a021; Culvenor & Smith *Aust J Chem* **10** 474 1957, DOI: 10.1071/CH9570474.] The **hydrochloride** has m 212-214° (from MeOH/Et₂O) and [α]_D²⁸ -38.4 (c 5, H₂O) [Adams & Gianturco *J Am Chem Soc* **78** 1922 1956, DOI: 10.1021/ja01590a041]. The **picrate** has m 230-231.5° (dec) [Adams et al. *J Am Chem Soc* **74** 5612 1952, DOI: 10.1021/ja01142a021]. [*Beilstein* **27** III/IV 6660.] Present in **Crotalinae** snakes (pit

vipers, rattlesnakes, copperhead, cottonmouth, adders), and as alkaloids in Seneceae (*Polygala senega*) and Retorseae plants. [Ruan et al. *Chem Res Toxicol* **6** 16 2014, PMID: 24836403; Culvenor et al. *Ann NY Acad Sci* **163** 837 1969, DOI: 10.1111/j.1749-6632.1969.tb24903.x].

Curcumin [bis(4-hydroxy-3-methoxycinnamoyl)methane, 1,7-bis(4-hydroxy-3-methoxyphenyl)(1*E*,6*E*)-1,6-heptadiene-3,5-one] [458-37-7] $C_{21}H_{20}O_6$, *M* 368.4, *m* 182°, 183°, 184-195°, 184-186°, *pK*₂₅ 9.30 (50% v/v aqueous EtOH), *pK*₁ 7.80, *pK*₂ 8.5 and *pK*₃ 9.0 (H₂O). Purify curcumin (principal cucuminoid of turmeric) by recrystallising it from MeOH, EtOH or acetic acid (yellow, orange-yellow prisms) and drying it *in vacuo*. It is best separated from related compounds by HPLC (see refs below). It is sparingly soluble in H₂O but soluble in AcOH and in alkaline buffer. In solution at pH 1-7 the colour is yellow and at pH 7-9 the colour is brownish-red or deep red. It was commercialised as a pH indicator under the name of '*curcuma paper*'. Curcumin is a phenolic substance which occurs naturally in *Curcuma domestica*, *C. xanthorrhiza*, *C. aromatica*, *C. longa*, and is used in Indian curry cooking as *turmeric*. It has many physiological properties including anti-oxidant, anti-inflammatory, potent anti-tumour, and is an inhibitor of several enzymes (EGFR tyrosine kinase, IκB kinase, nitric oxide synthase, cyclooxygenase, lipoxygenase). It penetrates cell membranes accumulating in plasma membranes, nuclear envelope and endoplasmic reticulum. Its UV-VIS has λ_{\max} (log ϵ)(EtOH) at 268 (4.09) and 430nm (4.74), (dioxane) at 265 (4.18) and 420nm (4.37) and (40% aqueous THF) at 429nm (4.780), with fluorescence maxima at λ_{excit} 433nm and λ_{emis} 511nm in 40% aqueous THF. Its IR (KBr) has ν_{\max} at 3400 (br), 1625, 1600, 1500, 1275, 1025, 960 cm^{-1} . The ¹H NMR [400MHz, CDCl₃] has δ at 6.06 (1H, s, H-4), 16.41 (1H, br s, OH), 7.57 (2H, d, *J* = 16.0Hz, H-2,6), 6.75 (2H, d, *J* = 16.0Hz, H-1,7), 7.32 (2H, d, *J* = 2.0Hz, H-2',2''), 9.64 (2H, s, OH-4',4''), 6.85 (2H, d, *J* = 8.1Hz, H-5',5''), 7.16 (2H, dd, *J* = 2.0, 8.1Hz, H-6',6''), 3.85 (6H, s, 3',3'' OMe) (from TMS); and the ¹³C NMR [100MHz, CDCl₃] has δ at 100 (C-4), 183.2 (C-3,5), 121.1 (C-2,6), 140.7 (C-1,7), 126.4 (C-1',1''), 111.5 (C-2',2''), 148.0 (C-3',3''), 149.4 (C-4',4''), 115.8 (C-5',5''), 123.0 (C-6',6''), 55.7 (OCH₃) (from TMS). It **complexes** strongly with boron, iron, copper and nickel, but weakly with calcium and magnesium. As a 0.1% solution in AcOH it is used for the spectrophotometric determination of Boron as a complex with λ_{\max} 550nm (ϵ 180,000). [Dyrssen et al. *Analyt Chim Acta* **60** 139 1972, DOI: 10.1016/S0003-2670(01)81893-6; Spicer & Strickland *Analyt Chim Acta* **18** 231 1958, DOI: 10.1016/S0003-2670(00)87133-0; Roughley & Whiting, *JCS Perkin Trans I* 2379 1973, DOI: 10.1039/P19730002379; Zsila et al. *Tetrahedron Asymm* **14** 2433 2003, DOI: 10.1016/S0957-4166(03)00486-5; cf. Cooke & Segal *Aust J Chem* **8** 107 1955, DOI: 10.1071/CH9550107; Beilstein **8** H 554, **8** I 757, **8** II 588, **8** III 4312, **8** IV 3697.] The **diacetyl derivative** has *m* 171-172° (from EtOH) and the **dibenzoyl derivative** has *m* 210° (from EtOH or C₆H₆*). The **dimethyl derivative** [1,7-bis(3,4-dimethoxyphenyl)(1*E*, 6*E*)-1,6-heptadiene-3,5-one] has *m* 128-130°, *M* 396.2; the UV has λ_{\max} at 427nm, and the ¹H NMR [200MHz, CDCl₃] has δ at 3.92 (6H, s, 2 x CH₃O-Ph), 3.93 (6H, s, 2 x CH₃O-Ph), 5.82 (1H, s, H-4), 6.45 (2H, d, *J* = 13Hz, H-1,2), 6.86 (6H, m, Arom-H), 7.57 (2H, d, *J* = 13Hz, H-6,7) [Nurfina et al. *Eur J Med Chem* **32** 321 1997, DOI: 10.1016/S0223-5234(97)89084-8].

5'-Methoxycurcumin [1-(4-hydroxy-3,5-dimethoxyphenyl)-7-(4-hydroxy-3-methoxyphenyl)-(1*E*,6*E*)-1,6-hepta-diene-3,5-one] *M* 398.4, has *m* 145-146° (from MeOH), and was isolated as a yellow powder from *C. xanthorrhiza* with UV λ_{\max} at 429nm in MeOH.

1-Hydroxycurcumin [1-hydroxy-1,7-bis(4-hydroxy-3-methoxyphenyl)-(6*E*-6-heptene-3,5-dione] *m* 84-88° was isolated from the same source, had UV with λ_{\max} at 372nm in MeOH and had a weaker antioxidant activity than the other curcumins towards the autoxidation of linoleic acid in a water-alcohol system [Masuda et al. *Phytochemistry* **31** 3645 1992, DOI: 10.1016/0031-9422(92)83748-N]. An optically active form, (**1*ξ***)-1-hydroxy-1,7-bis(4-hydroxy-3-methoxyphenyl)-(1*E*,6*E*)-1,6-heptadiene-3,5-one, has *m* 92.0-96.0°, with [α]_D +12.2 (c 0.06, EtOH, configuration unknown), *M* 386.4. Its UV has λ_{\max} nm (ϵ)(MeOH) at 230 (sh), 260 (21000), 283 (18000), 370 (56000); and its IR (CDCl₃) has ν_{\max} at 965, 1000, 1120, 1260, 1325, 1375, 1460, 1560, 1600, 1620, 1650, 2850, 2900, 3525 cm^{-1} . Its ¹H NMR [270MHz, CDCl₃] has δ at 2.65(1H, dd, *J* = 17, 3.5Hz), 2.93 (1H, dd, *J* = 17, 14Hz), 3.93 (3H, s), 3.95 (3H, s), 5.39 (1H, dd, *J* = 14, 3.5Hz), 5.60 (1H, s), 5.80 (1H, br s, disappeared on addition of D₂O), 5.86 (1H, br s, disappeared on addition of D₂O), 6.44 (1H, d, *J* = 16Hz), 6.90 (1H, d, *J* = 8Hz), 7.00 (4H, br s), 7.03 (1H, dd, *J* = 8, 2Hz), 7.30 (1H, d, *J* = 16Hz) and *R*_F (Silica gel 60 F₂₅₄, C₆H₆*/EtOAc 1:1) of 0.2. When this **hydroxyheptadiene** was heated at 130°/15mmHg/10 hours it gave **curcumin** in 79% yield. (**3*S*,5*S***)-1,7-Bis(4-hydroxy-3-methoxyphenyl)heptane-3,5-diol [colourless needles *m* 132-134°, with [α]_D -18.5 (c 0.26, EtOH), *R*_F 0.1] was also isolated from the rhizomes of *C. xanthorrhiza* (Zingiberaceae) and characterised; and (**3*R*,5*R***)-1-(4-hydroxyphenyl)-7-phenylheptane-3,5-diol

[colourless needles **m 109-111°**, with $[\alpha]_D^{25} +8.3$ (c 0.06, CDCl_3), R_F 0.3] was isolated from the rhizomes of *Alpinia officinarum* (Zingiberaceae) and characterised; the absolute configurations were determined from the circular dichroism (CD) spectra of the 3,5-bis-*p*-dimethylaminobenzoate esters of their methylated derivatives, and applying the *exciton chirality rule* [Uehara et al. *Chem Pharm Bull Jpn* **35** 3298 1987, DOI: org/10.1248/cpb.35.3298].

Desmethoxycurcumin [22608-11-3] $\text{C}_{20}\text{H}_{18}\text{O}_5$, **M 338.3**, **m 173-175°** (also reported are **168°** and **172°**) is a contaminant of curcumin, isolated from the same source as curcumin, and best purified by HPLC and crystallises from EtOH as an orange-yellow powder. Its UV-VIS has λ_{max} (log ϵ) (MeOH) at 419nm (4.706) and (40% aqueous THF) 425nm (4.806) with fluorescence maxima at λ_{excit} 428nm and λ_{emis} 505nm in 40% aqueous THF. The ^1H NMR [400MHz, CDCl_3] has δ at 5.97 (1H, s, H-4), 7.60 (2H, d, J = 16.0Hz, H-2,6), 6.69 (1H, d, J = 16.0Hz, H-1 or 7), 6.64 (1H, d, J = 16.0Hz, H-7 or 1), 7.34 (1H, d, J = 1.7Hz, H-2' or 2''), 6.90 (1H, d, J = 8.0Hz, H-2'' or 2'), 7.56 (1H, d, J = 8.0Hz, H-5'), 6.88 (1H, d, J = 8.0Hz, H-5''), 7.27 (1H, dd, J = 8, 1.7Hz, H-6'), 6.9 (1H, d, J = 8Hz, H-6''), 3.92 (3H, s, OCH_3) (from TMS); and the ^{13}C NMR [100MHz, CDCl_3] has δ at 101.6 (C-4), 184.4 and 184.5 (C-3,5), 121.1 and 122.3 (C-2,6), 141.4 and 141.0 (C-1,7), 128.2 and 127.7 (C-1',1''), 111.5 and 130.9 (C-2',2''), 148.8 and 116.8 (C-3',3''), 150.0 and 160.5 (C-4',4''), 116.2 and 116.8 (C-5',5''), 123.8 and 130.9 (C-6',6''), 56.3 (OCH_3) (from TMS).

Similarly **bis-desmethoxy-curcumin** [22608-12-4, 24939-16-0] $\text{C}_{20}\text{H}_{18}\text{O}_5$, **M 308.3**, **m 224°** (also reported are **216-218°** and **222°**) is a contaminant of curcumin, occurring in the same source, best purified by HPLC and crystallises as the *hydrate* in yellow plates from EtOH. Its UV has λ_{max} (log ϵ) (MeOH) at 414nm (4.675) and (40% aqueous THF) 420nm (4.702) with fluorescence maxima at λ_{excit} 425nm and λ_{emis} 501nm in 40% aqueous THF. The ^1H NMR [400MHz, CDCl_3] has δ at 6.03 (1H, s, H-4), 16.4 (1H, br s, 3-OH), 7.56 (2H, d, J = 15.9Hz, H-2,6), 7.56 (2H, d, J = 15.9Hz, H-1,7), 6.84 (2H, d, J = 8.2Hz, H-2',2''), 7.56 (2H, d, J = 8.2Hz, H-3',3''), 10.03 (2H, s, 4',4''-OH), 7.56 (2H, d, J = 8.2Hz, H-5',5''), 8.64 (2H, d, J = 8.2Hz, H-6',6'') (from TMS); and the ^{13}C NMR [100MHz, CDCl_3] has δ at 100.9 (C-4), 183.2 (C-3,5), 120.8 (C-2,6), 140.3 (C-1,7), 125.8 (C-1',1''), 130.3 (C-2',2''), 115.9 (C-3',3''), 159.8 (C-4',4''), 115.9 (C-5',5''), 130.3 (C-6',6'') (from TMS). [Jayaprakasha et al. *J Agric Food Chem* **50** 3668 2002, DOI: 10.1021/jf025506a; Inoue et al. *J Agric Food Chem* **56** 9328 2008, DOI: 10.1021/jf801815n, (HPLC, UV, fluorescence) Rouseff *J Food Sci* **53** 1823 1988, DOI: 10.1111/j.1365-2621.1988.tb07851.x; (MS) Jiang et al. *Rapid Commun Mass Spectrom* **20** 1001 2006, DOI: 10.1002/rcm.2401; (HPLC, MS) Jiang et al. *J Chromatogr A* **1111** 21 2006, DOI: 10.1016/j.chroma.2006.01.103; (HPLC, MS) Hiserodt et al. *J Chromatogr A* **740** 51 1996, DOI: 10.1016/0021-9673(96)00103-3; (spectroscopy) Haukvik et al. *Pharmazie* **65** 600 2010, DOI: 10.1691/ph.2010.0048; (IR) Tanaka et al. *J Agric Food Chem* **56** 8787 2008, DOI: 10.1021/jf801338e; (spectroscopy) Péret-Almeida et al. *Food Research International* **38** 1039 2005, DOI: 10.1016/j.foodres.2005.02.021; (cytotoxicity) Ishida et al. *Bioorg Med Chem* **10** 3481 2002, DOI: 10.1016/S0968-0896(02)00249-3; (anti-inflammatory, antioxidant and anti-amyloid activity) Yang et al. *J Biol Chem* **280** 5892 2005, PMID: 15590663.]

The keto-enol tautomerism of curcumin derivatives was shown to be a property that can be employed for amyloid detection in *Alzheimer's disease*, and may be exploited for the design of amyloid-binding agents for the therapy of the disease [Yanagisawa et al. *Biomaterials* **31** 4179 2010, DOI: 10.1016/j.biomaterials.2010.01.142]. Over the years, curcumin derivatives have been screened for a variety of diseases (see above) including anti-HIV, and although few have displayed very potent anticancer activity, they are still being considered for development as future anticancer agents. A review on their potential as anticancer agents has been published recently. [Agrawal & Mishra *Med Res Rev* **30** 818 2010, DOI: 10.1002/med.20188.] The above information under *Curcumin* was kindly supplied by Prof. Hiroyasu Taguchi, Shiga University of Medical Science, Molecular Neuroscience Research Centre, JAPAN.

Cytidine [65-46-3] $\text{C}_9\text{H}_{13}\text{N}_3\text{O}_5$, **M 243.2**, **m 210-220°(dec)**, **230° (dec)**, **251-252° (dec)**, $[\alpha]_{546}^{20} +37$ (c 9, H_2O), $[\alpha]_D^{20} +29$ (c 9, H_2O), **pK²⁵ 3.85**. Cytidine crystallises from 90% aqueous EtOH. It has also been converted to *sulfate* by dissolving (~200mg) in a solution of EtOH (10ml) containing H_2SO_4 (50mg), whereby the salt crystallises out. It is collected, washed with EtOH and dried for 5 hours at 120°/0.1mm. The *sulfate* has **m 225°**. The *free base* is obtained by shaking the salt solution with a weak ion-exchange resin, filtering, evaporating and recrystallising the residue from EtOH as before. [Fox & Goodman *J Am Chem Soc* **73** 3256 1956, DOI: 10.1021/ja01151a076; Fox & Shugar *Biochim Biophys Acta* **9** 369 1952, DOI: 10.1016/0006-3002(52)90181-9; see Prytsas & Sorm in *Synthetic Procedures in Nucleic Acid Chemistry* (Zorbach & Tipson Eds) **Vol 1** 404 1973.] [Beilstein **25** III/IV 3667.] It is a component of RNA.

Cytochalasin B (from dehydrated mould matter, Phomin) [14930-96-2] $C_{29}H_{37}NO_5$, M 479.6, m 218-221°, 215-223°. Purify it by MeOH extraction, reverse phase C18 silica gel batch extraction by selective elution with 1:1 v/v hexane/tetrahydrofuran, crystallisation, subjected to TLC and recrystallisation [Lipski et al. *Anal Biochem* **161** 332 1987, DOI: 10.1016/0003-2697(87)90459-3]. It is soluble in EtOH (3.6%), Me₂CO (1%), Me₂SO (37%) and Me₂NCHO (49%) at 24°, and can be recrystallised from the first two solvents. It interferes with cellular movement [Korn *Physiol Reviews* **62** 672 1982]. For synthesis see Haidle & Myers [*Proc Natl Acad Sci* **101**(33) 12048 2004, DOI: 10.1073/pnas.0402111101, PMID: 15208404]. **VERY TOXIC.**

Cytosine (4-amino-2-hydroxypyrimidine) [71-30-7] $C_4H_5N_3O$, M 111.1, m 313° (hydrate, dec), 320-325° (anhydrous, dec), pK_1^{25} 4.6, pK_2^{25} 12.1. Cytosine crystallises from H₂O as the *monohydrate* which loses water on heating above 100°. Its solubility in H₂O is 0.77%. Its UV has λ_{max} at 267nm (ϵ 6,100) in H₂O pH 8.8 and 275nm (ϵ 10,400) in 0.1N HCl. [Hilbert & Johnson *J Am Chem Soc* **52** 1152 1930, DOI: 10.1021/ja01366a051; Hilbert et al. *J Am Chem Soc* **57** 552 1935, DOI: 10.1021/ja01306a050; *Beilstein* **25** III/IV 3654.]

Cytosine-1- β -O-arabinofuranoside (Cytarabine) [147-94-4] $C_9H_{13}N_3O_5$, M 243.2, m ~220°(dec), 212-213.5°, $[\alpha]_D^{20}$ +155 (c 1, H₂O), pK^{25} 4.3. Purify cytarabine by recrystallisation from aqueous EtOH or a large volume of H₂O (its solubility at ~20° is 5%). It has λ_{max} at 212 and 279nm at pH 2, and 272nm at pH 12. It is an acute *leukaemic* agent. (October 2007). It is used in myeloid leukemia chemotherapy [Pigneux et al. *Haematologica* **92**(10) 1327 2007, DOI: 10.3324/haematol.11068; PMID: 18024370]; and has *antiviral activity* in humans [Lauter et al. *Antimicrob Agents Chemother* **6**(5) 598 1974, DOI: 10.1128/aac.6.5.598; PMID: 15825312; Walwick et al. *Proc Chem Soc (London)* **73** 84 1959, DOI: 10.1039/PS9590000073; *Beilstein* **25** III/IV 3669.]

Demeclocycline hydrochloride (7-chloro-6-demethyltetracycline hydrochloride, Clortetrin) [127-33-3 (free base), 64-73-3 HCl] $C_{21}H_{21}N_2O_8 \cdot HCl$, M 501.3, m 174-178°(dec, for sesquihydrate), $[\alpha]_D^{25}$ -258 (c 0.5, 0.1N H₂SO₄), pK^{25} 4.45 [H₂O-Me₂NCHO (1:1)]. Crystallise the salt from EtOH/Et₂O or H₂O and dry it in air [McCormick et al. *J Am Chem Soc* **79** 4561 1957, DOI: 10.1021/ja01573a089; Dobrynin et al. *Tetrahedron Lett* 901 1962, DOI: 10.1016/S0040-4039(00)70561-2]. [*Beilstein* **14** IV 2625.]

2'-Deoxyadenosine (adenine 2'-deoxyriboside) [16373-93-6] $C_{10}H_{13}N_5O_3$, M 251.2, m 187-189°, 189-191°, $[\alpha]_D^{20}$ -25 (c 0.5, H₂O), $[\alpha]_{589}^{25}$ -26, $[\alpha]_{310}^{25}$ -206 (c 0.5, H₂O), pK^{20} 3.79. Purify it by recrystallisation from H₂O (hydrated crystals; solubility of the *monohydrate* is 1.1% in H₂O at 20°). It has UV with λ_{max} at 258nm (pH 1), 260nm (pH 7) and 261nm (pH 13). [Ness & Fletcher *J Am Chem Soc* **81** 4752 1959, DOI: 10.1021/ja01526a083; Walker & Butler *Can J Chem* **34** 1168 1956, DOI: 10.1139/v56-153.] The 3',5'-*O*-diacetyl derivative has m 151-152° (from EtOAc/petroleum ether). [*Beilstein* **26** III/IV 3589.]

3'-Deoxyadenosine (Cordycepin, adenine 3'-deoxyriboside) [73-03-0] $C_{10}H_{13}N_5O_3$, M 251.2, m 225-226°, 225-229°, $[\alpha]_D^{20}$ -47 (H₂O), $pK_{Est} \sim 4.8$. 3'-Deoxyadenosine forms needles from EtOH, *n*-BuOH and *n*-PrOH, and a *monohydrate* from H₂O. It has UV with λ_{max} at 260nm (ϵ 14,600) in EtOH. The *picrate* has m 195°(dec, yellow crystals from H₂O). [Kaczka et al. *Biochim Biophys Res Commun* **14** 456 1964, DOI:10.1016/0006-291X(64)90086-5; Todd & Ulbricht *J Chem Soc* 3275 1960, DOI: 10.1039/JR9600003275; Lee et al. *J Am Chem Soc* **83** 1906 1961, DOI: 10.1021/ja01469a030; Walton et al. *J Am Chem Soc* **86** 2952 1964, DOI: 10.1021/ja01068a049; for synthesis see Norman & Reese *Synthesis* 304 1983, *Beilstein* **26** III/IV 3594.]

2'-Deoxycytidine monohydrate [951-77-9, 652157-52-3 (hydrate), 207121-53-7] $C_9H_{13}N_3O_4 \cdot H_2O$, M 245.2, m 119-200°, 207-209°, 213-215°, $[\alpha]_D^{25}$ +78 (c 0.4, N NaOH), $[\alpha]_D^{23}$ +57.6 (c 2, H₂O), pK^{25} 4.25. Purify 2'-deoxycytidine by recrystallisation from MeOH/Et₂O or EtOH and dry it in air. [For NMR see Miles *J Am Chem Soc* **85** 1007 1963, DOI: 10.1021/ja00890a043; for UV see Fox & Shugar *Biochim Biophys Acta* **9** 369 1952, DOI: 10.1016/0006-3002(52)90181-9.] The *hydrochloride* crystallises from H₂O/EtOH and has m 174°(dec, 169-173°). [Walker & Butler *Can J Chem* **34** 1168 1956, DOI: 10.1139/v56-153.] The *picrate* has m 208°(dec). [Fox et al. *J Am Chem Soc* **83** 4066 1961, DOI: 10.1021/ja01480a027; *Beilstein* **25** III/IV 3662.]

2'-Deoxycytidine 5'-monophosphoric acid (deoxycytidylic acid) [1032-65-1] $C_9H_{14}N_3O_7P \cdot H_2O$, M 325.2,

m 170-172°(dec), 179-180°, 183-184°(dec), 183-187°(dec), $[\alpha]_D^{21} +35$ (c 0.2, H₂O), pK₁ 4.6, pK₂ 6.6. Recrystallise the acid from H₂O or aqueous EtOH and dry it in a vacuum. [Volkin et al. *J Am Chem Soc* **73** 1533 1951 DOI: 10.1021/ja01148a037; for UV see Fox et al. *J Am Chem Soc* **75** 4315 1953, DOI: 10.1021/ja01113a051; IR: Michelson & Todd *J Chem Soc* 34 1954, DOI: 10.1039/JR9540000034; *Beilstein* **25** IV 3664.]

2'-Deoxyguanosine monohydrate (9-[2-deoxy-β-D-ribofuranosyl]guanidine) [961-07-9, 312693-72-4] **C₁₀H₁₃N₅O₄ · H₂O, M 285.3, m ca 200°(dec), $[\alpha]_D^{20} +37.5$ (c 2, H₂O), $[\alpha]_D^{14} -47.7$ (c 0.9, N NaOH), pK_{Est(1)} ~ 3.3, pK_{Est(2)} ~ 9.2.** 2'-Deoxyguanosine recrystallises from H₂O as the *monohydrate*. [Brown & Lythgoe *J Chem Soc* 1990 1950, DOI: 10.1039/JR9500001990; Levene & London *J Biol Chem* **81** 711 1929, <http://www.jbc.org/content/81/3/711>; Levene & London *J Biol Chem* **83** 793 1929, <http://www.jbc.org/content/83/3/793>; UV: Hotchkiss *J Biol Chem* **175** 315 1948, PMID: 18873306; ORD: Levendahl & James *Biochim Biophys Acta* **26** 89 1957, DOI: 10.1016/0006-3002(57)90058-6.] The **3',5'-di-O-acetyl derivative** crystallises from aqueous EtOH with **m 222°(dec)**, and $[\alpha]_D^{18} -38$ (c 0.3, 10% aqueous EtOH) [Hayes et al. *J Chem Soc* 808 1955, DOI: 10.1039/JR9550000808]. [*Beilstein* **26** III/IV 3897.]

2'-Deoxyinosine [890-38-0] **C₁₀H₁₂N₄O₄, M 252.2, m 206°(dec), 218-220°(dec), $[\alpha]_D^{25} -21$ (c 2, N NaOH), $[\alpha]_D^{21.5} -21$ (c 1, H₂O), pK_{Est(1)} ~ 8.9, pK_{Est(2)} ~ 12.4.** Purify 2'-deoxyinosine by recrystallisation from H₂O. Store it at -20°. [Brown & Lythgoe *J Chem Soc* 1990 1950, DOI: 10.1039/JR9500001990; UV: Macnutt *Biochem J* **50** 384 1952, DOI: 10.1042/bj0500384; *Beilstein* **26** III/IV 2086.]

5-Deoxy-5-(methylthio)adenosine [2457-80-9] **C₁₁H₁₅N₅O₃S, M 297.3, m 210-213°(dec), 211°, 212°, 213-214°, $[\alpha]_D^{20} -23.7$ (c 0.02, pyridine), $[\alpha]_D^{20} -8$ (c 1, 5% aqueous NaOH), $[\alpha]_D^{25} +15$ (c 0.4-1.0, 0.3N aqueous AcOH), pK_{Est} ~3.5.** Recrystallise it from H₂O and sublime it at 200°/0.004mm. [v. Euler & Myrbäck *Hoppe Seyler's Z physiol Chem* **177** 237 1928, Weygand & Trauth *Chem Ber* **84** 633 1951, DOI: 10.1002/cber.19510840714; Baddiley et al. *J Chem Soc* 2662 1953, DOI: 10.1039/JR9530002662.] The *hydrochloride* has **m 161-162°** [Kuhn & Henkel *Hoppe Seyler's Z Physiol Chem* **269** 41 1941]. The *picrate* has **m 183°(dec)** (from H₂O). [*Beilstein* **26** III/IV 3675.]

Deoxyribonucleic acid (from plasmids). Purify plasmid DNA by two buoyant density ultracentrifugations using ethidium bromide-CsCl. The fluorescent band of the desired DNA is collected. The ethidium bromide is extracted with Et₂O, and the DNA is dialysed against buffered EDTA and lyophilised. The purity is then assessed by PAGE (polyacrylamide gel electrophoresis). [Marmur & Doty *J Mol Biol* **5** 109 1962, DOI: 10.1016/S0022-2836(62)80066-7; Guerry et al. *J Bacteriol* **116** 1064 1973, PMID: 4583233.] See Introduction.

3'-Deoxythymidine {2',3'-dideoxythymidine, 1-[(2*r*)-5*c*-hydroxymethyltetrahydro(2*r*)-furyl]-5-methylpyrimidine-2,4-dione} [3416-05-5] **C₁₀H₁₄N₂O₄, M 226.2, m 145°, 149-151°, 155-156°, $[\alpha]_D^{26} +18$ (+20) (c 1, H₂O), pK_{Est} ~ 9.2.** Crystallise it from Me₂CO/MeOH. [Michelson & Todd *J Chem Soc* 816 1955, DOI: 10.1039/JR9550000816; *Beilstein* **24** III/IV 1297.] Used in DNA sequencing, inhibits at thymidine residues.

2'-Deoxyuridine [1-(β-D-erythro-2-deoxypentofuranosyl)-1*H*-pyrimidine-2,4-dione] [951-78-0] **C₉H₁₂N₂O₅, M 228.2, m 163°, 163-163.5°, 165-167° 167°, $[\alpha]_D^{26} +30$ (c 2, H₂O), $[\alpha]_D^{22} +50$ (c 1, N NaOH), pK₂₅ 9.3.** 2'-Deoxyuridine forms needles from absolute EtOH or 95% EtOH, and is dried *in vacuo*. [Dekker & Todd *Nature* **166** 557 1950, DOI: 10.1038/166557b0; Brown et al. *J Chem Soc* 3035 1958, DOI: 10.1039/JR9580003035; NMR Jardetzky *J Am Chem Soc* **83** 2919 1961, DOI: 10.1021/ja01474a030; Fox & Shugar *Biochim Biophys Acta* **9** 369 1952, DOI: 10.1016/0006-3002(52)90181-9; UV: MacNutt *Biochem J* **50** 384 1952, DOI: 10.1042/bj0500384; *Beilstein* **24** III/IV 1200.]

3'-Deoxyuridine {1-[(2*R*)-5*c*-hydroxymethyltetrahydro(2*r*)furyl]-5-methylpyrimidin-2,4-dione, 2',3'-dideoxythymidine} [7057-27-4, 3416-05-5] **C₉H₁₂N₂O₅, M 228.2, m 149-151°, $[\alpha]_D^{20} +18$ (c 1, H₂O), pK_{Est} ~ 9.3.** 3'-Deoxyuridine is recrystallised from Me₂CO/MeOH and is dried in a vacuum. [Michelson & Todd *J Chem Soc* 816 1955, DOI: 10.1039/JR9550000816.]

Desthiobiotin (4*R*-cis-5-methyl-2-oxo-4-imidazolidinehexanoic acid) [533-48-2] **C₁₀H₁₈N₂O₃, M 214.3, m**

156-158°, $[\alpha]_D^{20} +10.5$ (c 2, H₂O), $pK_{Est} \sim 2.8$. Dissolve desthiobiotin in 0.5% Na₂CO₃, filter, acidify with HCl to Congo Red, concentrate to a small volume (2-3 ml) to give fine needles, filter it off and recrystallise it twice from H₂O, **m 157-158°**. It also crystallises from 95% EtOH. The **methyl ester** crystallises from MeOH and sublimes at 100°/high vacuum, **m 69-70°**, with $[\alpha]_D^{27.5} +2.6$ (c 2, CHCl₃). [Melville et al. *Science* **98** 497 1943, DOI: 10.1126/science.98.2553.497; *J Am Chem Soc* **66** 1422 1944, DOI: 10.1021/ja01236a505; *Beilstein* **25** III/IV 1543.]

Di- and tri-carboxylic acids. These are separated by anion-exchange chromatography. [Bengtsson & Samuelson *Anal Chim Acta* **44** 217 1969, DOI: 10.1016/S0003-2670(01)81756-6.]

7,8-Dihydrofolic acid (7,8-dihydropteroyl-L-glutamic acid, DHFA) [4033-27-6] **C₁₉H₂₁N₇O₆, M 443.4, pK_1 2.0 (basic 10-NH), pK_2 2.89 (2-NH₂), pK_3 3.45 (α -CO₂H), pK_4 4.0 (basic 5N), pK_5 4.8 (γ -CO₂H), pK_6 9.54 (acidic 3NH).** DHFA is best purified by suspending (1g mostly dissolved) in ice-cold sodium ascorbate (300ml of 10% at pH 6.0, prepared by adjusting the pH of 30g of sodium ascorbate in 150ml of H₂O by adding 1N NaOH dropwise using a glass electrode till the pH is 6.0). This gives a clear solution at pH ~5. While stirring at 0°, add N HCl dropwise slowly (0.1ml/min) until the pH drops to 2.8 when white birefringent crystals separate. These are collected by centrifugation (1000xg for 5 minutes), washed 3x with 0.001N HCl also by centrifugation and decantation. The residue is then dried in a vacuum (0.02mm) over P₂O₅ (change the P₂O₅ frequently at first) and KOH at 25° in the dark. After 24 hours the solid reaches constant weight.

For the **assay of dihydrofolate reductases** (see [9002-03-3] 'Proteins, Enzymes...' in this Chapter): suspend ~66.5mg of DHFA in 10ml of 0.001M HCl containing 10mM dithiothreitol (**DTT** stock made from 154mg in 10ml H₂O making 0.1M), shake well and freeze in 400 μ l aliquots. Before use, mix 400 μ l of this suspension with 0.1M **DTT** (200 μ l, also made in frozen aliquots), and the mixture is diluted with 200 μ l of 1.5M Tris-HCl pH 7.0 and 1.2ml of H₂O (making a total volume of ~2ml) to give a clear solution. To estimate the concentration of DHFA in this solution, dilute 20 μ l of this solution to 1ml with 0.1M Tris-HCl pH 7.0 and read the OD at 282nm in a 1cm path length cuvette. ϵ at 282nm is 28,000M⁻¹cm⁻¹. [Futterman *Methods Enzymol* **6** 801 1963, DOI: 10.1016/0076-6879(63)06252-2; Reyes & Rathod *Methods Enzymol* **122** 360 1986, DOI: 10.1016/0076-6879(86)22194-1; *Beilstein* **26** III/IV 3934.]

DL-erythro-Dihydrosphingosine (dl-erythro-2-aminooctadecan-1,3-diol) [3102-56-5] **C₁₈H₃₉NO₂, M 301.5, m 85-86°, 85-87°, $pK_{Est} \sim 8.8$.** Purify it by recrystallisation from petroleum ether/EtOAc or CHCl₃. The (\pm)-**N-dichloroacetyl derivative** has **m 142-144°** (from MeOH). [Shapiro et al. *J Am Chem Soc* **80** 2170 1958, DOI: 10.1021/ja01542a034; Shapiro & Sheradsky *J Org Chem* **28** 2157 1963, DOI: 10.1021/jo01043a514.] The **D-isomer** [764-22-7] crystallises from petroleum ether/Et₂O and has **m 78.5-79°, $[\alpha]_{546}^{28} +6$ (CHCl₃/MeOH, 10:1);** its solubility in EtOH or in Me₂SO is 25mg/ml (warm). [Grob & Jenny *Helv Chim Acta* **35** 2106 1952, DOI: 10.1002/hlca.19520350641; Jenny & Grob *Helv Chim Acta* **36** 1454 1953, DOI: 10.1002/hlca.19530360632; *Beilstein* **4** I 448, **4** II 757, **4** III 854, **4** IV 1887.]

Dihydrostreptomycin sesquisulfate [5490-27-7] **C₂₁H₄₁N₇O₁₂. 3/2H₂SO₄, M 730.7, m 250°(dec), 255-265°(dec), $[\alpha]_D^{20} -92.4$ (c 1, H₂O), $pK_{Est(1)} \sim 9.5$ (NMe), $pK_{Est} \sim 13.4$ (guanidino).** The antibiotic crystallises from H₂O, MeOH, *n*-BuOH or methyl ethyl ketone. The crystals are not hygroscopic like the amorphous powder; however, both forms are soluble in H₂O but the **amorphous solid** is about 10 times more soluble than the crystals. The **free base** [128-46-1] **C₂₁H₄₁N₇O₁₂, M 583.6,** also crystallises from H₂O/Me₂CO and has $[\alpha]_D^{26} -92$ (a strong base, with aqueous solution pH 7.0, see pK_a). [Solomons & Regna *Science* **109** 515 1949, DOI: 10.1126/science.109.2838.515; Wolf et al. *Science* **109** 515 1949, DOI: 10.1126/science.109.2838.515-a; McGilveray & Rinehart *J Am Chem Soc* **77** 4003 1956, DOI: 10.1021/ja01095a054; Gardin et al. *J Am Vet Med Assoc* **183**, 434 1983, PMID: 6618969]. [*Beilstein* **18** III/IV 7538.] Active against bacteria and mycoplasma.

2,4-Dihydroxyimidazole (hydantoin, imidazolin-2,4-dione) [461-72-3] **C₃H₄N₂O₂, M 100.1, m 215°, 218-220°, 220°, pK^{25} 9.15, pK^{18} 9.16.** Well over the past hundred years many syntheses of hydantoin have been reported involving the reaction of alloxanic acid with HI [Baeyer *Justus Liebigs Ann Chem* **119** 126 1861, DOI: 10.1002/jlac.18611190115; and **130** 129 159 1864, DOI: 10.1002/jlac.18641300202], the cyclisation of urea derivatives [Harries & Weiss *Chem Ber* **33** 3418 1900, DOI: 10.1002/cber.190003303122; and *Justus Liebigs Ann Chem* **327** 355 366 1903, DOI: 10.1002/jlac.19033270307], the cyclisation of ethyl *N*-carbamoylglycinate,

using NaOMe/MeOH in 85% yield [Kaválek et al. *Coll Czech Chem Commun* **51** 375 1986, DOI: org/10.1135/cccc19860375], and more recently by the acid catalysed condensation of glycine with KNCO [El-Deeb et al. *Eur J Med Chem* **45** 2516 2010, DOI: 10.1016/j.ejmech.2010.02.038]. In a later patent, glycine (75g, 1 mole) and urea (140g, 2.3 moles) in H₂O (120ml) were stirred under reflux for 12 hours. The mixture was acidified by careful addition of concentrated H₂SO₄ while cooled in an ice bath, then boiled for 1 hour, cooled to 0-5°, the crystals were filtered off, washed with cold H₂O and dried. Recrystallisation from AcOH (10 parts) gave **pure hydantoin** (71g, 70%) m 220-220°. [Alkaloida US patent 4647694 1987.] Also purified from 2.5 parts of hot H₂O (charcoal), filtering, acidifying to pH ~6, filtering again, collecting the solid, washing with ice cold H₂O, drying, and recrystallising from EtOH (needles) or 10 parts of AcOH. Its solubility is 39.7g/L in H₂O and 3.24g/L in EtOH at 25°. Its UV has λ_{\max} (ϵ) at 215(400) and 245(60)nm in 0.01N EtOH/NaOH; 215(360) and 245(50)nm in EtOH/H⁺; and IR has ν_{\max} at 1697 (CO) and 1776 (CO) cm⁻¹ in a paraffin mull [West *J Biol Chem* **34** 187 1918, <http://www.jbc.org/content/34/1/187>; Crombie & Hooper *J Chem Soc* 3010 1955, DOI: 10.1039/JR9550003010; *Beilstein* **24** H 242, **24** I 287, **24** II 127, **24** III/IV 1034].

1,2-Dilauroyl-sn-glycero-3-phosphoethanolamine (\pm -dilauroyl- α -cephalin, **3-sn-phosphatidylethanolamine 1,2-didodecanoyl**) [59752-57-7] C₂₉H₅₈NO₈P, **M 579.8**, **m 210°**, **pK_{Est(1)} ~ 5.8 (PO₄H)**, **pK_{Est(2)} ~ 10.5 (NH₂)**. Recrystallise it from EtOH or tetrahydrofuran. Dry it in a vacuum and store it at -20°. [Bevan & Malkin *J Chem Soc* 2667 1951, DOI: 10.1039/JR9510002667; IR: Bellamy & Beecher *J Chem Soc* 728 1953, DOI: 10.1039/JR9530000728; *Beilstein* **4** IV 1417.]

3,7-Dimethyl-2,6-dioxopurine (Theobromine) [83-67-0] C₇H₈N₄O₂, **M 180.2**, **m 337°** (sublimes slowly at **290°** and finally melts at **~351°**), **pK₁⁴⁰ -0.16**, **pK₁²⁵ 9.96**. Recrystallise it from hot H₂O (solubility in w/v% is 0.06 at 15°; 1.25 at 100°). Forms salts with heavy metals, is a diuretic, vasodilator and cardiac stimulant. [Lister *Purines Part II. Fused Pyrimidines*, 1971, J.Wiley NY, ISBN 047138205-1; *Beilstein* **26** III/IV 2336.]

1,2-Dimyristoyl-sn-glycero-3-phosphocholine monohydrate (dimyristoyl-L- α -lecithin) [18194-24-6 (1H₂O)] C₃₆H₇₂NO₈P, **M 696.0 (H₂O)**, **[α]_D²⁴ +7 (c 8, EtOH-CHCl₃ 1:1 for α_1 form)**, **pK_{Est} ~ 5.8 (PO₄)**. It has three forms α_1 , α_2 and β . Recrystallise it from aqueous EtOH or EtOH/Et₂O. Its solubility at 22-23° in Et₂O is 0.03%, in Me₂CO it is 0.06% and in pyridine it is 1.3%. [Baer & Kates *J Am Chem Soc* **72** 942 1950, DOI: 10.1021/ja01158a078; Baer & Maurukas *J Am Chem Soc* **74** 158 1952, DOI: 10.1021/ja01121a039; for IR see Marinetti & Stotz *J Am Chem Soc* **76** 1347 1954, DOI: 10.1021/ja01634a047; *Beilstein* **4** IV 1463.] The **S-isomer** with 1H₂O recrystallises from 2,6-dimethylheptan-4-one with **m 226-227°** (sintering at 90-95°), and **[α]_D²⁰ -7 (c 6, MeOH/CHCl₃ 1:1)**. Store at -20°. [Baer & Martin *J Biol Chem* **193** 835 1951, PMID: 14907771.] (\pm)-**1,2-Dimyristoyl-sn-glycero-3-phosphoethanolamine** (dimyristoyl- α -cephalin) [998-07-2] C₃₃H₆₆NO₈P, **M 635.9**, **m 207°**, **pK_{Est(1)} ~ 5.8 (PO₄H)**, **pK_{Est(2)} ~ 10.5 (NH₂)**. Recrystallise the cephalin from EtOH [Bevan & Malkin *J Chem Soc* 2667 1951, DOI: 10.1039/JR9510002667]. The **R-isomer** has **m 195-196°** (sintering at 130-135°) after recrystallisation from CHCl₃/MeOH, and **[α]_D²⁶ +6.7 (c 8.5, CHCl₃/AcOH 9:1)**. Store it at -20°. [Baer *Can J Biochem Physiol* **35** 239 1957, DOI: 10.1139/o57-030; Baer et al. *J Am Chem Soc* **74** 152 1952, DOI: 10.1021/ja01121a038; *Beilstein* **4** IV 1463.]

S(-)-1,2-Dipalmitin (**S-1,2-dipalmitoyl-sn-glycerol**) [30334-71-5] C₃₅H₆₈O₅, **M 568.9**, **m 65-69°**, **68-69°** **[α]_D²⁰ -2.9 (c 8, CHCl₃)**, the **R(+)-isomer (2,3-dipalmitoyl-sn-glycerol)** [6076-30-8] has **m 67.5-68.2°**, **[α]_D²⁰ -2.8 (c 8, CHCl₃)**. Crystallise S(-)-1,2-dipalmitin from chloroform/petroleum ether (b 40-60°) ~1:1.5. [For the S(-)-isomer see Baer & Kates *J Am Chem Soc* **72** 942 1950, DOI: 10.1021/ja01158a078; Hanahan & Vercamer *J Am Chem Soc* **76** 1804 1954, DOI: 10.1021/ja01636a021; and for the R(+)-isomer see Tatttrie et al. *Arch Biochem* **78** 319 1958, DOI: 10.1016/0003-9861(58)90355-2; *Beilstein* **2** IV 1173.] The **racemate** [40290-32-2] is polymorphic with different IR spectra. Crystallisation from hexane, or other solvents, gives the higher melting form, **m 71.5-72.5°**. The melt then solidifies to the lower melting **α -form**, **m 49.7-50°**. The **β -form** has **m 61°** (**65-66°** is also reported). Keeping the lower melting forms at their melting temperatures for a while, converts to the higher melting form. Store them at -20°. [Howe & Malkin *J Chem Soc* 2663 1951, DOI: 10.1039/JR9510002663; Baer & Kates *J Am Chem Soc* **72** 942 1950, DOI: 10.1021/ja01158a078; *Beilstein* **2** IV 1173.]

R-Dipalmitoyl-sn-glycero-3-phosphatidic acid [7091-44-3] C₃₅H₆₉O₈P, **M 648.9**, **[α]_D²⁶ +4 (c 10, CHCl₃)**, **pK_{Est(1)} ~ 1.6**, **pK_{Est(2)} ~ 6.1**. Recrystallise the acid from Me₂CO at low temperature. At 21° it is soluble in

*C₆H₆ (4.2%), petroleum ether (0.01%), MeOH (2%), EtOH (2.5%), AcOH (1.3%), Me₂CO (1.76%), and Et₂O (1.5%). [Baer *J Biol Chem* **189** 235 1951, PMID: 14832235.] The *sodium salt* [71065-87-7] decomposes on heating, and its solubility ratio in CHCl₃/MeOH/AcOH is 4:1:0.1. Store the acid and salt at -20°.

R-1,2-Dipalmitoyl-sn-glycero-3-phosphocholine monohydrate (dipalmitoyl- α -L-lecithin) [63-89-8] C₄₀H₈₀NO₈P. H₂O, M 752.1, m sinters at 120°, [α]_D²⁵ +7.0 (c 5.6, absolute CHCl₃), pK_{Est} ~ 5.8 (PO₄). It has three crystalline forms α_1 , α_2 and β' which changes at 60-70° and at 229°, respectively. In order to obtain a fine powder, ~2 g are dissolved in CHCl₃ (15ml) and petroleum ether (b 35-60°) is added; the solution is evaporated to dryness *in vacuo* at <20° and then dried at 0.1mm over CaCl₂. Store at -20° in a sealed container. [Baer & Maurukas *J Am Chem Soc* **74** 158 1952, DOI: 10.1021/ja01121a039; Baer & Kates *J Biol Chem* **185** 615 1950, PMID: 14774404; Beilstein **4** IV 1463.]

dl- β -Dipalmitoylphosphatidyl choline [2797-68-4, 2644-64-6] C₄₀H₈₀NO₈P, M 734.0, m 230-233°, pK_{Est} ~ 5.8 (PO₄). Recrystallise the choline from chloroform and dry it for 48 hours at 10⁻⁵ Torr [O'Leary & Levin *J Phys Chem* **88** 1790 1984, DOI: 10.1021/j150653a025].

1,2-Distearoyl-sn-glycerol [1429-59-0, 10567-21-2] C₃₉H₇₆O₅, M 625.0. The *dl*-form recrystallises from CHCl₃/petroleum ether (b 40-60°), m 59.5° (α form) and 71.5-72.5° (β form). Recrystallisation from solvents such as EtOH, MeOH, toluene, Et₂O gives the higher melting form, and resolidification gives the lower melting form. [IR: Chapman *J Chem Soc* 4680 1958, DOI: 10.1039/JR9580004680; 2522 1956, DOI: 10.1039/JR9560002522.] The *S-isomer* is recrystallised from CHCl₃/petroleum ether and has m 76-77°, with [α]_D²⁴ -2.8 (c 6, CHCl₃). Store them at -20° in sealed containers. [Baer & Kates *J Am Chem Soc* **72** 942 1950, DOI: 10.1021/ja01158a078; Beilstein **2** IV 1231.]

1,2-Distearoyl-sn-glycero-3-phosphoethanolamine (distearoyl- α -cephalin) [1069-79-0] C₄₁H₈₂NO₈P, M 748.1, m 180-182° (*R*-form, sintering at 130-135°), m 196° (\pm form), pK_{Est(1)} ~ 5.8 (PO₄H), pK_{Est(2)} ~ 10.5 (NH₂). The *R*-form is recrystallised from CHCl₃/MeOH, and the \pm -form is recrystallised from EtOH. Store at -20°. [Bevan & Malkin *J Chem Soc* 2667 1951, DOI: 10.1039/JR9510002667; Baer *Can J Biochem* **52** 570 1974, DOI: 10.1139/o74-082; Baer *Can J Biochem Physiol* **34** 288 1956, DOI: 10.1139/y56-033; Beilstein **4** IV 1420.]

(-)-Domoic acid [4-(2-carboxyhexa-3,5-dienyl)-3-carboxymethylproline] [14277-97-5] C₁₅H₂₁NO₆, M 311.3, m 215°, 217°, b 607.2°/760mm, [α]_D²⁰ -108 (c 1, H₂O), pK₁ 2.20 (2-CO₂H), pK₂ 3.72 (CO₂H), pK₃ 4.93 (3-CH₂CO₂H), pK₄ 9.82 (NH). Domoic acid (~300 mg) is purified on a Dowex1 column (3.5 x 40 mm, 200-400 mesh, acetate form), washed with H₂O until neutral, then eluted with increasing concentrations of AcOH (8L) from 0 to 0.25M. The fraction containing domoic acid (in 50ml) is collected, evaporated to dryness under reduced pressure and recrystallised from aqueous EtOH. It is a *glutamate* and a *Kainate* receptor agonist. A kainic acid neurotoxin that causes shellfish poisoning-type *amnesia*. [Impellizzeri et al. *Phytochemistry* **14** 1549 1975, DOI: 10.1016/0031-9422(75)85349-0; Takemoto & Daigo *Arch Pharm* **293** 627 1960, DOI: 10.1002/ardp.19602930608; Beilstein **22/4** V 371.] For risk assessment of the amnesic shellfish poison, domoic acid, on animals and humans see Kumar et al. [*J Environ Biol.* **30**(3) 319 2009, PMID: 20120452].

Enniatin A [11113-62-5, 2503-13-1] C₃₆H₆₃N₃O₉, M 681.9, m 122-122.5°, [α]_D¹⁸ -92 (c 0.9, CHCl₃). It is a cyclic peptidic ester antibiotic which is recrystallised from EtOH/water but is deactivated in alkaline solution. It is soluble in MeOH (10mg/ml), EtOH (10mg/ml), Me₂SO (10mg/ml), in DMF and partly in H₂O. Store at -20°. [Ovchinnikov & Ivanov in *The Proteins* (Neurath and Hill eds) Academic Press, NY, Vol V pp. 365 and 516 1982; Sy-Cordero et al. *J Antibiot* **65** 541 2012, DOI: 10.1038/ja.2012.71.] Ammonium ion binding ionophore [Ovchinnikov et al. *Int J Peptide & Protein Res* **6**(6) 465-498 1974, DOI: 10.1111/j.1399-3011.1974.tb02407.x].

(-)-Ephedrine (1R,2S-2-methylamino-1-phenylpropanol) [299-42-3] C₁₀H₁₅NO, M 165.2, m ~34°, 36°, 38.1°, 40°, b 126-129°/7mm, 225-227°/760mm, d₄²² 1.0085, [α]_D²⁰ -47 and [α]_D²⁶ -42 (c 4, 3% HCl), [α]_D^{22.5} +15.1 (c 0.8, H₂O), -9.36 (c 3, MeOH), pK₂₂ 9.58 (pK₂₅ 8.84 in 80% aqueous methoxyethanol). Purify (-)-ephedrine by vacuum distillation (dehydrates) and forms waxy crystals or granules, and may pick up 0.5 H₂O.

The presence of H₂O raises its melting point to 40°. [Moore & Tabern *J Am Pharm Soc* **24** 211 1935, DOI: 10.1002/jps.3080240306.] The **anhydrous base** crystallises from dry ether [Fleming & Saunders *J Chem Soc* 4150 1955, DOI: 10.1039/JR9550004150]. It gradually decomposes on exposure to light and is best stored in an inert atmosphere in the dark (preferably at -20°). Its solubility in H₂O is 5%, in EtOH it is 1% and it is soluble in CHCl₃, Et₂O and mineral oils. It has pK_a values in H₂O of 10.25 (0°) and 8.69 (60°) [Everett & Hyne *J Chem Soc* 1636 1958, DOI: 10.1039/JR9580001636; Prelog & Häfliger *Helv Chim Acta* **33** 2021 1950, DOI: 10.1002/hlca.19500330708] and pK_a²³ 8.84 in 80% aqueous methoxyethanol [Simon *Helv Chim Acta* **41** 1835 1958, DOI: 10.1002/hlca.19580410639]. The **hydrochloride** has **m 220°** (from EtOH/Et₂O) and [α]_D²⁰ -38.8 (c 2, EtOH). [IR: Chatten & Levi *Anal Chem* **31** 1581 1959, DOI: 10.1021/ac60153a045.] The **anhydrous base** crystallises from Et₂O [Fleming & Saunders *J Chem Soc* 4150 1955, DOI: 10.1039/JR9550004150]. [Beilstein **13** H 373, **13**, III 1720, **13** IV 1879.] It is a **sympathomimetic** amine. [Kobayashi, et al. *Anesth Analg* **97** 1239 2003, DOI: 10.1213/01.ANE.0000092917.96558.3C].

(+)-Ephedrine hydrochloride (1S-2R-2-methylamino-1-phenylpropan-1-ol hydrochloride) [24221-86-1] C₁₀H₁₅NO. HCl, **M 201.7**, has **m 216-219°**, [α]_D²⁰ +34 (c 11.5, H₂O). Recrystallise the hydrochloride from EtOH/Et₂O. The **free base** crystallises from *C₆H₆ with **m 40-41°** (Skita et al. *Chem Ber* **66** 974 1933, DOI: 10.1002/cber.19330660713]. [Beilstein **13** H 254, **13** II 375, **13** III 1721, **13** IV 1879.]

(-)-Ephedrine hydrochloride (1R-2S-2-methylamino-1-phenylpropan-1-ol hydrochloride) [50-98-6] has **m 218°**, [α]_D²⁰ -48 (c 5, 2M HCl). It crystallises from water. See **free base** above. [Beilstein **13** H 254.]

α-Erythroidine (3R,5S,12S-1,2,6,7-Tetradehydro-12,17-dihydro-3β-methoxy-16(15H)-oxaerythrinan-15-one) [466-80-8] C₁₆H₁₉NO₃, **M 273.3**, **m 52-55°, 58-60°**, [α]_D²⁷ +136 (c 0.5, H₂O), pK_{Est(3)} ~7.4 Recrystallise α-erythroidine from pentane. It is best prepared freshly from the more stable hydrochloride. The **hydrochloride** (1.3g) in H₂O (20ml) is basified with NaHCO₃ to pH ~8 and extracted with *C₆H₆ (6 x 10ml). The combined extracts are evaporated to a small volume and refrigerated. The **free base** separates as white hygroscopic crystals **m 52-55°**, which can be recrystallised from pentane. Although stable in solution, the crystals turn brown on exposure to air.

α-Erythroidine hydrochloride is best purified by dissolving 10g in H₂O (100ml), adjusting the pH to 8 with aqueous NaHCO₃, extracting with *C₆H₆ (4 x 20ml), evaporating to 20ml, passing through activated Al₂O₃ and eluting with *C₆H₆. The eluate is evaporated to a small volume and the crystals are collected, dissolved in EtOH and dry HCl gas is passed through to give the **pure hydrochloride**. When recrystallised from EtOH, it has **m 226-228°(dec)**, and [α]_D³² +118 (c 0.5, H₂O). It has UV with λ_{max} at 224nm (ε 35,500); compare with β-erythroidine hydrochloride below. [Boelkeleide & Grundon *J Am Chem Soc* **75** 2563 1953, DOI: 10.1021/ja01107a006; for conversion of **α to β forms** see Boelkeleide & Morrison *J Am Chem Soc* **80** 3905 1958, DOI: 10.1021/ja01548a027; Hill & Scheerer *J Org Chem* **27** 921 1962, DOI: 10.1021/jo01050a056; Boelkeleide & Wenzinger *J Org Chem* **29** 1307 1964, DOI: 10.1021/jo01029a008; Beilstein **27** III/IV 3569.]

β-Erythroidine (3R,5S) [466-81-9] C₁₆H₁₉NO₃, **M 273.3**, **m 97-99°, 99.5-100°**, [α]_D²⁵ +89 (c 0.5, H₂O). Purify it like the **α-isomer** but recrystallise it from EtOH. The **free base** is unstable in air and light, but the hydrochloride is more stable and best stored as such. The **methiodide** has **m 211°** (prisms from EtOH).

β-Erythroidine hydrochloride can be obtained from the **α-isomeric salt** as follows: The α-hydrochloride (1.2g) in 10% aqueous NaOH (12ml) is refluxed for 3 hours under N₂, cooled in ice and concentrated HCl added to pH 2. After standing for 3 hours, NaHCO₃ is carefully added to pH 7, the solution is extracted with CHCl₃ (5x), and the extracts are dried, filtered, and concentrated to give an oil which on seeding gives β-erythroidine **m 97-99°**. When dissolved in EtOH and dry HCl gas is passed through the solution, the **pure β-erythroidine hydrochloride** crystallises out with **m 230.5-231.5°**, [α]_D²⁵ +10 (c 0.5, H₂O). It is a **muscle relaxant** with a curare-like-action and is **more active** than the α-isomer. [Koniuszy & Folkers *J Am Chem Soc* **72** 5579 1950, DOI: 10.1021/ja01168a061; Boelkeleide et al. *J Am Chem Soc* **75** 2550 1953, DOI: 10.1021/ja01107a004; Boelkeleide & Morrison *J Am Chem Soc* **80** 3905 1958, DOI: 10.1021/ja01548a027; Boelkeleide & Wenzinger *J Org Chem* **29** 1307 1964, DOI: 10.1021/jo01029a008; Berger & Schwartz *J Pharmacol Exp Therap* **93** 362 1948, PMID: 18882142; Beilstein **4** IV 3568.]

Erythromycin A [114-07-8] C₃₇H₆₇NO₁₃, **M 733.9**, **m 133-135°(dec), 135-140°, 137-140°**, [α]_D²⁰ -75 (c 2, EtOH), pK₂₅ 8.9. It is recrystallised from H₂O to form **hydrated** crystals which melt at **ca 135-140°**, resolidify and melt again at **190-193°**. The melting point after drying at 56°/8mm is that of the **anhydrous** material and is

at **137-140°**. Its solubility in H₂O is ~2mg/ml. The **hydrochloride** has **m 170°, 173°** (from aqueous EtOH, EtOH/Et₂O). [Flynn et al. *J Am Chem Soc* **76** 3121 1954, DOI: 10.1021/ja01641a005; constitution: Wiley et al. *J Am Chem Soc* **79** 6062 1957, DOI: 10.1021/ja01579a059]. [*Beilstein* **18/10** V 398.] A polyketide antibiotic [Washington & Wilson *Mayo Clinic Proceedings* **60** 189 1985, DOI: 10.1016/S0025-6196(12)60219-5].

Ethidium bromide [1239-45-8] C₂₁H₂₀N₃. Br, **M 394.3, m 260-262°**. Crystallise the purple-red solid from MeOH or EtOH [Lamos et al. *J Am Chem Soc* **108** 4278 1986, DOI: 10.1021/ja00275a007]. Its solubility in H₂O is 1%. Nucleic acid intercalating and colouring agent (used in gels). [*Beilstein* **22/11** V 352.] **CARCINOGEN**.

Ethyl 7-acetoxycoumarin-3-carboxylate [13207-77-3] C₁₄H₁₂O₆, **M 276.3, m 147-148°, 151°, 152°, 152-154°, pK²⁵ 7.04**. Purify the ester by recrystallisation from EtOH or aqueous EtOH, dry it *in vacuo* and store it at 2-8°. It is a good **fluorogenic substrate for esterases** that hydrolyse it to the acid, which acts as an indicator. The fluorescence has λ_{ex} 386nm, λ_{em} 445nm in 0.1 M Tris buffer pH 8.0. It has been prepared by refluxing 2-imino-3-ethoxycarbonyl-7-hydroxy-2H-chromene with Ac₂O for 1 hour, evaporating and recrystallising the residue from aqueous MeOH [Yasuda & Midorikawa *Bull Chem Soc Jpn* **39** 1754 1966, DOI: 10.1246/bcsj.39.1754]. Alternatively, it can be prepared by acetylation of 7-hydroxy-3-carbethoxycoumarin by the Ac₂O/pyridine method, and crystallises from EtOH in plates, **m 152°**. An attempted Fries rearrangement by heating the ester with anhydrous AlCl₃ at 16-170° failed, but gave **7-hydroxy-3-carboxycoumarin, m 271° (dec)** (from H₂O). [Shah & Shah *J Org Chem* **19** 1681 1954, DOI: 10.1021/jo01375a018; see also Boehm *Archiv der Pharmazie (Weinheim)* **271** 490 1933, DOI: 10.1002/ardp.19332710808].

Farnesol (*trans-trans-3,7,11-trimethyl-2,6,10-dodecatrien-1-ol*) [106-28-5, 4602-84-0 (*trans-trans, E,E*)] C₁₅H₂₆O, **M 222.4, b 111°/0.35mm, 126-127°/0.5mm, 142-143°/2mm, d₄²⁰ 0.8871, n_D²⁵ 1.4870**. The main impurity is the *cis-trans* isomer. Purify it by gas chromatography using a 4ft x 0.125in 3%OV-1 column at 150°. [Corey & Yamamoto *J Am Chem Soc* **92** 6637 1970, DOI: 10.1021/ja00725a044; Popjak et al. *J Biol Chem* **237** 56 1962, PMID: 14487886.] It has also been fractionated through a 14-in Podbielniak column at 11°/0.35mm. Alternatively, it has been purified by gas chromatography using SF96 silicone on Fluoropak columns or Carbowax 20M on Fluoropak or base-washed 30:60 firebrick (to avoid decomposition, prepared by treating the firebrick with 5N NaOH in MeOH and washed with MeOH to pH 8) at 210° with Helium carrier gas at 60 ml/min flow rate. It distils at high vacuum, and should be stored in sealed containers under N₂ in the dark. The **diphenylcarbamoyl derivative** has **m 61-63°** (from MeOH) and has an IR band at 3500 cm⁻¹. It has the scent of flowers. [Bates et al. *J Org Chem* **28** 1086 1963, DOI: 10.1021/jo01039a054; *Beilstein* **1** IV 2335.] **Farnesyl acetate** (*trans-trans-3,7,11-trimethyl-2,6,10-dodecatrien-1-yl acetate*) [4128-17-0 (*trans-trans, E,E*); 29548-30-9 (*isomeric mixture*)] C₁₇H₂₈O₂, **M 264.4, has b 115-125°/0.3mm, 167-169°/0.3mm, d₄²⁰ 0.91, n_D²⁰ 1.4870**. Purify and store farnesyl acetate as for the alcohol above. [*Beilstein* **2** II 154, **2** III 303.] Used in the perfumery. **Farnesyl pyrophosphate** [13058-04-3, *E,E*: 372-97-4] C₁₅H₂₈O₇P₂, **M 382.3, has pK_{Est(1)}~<2, pK_{Est(2)}~<2, pK_{Est(3)}~3.95, pK_{Est(4)}~6.26**. Purify the it by chromatography on Whatman No3 MM paper in a system of isopropanol-isobutanol/ammonia/water (40:20:1:30) (v/v). Store it as the Li or NH₄ salt at 0°. See geranyl-geranyl pyrophosphate below. Intermediate in the biosynthesis of terpenes, sterols and terpenoids substances.

Fenretinide [4-HPR, *N*-(4-hydroxyphenyl)retinamide] [65646-68-6] C₂₆H₃₃NO₂, **M 391.5.6, m 173-175°**. It is prepared from all-*trans*-retinoic acid [3.0g; see 302-79-4 above] and redistilled PCl₃ (0.92g) in dry *C₆H₆ (50ml) which are stirred for 2.5 hours, and during 20 minutes. This mixture is added to a solution of *p*-aminophenol (5.46g) in DMF (16ml) and Et₂O (2ml) with stirring under N₂ and ice-cooling, then stirred further at ~25°/3hrs followed by 50°/1hr. This is then diluted with Et₂O (150ml), the ethereal layer is extracted with cold 5% aqueous HCl (2 x 25ml), H₂O (25ml), dried (Na₂SO₄), filtered and evaporated to give a dark yellow solid which crystallises from MeOH (6ml/g), then 1:1.7 CHCl₃/*n*-hexane (8ml/g) to provide **fenretinide m 162-163°**. [Gander & Gurney US Patent 4190594 A to Johnson & Johnson 26 Feb 1980]. **Note** that it is polymorphic and has various forms, e.g. from aqueous EtOH the crystals have **m 173-175°**, and from MeOH the crystals have **m 178-181°**. The UV has λ_{max} at 362nm (ϵ 47900) in MeOH, and 370nm (ϵ 44500) in CHCl₃. [Synthesis and properties: Shealy et al. *J Pharm Sci* **73** 745 1984, DOI: 10.1002/jps.2600730610.] **Fenretinide** is a synthetic retinoid agonist agent, with anti-proliferative, anti-oxidant, anti-cancer activity with a **long half life in vivo**. Its **apoptotic action** appears different from that of *classical* retinoids. [Ulukaya & Wood *Cancer*

Treat Rev **25** 229 1999, DOI: 10.1053/ctrv.1999.0127.]

Fentanyl [*N*-(4-phenethyl-1-piperidiny)-*N*-phenylpropionamide] [437-38-7] $C_{22}H_{28}N_2O$, **M 336.5**, **m 83-84°**, **87.5°**, **pK_{Est}²⁵ ~10.0**. The syntheses of this drug and related compounds are described in a patent by Janssen [USP 3164600 A, Jan 5 1965]. -Purify it by dissolving in di-*iso*-propyl ether, filtering several times until clear, concentrate to a small volume and cool (2hrs/~0°) whereby **Fentanyl** crystallises out. *Alternatively*, recrystallise from di-*iso*-propyl ether, hexone or petroleum ether. It is a **strong organic base** (see pK_a) and is best kept as its salts. **Fentanyl citrate** [990-73-8] $C_{22}H_{28}N_2O \cdot C_6H_8O_7$, **M528.6**, **m 149-151°**, has a solubility in H₂O at ~25° of 2.5g/100ml, and is soluble in MeOH, but poorly soluble in CHCl₃. It has a bitter taste and is sold under the trade name of **Sublimaze**, among other names. It is a potent **opioid analgesic**, ~80-100 times **more potent than morphine** [57-27-2] and ~40-50 times the potency of pure pharmaceutical grade **heroin**. It is a **controlled substance** and a medical prescription. Used for severe trauma, strong chronic pain and apparently less addictive than morphine. Related useful analgesics are **Alfentanil** (ultra-short acting, 5-10min) and **Sufentanil** (5-10 times more potent than **Fentanil**). [Hess et al. *Eur J Clin Pharmacol* **4** 137 1972, DOI: 10.1007/BF00561135, PMID: 4655287; Stanley *J Pain Symptom Manage* **7** (3 Suppl, Issue 3) S3-7 1992, DOI: 10.1016/0885-3924(92)90047-L, PMID: 1517629; Black *J Med Chem.* **48** 1687 2005, DOI: 10.1021/jm040195b; PMID: 15771410.]

Flavin adenine dinucleotide (di-Na, 2H₂O salt, FAD) [146-14-5, 84366-81-4 (anhydrous)] $C_{27}H_{33}N_9O_{15}P_2$, **M 865.6** (free acid), **865.6**, **[α]₅₄₆ -54 (c 1, H₂O)**, **pK_a 1.13**, **pK_b 12.87**. Small quantities of FAD are purified by paper chromatography using *tert*-butyl alcohol/water, cutting out the main spot and eluting with water. Larger amounts can be precipitated from water as the **uranyl complex** by adding a slight excess of uranyl acetate to a solution at pH 6.0, dropwise and with gentle stirring. The solution is set aside overnight in the cold, and the precipitate is centrifuged off, washed with small portions of cold EtOH, then with cold peroxide-free diethyl ether. It is dried in the dark under vacuum over P₂O₅ at 50-60°. For **recovery**, the uranyl complex is suspended in water, and, after adding sufficient 0.01M NaOH to adjust the pH to 7, the precipitate of uranyl hydroxide is removed by centrifugation and the supernatant is lyophilised [Huennekens & Felton *Methods Enzymol* **3** 950 1957, DOI: 10.1016/S0076-6879(57)03484-9]. It can also be crystallised from water. It should be kept in the dark. More recently it was purified by elution from a DEAE-cellulose (Whatman DE 23) column with 0.1M phosphate buffer pH 7, and the purity was checked by TLC. [Holt & Cotton, *J Am Chem Soc* **109** 1841 1987, DOI: 10.1021/ja00240a039; *Beilstein* **26** III/IV 3632.] Cofactor of oxido-reductase enzymes.

Flavin mononucleotide (Na, 2H₂O salt, riboflavin-5'-phosphate [Na salt, 2H₂O], FMN) [130-40-5, 6184-17-4 (Na salt)] $C_{17}H_{21}N_4O_9P$, **M 456.3** (free acid), **514.4**, **pK₁ 2.1 (PO₄H₂)**, **pK₂ 6.5 (PO₄H⁻)**, **pK₃ 10.3 (CONH)**, **fluorescence λ_{max} 530nm (870nm for reduced form)**. Purify FMN by paper chromatography using *tert*-butanol/water, cutting out the main spot and eluting it with water. It can also be purified by adsorption onto an apo-flavodoxin column, followed by elution and freeze drying. It crystallises from aqueous acidic solution. [Mayhew & Strating *Eur J Biochem* **59** 539 1975, DOI: 10.1111/j.1432-1033.1975.tb02480.x; *Beilstein* **26** III/IV 2555.] Prosthetic group for NADH(P) oxido-reductases.

5-Fluorouridine (5-FU, 5-fluoro-1-β-D-ribofuranosyl-1H-pyrimidine-2,4-dione) [316-46-1, 77180-80-1] $C_9H_{11}FN_2O_6$, **M 262.2**, **m 180-182°**, **182-184°**, **[α]_D²⁰ +18 (c 1, H₂O)**, **pK_{Est(1)} ~ 8.0**, **pK_{Est(2)} ~ 13**. 5-Fluorouridine is recrystallised from EtOH/Et₂O and dried at 100° in a vacuum. It has UV with λ_{max} at 269nm (pH 7.2, H₂O), 270nm (pH 14, H₂O). [Liang et al. *Mol Pharmacol* **21** 224 1982, PMID: 6813676; *Beilstein* **24** III/IV 1231.] It is an antitumour agent.

Folic acid (FA, pteroyl-S-glutamic acid) [59-30-3, 75708-92-8 (2H₂O)] $C_{19}H_{19}N_7O_6$, **M 441.4**, **m >250°(dec)**, **[α]_D²⁵ +23 (c 0.5, 0.1N NaOH)**, **pK₁ 2.35 (protonation N10)**, **pK₂ 2.75 (protonation N1)**, **pK₃ 3.49 (α-CO₂H)**, **pK₄ 4.65 (γ-CO₂H)**, **pK₅ 8.80 (acidic N3)**. If paper chromatography indicates impurities, then recrystallise it from hot H₂O or from dilute acid [Waller et al. *J Am Chem Soc* **70** 19 1948, DOI: 10.1021/ja01181a006]. Impurities are removed by repeated extraction with *n*-BuOH of a neutral aqueous solution of folic acid (by suspending in H₂O and adding N NaOH dropwise till the solid dissolves, then adjusting the pH to ~7.0-7.5) followed by precipitation with acid, filtration, or **better** collected by centrifugation and recrystallisation from hot H₂O (fine yellow crystals). [Blakley *Biochem J* **65** 331 1957, DOI: 10.1042/bj0650331; Khalifa et al. *Helv Chim Acta* **61** 2739 1978, DOI: 10.1002/hlca.19780610742.]

Chromatography on cellulose followed by filtration through charcoal has also been used to obtain *pure acid*. [Sakami & Knowles *Science* **129** 274 1959, DOI: 10.1126/science.129.3344.274.] Its UV has λ_{\max} at 247 and 296nm (ϵ 12,800 and 18,700) in H₂O pH 1.0; 282 and 346nm (ϵ 27,600 and 7,200) in H₂O pH 7.0; 256, 284 and 366nm (ϵ 24,600, 24,500 and 86,00) in H₂O pH 13 [Rabinowitz in *The Enzymes* (Boyer et al. Eds), **2** 185 1960]. [Beilstein **26** III/IV 3944.] It is an important *food additive* and essential vitamin precursor of the cofactor dihydrofolic acid, necessary for thymidine synthesis and DNA; and the development of the neuronal tube in foetuses where deficiency leads to incomplete sealing of the tube and subsequently *Spina bifida*.

3-Formylchromone (4-oxo-4H-1-benzopyran-3-carbaldehyde) [17422-74-1] C₁₀H₆O₃, M **174.2**, m **151-153°**, **152°**, **152-153°**. This useful precursor for heterocyclic compounds [Sabitha *Aldrichim Acta* **29** 12 1996] is best prepared by a Vilsmeier-Haack reaction. Tetrachloropyrophosphate (80ml) is added dropwise to a solution of *o*-hydroxyacetophenone (25g, 0.184mole) in DMF (80ml) at -20° to 20° during 10 minutes. The mixture is stirred at ~25° for 13 hours, poured into ice-water and the solid is filtered off, washed with H₂O, then EtOH, dried *in vacuo*, and recrystallised from Me₂CO to give white crystals of *pure 3-formylchromone* (19.6g, 61%). Alternatively, DMF/POCl₃ is used. It has IR bands with ν_{\max} at 1605 (CHO), 1650 (C=O) and 1620 (C=C) cm⁻¹, the ¹H NMR [60MHz, DMSO-*d*₆, TMS] has δ at 10.16 (s, CHO), 8.9 (s, H-2) and 8.3-7.4 (m, H-5,6,7 and 8), and the MS *m/z*(%) has peaks at 174(4), 147(10), **146**(100), 121(8), 120(24), 118(3), 105(6), **104**(34), 92(17), 90(13), 89(11), **76**(13), 64(12), 63(18), 53(24), fragments in bold are characteristic of the 4-oxo-4H-1-benzopyran-3-carbaldehyde molecule. A large number of substituted derivatives have been prepared in this way and some, particularly the *6-bromo-derivative*, showed relatively *strong anti-anaphylactic activity* with low LD₅₀, and decreased gastric acid volume. [Nohara et al *Tetrahedron* **30** 3553 1974, DOI: 10.1016/S0040-4020(01)97034-6; Klutcho et al. US Pat 4,098,799 1978, *Chem Abs* **90** 22803 1979.]

Fructose-1,6-diphosphate (trisodium salt) [38099-82-0] C₆H₁₁Na₃O₁₂P₂·xH₂O, M **406.1**, pK₁²⁵ **1.48**, pK₂²⁵ **6.14**, pK₃²⁵ **6.29**, pK₄²⁵ **6.93** (free acid). Fructose-1,6-diphosphate is best purified *via* the acid strychnine salt which is stable for several months. To remove the strychnine, dissolve 5g in H₂O (150ml), and add 5N NaOH (or KOH to obtain the K salt) to pH 8.3 (phenolphthalein) with vigorous stirring. Remove the precipitate by centrifugation, wash with cold H₂O (2x 25ml), extract with CHCl₃ until the extract is free of strychnine (*ca* 8 to 10 times, Mandelein spot test). Freeze-dry the aqueous solution to give the Na salt which is *hygroscopic*. It has been recrystallised from aqueous EtOH as the *octahydrate*, m **125°**, [α]_D²⁰ **+2.6** (c **1**, H₂O). A neutral solution of the salt keeps well in a frozen state for over several months. Store it at -20°. [Sable *Biochemical Preparations* **2** 52 1952, Stumpf *J Biol Chem* **182** 261 1950, <http://www.jbc.org/content/182/1/261>; for estimation of hexose diphosphate see Kahan *Arch Biochem Biophys* **48** 331 1954, DOI: 10.1016/0003-9861(54)90348-3]. The *calcium salt* can be partially purified by dissolving in ice-cold N HCl (1g per 10ml) and re-precipitating by dropwise addition of 2N NaOH: the precipitate and supernatant are heated on a boiling water bath for a short time, then filtered, and the precipitate is washed with hot water. The *magnesium salt* can be precipitated from a cold aqueous solution by adding four volumes of EtOH. The *tetramethyl ester* is an oil with n_D^{10} **1.4648**, [α]_D¹⁸ **+20.2** (c **0.4**, MeOH). [Schulbach & Rauchenberger *Chem Ber* **60** 1178 1927, DOI: 10.1002/cber.19270600536; Beilstein **1** IV 4424.]

Fructose-6-phosphate (Neuberg ester, β -D-form) [643-13-0] C₆H₁₃O₉P, M **260.1**, [α]_D²¹ **+2.5** (c **3**, H₂O), pK₁²¹ **0.97**, pK₂²¹ **6.11**, (pK₂²⁵ **5.84**). Crystallise fructose-6-phosphate as the *barium salt* from water by adding 4-volumes of EtOH. The barium can be removed by passage through the H⁺ form of a cation exchange resin, and the *free acid* is collected by freeze-drying. Alternatively, the Ba salt is dissolved in H₂O, and one equivalent of Na₂SO₄ is added in small portions with stirring, filter off BaSO₄ and freeze dry to give the *Na salt*. The 6-phosphate hydrolyses more slowly than the 1-phosphate and considerably slower than pyrophosphoric acid (10² times) and triphosphoric acid (10³ times). [For hydrolysis see Friess *J Am Chem Soc* **74** 5521 1954, DOI: 10.1021/ja01141a510; Tankó & Robinson *Biochem J* **29** 961 1935, DOI: 10.1042/bj0290961; Neuberg *Biochem Zeitschrift* **88** 432 1918, pKa: Meyerhof & Lohmann *Biochem Zeitschrift* **185** 113, 131 1927, Neuberg et al. *Arch Biochem* **3** 33, 40 1944, Beilstein **1** I 464, **1** IV 4423, H **31** 537.]

Gangcyclovir [9-{(1,3-dihydroxy-2-propoxy)methyl}guanine; 2-amino-1,9-[(2-hydroxy-1-hydroxy-methyl)-ethoxymethyl]-6H-purin-6-one; Cytovene; Cymeva(e)n(e)] [82410-32-0] C₉H₁₃N₅O₄, M **255.2**,

m >290°(dec), >300°(dec), monohydrate m 248-249°(dec), $pK_{Est(1)} \sim -1.1$, $pK_{Est(2)} \sim 4.1$, $pK_{Est(3)} \sim 9.7$. Recrystallise gangcyclovir from MeOH. *Alternatively*, dissolve ~90g of it in 700ml of H₂O, filter and cool (*ca* 94% recovery). It has UV with λ_{max} (MeOH) at 254nm (ϵ 12,880), 270sh nm (ϵ 9,040); its solubility in H₂O at 25° is 4.3mg/ml at pH 7.0. [Ogilvie et al. *Can J Chem* **60** 3005 1982, DOI: 10.1139/v82-430; Ashton et al. *Biochem Biophys Res Commun* **108** 1716 1982, DOI: 10.1016/S0006-291X(82)80109-5; Martin et al. *J Med Chem* **26** 759 1983, DOI: 10.1021/jm00359a023.] It is an antiviral drug.

Geraniol (*trans*-3,7-Dimethyl-2,6-octadien-1-ol) [*trans*, *E*-form 106-24-1] C₁₀H₁₈O, M 154.2, b 114-115°/12mm, 229-230°/757mm, d_{15}^{15} 0.889, n_D^{20} 1.4628. Purify geraniol by fractional distillation preferably at high vacuum. It has a sweet rose odour when pure, and the UV has λ_{max} at 190-195nm (ϵ 18,000). [Beilstein **1** H 457, **1** IV 2277.] It is an important biosynthetic intermediate, is present in rose, geranium, lemon and other essential oils, used in perfumery and as a flavouring agent.

Geranyl acetate [*trans*, *E*-form 105-87-3; 16409-44-2] C₁₂H₂₀O₂, M 196.3, m <25°, b 118°/12mm, 138°/25mm, 236-242°/760mm(dec), d_{15}^{15} 0.9174, n_D^{15} 1.4766. Purify the fragrant smelling geranyl acetate by fractional distillation at as high a vacuum as possible. It is very soluble in EtOH but insoluble in H₂O. [Beilstein **2** H 140, **2** I 65, **2** II 153, **2** III 299, **2** IV 204.] It has the scent of flowers and is used in food and cosmetics see *Food and Cosmetics Toxicology* **12** 885 1974, DOI: 10.1016/0015-6264(74)90167-9.

Geranylgeranyl pyrophosphate [6699-20-3 (NH₄ salt), 64732-91-8] C₂₀H₃₆O₇P₂, M 450.5, $pK_{Est(1)} \sim <2$, $pK_{Est(2)} \sim <2$, $pK_{Est(3)} \sim 3.95$, $pK_{Est(4)} \sim 6.26$. Purify the pyrophosphate by countercurrent distribution between two phases of a butanol/isopropyl ether/ammonia/water mixture (15:5:1:19) (v/v), or by chromatography on DEAE-cellulose (linear gradient of 0.02M KCl in 1mM Tris buffer, pH 8.9). *Alternatively*, purify it through a column of Dowex 1-x8 (formate form previously washed with MeOH) and eluted with a linear gradient of 0.053–0.43M ammonium formate in a total volume of 300ml of MeOH. The purity can be checked by TLC on Silica gel G on buffered plates (pH 6.5), eluted with CHCl₃/MeOH/H₂O (60:40:9) and developed with I₂ vapour. Store it as a powder at 0°. [Altman et al. *J Am Chem Soc* **94** 3257 1972, DOI: 10.1021/ja00764a073.] It is more stable as the *di(tri-n-butylammonium)hydrogen phosphate salt* which can be obtained from the acid by evaporation in a rotary evaporator below 32° [Upper & West *J Biol Chem* **242** 3285 1967, PMID: 4291475]. [Gregonis & Rilling *Biochemistry* **13** 1538 1974, DOI: 10.1021/bi00704a033; Gregonis & Rilling *Biochem Biophys Res Commun* **54** 449 1973, DOI: 10.1016/0006-291X(73)90942-X; for medical aspects see Wiemer et al. [*Clin Pharmacol Therap* **90**(6) 804 2011, DOI: 10.1038/clpt.2011.215. PMID: 22048229].

Geranyl pyrophosphate [*E*-form 763-10-0 tri-(NH₄ salt), *Z*-form 16751-02-3] C₁₀H₂₀O₇P₂, M 314.2, $pK_{Est(1)} \sim <2$, $pK_{Est(2)} \sim <2$, $pK_{Est(3)} \sim 3.95$, $pK_{Est(4)} \sim 6.26$. Purify the pyrophosphate by paper chromatography on Whatman No 3 MM paper in a system of isopropyl alcohol/isobutyl alcohol/ammonia/water (40:20:1:39), R_F 0.77-0.82. Store it in the dark as the *ammonium salt* at 0°. The *E*-form crystallises in platelets from aqueous Me₂CO, m ~120°. It dissolves in dry MeCN. *Alternatively*, purify it through a column of Dowex AG 1x8(200-400mesh) equilibrated with 50mM NH₄ formate, and elute with MeOH/H₂O/NH₄OH (95:5:05), then freeze-dry. [Dixit et al. *J Org Chem* **46** 1967 1981, DOI: 10.1021/jo00322a060; Beilstein **1** IV 3580.] It is an intermediate in the HMG-CoA reductase pathway.

Gliotoxin (3*R*-6*t*-hydroxy-3-hydroxymethyl-2-methyl-(5*at*)-2,3,6,10-tetrahydro-5*aH*-3,10*ac*-epidisulfido-[1,2-*a*]-indol-1,4-dione) [67-99-2] C₁₃H₁₄N₂O₄S₂, M 326.4, m 191-218°(dec), 220°(dec), 221°(dec), $[\alpha]_D^{20}$ -254 (c 0.6, CHCl₃), $[\alpha]_D^{25}$ -270 (c 1.7, pyridine). Purify gliotoxin by recrystallisation from MeOH. Its solubility in CHCl₃ is 1%. The *dibenzoyl derivative* has m 202° (from CHCl₃/MeOH). [Glister & Williams *Nature* **153** 651 1944, DOI: 10.1038/153651a0; Elvidge & Spring *J Chem Soc Suppl* **135** 1949, DOI: 10.1039/JR949000S135; Johnson et al. *J Am Chem Soc* **65** 2005 1943, DOI: 10.1021/ja01250a051; Bracken & Raistrick *Biochem J* **41** 569 1947, DOI: 10.1042/bj0410569; for a review see Waring & Chai *Aust J Chem* **68** 178 2015, DOI: 10.1071/CH14482.] It is a highly bioactive mycotoxin produced by various fungi.

Glucose-1-phosphate (G-1-P) (Cori ester) [59-56-3] C₆H₁₃O₉P, M 260.1, $[\alpha]_D^{25}$ +120 (c 3, H₂O), $[\alpha]_D^{20}$ +78 (c 4, H₂O of di-K salt), pK_1 1.11, pK_2 6.13 [pK^{25} 6.50]. Two litres of a 5% aqueous solution of the phosphate are purified by adjusting the pH to 3.5 with glacial acetic acid (+ 3g of charcoal) and filtering. An

equal volume of EtOH is added, the pH is adjusted to 8.0 (glass electrode) and the solution is stored at 3° overnight. The precipitate is filtered off, dissolved in 1.2L of distilled water, filtered and an equal volume of EtOH is added. After standing at 0° overnight, the crystals are collected at the centrifuge and washed with 95% EtOH, then absolute EtOH, ethanol/diethyl ether (1:1), and diethyl ether. [Sutherland & Wosilait, *J Biol Chem* **218** 459 1956, PMID: 13278353.] Its **barium salt** can be crystallised from water and EtOH. Heavy metal impurities are removed by passage of an aqueous solution (*ca* 1%) through an Amberlite IR-120 column (in the appropriate H⁺, Na⁺ or K⁺ forms). **Di-K salt** crystallises as a **dihydrate** from EtOH. [see McGready *Biochemical Preparations* **4** 63 1955.] [Beilstein **17/8** V 247.] This is the α -form, the β -form occurs in microbes.

Glucose-6-phosphate (G-6-P) (Robison ester) [acid 56-73-5; Ba salt 58823-95-3; Na salt 54010-71-8] **C₆H₁₃O₉P**, M 260.1, m 205-207°(dec) mono Na salt, [α]_D²⁰ +41 (c 5, H₂O), pK₁ 1.65, pK₂ 6.11, pK₃²⁵ 11.71 [-C₁(OH)O⁻]. It can be freed from metal impurities as described for glucose-1-phosphate. The solubility of the Na salt is 5% in H₂O at 20°. Its **barium salt** can be purified by solution in dilute HCl and precipitation by neutralising the solution. The precipitate is washed with small volumes of cold water and dried in air. Alternatively, the barium salt is dissolved in H₂O, 4 volumes of EtOH are added, the precipitate is collected, washed with 90% EtOH, absolute EtOH, 75% EtOH/25% Et₂O, 25% EtOH/75% Et₂O and finally dry Et₂O. The **dry pure Barium salt** has [α]_D²⁵ +17.9° (c 1, H₂O). [Beilstein **1** IV.] **G-6-P** is relatively more stable to hydrolysis (12% hydrolysis in N HCl/100° in 4 hours) than G-1-P (45% hydrolysis in N HCl/20° in 20 hours). [Lardy & Fischer *Biochemical Preparations* **2** 39 1952.]

L- α -Glycerol phosphocholine (Cadmium Chloride)_x complex [64681-08-9] **C₈H₂₀NO₆P. CdCl₂. M 257.2 + (183.3)_x, pK_{Est} ~ 5.5. Glycerol phosphocholine is purified *via* the CdCl₂ complex which is recrystallised four times from 99% EtOH by standing at 0° for 1 hour. The white precipitate is collected, washed with EtOH, Et₂O and dried in a vacuum. The amorphous Cd complex can be converted to the crystalline form [**C₈H₂₀O₆NP.CdCl₂.3H₂O**] by dissolving 34.4g in H₂O (410ml), and 99% EtOH (1650ml total) is added slowly with stirring and allowing the clear solution to stand at 25° for 12 hours, then at 5° for 12 hours. The crystalline complex is filtered off, washed with cold 80% EtOH and dried in air. **Glycerol phosphocholine** is recovered from the complex by dissolving in H₂O (2% solution) and passing it through an ion-exchange column (4.9 x 100cm, of 1 volume IRC-50 and 2 volumes of IR-45). The effluent is concentrated to a thick syrup at 45°. It is dried further at 50°/P₂O₅/48 hours. The vitreous product (~8.25g) is then dissolved in 99% EtOH (50ml), and the clear solution is cooled to 5°, whereby crystals begin to appear, and crystallisation is completed at -15°/16 hours. The crystals are filtered off, washed with 99% EtOH, and Et₂O then dried at 50° *in vacuo* over P₂O₅. It can be recrystallised from 99.5% EtOH (long prisms). It is **hygroscopic** and must be handled in a H₂O-free atmosphere [Tattie & McArthur *Biochemical Preparations* **6** 16 1958, Baer & Kates *J Am Chem Soc* **70** 1394 1948, DOI: 10.1021/ja01184a031; Abrahamsson & Pascher *Acta Cryst* **21** 79, 87 1966, DOI: 10.1107/S0365110X66002366].**

Hematin (ferrihaeme hydroxide) [15489-90-4] **C₃₄H₃₃N₄O₅Fe**, M 633.5, m 200°(dec), pK_{Est} ~ 4. Crystallise the bluish-black or brownish-black solid from pyridine. Dry it at 40° *in vacuo*. It is soluble in dilute aqueous NaOH or KOH and has λ_{\max} (2.5N NaOH) at 580nm (ϵ 10500). [Beilstein **26** III/IV 3047.]

Hesparin IX (8,13-bis(1-hydroxyethyl)-3,7,12,17-tetramethyl-21H-23H-porphin-2,18-dipropionic acid, 3,3'-[7,12-bis(1-hydroxyethyl)-3,8,13,17-tetramethyl-porphyrin-2,18-diyl]-dipropionic acid) [14459-29-1] **C₃₄H₃₈N₄O₆**, M 598.7, pK_{Est} ~4.8, 3.67. Purify it by dissolving it in EtOH and adding H₂O or Et₂O to give deep red crystals. It has also been recrystallised from MeOH. Its UV has λ_{\max} at 615.5, 565, 534.4 and 499.5nm in 0.1N NaOH, and 597, 619, 634, 653, 683 and 701nm in 2N HCl [Falk *Porphyrins and Metalloporphyrins* Elsevier, NY, p 175 1964, LCCCN0 63-19821.] It is used in the affinity chromatographic purification of Heme proteins [Olsen *Methods Enzymol* **123** 324 1986, DOI: 10.1016/S0076-6879(86)23038-4]. The **dihydrochloride** [17696-69-4] **M 671.6**, crystallises from EtOH/HCl/Et₂O and is partly soluble in H₂O and 5w/v% in pyridine. The **O-methyl-dimethyl ester** has m 203-206° (from CHCl₃/MeOH), and the **O,O'-dimethyl-dimethyl ester** has m 145° (from CHCl₃/MeOH). [Paul *Acta Chem Scand* **5** 389 1951, DOI: 10.3891/acta.chem.scand.05-0389; Beilstein **26** III/IV 3157, and 3158 for the HCl.] **Hematoporphyrin dimethyl ester** [33070-12-1, 5594-29-6] **C₃₆H₄₂N₄O₆**, M 626.7, has m 212°, 212.2-217.2°. It crystallises from CHCl₃/MeOH [Beilstein **26** III/IV 3157.]

Hematoxylin (\pm -11*bc*-7,11*b*-dihydroindeno[2,1-*c*]-chromen-3,4-6*ar*-9,10-pentaol) [517-28-2] $C_{16}H_{14}O_6$, **M 302.3**, **m 200°(dec)**, **210-212°(dec)**. Hematoxylin crystallises from H_2O (as *trihydrate*) in white-yellow crystals which become red on exposure to light and then melt at 100-120°. It can be recrystallised from $Me_2CO/*C_6H_6$. It has also been recrystallised from dilute aqueous $NaHSO_3$ until colourless and is soluble in alkali, borax and glycerol. Store it in the dark below 0°. [Morsingh & Robinson *Tetrahedron* **26** 281 1970, DOI: 10.1016/0040-4020(70)85029-3; Dann & Hofmann *Chem Ber* **98** 1498 1965, DOI: 10.1002/cber.19650980524; *Beilstein* **17/8** V 469.] Used extensively for staining plant and animal tissue. [Puchtler et al. *Histochemistry* **85** 353 1986, DOI: 10.1007/BF00982665.] It has one more phenolic group than *Brazilin* (474-07-7, *Brazilein*).

Hemin (ferriprotoporphyrin IX chloride) [16009-13-5] $C_{34}H_{32}ClN_4O_4Fe$, **M 652.0**, **m sinters at 240°**, **pK_{Est} ~4.8**. Hemin is purified by recrystallisation from AcOH. Also, hemin (5g) is shaken in pyridine (25ml) till it dissolves, then $CHCl_3$ (40ml) is added, the container is stoppered and shaken for 5 minutes (releasing the stopper occasionally). The solution is filtered under slight suction, and the flask and filter are washed with a little $CHCl_3$ (15ml). During this period, AcOH (300ml) is heated to boiling, and saturated aqueous NaCl (5ml) and concentrated HCl (4ml) are added. The $CHCl_3$ filtrate is poured in a steady stream, with stirring, into the hot AcOH mixture and set aside for 12 hours. The crystals are filtered off, washed with 50% aqueous AcOH (50ml), H_2O (100ml), EtOH (25ml), Et₂O and dried in air. [Fischer *Org Synth Coll Vol* **3** 442 1955, DOI: 10.15227/orgsyn.021.0053; *Beilstein* **26** III/IV 3048.] It is used in the treatment of *porphiria*.

α -1,2,3,4,5,6-Hexabromocyclohexane (NSC 7908, **Jak2 inhibitor**, **α -1*r*,2*t*,3*t*,4*c*,5*t*,6*c*-hexabromocyclohexane**) [1837-91-8] $C_6H_6Br_6$, **M 557.5**, **m 222°**, **216-218°**, **218-223°**, **224-225°**, **b 388.9° /760mm (predicted)**, **d²⁵ 2.90g/cm³**, **n_D²⁰ 1.70**. Although eight stereoisomers are possible, the least adaptable isomers should be the ones where 1,3-diaxial bromine atoms in the cyclohexane ring, which would be dis-favoured due to strong non-bonded interactions. The most favourable configuration, and consequently more stable, is the one where the cyclohexane ring is in the *chair* conformation with all the six bromine atoms holding *equatorial* orientations. α -1,2,3,4,5,6-Hexabromocyclohexane has been prepared in 86% yield by prolonged exposure of Br_2 (50g, 16ml) in $*C_6H_6$ (100ml) to sunlight and isolating successive crops, combining them and recrystallising them from toluene to give white cubes with **m 216-218°**. The yield is affected by the light source and strength — stronger light greatly decreases the yield. A series of reactions occur here and were unravelled by Bolton et al. [*JCS Perkin Trans 1* 893 1984, DOI: 10.1039/P19840000893.] who showed that it was the isomer where four Br atoms on consecutive carbon atoms of the cyclohexane ring are in the *equatorial* orientation, while the fifth and sixth adjacent Br atoms are in the *axial* configuration. It used to be referred to as ‘ α -benzene hexabromide’ and its structure was confirmed by X-ray crystallography (130K), and by ¹H and ¹³C NMR spectroscopy to be *r-1, t-2, t-3, c-4, t-5, c-6-hexabromocyclohexane* (where *r* = reference point, *t* = trans and *c* = cis) [Carman et al. *Aust J Chem* **47** 1395 1994, DOI: 10.1071/CH9941395].

β -1,2,3,4,5,6-Hexabromocyclohexane [87-82-1, 3015-41-0] is apparently the thermodynamically stable isomer which has been known for some time. It was prepared by Orndorff and Howells [*J Am Chem Soc* **18** 312 189], and repeated by Hendricks and Bilicke [*J Am Chem Soc* **48** 3007 1928, DOI: 10.1021/ja01691a001] who added slowly, and successively, small amounts of Br_2 to a layer of $*C_6H_6$ floating on a 1% solution of NaOH at a temperature near 0°. The **α - and β - isomers** were isolated from the solid that separated from the lower layer by boiling with EtOH in which they have limited solubility. The **α - form** was separated by extraction with $CHCl_3$ leaving the **β -form** behind. The latter was recrystallised from boiling xylene which gave a 0.1% yield of small crystals, 1mm thick, which showed a crystal development of [111] and [110]. These crystals had a sharp **melting point at 253°**. The latter authors have determined the X-ray structure of the crystals of this **β -isomer** which showed that they are of the point group C_{3i} , which require a three-fold axis and a centre of symmetry, consistent with a cyclohexane ring with **all the bromine atoms** in the *equatorial* orientation.

α -1,2,3,4,5,6-Hexabromocyclohexane has limited solubility in most solvents but can be recrystallised from toluene and is extractable from the **β -isomer** with hot $CHCl_3$ (see above). To obtain higher solubility make a solution in DMSO (5mg/ml) by warming at 37° with shaking in an ultrasonic bath for several minutes. This stock solution can be stored below -20° for several months. The **enantiomers** of this isomer have been separated by gas chromatography (see Kallenborn et al. *ChemInform Abstract* **22** (Issue 20) page no, May 21, 1991, on line 23 Aug 2010, DOI: 10.1002/chin.199120281].

Sayeski and coworkers [Sandberg et al. *J Med Chem* **48** 2526 2005, DOI: 10.1021/jm049470k, PMID: 15801842] have identified this **α -isomer** as a small molecule inhibitor of Jak2 tyrosine kinase autophosphor-

ylation which specifically *inhibits* ligand-dependent Jak2 activation.

(+)-Hydroquinidine anhydrous (9S-6'-methoxy-10,11-dihydrocinchonan-9-ol) [1435-55-8] $C_{20}H_{26}N_2O_2$, **M 326.4, m 168-169°, 169°, 169-170°, 171-172°, $[\alpha]_D^{20} +231$ (c 2, EtOH), +299° (c 0.82, 0.1N H_2SO_4), $pK_{Est} \sim 8.8$.** (+)-Hydroquinidine forms needles from EtOH and plates from Et_2O . It is slightly soluble in Et_2O and H_2O but readily soluble in hot EtOH. The **hydrochloride** [1476-98-9] has **m 273-274° (dec, darkening at $\sim 270^\circ$), and $[\alpha]_D^{26} +183.9$ (c 1.3, MeOH);** and is very soluble in MeOH and $CHCl_3$, but less soluble in H_2O , EtOH and still less soluble in dry Me_2CO . [Heidelberger & Jacobs *J Am Chem Soc* **41** 817 1919, DOI: 10.1021/ja02226a015; King *J Chem Soc* 523 1946, DOI: 10.1039/JR9460000523.] [Kyker & Lewis *J Biol Chem* **157** 707 1945, <http://www.jbc.org/content/157/2/707>; Emde *Helv Chim Acta* **15** 557 1932, DOI: 10.1002/hlca.19320150156; *Beilstein* **23/13** V 340.] It is a useful base for **optical resolutions** of chiral acids.

Hydroquinine [(8 α ,9R)-10,11-dihydro-6'-methoxycinchonan-9-ol] [522-66-7] $C_{20}H_{26}N_2O_2$, **M 326.4, m 168-171°, 171.5°, $[\alpha]_D^{16} -143.5$ (-148) (c 1.1, EtOH), pK^{15} 5.33 and 8.87.** Recrystallise hydroquinine from EtOH, Et_2O or $*C_6H_6$. The **hydrochloride** [1668-97-9] $C_{20}H_{26}N_2O_2 \cdot HCl \cdot 0.5H_2O$, crystallises from H_2O and decomposes at **206-208°** (after initial rapid heating to 200°). It is soluble in H_2O , EtOH and Me_2CO and has $[\alpha]_D^{21} -123.9$ (c 1.11 H_2O). [Rabe & Schultz *Chem Ber* **66** 120 1933, DOI: 10.1002/cber.19330660208; for synthesis see Heidelberger & Jacobs *J Am Chem Soc* **41** 817 1919, DOI: 10.1021/ja02226a015; *Beilstein* **23** III/IV 3193, **23/13** V 340.] It is a useful base for **optical resolutions** of chiral acids. The **racemate** has **m 175-175.5°** (from EtOH and drying at 100°/0.1mm).

5-Hydroxycreatinine (creatol, CTL, 2-amino-5-hydroxy-1-methylimidazolidin-4-one) [133882-98-1] $C_4H_7N_3O_2$, **M 129.1, m 190° (191°, dec) for the hydrochloride, pK^{25} 4.2.** Creatol is one of three metabolites of creatinine [60-27-5] which was isolated from inflamed rabbit skin tissues [Ienaga et al. *Tetrahedron Lett* **28** 4587 1987, DOI: 10.1016/S0040-4039(00)96571-7], and the urine of uraemic rats (adenine induced chronic renal failure) and uraemic patients. The other two metabolites are 1-methylhydantoin (1-methylimidazolidine-2,4-dione [616-04-6]) and 5-hydroxy-1-methylhydantoin [1-methyl-5-hydroxyimidazolidine-2,4-dione [84210-26-4]. [Ienaga et al. *JCS Chem Commun* 509 1991, DOI: 10.1039/C39910000509.] Urine from uraemic rats (44ml, collected after 24 hours of administration of creatinine) was applied onto a PK216 ion exchange column (H^+ form) that was washed with H_2O , then eluted with aqueous NH_3 ($2mol\ dm^{-3}$) and evaporated *in vacuo*. The residue, in H_2O , was applied onto a Biolex-70 ion exchange column (H^+ form) that was washed with H_2O , then eluted with aqueous AcOH ($0.1mol\ dm^{-3}$), and the eluate was evaporated *in vacuo* to give **creatol** (0.66mg, 0.11%, purity checked by ODS reverse phase HPLC). [The H_2O wash from the PK216 column gave the other two metabolites (1.5mg, 0.35 and 0.87mg, 0.15% respectively) which were separated on a silica gel column with EtOAc elution. **Creatol** has 1H NMR (D_2O , *tert*-BuOH as internal standard) with δ at 3.11 (s, in $N^{12}CH_3$ creatol; but d, in $N^{13}CH_3$ labelled creatol with $J = 143$ Hz) and 5.11 (s, H-5); and ^{13}C NMR (D_2O , dioxane as internal standard) with δ at 30 (NCH_3), 81 (C-5), 160 (C-2) and 174 (C-4). [Ienaga et al. *JCS Chem Commun* 509 1991, DOI: 10.1039/C39910000509.] The **natural enantiomer** most probably has the **S configuration** at C-5 [see Ienaga et al. *JCS Chem Commun* 509 1991, DOI: 10.1039/C39910000509]. [Nakamura & Ienaga, Glycocyamidine derivatives, *Japanese Kokai Tokyo Koho*, JP 2957217 1989, US patent 4957936 1990.] A urinary metabolite indicator of renal disfunction. Data kindly supplied by Dr Kazu Ienaga, Nippon Zoki Pharmaceutical Co Ltd, Osaka, Japan.

R-(-)-2-Hydroxy-3,3-dimethyl- γ -butyrolactone (3-hydroxy-4,4-dimethyl-4,5-dihydrofuran-2-one, D-pantolactone) [599-04-2] $C_6H_{10}O_3$, **M 130.1, m 89-91°, 90.5-91.5°, 91°, 92-93°, b 120-122°/15mm, $[\alpha]_D^{20} -28^\circ$ (c 5, MeOH), $[\alpha]_D^{20} -51$ (-49.1) (c 3, H_2O).** Recrystallise the lactone from Et_2O /petroleum ether, diisopropyl ether or $*C_6H_6$ /petroleum ether and sublime it at 25°/0.0001mm. It hydrolyses readily to the hydroxy-acid and racemises when heated above 145°. Store at 2-8°. The **Brucine salt** has **m 211-212°** (from EtOH). [Kuhn & Wieland *Chem Ber* **73** 1134 1940, DOI: 10.1002/cber.19400731020; and Stiller et al. *J Am Chem Soc* **62** 1779 1940, DOI: 10.1021/ja01864a037; Beutel & Tishler *J Am Chem Soc* **68** 1463 1946, DOI: 10.1021/ja01212a020; *Beilstein* **18/1** V 22.] The **racemate** [79-50-5], **m 74-78°, 80°, b 130°/18mm**, purified in the same way, is **hygroscopic** and its **resolution** has been described [Major & Finkelstein *J Am Chem Soc* **63** 1368 1941, DOI: 10.1021/ja01850a068; see also preceding references].

(±)-5-Hydroxy-1-methyl-imidazolidine-2,4-dione (HMH) [84210-26-4] $C_4H_6N_2O_3$, M 130.1, m 138°, pK 8.64. The racemic compound was obtained by oxidation of *N*-1-methylhydantoin with $Pb(OAc)_4$ in $*C_6H_6$ (60°, 24 hours), isolation of the 5-acetoxy derivative (*via* a short silica-gel column chromatography; eluting with $*C_6H_6$ /hexane), followed by hydrolysis with aqueous 0.3M H_2SO_4 (~25°, 40 minutes). Neutralisation, followed by evaporation, and silica-gel TLC (EtOAc) provided pure **racemate m 138°**, (64% yield) with the expected 1H NMR spectrum, EI MS and elemental analysis. It is a **plant growth regulator** affecting the flowering period of cut white chrysanthemums at concentrations as low as 10^{-6} M. [Ienaga et al. *Tetrahedron Lett* **28** 4587 1987, DOI: 10.1016/S0040-4039(00)96571-7.] The **natural metabolite N-1-methylhydantoin (MH [616-04-6]**, see below) was isolated from rabbit urine by diluting the urine two-fold with MeOH and evaporating. The residue was extracted with MeOH/EtOAc (1:6), filtered, the filtrate was concentrated, subjected to silica-gel column chromatography and eluted with MeOH/ $CHCl_3$ (1:9). The first eluate gave the pure crystalline **5-hydroxy compound (HMH, 28% of total)** that had **m 147-150°** and $[\alpha]_D^{25} -5.1$ (c 1.0, MeOH). The expected 1H NMR [$Eu(tfc)_3$] spectrum indicated that its enantiomeric excess was 56% (i.e. a 78:22 mixture). Some racemisation of the 5-hydroxy compound may well have occurred during isolation. The synthetic **racemate** was resolved by conversion into the diastereoisomeric Bz-L-proline ester (1:1, *R:S*) [with 1-(3-diethylaminopropyl)-3-ethylcarbodiimide HCl in the presence of DMAP in MeCN at 0°/2 hours) followed by purification through a silica-gel column and eluting with EtOAc/hexane (7:3)] in 56% yield. Fractional recrystallisation from Me_2CO gave pure 5-(*R*)-(N-Bz-L-prolyloxy)-1-methylimidazolidine-2,4-dione (30%), **m 189-192°**, $[\alpha]_D^{25} -2.1$ (c 1.0, MeOH) $\{^1H$ NMR, $DMSO-d_6$ has δ at 1.8-2.0 (m, 3H), 2.2-2.4 (m, 1 H), [2.52 (s) + 2.75 (s), 3H], 3.4-3.55 (m, 2 H), [4.35 (dd, *J* = 4, 9 Hz) + 4.46 (dd, *J* = 4, 9 Hz) 1H], [5.03 (s) + 5.10 (s) 2H], [6.90 (s) + 6.10 (s) 1H], 7.25-7.45 (m, 5H, aromatic) and 11.21 (br s, NH)}; and when these crystals were subjected to X-ray crystallography they revealed that the absolute configuration at C5 of the *imidazolidine-2,4-dione* part of the molecule was **R**, and that C5 of the *natural metabolite* was therefore **S** when a comparison of the HPLC of the mixture of prolyl-esters from the natural metabolite was made. The residue from evaporation of the mother liquors was recrystallised from EtOAc and gave the diastereoisomeric **5-(S)-N-Bz-L-prolyl ester** (8%) with **m 147-150°**, and $[\alpha]_D^{25} -68.9$ (c 1.0, MeOH) $\{^1H$ NMR, $DMSO-d_6$ has δ at 1.7-2.0 (m, 3H), 2.2-2.4 (m, 1 H), [2.63 (s) + 2.73 (s), 3H], 3.35-3.55 (m, 2 H), [4.36 (dd, *J* = 4, 9 Hz) + 4.47 (dd, *J* = 4, 9 Hz) 1H], [5.04 (d, *J* = 13Hz) + 5.07 (d, *J* = 13Hz) 1H], [5.07 (d, *J* = 13Hz) + 5.13 (d, *J* = 13Hz) 1H], [6.05 (s) + 6.12 (s) 1H], 7.25-7.45 (m, 5H, aromatic) and 11.25 (br s, NH)}. Hydrolysis of these esters gave **racemic** 5-hydroxyimidazoline-2,4-dione, however, owing to facile racemisation. [Ienaga et al. *JCS Perkin Trans I* 1153 1989, DOI: 10.1039/P19890001153.] The **racemate** and the **optically active** form of **5-hydroxy-1-methylhydantoin** have the same 1H NMR (400MHz, Me_2CO-d_6 , TMS) with δ at ~5.1 (d, C-5) and ~5.9 (d, 1-NMe). Data kindly supplied by Dr Kazu Ienaga, Nippon Zoki Pharmaceutical Co Ltd, Osaka, Japan. These are urinary metabolites and **indicators of renal health**.

(-)-Inosine {9-[(2*R*,3*R*,4*S*,5*R*)-3,4-dihydroxy-5-(hydroxymethyl)oxolan-2-yl]-6,9-dihydro-3*H*-purin-6-one} [58-63-9] $C_{10}H_{12}N_4O_5$, M 268.2, m 90° (dihydrate), 218° (dec, anhydrous), $[\alpha]_{546}^{20} -76$ (c 1, 0.1M NaOH), pK₁²⁵ 1.06, pK₂²⁵ 8.96, pK₃²⁵ 11.36. (-)-Inosine forms **anhydrous** crystals from aqueous 80% EtOH but the **dihydrate** from H_2O . [Beilstein **31** H 25, **26** III/IV 2087.] Inosine is commonly found in tRNAs.

myo-Inositol (cyclohexane[1*r*,2*c*,3*c*,4*t*,5*c*,6*t*]-hexol) [87-89-8] $C_6H_{12}O_6$, M 180.2, m 218° (di-hydrate), 225-227°, 226-230°. Recrystallise *myo*-inositol from aqueous 50% ethanol or H_2O forming a **dihydrate**, or **anhydrous** crystals from AcOH. The dihydrate is **efflorescent** and becomes anhydrous when heated at 100°. The anhydrous crystals are not hygroscopic. Its solubility in H_2O at 25° is 14%, at 60° it is 28%; it is slightly soluble in EtOH but insoluble in Et_2O . [Ballou & Anderson *J Am Chem Soc* **75** 648 1953, DOI: 10.1021/ja01099a039; Anderson & Wallis *J Am Chem Soc* **70** 2931 1948, DOI: 10.1021/ja01189a030; Beilstein **6** II 1157, **6** IV 7919.] Combined GC-MS study of many organic-rich marine sediments identified *myo*-inositol, *chiro*-inositol and *scyllo*-inositol, with the *myo*-isomer as the most abundant. [White & Miller *Science* **193** (issue 4256) 885 1976, DOI: 10.1126/science.193.4256.885]. *myo*-Inositol is useful in the treatment of several medical problems, e.g. bulimia, panic disorder, obsessive compulsive disorder (OCD), unipolar and bipolar depression, and is used as a food supplement [Clemments & Damell *Am J Clin Nutr* **33**(9) 1954 1980, PMID: 7416064].

Inositol monophosphate [15421-51-9] $C_6H_{13}O_9P$, M 260.1, m 195-197°(dec). Crystallise the phosphate from

water, and EtOH. Recrystallise 1g by dissolving it in 3ml of H₂O and adding slowly 15ml of commercial EtOH, filter the crystals, wash with a little EtOH then Et₂O and dry it in a vacuum. Although hydrolysed by boiling 6N HCl (~14 hours), it is rather resistant to boiling strong alkali. [McCormick & Carter *Biochemical Preparations* **2** 65 1952, Posternak *Helv Chim Acta* **12** 1165 1929, DOI: 10.1002/hlca.192901201125] It is involved in *trans-membrane signaling* when bound to lipids.

5-Iodouridine (5-iodo-1-[β-D-ribofuranosyl]-pyrimidine-2,4(1*H*)-dione) [1024-99-3] C₉H₁₁IN₂O₆, M 370.1, m 205-207°(dec), 205-208°(dec), 210-215°(dec), [α]_D²⁰ -23.5 (c 1, H₂O), pK²⁰ 8.5. Recrystallise 5-iodouridine from H₂O and dry it *in vacuo* at 100°. Its UV has λ_{max} at 289nm (0.01N HCl) and 278nm (0.01N NaOH). [Prusoff et al. *Cancer Res* **13** 221 1953, *Beilstein* **24** III/IV 1235.] It can be incorporated into RNA and used for X-ray crystallography of the RNA [Golden et al. *RNA* **2** 1295 1996, PMID: 8972777].

Isopentenyl pyrophosphate [358-71-4] C₅H₁₂O₇P₂, M 246.1, pK_{Est(1)} ~2, pK_{Est(2)} ~2, pK_{Est(3)} ~3.95, pK_{Est(4)} ~6.26. Purify the pyrophosphate by chromatography on Whatman No 1 paper using *tert*-butyl alcohol/formic acid/water (20:5:8, R_F 0.60) or 1-propanol/ammonia/water (6:3:1, R_F 0.48). Alternatively, purify it by chromatography on a DEAE-cellulose column or a Dowex-1 (formate form) ion-exchanger using formic acid and ammonium formate as eluents. A further purification step is to convert it to the *monocyclohexylammonium salt* by passage through a column of Dowex-50 (cyclohexylammonium form) ion-exchange resin. Also converted into its *lithium salt*. It is the primary compound in mevalonate biosynthesis. [Wiemer et al. *Current topics in medicinal chemistry* **10**(18) 1858 2010, PMID: 20615187.]

Kanamycin B (Bekanamycin, 4-*O*-[2,6-diamino-2,6-dideoxy-α-D-glucopyranosyl]-6-*O*-[3-amino-3-deoxy-α-D-glucopyranosyl]-2-deoxystreptamine) [4696-76-8, 29701-07-3 (sulfate salt)] C₁₈H₃₇N₅O₁₀·H₂SO₄, M 483.5 (free base), 587.6 (salt), m 170-179°(dec), 178-182°(dec), [α]_D¹⁸ +130 (c 0.5, H₂O), pK²⁵ 7.2. A small quantity of kanamycin B (24mg) can be purified on a small Dowex-1 x 2 column (6 x 50mm); the required fraction is evaporated to dryness and the residue crystallised from EtOH containing a small amount of H₂O. [Umezawa et al. *Bull Chem Soc Jpn* **42** 537 1969, DOI: 10.1246/bcsj.42.537.] It has been crystallised from H₂O by dissolving ~1g in H₂O (3ml), adding Me₂NCHO (3ml) and setting aside at 4° overnight. The needles are collected and dried to constant weight at 130°. It has also been recrystallised from aqueous EtOH. It is slightly soluble in CHCl₃ and isoPrOH. [IR: Wakazawa et al. *J Antibiot* **14A** 180, 187 1961, Ito et al. *J Antibiot* **17 A** 189 1964; for total synthesis of aminoglycosides see *J Antibiot* **66** 107 2013, DOI: 10.1038/ja.2012.126; *Beilstein* **18** III/IV 7631.] Antibiotic, potent against Gram-positive bacteria, affecting ribosomal RNA synthesis.

Ketamine [2-(2-chlorophenyl)-2-(methylamino)cyclohexanone)] [6740-88-1] C₁₃H₁₆ClNO, M 237.7, m 92-93°, pK²⁵ 7.50. Its preparation has been described by Stevens [US patent 3254124 A to Parke Davis & Co, May 31 1966] which required a rearrangement with ring expansion. The steps in one variant of the synthesis involved preparing *o*-chlorophenyl 1-hydroxycyclopentyl ketone *N*-methylimine [from cyclopentylMgBr + *o*-chlorobenzonitrile → *o*-chlorophenyl cyclopentyl ketone, b 96-97°/0.3mm, n_D²⁰ 1.5452 → *o*-chlorophenyl 1-bromocyclopentyl ketone, b 111-114°/0.1mm + MeNH₂ at 50°/1hr in a bomb → *o*-chlorophenyl 1-hydroxycyclopentyl ketone *N*-methylimine, m 62°) and heating it in excess MeNH₂ in an autoclave at ~200°/5hrs, evaporating to dryness (fume cupboard, and absorb MeNH₂ in aqueous HCl), adding *iso*-PrOH/HCl and isolating *ketamine hydrochloride* [1867-66-9] M 274.2, as white crystals m 262-263° after recrystallisation from EtOH-Et₂O. Its solubility in H₂O is 20w/v% at ~25°. *Ketamine* is obtained by basifying the hydrochloride with dilute aqueous NaOH (~0.5N, to pH >10), extracting with Et₂O, drying, evaporating and recrystallising the residue from pentane-Et₂O. The UV of the neutral species (0.01N NaOH in 95% MeOH, see pK_a above) has λ_{max}(nm)(E_{1%}^{1cm}) at 261(10.7), 268(9.8), 276(7.0) and 301(5.0). [For rearrangement of α-aminoketones see Stevens *J Am Chem Soc* **84** 2272 1962, DOI: 10.1021/ja00870a062].

Optical resolution: Ketamine has one asymmetric centre and has been resolved into its optical isomers *S*(-)-ketamine (*Esketamine*) [33643-46-8] and *R*(+)-ketamine (*Arketamine*) [33643-49-1] which is described by Steiner et al. [patent DE 1961665 A1 to Uetikon Chemie GmbH, Nov 20, 1997; see also Hudyama et al. German Patent 2,062,620 1970, *Chem Abstr* **75** 119119x 1971]. Only the preparation of *Esketamine* is provided here because it is ~3 times more active than its enantiomer, 2 times more active than the *racemate* and is available commercially; the *R*-enantiomer has not been marketed as an enantio pure drug for clinical use. Thus *RS*-ket-

amine (50g, 0.21 mol) in boiling Me₂CO (613ml) is treated with L(+)-tartaric acid (31.5g, 0.21 mol), and to obtain a clear solution H₂O (40ml) is added at the boiling point and the hot solution is filtered. After seeding, the solution is cooled to ~25°, the crystals are filtered off, dried first at ~25° then at 50-60° to give the crude (-)-base-(+) acid diastereomeric tartrate [**m 161°**, $[\alpha]_D^{20} +26.1$ (c 2, H₂O)]. This is recrystallised from a hot Me₂CO (1226ml) and H₂O (90ml) mixture, which on cooling to ~25° with stirring for 4 hours gives pure diastereomeric tartrate (38.8g, 95%) after drying as before, which has m 175.3°, $[\alpha]_D^{20} +68.9$ (c 2, H₂O). The (-)-base-(+) acid salt (38.8g) in aqueous NaOH (420ml, pH>10) is stirred with Et₂O (540ml), the ethereal extract is washed with H₂O and brine, dried (Na₂SO₄), filtered and evaporated to dryness to give colourless crystals [21.5g, 86%; m **118.9°**, $[\alpha]_D^{20} -55.8$ (c 2, H₂O)]. The crystals (10.75g) are recrystallised from hot cyclohexane (43ml), which after stirring with cooling at 10° during 1 hour gave pure *S*(-)-ketamine (10.3g, 96%) with m **120°** (also **120-122°**) and $[\alpha]_D^{20} -56.8$ (c 2, EtOH).

When *S*(-)-ketamine (5g) is dissolved in *iso*-PrOH (50ml) at ~50°, filtered through kieselguhr, then HCl gas is passed through the solution at 50-60° until the pH is 0-1 and cooled to ~20°; the crystals are collected washed with cold *iso*-PrOH (5ml), dried overnight in an oven at 50° provided *pure S*(+)-ketamine hydrochloride [33795-24-3] (5.09g, 88%) $[\alpha]_D^{25} +92.5$ (c 2, H₂O). [Note change in sign of the optical rotation].

R(+)-ketamine and *R*(-)-ketamine hydrochloride [33795-24-3] can be obtained in a similar way but using D(+)-tartaric acid. There is much confusion in the literature about the absolute configuration most probably because the free base has the opposite sign of rotation from that of its hydrochloride, and because it is the (+)-hydrochloride that is generally used as the drug since it is 3 times more active than the (-)-hydrochloride. Also when the name Ketamine is used, it is the *hydrochloride drug* that is usually intended. This was put to rest when Groth and coworkers [Ratti-Moberg *Acta Chem Scand* **45** 108 1991, DOI: 10.3891/acta.chem.scand.45-0108] determined the crystal structure of the *R,R*-tartrate salt of levo-ketamine showing that (-)-ketamine has the *S*- absolute configuration, and that its hydrochloride has a positive rotation for the sodium D line(s) at ~25°. Ketamine is a controlled anti-depressant drug that requires a medical prescription for use. It is an anaesthetic (dissociative) and a *non-competitive NMDA* (*N*-methyl-D-Aspartate) *receptor antagonist*, with the *S*(+)-HCl having three times the activity of its enantiomer. [Hashimoto in Zhang et al. *Pharmacol Biochem Behavior* **116** 137 2014, DOI: 10.1016/j.pbb.2013.11.033; review: Morgan et al. *Addiction* **107** 27 2012, DOI: 10.1111/j.1360-0443.2011.03576.x; Anis et al. *Br J Pharmacol* **79** 565 1983, PMCID: 2044888; Liu et al. *Anesth Analg* **92** 1173 2001, PMID: 11323343.]

Lansoprazole {2-[4-(2,2,2-trifluoroethoxy)-3-methyl-2-pyridylmethylsulfanyl]-1*H*-benzimidazole, Zoton, Takepron) [103577-45-3] C₁₆H₁₄F₃N₃O₂S, M369.4, m 178-182°(dec), pKa similar to Omeprazole. It is a drug related to *Omeprazole* (see below) with similar structure except that the 5-methyl group is missing, the 4-methoxy group is replaced by a 2,2,2-trifluoroethoxy group in the pyridine ring and the 5-methoxy group is missing in the benzimidazole ring. Although it exhibits similar properties to Omeprazole, Katsuki et al. [*Pharm Res* **13** 611-615 1996, DOI: 10.1023/A:1016062508580] found that the enantiomers of Lansoprazole show different pharmacokinetic profiles in humans. The *R*(+) isomer is more effective as a *proton pump inhibitor* due to its higher bioavailability and less adverse side effects.

Racemic Lansoprazole can be optically resolved, via the *diastereomeric* (1:2) inclusion compounds with *R*(+)-1,1,2-triphenyl-1,2-ethane diol. Thus, according to the procedure of Sun Young Jan et al. [WO2010068049 A2, 2010], a mixture of (±)-Lansoprazole (40.0g, 1mol) and *R*(+)-1,1,2-triphenyl-1,2-ethane diol (**TED**, [9506-46-4] 62.9g, 2mol) suspended in CHCl₃ (640ml) are boiled under reflux until a clear solution is formed, then slowly cooled to ~25° followed by stirring for 4 hours. The precipitate formed is filtered off, washed with cold CHCl₃, and dried at 40° to give the (+)(+)-1:2-inclusion complex (41.1g 80%, 97.2%ee) as a white crystalline powder, m **147.8-149.9°**, $[\alpha]_D^{25} +174.32$ (c 1, MeOH) 97.2% optical purity. Purification to >99.9% optical purity can be achieved by suspending the product (15g) in CHCl₃ (120ml) refluxing until clear, then hexane (180ml) is added, cooled slowly to ~25°, stirred for 1 hour, and the precipitate is collected, and dried at 40° to give *pure* (+)(+)-1:2-inclusion complex (13g, 87%, 99.9%ee) as a white crystalline powder, m **149.8-151.5°**, $[\alpha]_D^{25} +182.76$ (c 1, MeOH). Its IR (KBr) and ¹H NMR (CDCl₃) have been reported. Other solvents used for complex formation are MeOH/di-disopropyl ether (20:10, yield 70%, 96%ee), EtOH/di-disopropyl ether (20:10, yield 76%, 94.5%ee), MeOH/H₂O (25:5, yield 62%, 97.3%ee), *iso*PrOH/H₂O (25:5, yield 78%, 94.5%ee), Me₂CO/H₂O (20:5, yield 70%, 95.2%ee), and MeCN/H₂O (20:5, yield 62%, 95.5%ee).

R(+)-Lansoprazole is obtained from the (+)(+)-inclusion complex above (10.0g, 10.5mmol, >99.9%ee) by dis-

solving in CH_2Cl_2 (155ml), adding 0.2N KOH (105ml, 21.0mmol) and stirring. The aqueous layer is collected, diluted with MeOH (50ml) and the pH of the mixture is adjusted to 6.5 with AcOH, then stirred for 2 hours at $\sim 25^\circ$. The precipitate is collected, washed with H_2O , then dissolved in EtOAc (80ml); hexane (80ml) is added, stirred for 1hr and the crystalline precipitate is collected, dried at 40° to give **pure *R*-(+)-Lansoprazole $1.5\text{H}_2\text{O}$** (3.4g, 89%, 99.9%ee). This has **m 80-90 $^\circ$, $[\alpha]_{\text{D}}^{25} +154.7$ (c 1, CHCl_3)** [H_2O content 6.8%]; and its IR (KBr) and ^1H NMR (CDCl_3) have been reported. Also *R*-(+)-TED can be recovered by washing the above CH_2Cl_2 layer with H_2O (100ml), drying (anhydrous MgSO_4), filtering, evaporating the solvent to dryness and recrystallising the residual solid from EtOAc/hexane (1:10 v/v) to give *R*-(+)-1,1,2-triphenyl-1,2-ethanediol (5.5%, 90% recovery), **m 127-130 $^\circ$, $[\alpha]_{\text{D}}^{25} +205.5$ (c 1, MeOH, 99.8%ee)**; its IR (KBr) have been reported. Alternatively, to obtain the **anhydrous** drug, the (+)(+)-inclusion complex (10.0g, 10.5mmol, >99.9%ee) is dissolved in CH_2Cl_2 (155ml), stirred with 0.25N NaOH (105ml, 21.0mmol), the aqueous layer is isolated, mixed with EtOAc (200ml), the pH of the mixture is adjusted to 7.0 with hydrochloric acid whereby the neutral species remain in the EtOAc layer which is washed with brine (50ml), dried (MgSO_4), and evaporated to provide the **anhydrous *R*-(+)-drug** (3.5g, 91%) as an **amorphous powder m 76.8-77.2 $^\circ$, $[\alpha]_{\text{D}}^{25} \sim +156$ (c 1, MeOH, 99.9%ee)**.

NOTE: that from the original resolution the (-)(+)-inclusion complex of high optical purity can be isolated, and optically **pure *S*-(-)-Lansoprazole** can be obtained; **OR better**, by using the *S*-(-)-1,1,2-triphenyl-1,2-ethanediol [108998-83-0] $\text{C}_{20}\text{H}_{18}\text{O}_2$, M 290.4, **m 125-127 $^\circ$, $[\alpha]_{\text{D}}^{20} -214$ (c 1, EtOH)** as resolving agent, the **(-)(-)-1,2-inclusion complex** can be produced from which the ***S*-(-)-drug** can be obtained using exactly the same quantities of reagents and solvents as used for their antipodes above, without the necessity of exploring new reagents and solvents.

[For mechanism of action of Lansoprazole see also Nagaya et al. *J Pharmacol Exp Ther* **252** 1289 1990, PMID: 2156997; for the synthesis of the (\pm)-drug see Nohara & Maki to Takeda US Pat 4628098 A and EP 174726; and Clarke & Slemmon US Pat 5470983 A 1995 above; and for the process of preparing *R*-(+)-1,1,2-triphenyl-1,2-ethane diol see Sun et al. to Hanmi Pharm Co WO 2010068049 A2 and A3 2010.]

Lecithin (1,2-diacylphosphatidylcholine mixture) [8002-43-5] **M ~600-800, amorphous**. Lecithin from hen egg white is purified by solvent extraction and chromatography on alumina. It is suspended in H_2O and kept frozen until required [Lee & Hurst *J Am Chem Soc* **106** 7411 1984, DOI: 10.1021/ja00336a020; Singleton et al. *J Am Oil Chem Soc* **42** 53 1965, DOI: 10.1007/BF02558256]. For purification of commercial egg lecithin, see Pangborn [*J Biol Chem* **188** 471 1951, <http://www.jbc.org/content/188/2/471>; PMID: 14824134].

Leucopterin (2-amino-5,8-dihydropteridine-4,6,7(1H)-trione) [492-11-5] $\text{C}_6\text{H}_5\text{N}_5\text{O}_3$, **M 195.1, m >300 $^\circ$ (dec), $\text{pK}_1^{20} -1.66$, $\text{pK}_2^{20} 7.56$, $\text{pK}_3^{20} 9.78$, $\text{pK}_4^{20} 13.6$** . Leucopterin is purified by dissolving in aqueous NaOH, stirring with charcoal, filtering and precipitating by adding aqueous HCl, then drying at 100° in a vacuum. It separates with 0.5 mole of H_2O . Its solubility in H_2O is 1g/750 litres [Albert et al. *J Chem Soc* 4219 1952, DOI: 10.1039/JR9520004219; Albert & Wood *J Appl Chem (UK)* **2** 591 1952, DOI: 10.1002/jctb.5010021005; Pfeleiderer *Chem Ber* **90** 2631 1957, DOI: 10.1002/cber.19570901129]. [*Beilstein* **26** III/IV 4017.]

DL- α -Lipoamide (\pm -6,8-thioctic acid amide, 5-[1,2]-dithiolan-3-ylvaleric acid amide) [940-69-2; 3206-73-3] $\text{C}_8\text{H}_{15}\text{NOS}_2$, **M 205.3, 242.2, m 124-126 $^\circ$, 126-129 $^\circ$, 130-131 $^\circ$** . DL- α -Lipoamide is recrystallised from EtOH and has UV with λ_{max} at 331nm in MeOH. [Reed et al. *J Biol Chem* **232** 143 1958, PMID: 13549405; IR: Wagner et al. *J Am Chem Soc* **78** 5079 1956, DOI: 10.1021/ja01600a069; *Beilstein* **19/7** V 238.] **IRRITANT**.

DL- α -Lipoic acid (\pm -6,8-thioctic acid, 5-[1,2]-dithiolan-3-ylvaleric acid) [1077-28-7] $\text{C}_8\text{H}_{14}\text{O}_2\text{S}_2$, **M 206.3, m 59-61 $^\circ$, 60.5-61.5 $^\circ$ and 62-63 $^\circ$, b 90 $^\circ$ /10 $^{-4}$ mm, 150 $^\circ$ /0.1mm, $\text{pK}^{25} 4.7$** . It forms yellow needles from cyclohexane or hexane, has been distilled at high vacuum, and sublimes at $\sim 90^\circ$ and very high vacuum. It is insoluble in H_2O but dissolves in alkaline solution. [Lewis & Raphael *J Chem Soc* 4253 4263 1962, DOI: 10.1039/JR9620004253; Soper et al. *J Am Chem Soc* **76** 4109, DOI: 10.1021/ja01645a016; Reed & Niu *J Am Chem Soc* **77** 416 1955, DOI: 10.1021/ja01607a057; Tsuji et al. *J Org Chem* **43** 3606 1978, DOI: 10.1021/jo00412a044; Calvin *Fed Proc USA* **13** 697 1954, PMID: 13210461.] The ***S*-benzylisothiuronium salt** has **m 153-154 $^\circ$** (evacuated capillary, from MeOH), 132-134 $^\circ$, 135-137 $^\circ$ (from EtOH). The ***d*- and *l*-forms** have **m 45-47.5 $^\circ$ and $[\alpha]_{\text{D}}^{23} \pm 113$ (c 1.88, $^*\text{C}_6\text{H}_6$)**; and have UV in MeOH with λ_{max} at 330nm (ϵ 140). [*Beilstein* **19/7** V 237.] The reduced form, (\pm)-6,8-dimercaptooctanoic acid, [7516-48-5] **M 208.3**, is a light

yellow liquid which is sold in sealed ampoules.

D-Luciferin (firefly luciferin, *S*-2[6-hydroxybenzothiazol-2-yl]-4,5-dihydrothiazol-4-carboxylic acid), [2591-17-5] $C_{11}H_{18}N_2O_3S_2$, **M 280.3**, **m 189.5-190°(dec)**, **196°(dec)**, **201-204°**, **205-210°(dec)**, **browning at 170°**, $[\alpha]_D^{22}$ -36 (c 1.2, DMF), $pK_{Est(1)} \sim 1.2$ (benzothiazole-N), $pK_{Est(2)} \sim 1.6$ (thiazolidine-N), $pK_{Est(3)} \sim 6.0$ (CO_2H), $pK_{Est(4)} 8.5$ (6OH). D-Luciferin crystallises as pale yellow needles from H_2O , or MeOH (83mg/7ml). It has UV with λ_{max} at 263 and 327nm (log ϵ 3.88 and 4.27) in 95% EtOH; and fluorescence (H_2O) with λ_{ex} 328nm and λ_{em} 532nm. The **Na salt** has a solubility of 4mg in 1 ml of 0.05M glycine. Store it at -20°. [White et al. *J Am Chem Soc* **83** 2402 1961, DOI: 10.1021/ja01471a051; **85** 337 1963, DOI: 10.1021/ja00886a019; for UV and IR see Bitler & McElroy *Arch Biochem* **72** 358 1957, DOI: 10.1016/0003-9861(57)90212-6; Review: Cormier et al. *Fortschr Chem Org Naturst* **30** 1 1973, PMID: 4156520; *Beilstein* **27** III/IV 8934.] It is a substrate for *firefly luciferase* ($K_m \sim 2\mu M$) being involved in bioluminescence, and is a useful biological tag.

Lumichrome (7,8-dimethylalloxazine) [1086-80-2] **M 242.2**, **m >290°**, $pK_{Est(1)} \sim -0.1$ (basic), $pK_{Est(2)} \sim 9.9$ (acidic). Recrystallise lumichrome twice from glacial AcOH and dry it at 100° in a vacuum. [Cresswell & Wood *J Chem Soc* 4768 1960, DOI: 10.1039/JR9600004768; *Beilstein* **26** III/IV 2538.]

Lumiflavin (7,8,10-trimethylbenzo[g]pteridine-2,4(3*H*,10*H*)-dione) [1088-56-8] $C_{13}H_{12}N_4O_2$, **M 256.3**, **m 330°(dec)**, **340°(dec)**, $pK^{25} 10.2$. Lumiflavin forms orange crystals upon recrystallisation from 12% aqueous AcOH, or from formic acid. It sublimes at high vacuum. It is freely soluble in $CHCl_3$, but not very soluble in H_2O and most organic solvents. In H_2O and $CHCl_3$ it has a green fluorescence. Its UV has λ_{max} at 269, 355 and 445nm (ϵ 38,800, 11,700 and 11,800, respectively) in 0.1N NaOH, and 264, 373 and 440nm (ϵ 34,700, 11,400 and 10,400, respectively) in 0.1N HCl, while the UV in $CHCl_3$ has λ_{max} at 270, 312, 341, 360, 420, 445 and 470nm. [Hemmerich et al. *Helv Chim Acta* **39** 1242 1956, DOI: 10.1002/hlca.19560390511; Holiday & Stern *Chem Ber* **67** 1352 1934, DOI: 10.1002/cber.19340670812; Yoneda et al. *Chem Pharm Bull Jpn* **20** 1832 1972, DOI: org/10.1248/cpb.20.1832; Birch & Moye *J Chem Soc* 2622 1958, DOI: 10.1039/JR9580002622; Kuhn & Moruzzi *Chem Ber* 67 888 1934, DOI: 10.1002/cber.19340670539; *Beilstein* **26** III/IV 2539.]

Magnesium protoporphyrin dimethyl ester [14724-63-1] $C_{36}H_{36}N_4O_4Mg$, **M 580.7**, **m 214-217°** (others found **m 228-230°**). The Mg complex can be prepared from protoporphyrin dimethyl ester (see below) and $Mg(ClO_4)_2$ in boiling pyridine under N_2 for 3-4 hours until the band at 630nm (free porphyrin) is absent. Filter, wash the insolubles with Et_2O until the filtrate is colourless. Evaporate the solvent *in vacuo* at 50-60° to a very small volume, then add excess peroxide free Et_2O in a separating funnel, shake with H_2O (2x), evaporate the Et_2O , and remove the H_2O and pyridine by evaporating (azeotropically) with $*C_6H_6$ (4x). Purify the residue by dissolving the Mg complex in as little hot $*C_6H_6$ (50ml for 800mg) as possible, and add cold petroleum ether (b 30-60°), leave at room temperature until crystallisation begins, then further in a refrigerator to give twinned prisms, **m >330°**. Its IR has ν_{max} at 3080(w $CH=CH_2$), 1610, 1698, 1740 (s CO_2Me) cm^{-1} , but no NH. Its UV (Et_2O) has λ_{max} (ϵ) at 588nm (19,000), 550nm (19,300), 417nm (Soret, 252,000) [and 340nm (20,550)]. [Ramsey *Biochemical Preparations* **3** 39 1953, Fuhrhop & Granick *Biochemical Preparations* **13** 55 1971.] Even when pure, Mg protoporphyrin dimethyl ester is difficult to crystallise although crystals can be found to form on a glass slide under a microscope. For analysis, a portion of the pink powder is washed on the filter with dry Et_2O . The filtrate consists of a colloidal solution (a slight residue remains on the filter paper). To the filtrate is now added low boiling light petroleum and a precipitate of plates and highly twinned crystals results. These are centrifuged down, washed with this petroleum ether and dried *in vacuo*. The UV of the red solution in Et_2O is the same as above. [Granick *J Biol Chem* **175** 333 1948, PMID: 18873307.]

6-Mercaptopurine-9- β -D-ribofuranoside [574-25-4] $C_{10}H_{12}N_4O_4S$, **M 284.3**, **m 208-210°(dec)**, **210-211°(dec)**, **221-223°(dec)**, **220-223°(dec)**, **222-224°(dec)**, $[\alpha]_D^{25}$ -73 (c 1, 0.1N NaOH), $pK^{25} 7.56$. Recrystallise the riboside from hot H_2O or EtOH. It has UV with λ_{max} (H_2O) at 322nm (pH 1), 320 nm (pH 6.7) and 310nm (pH 13). It is soluble in DMSO. [IR: Johnson et al. *J Am Chem Soc* **80** 699 1958, DOI: 10.1021/ja01536a044; UV: Fox et al. *J Am Chem Soc* **80** 1669 1958, DOI: 10.1021/ja01540a041; *Beilstein* **26** III/IV 2100.] It is a substrate for adenosine deaminase.

R(-)-Methadone (Levomethadone, 6-dimethylamino-4,4-diphenylheptan-3-one) [125-58-6] $C_{21}H_{27}NO$, **M 309.4**, **m 98-100°**, $[\alpha]_D^{20} -32$ (c 1.8, EtOH), see below for pKa. This pharmacologically active (against narcotic addiction) enantiomer was obtained by optical resolution (using D-tartaric acid) of the racemate, and was purified by precipitation of the hydrochloride from aqueous solution at pH <6, dried and recrystallised from propan-2-ol. The **R-hydrochloride** [5967-73-7], when recrystallised from propan-2-ol, has **m 245-246°**, and $[\alpha]_D^{20} -169$ (c 2, EtOH). The **S-(+)-enantiomer** [5653-8-5] also recrystallises from propan-2-ol and has a recorded **m** of **100-101°**, and $[\alpha]_D^{25} +26$ (c 1.2, H₂O). The **S-hydrochloride** [15284-15-8], when crystallised from propan-2-ol, has **m 243-244°**, and $[\alpha]_D^{20} +169$ (c 2, EtOH). [Larsen et al. *J Am Chem Soc* **70** 4194 1948, DOI: 10.1021/ja01192a065; Brode & Hill *J Org Chem* **13** 191 1948, DOI: 10.1021/jo01160a004; Schultz et al. *J Am Chem Soc* **69** 2454 1947, DOI: 10.1021/ja01202a061; Easton et al. *J Am Chem Soc* **69** 2941 1947, DOI: 10.1021/ja01204a006; Winter & Flataker *J Pharmacol Exp Ther* **98** 305 1950, PMID: 15428991; Beilstein **14** III 278, **14** III/IV 300.] [Strain & Stitzer *The Treatment of Opioid Dependence* JHU Press p 63 2005, ISBN 978-0-8018-8303-3].

(±)-Methadone hydrochloride (6-dimethylamino-4,4-diphenylheptan-3-one HCl) [1095-90-5] $C_{21}H_{27}NO \cdot HCl$, **M 345.9**, has **m 241-242°**, $pK_1^{25} 8.94$, $pK_2^{20} 10.12$ (free base). The salt (see above) crystallises from EtOH, or EtOH/Et₂O. [see methadone references.]

Methoxantin coenzyme (PQQ, pyrrolo quinoline quinone, 2,7,9-tricarboxy-1H-pyrrolo-[2,3-f]-quinoline-4,5-dione, 4,5-dihydro-4,5-dioxo-1H-pyrrolo[2,3-f]quinoline-2,7,9-tri-carboxylic acid) [72909-34-3] $C_{14}H_6N_2O_8$, **M 330.2**, **m 220°(dec)**. Efflorescent yellow-orange needles of PQQ are formed on recrystallising from H₂O by addition of Me₂CO, or better from a supersaturated aqueous solution, as it forms an **acetone adduct**. [Salisbury et al. *Nature* **280** 843 1979, DOI: 10.1038/280843a0.] It has also been purified by passage through a C-18 reverse phase silica cartridge or a silanised silica gel column in aqueous solution whereby methoxantin remains behind as a **red-orange band** at the origin. This band is collected and washed thoroughly with dilute aqueous HCl (pH 2) and is then eluted with MeOH/H₂O (7:3) and evaporated *in vacuo* to give the coenzyme as a red solid. It has also been purified by dissolving it in aqueous 0.5M K₂CO₃ and acidifying to pH 2.5 whereby PQQ precipitates as a deep red solid which is collected and dried *in vacuo*. Methoxantin elutes at 3.55 retention volumes from a C18 μ Bondapak column using H₂O/MeOH (95:5) + 0.1% AcOH pH 4.5. It has UV with λ_{max} at 247 and 330nm (shoulder at 270nm) in H₂O, and λ_{max} at 250 and 340nm in H₂O at pH 2.5. With excitation at λ_{ex} 365nm it has a λ_{em} at 483nm. The ¹³C NMR has δ : 113.86, 122.76, 125.97, 127.71, 130.68, 137.60, 144.63, 146.41, 147.62, 161.25, 165.48, 166.45, 173.30 and 180.00.

When **dissolved in** a solution of 10% aqueous MeCO and adjusted to pH 9 with aqueous NH₃ then kept at 25° for 30 minutes, the **acetone adduct** is formed, which has UV with λ_{max} at 250, 317 and 360nm (H₂O, pH 5.5), and with λ_{ex} at 360nm it fluorescence with λ_{em} at 465nm; and the ¹³C NMR [(CD₃)₂SO, TMS] has been reported. It also forms a **methanol adduct**.

When it is reacted with Me₂SO₄/K₂CO₃ in dry Me₂NCHO at 80° for 4 hours, it forms the **trimethyl ester** which has **m 265-267°(dec)** [**260-263°(dec)** also reported] and after recrystallisation from hot MeCN it forms orange crystals with UV that has λ_{max} at 252 and 344nm (H₂O), and 251, 321 and 373nm (in MeOH; MeOH adduct?). [Duine et al. *Eur J Biochem* **108** 187 1980, DOI: 10.1111/j.1432-1033.1980.tb04711.x; Duine et al. *Adv Enzymology* **59** 169 1987, DOI: 10.1002/9780470123058.ch4; Corey & Tramontano *J Am Chem Soc* **103** 5599 1981, DOI: 10.1021/ja00408a067; Gainor & Weinreb *J Org Chem* **46** 4317 1981, DOI: 10.1021/jo00334a053; Hendrickson & de Vries *J Org Chem* **47** 1148 1982, DOI: 10.1021/jo00345a057; MacKenzie et al. *JCS Chem Commun* 1372 1983, DOI: 10.1039/C39830001372.]

Methyl benzylpenicillinate [653-89-4] $C_{17}H_{20}N_2O_4S$, **M 336.4**, **m 97°**, $[\alpha]_D^{20} +328$ (c 1, MeOH), $[\alpha]_D^{20} +286$ (c 1, CHCl₃). Crystallise the ester once from Et₂O, and once from CCl₄ or EtOAc/hexane. [Sheehan & Henery-Logan *J Am Chem Soc* **84** 2983 1962, DOI: 10.1021/ja00874a029; Review: Chain et al. *Antibiotics* (Oxford University Press) **2** 1949; Kumler et al. *J Am Chem Soc* **71** 3382 1949, DOI: 10.1021/ja01178a032]

1-Methylimidazolidine-2,4-dione (1N-methylhydantoin, MH) [616-04-6] $C_4H_6N_2O_2$, **M 114.1**, **m 158°, 155-156°, 157-159°, and 184-185° (dimorphic ?)**, $pK^{25} 9.20$. It is purified by dissolving it in the minimum volume of H₂O, extracting with EtOAc, the extract is dried and evaporated to dryness. The residue, in EtOAc is subjected to column chromatography (silica gel with EtOAc as eluent), or silica gel TLC [MeOH/CHCl₃ (1:9)] and recrystallised from EtOH (elongated plates). It is identified by ¹H NMR and MS. The IR has ν_{max} at 1712

(2CO) and 1761 (4CO) cm^{-1} . The **3-acetate** has **m 134-135°**. [Miller & Robson *J Chem Soc* 1910 1938, DOI: 10.1039/JR9380001910; West *J Biol Chem* **34** 187 1918, <http://www.jbc.org/content/34/1/187>; Ienaga et al. *Tetrahedron Lett* **28** 4587 1987, Ienaga et al. *JCS Perkin Trans I* 1153 1989, DOI: 10.1039/P19890001153; Beilstein **24** H 44.] Data by Dr Kazu Ienaga, Nippon Zoki Pharmaceutical Co Ltd, Osaka, Japan.

(±)-5-Methylimidazoline-2,4-dione (5-methylhydantoin) [616-03-5] $\text{C}_4\text{H}_6\text{N}_2\text{O}_2$, **M 114.1**, **m 145°, 148-152°, 149-151°, 150°, pK²⁵ 9.07**. Purify it by recrystallisation from H_2O , and dry it *in vacuo*. [Ienaga et al. *Tetrahedron Lett* **28** 4587 1987, DOI: 10.1016/S0040-4039(00)96571-7; Ienaga et al. *JCS Perkin Trans I* 1153 1989, DOI: 10.1039/P19890001153; IR: Burland & Christian *Can J Chem* **35** 444 1957, DOI: 10.1139/v57-064; Beilstein **24** H 279, **24** I 305, **24** II 155, **24** III/IV 1083.] Data supplied by Dr Kazu Ienaga (see above).

5-Methyltetrahydrofolic acid disodium salt (prefolic A) [68792-52-9] $\text{C}_{20}\text{H}_{23}\text{N}_7\text{O}_6 \cdot 2\text{Na}$, **M 503.4**, **pK₁ 2.4 (N10 protonation), pK₂ 2.7 (pyrimidine N1 protonation), pK₃ 3.5 ($\alpha\text{-CO}_2\text{H}$), pK₄ 4.9 ($\gamma\text{-CO}_2\text{H}$), pK₅ 5.6 (N5-Me), pK₆ 8.5 (3NHCO acidic)**. First check the purity by measuring the UV at pH 7.0 (use phosphate buffer), and it should have λ_{max} at 290nm and λ_{min} at 245nm with a ratio of A_{290}/A_{250} of 3.7. This ratio goes down to 1.3 as oxidation to the dihydro derivative occurs. The latter can be reduced back to the tetrahydro compound by reaction with 2-mercaptoethanol at room temperature. If oxidation has occurred, then the compound should be chromatographed on DEAE-cellulose (~0.9 milliequiv/g, in AcO^- form) in $(\text{NH}_4)_2\text{CO}_3$ (1.5 M) and washed with 1M NH_4OAc containing 0.01M mercaptoethanol till free from UV absorption and then washed with 0.01M mercaptoethanol. All is done in a nitrogen atmosphere. The reduced folate is then eluted with a gradient between 0.01M mercaptoethanol and 1M NH_4OAc containing 0.01M mercaptoethanol, and the fractions with absorption at 290nm are collected. These are evaporated under reduced pressure at 25°, and traces of NH_4OAc and H_2O are removed at high vacuum/25° (~24-48 hours). The residue is dissolved in the minimum volume of 0.01M mercaptoethanol, and an equivalent of NaOH is added to convert the acid to the diNa salt and evaporated to dryness at 25°/high vacuum. The **pure product** has λ_{max} 290nm (ϵ 32,000) in pH 7.0 buffer. [Sakami *Biochemical Preparations* **10** 103 1963; for a convenient preparation of (±)-5-MeTHF and its Ca and Ba salts in quantity see Blair & Saunders *Analyt Biochem* **34** 376 1970, DOI: 10.1016/0003-2697(70)90122-3] It is a biological methylating agent [Banerjee & Snyder *Science* **182** 74 1973, DOI: 10.1126/science.182.4107.74].

Metoclopramide [Reglan, **4-amino-5-chloro-2-methoxy-N-(β-diethylaminoethyl)benzamide**] [364-62-5] $\text{C}_{14}\text{H}_{22}\text{ClN}_3\text{O}_2$, **M 299.8**, **m 146.5-148°, 147.3°, pK₁ 0.6 (ArNH), pK₂ 9.3 (Alkyl-tert-N)**. It can be prepared from 5-chloro-2-methoxy-4-nitrobenzoic acid [35633-81-4], in several steps from *o*-toluidine, see e.g. Büchi et al. *Helv Chim Acta* **34** 1002 1951, DOI: 10.1002/hlca.19510340404). It is converted to the acid chloride which is reacted with *N,N*-diethylethylenediamine to give the respective *N*-(diethylaminoethyl) 5-chloro-2-methoxy-4-nitrobenzamide which is reduced catalytically to **Metoclopramide**. [Thominet US Patent 3177252, 1965 to Soc d'Etudes Sci Ind de l'Ile-de-France.] This base is purified by recrystallisation from aqueous EtOH containing a small amount of NH_3 in order to avoid contact with CO_2 (to form a carbonate salt as it is a strong base) and drying in a desiccator over solid KOH. The solubility of the **free base** in w/v at ~25° in H_2O is 0.01%, in $^*\text{C}_6\text{H}_6$ is 0.10%, in absolute EtOH is 1.90%, in 95% aq EtOH is 2.30% and in CHCl_3 is 6.60%. It is best to convert it into the **monohydrochloride monohydrate (Clopromate, Maxolon, Meclopran)** [54143-57-6] which is a white crystalline solid **m 182.5-184°** obtained by dissolving the **free base** in EtOH (3 volumes) to which is added 5N alcoholic HCl (1.5 equivalents), filtering off the salt and drying it in a vacuum desiccator over KOH. Similarly, the **dihydrochloride monohydrate (Emetid, Gastronerton, Primperan)** [7232-21-5] **m 145°(dec)**, is prepared as for the **monohydrochloride** except that a large excess of HCl is added. It is stable in acidic media but unstable in strong alkaline solutions. The solubility in w/v at ~25° in CHCl_3 is 0.10%, in $^*\text{C}_6\text{H}_6$ is 0.10%, in absolute EtOH is 6.0%, in 95% aqueous EtOH is 9.0% and in H_2O it is 48.0%.

Metoclopramide is a useful **anti-emetic**, and used for alleviating nausea (also in radiation sickness), vomiting and gastric reflux. It appears to bind to dopamine D_2 receptors with nanomolar affinity (K_i 28.8 nM), and is a **mixed 5-HT₃ receptor antagonist/5-HT₄ receptor agonist**. [Justin-Besançon & Laville *Comptes Rendus des Séances de la Société de Biologie et de ses Filiales* (in French) **158** 723 1964. PMID: 14186927; review: Hibbs & Lorch *Pediatrics* **118** 746 2006, PMID: 16882832; Keabian & Calne *Nature* **277** 93 1979, DOI: 10.1038/277093a0; Tonini et al. *Pharmacol Res* **31** 257 1995, PMID: 7479521.]

(±)-Mevalonic acid lactone [674-26-0] $\text{C}_6\text{H}_{10}\text{O}_3$, **M 130.2**, **m 28°, b 145-150°/5mm**. Purify the lactone *via*

the *dibenzyl-ethylenediammonium salt* (**m** 124-125°) [Hofmann et al. *J Am Chem Soc* **79** 2316 1957, DOI: 10.1021/ja01566a080], or by chromatography on paper or on a Dowex-1 (formate) column. [Bloch et al. *J Biol Chem* **234** 2595 1959, PMID: 13801508.] Store it as the *N,N'*-dibenzylethylenediamine (DBED) salt, or as the lactone in a sealed container at 0°. [Beilstein **18/1** V 19.]

Mevalonic acid 5-phosphate [1189-94-2] $C_6H_{13}O_7P$, **M** 228.1, $pK_{Est(1)} \sim 1.5$ (PO_4H_2), $pK_{Est(2)} \sim 4.4$ (CO_2H), $pK_{Est(3)} \sim 6.31$ (PO_4H^-). Purify the acid by conversion to the *tricyclohexylammonium salt* (**m** 154-156°) by treatment with cyclohexylamine. Recrystallise the salt from water/acetone at -15°. Alternatively, the phosphate is chromatographed through an ion-exchange resin or paper (Whatman No 1) in a system of isobutyric acid/ammonia/water (66:3:30; R_F 0.42). See following entry. Store it as the cyclohexylammonium salt.

Mevalonic acid 5-pyrophosphate [1492-08-6] $C_6H_{14}O_{10}P_2$, **M** 308.1, $pK_{Est(1)} \sim 2$, $pK_{Est(2)} \sim 2$, $pK_{Est(3)} \sim 3.95$ (PO_4), $pK_{Est(4)} \sim 4.4$ (CO_2H), $pK_{Est(5)} \sim 6.26$ (PO_4). Purify the pyrophosphate by ion-exchange chromatography on Dowex-1 formate [Bloch et al. *J Biol Chem* **234** 2595 1959, PMID: 13801508], DEAE-cellulose [Skilletar and Kekwick, *Anal Biochem* **20** 171 1967, DOI: 10.1016/0003-2697(67)90275-8], or by paper chromatography [Rogers et al. *Biochem J* **99** 381 1966, DOI: 10.1042/bj0990381]. Likely impurities are ATP and mevalonic acid phosphate. Store it as a dry powder or in a slightly alkaline (pH 7-9) solution at -20°.

Mexiletine hydrochloride [Mexitil, MEX, 1-(2,6-dimethylphenoxy)-2-propanamine hydrochloride] [5370-01-4] $C_{11}H_{17}NO$. HCl, **M** 215.7, **m** 203-205°, **pKa** 9.0. The synthesis of the *racemate* is described in a patent by Köppe et al. [US 3954872 1976 to Boehringer] where starting from 2,6-dimethylphenol, the 1-(2,4-dimethylphenoxy)propan-2-one oxime is obtained and reduced catalytically with Raney Ni in MeOH at 60°/5 atmospheres in a pressure bomb until the desired volume of H_2 is absorbed, cooled, filtered and evaporated to dryness to give the *free base* [31828-71-4] as an oil. This oil is dissolved in EtOH and acidified with ethereal HCl and cooled; the solid is collected and recrystallised by dissolving in the minimum volume of hot EtOH and adding excess dry Et_2O to give **Mexiletine hydrochloride m** 203-205° (51.5% yield) as off-white crystals with 1H NMR (200MHz, $DMSO-d_6$, TMS) which has δ at 1.35 (d, 3H, $J = 6.6$ Hz, CH_3-CH), 2.23 (s, 6H, 2 CH_3-Ar), 3.5-3.58 (m, 1H, $CH_2-CH-CH_3$), 3.81 (d, 2H, $J = 5.2$ Hz, $O-CH_2-CH$), 6.87-7.04 (m, 3H, $Ar-H$), 8.49 (br s, 3H, NH_3^+) ppm [from *Chirality* **6** 590 1994, DOI: 10.1002/chir.530060713]. The compound has one asymmetric centre and could be resolved into its optical antipodes using di-*O*-benzoyl-L-tartaric acid or D-3-bromocamphor-8-sulfonic acid. The (-)-*R*-mexiletine has been found to be more potent than the (+)-*S*-enantiomer in *experimental arrhythmias* [Turgeon et al. *J Pharm Pharmacol* **43** 630 1991, DOI: 10.1111/j.2042-7158.1991.tb03552.x], in binding studies on cardiac sodium channels [Hill et al. *Mol Pharmacol* **34** 659 1988, PMID: 2848186], and a more potent blocker of Na currents in adult frog skeletal muscle fibres [De Luca et al. Naunyn Schmiedeberg's *Arch Pharmacol* **352** 653 1995, DOI: 10.1007/BF00171325; Luca et al. Naunyn Schmiedeberg's *Arch Pharmacol* **356** 777 1997, PMID: 9453464]. However, the reverse is true where the (+)-*S*-enantiomer is more effective than the (-)-*R*-enantiomer as an *inhibitor of chlorethylene-resistant amine oxidase (CRAO)* [Clarke et al. *Biochem Pharmacol* **31** 27 1982, DOI: 10.1016/0006-2952(82)90231-3], and in the treatment of *allodynia* [Zeitlin US Patent 97,7867,49, 1999, *Chem Abstr* **129** 153250 1999]. Various methods have been devised to obtain the two enantiomers: the procedure of G. Lentini and coworkers [Carocci et al. *Chirality* **12** 103 2000, DOI: 10.1002/(SICI)1520-636X] requires three steps starting from commercially available *S*(-)- and *R*(+)- propylene oxides in acceptable overall yields (24-34%) with 96% 'ee' and 93% 'ee' respectively, and checked by TLC on Kieselgel 60F₂₅₄) with $[\alpha]_D^{20}$ -1.1 (c 5, $CHCl_3$), and by the 1H NMR and MS which has m/z 180 M^+ (26), 122 (100). Similarly *R*(+)-propylene oxide gives ***R*(+)-1-(2,6-dimethylphenoxy)-2-propanol** in 57% yield with $[\alpha]_D^{20}$ +0.9 (c 5, $CHCl_3$) with the same spectra. ***S*(-)-1-(2,6-dimethylphenoxy)-2-propanol**, phthalimide and PhP_3 in THF under N_2 is treated dropwise with a solution of DEAD in THF, stirred overnight, concentrated *in vacuo*, Et_2O is added to the residue, the solid is filtered off and the filtrate is evaporated *in vacuo*, followed with purification by flash chromatography to give ***R*(-)-1-(2,6-dimethylphenoxy)-2-propyl N-phthalimide** (68% yield: **NOTE** inversion of configuration at asymmetric centre) with $[\alpha]_D^{20}$ -55 (c 2.2, $CHCl_3$) and 1H NMR (90MHz, $CDCl_3$ TMS) has δ at 1.55 (d, 3H, $J = 7.5$ Hz, CH_3-CH), 2.2 (s, 6H, 2 CH_3-Ar), 3.9 (dd, 1H, $J = 9$ Hz, $J = 6$ Hz, $OCHH$), 4.39 (apparent t, 1H, $J = 6$ Hz, $OCHH$), 4.85 (apparent br sextet, 1H, $J = 7.5$ Hz, CH_3CH), 6.9 (br s, 3H, PhO), 7.6-8.0 (m, 4H, Ar) ppm; and

MS has m/z 309 M^+ (5), 188 (100). Similarly, the *S*(+)-1-(2,6-dimethylphenoxy)-2-propyl *N*-phthalimide (with inversion of configuration) is obtained in 84% yield with $[\alpha]_D^{20} +55$ (c 2.5, $CHCl_3$) and same spectra as its enantiomer. Removal of the phthalimido group is achieved in the usual way, e.g. to the *R*(-)-phthalimide derivative in MeOH are added 2 equivalents of AcOH and 2 equivalents of $NH_2NH_2 \cdot H_2O$, stirred for 6 hours, the phthalazine-1,4-dione that separated is filtered off, washed twice with EtOAc, the combined filtrates are dried (Na_2SO_4), filtered, and evaporated to provide *R*(-)-**Mexiletine** [81771-86-0] in 66% yield as a colourless oil with $[\alpha]_D^{20} -2.7$ (c 4.7, $CHCl_3$), MS: m/z 179 M^+ (10), 44 (100). *R*(-)-**Mexiletine hydrochloride** **m 204-205°**, $[\alpha]_D^{20} -2.5$ (c 5, MeOH), is prepared by dissolving the *free base* in a small volume of aqueous 1M HCl, removing the H_2O azeotropically (e.g. with toluene) and the crude salt is recrystallised from EtOH-Et₂O to give analytically pure *hydrochloride*. The enantiomeric excess ('ee') is determined by HPLC analysis of *N*-acetyl-*R*(-)-**Mexiletine**. The acetate is prepared by treating a solution of the *free base* in dry THF with 2-equivalents of Ac_2O and 2-equivalents of Et_3N , stirring at $\sim 25^\circ$ for 8 hours, evaporated *in vacuo*, the residue is dissolved in EtOAc, washed with $NaHCO_3$ then brine, dried (Na_2SO_4), filtered and evaporated. The *acetate* is shown by HPLC (CSP: on CHIRACEL OD/R, with 9:1 MeOH- H_2O mobile phase at 0.4ml/min flow rate) to have 96% ee purity. Similarly *S*(+)-**Mexiletine** [9491-72-7] $[\alpha]_D^{20} +2.5$ (c 4.9, $CHCl_3$), and the *S*(+)-**Mexiletine hydrochloride** $[\alpha]_D^{20} +2.2$ (c 5, MeOH), are obtained; and the *S*(+)-**Mexiletine acetate** that is obtained in this case crystallised from 1:1 EtOAc-petroleum ether, and had 1H NMR (200MHz, $CDCl_3$, TMS) with δ at 1.38 (d, 3H, $J = 6.8Hz$, CH_3-CH), 2.01 (s, 3H, CH_3-CO), 2.24 (s, 6H, 2 CH_3-Ar), 3.65-3.80 (d, 2H, $O-CH_2-CH$), 4.35 (m, 1H, $CH_2-CH-CH_3$), 6.15 (br s, 1H, NH -acetyl) ppm with 96% 'ee' purity as determined by using the $Eu(hfc)_3$ chemical shift reagent and by CSD HPLC (see above). An earlier different synthesis of chiral *Mexiletine* by Franchini and coworkers [*Chirality* **6** 590 1994, DOI: 10.1002/chir.530060713] starts with the reaction of *S*(+)-3-bromo-2-methyl-1-propanol, of known absolute configuration, and sodium 2,6-dimethylphenolate. It provides *R*(-)-**Mexiletine hydrochloride** in four further steps without involving the chiral centre. It also establishes the absolute configuration of the drug, which is further confirmed by X-ray crystallographic analysis of the related *S*(+)-**Mexiletine hydrobromide** prepared from an optical resolution of racemic *Mexiletine* L(-)-dibenzoyl-tartrate. For optical resolution of *Mexiletine* enantiomers using di-*O*-*p*-toluoyltartrate salts see Turgeon et al. [*J Pharm Pharmacol* **43** 630 1991, DOI: 10.1111/j.2042-7158.1991.tb03552.x]. The syntheses and pharmacological evaluation as selective Na^+ channel blockers of a variety of optically active *Mexiletine* analogues have been described [Franchini et al. *J Med Chem* **46** 5238 2003, DOI: 10.1021/jm030865y]. *R*(-)-**Mexiletine** [81771-86-0] {with reported IR, 1H NMR (200MHz, $CDCl_3$, TMS) and MS reported} has also been prepared enantiomerically pure (ee 98%, by DIACEL CHIRACEL HPLC of the *N*-acetyl derivative, see above) using a *hydrolytic kinetic resolution* of (\pm) -2-[(2,6-dimethylphenoxy)methyl]oxirane with Jacobsen's catalyst *R,R*-Salen-Co(III) (see 'Catalysts Part-2' in Chapter 5) [Sasikumar et al. *Tetrahedron: Asymmetry* **20** 2814 2009, DOI: 10.1016/j.tetasy.2009.11.014]. *Mexiletine* is an *antiarrhythmic (Class 1B) drug* that is also used to treat *refractory pain* and *muscle stiffness* from *myotonic dystrophy* (Steinert's disease) or *congenital mitonia* (Thomsen disease).

Mithramycin A (Aureolic acid, Plicamycin) [18378-89-7] $C_{52}H_{76}O_{24}$, **M 1085.2, m 164-167°, 180-183°, $[\alpha]_D^{20} -51$ (c 0.3, EtOH), $pK_{Est} \sim 9.2$** . Purify the antibiotic *mithramycin A* by crystallisation from $CHCl_3$. It is soluble in MeOH, EtOH, Me_2CO , EtOAc, Me_2SO and H_2O , and moderately soluble in $CHCl_3$, but is slightly soluble in $*C_6H_6$ and Et_2O . It is a fluorescent antitumour agent used in flow cytometry. Store it at -20° . [Thiem & Meyer *Tetrahedron* **37** 551 1981, DOI: 10.1016/S0040-4020(01)92428-7; for NMR see Yu et al. *Nature* **218** 193 1968, DOI: 10.1038/218193a0; *Beilstein* **17/1** V 672.] It is an *inhibitor of DNA/RNA*

Mitomycin C {[1a*S*-(1*α*,8*β*,8*α*,8*β*)-6-amino-8-[(aminocarbonyl)oxy]methyl]-1,1*a*,2,8,8*a*,8*b*-hexahydro-8*a*-methoxy-5-methylzirino[2',3':3,4]pyrrolo[1,2-*a*]indole-4,7-dione} [50-07-7] $C_{15}H_{18}N_4O_5$, **M 334.4, m >360°, $pK_{Est(2)} \sim 8.0$** . Mitomycin C forms blue-violet crystals from $*C_6H_6$ /petroleum ether and is soluble in Me_2CO , MeOH and H_2O , moderately soluble in $*C_6H_6$, CCl_4 and Et_2O but insoluble in petroleum ether. It has UV with λ_{max} at 216, 360 and a weak peak at 560nm in MeOH. [Stevens et al. *J Med Chem* **8** 1 1965, DOI: 10.1021/jm00325a001; Shirahata & Hirayama *J Am Chem Soc* **105** 7199 1983, DOI: 10.1021/ja00362a046; *Beilstein* **25** III/IV 516.] **Anticancer agent** [see 'Effect of Mitomycin C on Bladder Cancer: A 7 Years of Followup' Tolley et al. *J Urology* **155** 1233 1996, DOI: 10.1016/S0022-5347(01)66226-8].

(-)-Morphine (H_2O) (5*α*,6*α*)-7,8-didehydro-4,5-epoxy-17-methyl-3,6-diol [57-27-2] $C_{17}H_{19}NO_3 \cdot H_2O$, **M**

285.3 (anhydrous), **302.2**, **m** 230°(dec), **254°**(dec, rapid heating), **255°**, **260°**(Kofler block), d_4^{20} **1.32**, $[\alpha]_D^{23}$ **-130.9** (MeOH), $[\alpha]_D^{25}$ **-133** (MeOH), **pK₁ 8.31** (NMe), **pK₂ 9.51** (OH). Morphine is among the three more abundant (9-14% depending on source) narcotic alkaloids isolated from opium poppy seed pods; the other two being *codeine* (3-methoxymorphine, 1-3%) and *thebaine* (3,6-dimethoxy derivative of morphine with double bonds at C₅=C₆ and C₈=C₁₄, 0.3 to 1.5%) [Robinson & Gulland numbering see Bergel & Morrison *Quart Rev Chem Soc* **2** 349 1948, DOI: 10.1039/QR9480200349]. Crystallise the narcotic from MeOH or anisole (short orthorhombic crystals). It *dehydrates* at 130°, and sublimes at bath temperature 190-200°/0.2mm. A *metastable form* melts at ~197°. Its solubility in H₂O is 0.2g/L at 20° and 0.9g/L at 100°; 60g/L in dilute HCl or H₂SO₄ at 20°; and in EtOH it is 5g/L at 20° and 10g/L on boiling. The solubility (w/v) at ~25° in CHCl₃ is 0.08%, in Et₂O is 0.016%, in amyl alcohol is 0.7%, and in boiling EtOAc is 10%. Store it away from light as it darkens in its presence, but with little loss of activity. It is basic due to the 'aliphatic type' NMe group and acidic due to its phenolic OH group (see pK_a's), forms salts readily and is useful for the *optical resolution* of acids. The *stypnate* has **m** **189°** (from aqueous EtOH).

Morphine sulfate (2:1) [anhydrous 64-31-3; 5H₂O 6211-15-0] [C₁₇H₁₉NO₃]₂ 5H₂SO₄, **M** **668.8**, has $[\alpha]_D^{25}$ **-109** (**c** **4** anhydr, H₂O). The *pentahydrate* crystallises from hot H₂O (solubility is 6.5w/w% at ~25°, 143w/w% at 80°) and loses 1H₂O at ~25°, 3H₂O at 100°, all H₂O at 130°, and decomposes at ~250°. Its solubility in EtOH is 0.18w/w% at ~25° and 0.42w/w% at 60°. Store away of light as it darkens, but without loss of activity.

Morphine acetate has $[\alpha]_D^{15}$ **-77** (H₂O) forms a *trihydrate* from H₂O (solubility w/v is 40% at 100°), and the solubility in EtOH is 4.5w/v% at ~25°, 50w/v% at 60°, and in CHCl₃ it is 0.21w/v% at ~25°. **Morphine hydrochloride** [52-26-6] $[\alpha]_D^{25}$ **-113** (**c** **2.2** anhydr, H₂O) crystallises as a *trihydrate* from H₂O which loses H₂O, becomes yellow and melts at ~200°(dec). [Beilstein **27** II 118, **27** III/IV 2223.]

Morphine and its salts are strong *analgesic* substances, are *addictive narcotics* with *strong withdrawal effects*, rapid acting, cause respiratory depression and death when used in excess, and are *controlled drugs* that require a medical prescription for use [Narcotic Drugs 2014 INTERNATIONAL NARCOTICS CONTROL BOARD. 2015, ISBN 9789210481571].

The *second* abundant opiate alkaloid in poppy seedpods is *Codeine* (3-methylmorphine, 5 α ,6 α)-7,8-didehydro-4,5-epoxy-3-methoxy-17-methyl-6-ol) [76-57-3] C₁₈H₂₁NO₃, **M** **299.4**, **m** **151-154°**, **154-156°**, d_4^{20} **1.32**, $[\alpha]_D^{15}$ **-112** (**c** **2** anhydr, CHCl₃), $[\alpha]_D^{15}$ **-136** (**c** **2** anhydr, EtOH), **pK₂₅ 8.21**. Purify it by recrystallisation from H₂O or aqueous EtOH (rods or octahedral plates) to give the *monohydrate* **m** **157-158.5°**, which has $[\alpha]_D^{25}$ **-136** (**c** **2.8**, EtOH), and is dried at 80°. The dried base sublimes at bath temperature 140-150°/~1.5mm. If it contains morphine (which has a phenolic group see pK_a above) then dissolve it in *C₆H₆ (solubility of codeine is 7.7w/v% at ~20°) extract this impurity into aqueous Na₂CO₃, dry the organic extract, evaporate and recrystallise. Its solubility (w/v) in H₂O is 0.83%(20°) and 1.67%(80°); in EtOH it is 50%(25°) and 83.3%(~78°); in Et₂O it is 5.6%(~25°); and is very soluble in CHCl₃ (200% at ~25°). Evaporation of a CHCl₃ extract gives a colourless glass which crystallises on scratching. **Codeine hydrobromide** [125-25-7] crystallises in needles from H₂O as the *dihydrate* (PubChem CID 517281), and *effervesces* at **151-160°** (**m** **190-192°** also reported) solidifies and remelts with extensive decomposition at 273-278°. It sublimes at 100°/0.03mm. [Gates *J Am Chem Soc* **75** 4340 1953, DOI: 10.1021/ja01113a511; Dauben et al. *J Org Chem* **44** 1567 1979, DOI: 10.1021/jo01323a045; Beilstein **27** II 136, **27** III/IV 2228.] **Codeine hydrochloride** [1422-07-7] crystallises in small needles from H₂O (solubility in w/v is 5% at ~25° and 100% at boiling) as the *dihydrate* with **m** **~287°**(dec) and has $[\alpha]_D^{15}$ **-108.2** (H₂O). Its solubility in EtOH is 0.56w.v% at ~25°. **Codeine sulfate** [1420-53-7] [C₁₈H₂₁NO₃]₂·H₂SO₄, crystallises in small crystals from H₂O (solubility in w/v is 3.3% at ~25° and 14.7% at 80°) as the *pentahydrate* with **m** **278°**(anhydrous) and has $[\alpha]_D^{15}$ **-100.9** (**c** **3**, H₂O). Its solubility in EtOH is very low at 0.076w/v% at ~25°. Like morphine and its salts, *codeine* and its salts are *light sensitive* and darken, although their activity is barely diminished. They should be stored in the dark. *Codeine* is also an *opiate analgesic* but much milder than morphine and less addictive, so much so that in some countries it is dispensed without a medical prescription. It is used in cough medicines and sometimes mixed with *paracetamol* or *ibuprofen*, and is a *muscle relaxant*. Its apparently weaker action is because it is demethylated to morphine during metabolism and people who do not have the required demethylating enzymes suffer from morphine side effects. See below for chemical syntheses

The *third* abundant opiate alkaloid in poppy seedpods (see above) is *Thebaine* (6,7,8,14-tetradehydro-4.5 α -epoxy-3,6-dimethoxy-17-methylmorphinan, *Paramorphine*) [115-37-7] C₁₉H₂₁NO₃, **M** **311.4**, **m** **~193°**(rapid heating), $[\alpha]_D^{23}$ **-230** (**c** **2**, CHCl₃), $[\alpha]_D^{15}$ **-219** (**c** **2**, EtOH), **pK₂₅ 8.15**. Purify by sublimation at 170-180°/atm or in a vacuum to give rectangular plates. It is poorly soluble in H₂O (solubility is 0.068w/v% at

15°), but is more soluble in CHCl_3 (7.7w/v% at ~25°), EtOH (6.7w/v% at boiling), C_6H_6 (4.0w/v% at ~25°), and Et_2O (0.5w/v% at ~25°). **Thebaine hydrochloride** [850-57-7] $\text{C}_{19}\text{H}_{21}\text{NO}_3 \cdot \text{HCl}$, **M 347.8, decomposes on heating emitting toxic NO_x and HCl fumes, $[\alpha]_{\text{D}}^{25} -164$ (c 2, EtOH).** Purify by recrystallisation from EtOH (colourless prisms). Its solubility in H_2O is 8.3w/v% at ~25°. Unlike morphine and its compounds, **thebaine compounds** are **opiates** with stimulatory rather than depressant effects on the nervous system, and at **high doses cause convulsions** not unlike strychnine. The synthetic **(+)-thebaine exhibits analgesic effects** unlike the natural (-)-thebaine which is inactive in this respect. It is not used therapeutically but is important commercially as it is used for preparing several physiologically active compounds such as **oxycodone** [76-42-2] and **naloxone** [465-65-6]. Many **chemical syntheses** of natural (-)-morphine, its enantiomer (+)- morphine as well as the racemate (see below) have been reported, but none are cheaper to produce than the drug from natural sources. [Beyerman et al. 'A convenient synthesis of codeine and morphine' *Recl Trav Chim Pays-Bas* **95** 24 1976, DOI: 10.1002/recl.19760950107; Beyerman et al. 'Synthesis of racemic and optically active codeine and morphine via the N-formylnordihydro-thebainones' *Recl Trav Chim Pays-Bas* **97** 127 1978, DOI: 10.1002/recl.19780970504; Gates & Tschudi 'The Synthesis of Morphine' *J Am Chem Soc* **78** 1380 1956, DOI: 10.1021/ja01588a033; Rice 'Synthetic opium alkaloids and derivatives. A short total synthesis of (+/-)-dihydrothebainone, (+/-)-dihydrocodeinone, and (+/-)-nordihydrocodeinone as an approach to a practical synthesis of morphine, codeine, and congeners' *J Org Chem* **45** 3135 1980, DOI: 10.1021/jo01303a045; Evans & Mitch 'Studies directed towards the total synthesis of morphine alkaloids' *Tetrahedron Lett* **23** 285 1982, DOI: 10.1016/S0040-4039(00)86810-0; Toth et al. 'Studies on the total synthesis of (dl)-morphine' *J Org Chem* **53** 4694 1988, DOI: 10.1021/jo00255a008; Parker & Fokas 'Convergent synthesis of (+/-)-dihydroisocodeine by the tandem radical cyclisation strategy and a formal synthesis of (+/-)-morphine' *J Am Chem Soc* **114** (24): 9688 1992, DOI: 10.1021/ja00050a075; Hong et al. 'Asymmetric synthesis of either enantiomer of opium alkaloids and morphinans, total synthesis of (-)- and (+)-dihydrocodeinone, and (-)- and (+)-morphine' *J Am Chem Soc* **115** 11028 1993, DOI: 10.1021/ja00076a086; Mulzer et al. 'Formal Total Synthesis of (-)-Morphine by Cuprate Conjugate Addition' *Angew Chem Int Ed* **35** 2830 1996, DOI: 10.1002/anie.199628301; White et al. 'Asymmetric Total Synthesis of (+)-Codeine via Intramolecular Carbenoid Insertion' *J Org Chem* **64** 7871 1999, DOI: 10.1021/jo990905z; Taber et al. 'Synthesis of (-)-Morphine' *J Am Chem Soc* **124** 12416 2002, DOI: 10.1021/ja027882h, PMID: 12381175; Trost & Tang 'Enantioselective Synthesis of (-)-Codeine and (-)-Morphine' *J Am Chem Soc* **124** 14542 2002, DOI: 10.1021/ja0283394. PMID: 12465957; Uchida et al. 'Total Synthesis of (±)-Morphine' *Org Lett* **8** 5311 2006, DOI: 10.1021/ol062112m, PMID: 17078705; Varin et al. 'Diastereoselective Total Synthesis of (±)-Codeine' *Chemistry – A European Journal* **14** 6606 2008, DOI: 10.1002/chem.200800744; Stork et al. 'Regiospecific and Stereoselective Syntheses of (±) Morphine, Codeine, and Thebaine' *J Am Chem Soc* **131** 11402 2009, DOI: 10.1021/ja9038505, PMID: 19624126

Muramic acid [**R-2(2-amino-2-deoxy-D-glucose-3-yloxy)-propionic acid**] [1114-41-6] $\text{C}_9\text{H}_{17}\text{NO}_7$, **M 251.2, m 145-150°(dec), 152-154°(dec), 155°(dec), $[\alpha]_{\text{D}}^{25} +109$ (c 2, H_2O), +165.0° (extrapolated to 0 time) → +123° [after 3 hours (c 3, $\text{H}_2\text{O})$], $\text{pK}_{\text{Est}(1)} \sim 3.8$ (CO_2), $\text{pK}_{\text{Est}(2)} \sim 7.7$ (NH_2).** Muramic acid crystallises from H_2O or aqueous EtOH as the **monohydrate** which loses H_2O at 80° *in vacuo* over P_2O_5 . It sometimes contains some NaCl. It has been purified by dissolving 3.2g in MeOH (75ml), filtering from some insoluble material, concentrating to ~10ml and refrigerating. The colourless crystals are washed with absolute MeOH. This process does not remove NaCl; to do so, the product is recrystallised from an equal weight of H_2O to give a low recovery yield of **very pure acid** (0.12g). On paper chromatography 0.26µg give one ninhydrin positive spot after development with 75% phenol (R_F 0.51) or with *sec*-BuOH/ $\text{HCO}_2\text{H}/\text{H}_2\text{O}$ (7:1:2) (R_F 0.30). [Matsushima & Park *Biochemical Preparations* **10** 109 1963, *J Org Chem* **27** 3581 1962, DOI: 10.1021/jo01057a045.] The acid has also been purified by dissolving 990mg in 50% aqueous EtOH (2ml), cooling, collecting the colourless needles on a sintered glass funnel and drying over P_2O_5 at 80°/0.1mm to give the **anhydrous acid**. [Lambert & Zilliken *Chem Ber* **93** 2915 1960, DOI: 10.1002/cber.19600931224.] Alternatively, the acid is dissolved in a small volume of H_2O , neutralised to pH 7 with ion-exchange resin beads (IR4B in OH^- form), filtered, evaporated and dried. The residue is recrystallised from 90% EtOH (v/v) and dried as above for 24 hours. [Strange & Kent *Biochem J* **71** 333 1959, DOI: 10.1042/bj0710333.] The **N-acetyl derivative (NAMA, R-2-(acetyl-amino)-3-O-(1-carboxyethyl)-2-deoxy-D-glucose, R-2(2-acetyl-amino-2-deoxy-D-glucose-3-yloxy)-propionic acid**] [10597-89-4], **M 292.3, has m ~125° (dec) and $[\alpha]_{\text{D}}^{20} +41.2$ after 24 hours (c 1.5, H_2O), $\text{pK}_{\text{Est}} \sim 3.6$.** [Watanabe & Saito *J Bacteriol* **144** 428 1980, PMID: 7419493; Beilstein **4** IV 2029.] It is a component of bacterial cell walls.

Muscimol (pantherine, 5-aminoethyl-3[2*h*]-isoxazolone) [2763-96-4] $C_4H_6N_2O_2$, **M** 114.1, **m** 170-172°(dec), 172-174°(dec), 172-175°, 175°, 176-178°(dec), 184-185°, $pK_{Est(1)} \sim 6$ (acidic, ring 2-NH), $pK_{Est(2)} \sim 8$ ($CH_2CH_2NH_2$). Recrystallise muscimol from MeOH/tetrahydrofuran or EtOH and sublime it at 110-140° (bath) at 10^{-4} mm to give a yellow spot with ninhydrin which slowly turns purple [for NMR see Bowden et al. *J Chem Soc (C)* 172 1968, DOI: 10.1039/J39680000172]. It can also be purified by dissolving in the minimum volume of hot H_2O and adding EtOH dropwise until cloudy, cool, and colourless crystals separate; its IR has ν_{max} at 3445w, 3000-2560w br, 2156w, 1635s and 1475s cm^{-1} . [For NMR see Jager & Frey *Justus Liebigs Ann Chem* 817 1982, DOI: 10.1002/jlac.198219820423.] Alternatively, it has been purified by two successive chromatographic treatments on Dowex-1 x 8, with the first elution with 2M AcOH and a second with a linear gradient between 0–2M AcOH, evaporating the desired fractions and recrystallising the residue from MeOH. [McCarry & Savard *Tetrahedron Lett* 22 5153 1981, DOI: 10.1016/S0040-4039(01)92445-1; Nakamura *Chem Pharm Bull Jpn* 19 46 1971, DOI: org/10.1248/cpb.19.46.] It is a psychoactive ingredient of a poisonous mushroom (*Amanita muscaria*).

Mycophenolic acid (6-[1,3-dihydro-7-hydroxy-5-methoxy-4-methyl-1-oxoisobenzofuran-6-yl]-4-methylhex-4-enoic acid) [24280-93-1] $C_{17}H_{20}O_6$, **M** 320.3, **m** 141°, 141-143°, $pK_{Est(1)} \sim 2.5$ (CO_2H), $pK_{Est(2)} \sim 9.5$ (phenolic OH). Purify the acid by dissolving it in the minimum volume of EtOAc, applying onto a silica gel column (0.05-0.2 mesh) and eluting with a mixture of EtOAc/ $CHCl_3$ /AcOH (45:55:1) followed by recrystallising from heptane/EtOAc, from aqueous EtOH or from hot H_2O and drying *in vacuo*. It is a weak dibasic acid, moderately soluble in Et_2O , $CHCl_3$ and hot H_2O but weakly soluble in C_6H_6 and toluene. [Birch & Wright *Aust J Chem* 22 2635 1969, DOI: 10.1071/CH9692635; Canonica et al. *JCS Perkin Trans 1* 2639 1972, DOI: 10.1039/P19720002639; Birkinshaw et al. *Biochem J* 50 630 1952, DOI: 10.1042/bj0500630; Beilstein 18 II 393, 18 III/IV 6513.] It is an immunosuppressive agent of use in organ transplant.

Myricetin (Cannabiscetin, 3,3',4',5,5',7-hexahydroxyflavone) [529-44-2] $C_{15}H_{10}O_8$, **M** 318.2, **m** >300°, 357°(dec) (polyphenolic $pK_{Est} \sim 8-11$). Recrystallise myricetin from aqueous EtOH (**m** 357° dec, as *monohydrate*) or Me_2CO (**m** 350° dec, with one molecule of Me_2CO) as yellow crystals. It is almost insoluble in $CHCl_3$ and AcOH. The *hexaacetate* has **m** 213°. [Hergert *J Org Chem* 21 534 1956, DOI: 10.1021/jo01111a013; Spada & Cameroni *Gazzetta* 86 965, 975 1956, Kalff & Robinson *J Chem Soc* 127 181 1925, DOI: 10.1039/CT9252700181; Beilstein 18/5 V 670.] This is a polyphenolic flavonoid antioxidant.

Nalidixic acid (1-ethyl-7-methyl-1,8-naphthyridin-4-one-3-carboxylic acid) [389-08-2] $C_{12}H_{12}N_2O_3$, **M** 232.3, **m** 226.8-230.2°, 228-230°, 229-230°, pK^{25} 6.0. Nalidixic acid crystallises from H_2O or EtOH as a pale buff powder. It is soluble at 23° in $CHCl_3$ (3.5%), toluene (0.16%), MeOH (0.13%), EtOH (0.09%), H_2O (0.01%) and Et_2O (0.01%). It inhibits nucleic acid and protein synthesis in yeast [Leshner et al. *J Med Chem* 5 1063 1962, DOI: 10.1021/jm01240a021] and is a synthetic quinolone antibiotic.

Naloxone hydrochloride hydrate $C_{19}H_{21}NO_4 \cdot HCl \cdot H_2O$, (Narcan, 1-*N*-propenyl-7,8-dihydro-14-hydroxymorphinan-6-one hydrochloride) [357-08-4 anhydrous, 51481-60-8 hydrate] **M** 399.9, **m** 200-205°, $[\alpha]_D^{20}$ -164 (c 2.5, H_2O), $pK_{Est(1)} \sim 6$ (*N*-propenyl), $pK_{Est(2)} \sim 9.6$ (phenolic OH). This opiate antagonist has been recrystallised from EtOH/ Et_2O or H_2O . It is soluble in H_2O (5w/v%) and EtOH but insoluble in Et_2O . The *free base* [465-65-6] has **m** 184° (177-178° also reported) after recrystallisation from EtOAc, and $[\alpha]_D^{20}$ -194.5 (c 0.93, $CHCl_3$). [Olofson et al. *Tetrahedron Lett* 1567 1977, DOI: 10.1016/S0040-4039(01)93104-1; Gold et al. *Med Res Rev* 2 211 1982, DOI: 10.1002/med.2610020302.] It blocks the effects of opioids when overdosed.

Naltrexone hydrochloride dihydrate (1-*N*-cyclopropylmethyl-7,8-dihydro-14-hydroxy-morphinan-6-one hydrochloride) [16676-29-2] $C_{20}H_{23}NO_4 \cdot HCl \cdot 2H_2O$, **M** 413.9, **m** 274-276°, $[\alpha]_D^{20}$ -173 (c 1, H_2O), $pK_{Est(1)} \sim 6$ (*N*-cyclopropylmethyl), $pK_{Est(2)} \sim 9.6$ (phenolic OH). This narcotic antagonist has been purified by recrystallisation from MeOH and dried in air. The *free base* [465-65-6] has **m** 168-170° after recrystallisation from Me_2CO . [Cone et al. *J Pharm Sci* 64 618 1975, DOI: 10.1002/jps.2600640409; Gold et al. *Med Res Rev* 2 211 1982, DOI: 10.1002/med.2610020302.] Blocks the effect of opioids.

1-Naphthyl phosphate disodium salt [2183-17-7, 207569-06-0] $C_{10}H_7O_4P \cdot 2Na$, **M** 268.1, pK_1^{26} 0.97, pK_2^{26}

5.85 (for free acid). Purify the salt through an acid ion-exchange column (in H^+ form) to give the *free acid* [1136-89-6], M 224.2, which is obtained by freeze drying and recrystallising from $Me_2CO/*C_6H_6$, or by adding 2.5 volumes of hot $CHCl_3$ (or 20 parts of boiling $*C_6H_6$) to a hot solution of 1 part acid and 1.2 parts Me_2CO and cooling (**m 155-157°**, **157-158°**). The acid is dissolved in the minimum volume of H_2O to which 2 equivalents of NaOH are added and then freeze dried, or by adding the equivalent amount of MeONa in MeOH to a solution of the acid in MeOH and collecting the *Na salt*, washing with cold MeOH, then Et_2O , and drying in a vacuum. [Friedman & Seligman *J Am Chem Soc* **72** 624 1950, DOI: 10.1021/ja01157a505; Chanley & Feageson *J Am Chem Soc* **77** 4002 1955, DOI: 10.1021/ja01620a015.] The *monosodium salt* [1136-89-6] is similarly prepared but with 1 equiv of NaOH. The phosphate group hydrolyses at pH 1.1-5.85 at 70°. [Beilstein **6** IV 4226.] A *substrate* for *alkaline phosphatase* [Gomori *Methods Enzymol* **4** 381 1957, DOI: 10.1016/0076-6879(57)04066-5; **128** 212 1968], and *prostatic phosphatase* [Babson *Clin Chem* **30** 1418 1984, PMID: 6744605].

2-Naphthyl phosphate monosodium salt [14463-68-4] $C_{10}H_8O_4P.Na$, M 246.2, m 177-178°, 296° (sintering at 228°), pK_1^{26} 1.28, pK_2^{26} 5.53, pK_3^{26} 6.57 (for free acid). The *free acid* [41845-15-2] is purified as for the preceding 1-isomer and has m 176-177° (also 177-178°) after several recrystallisations by adding 2.5 volumes of hot $CHCl_3$ to a hot solution of 1 part of acid in 1.3 volumes of Me_2CO as for the 1-isomer above. It is neutralised with one equivalent of NaOH and freeze dried or prepared as the 1-isomer above. Its solubility in H_2O is ~5%. It also forms a **0.5 Na. 1 H_2O salt** which has m 203-205° (244° also reported). [Friedman & Seligman *J Am Chem Soc* **72** 624 1950, DOI: 10.1021/ja01157a505; Chanley & Feageson *J Am Chem Soc* **77** 4002 1955, DOI: 10.1021/ja01620a015; Beilstein **6** IV 4285.]

D(+)-Neopterin [2009-64-5] $C_9H_{11}N_5O_4$, M 253.2, m >300°(dec), $[\alpha]_{546}^{20}$ +64.5 (c 0.14, 0.1M HCl), $[\alpha]_D^{25}$ +50.1 (c 0.3, 0.1N HCl), pK_1 2.23 (basic), pK_2 7.89 (acidic). Purification is as for biopterin. Also purify it on a Dowex-1 x 8 (formate form) column and elute with 0.03M ammonium formate buffer pH 8.0 then pH 7.2. The fluorescent neopterin fraction is evaporated under reduced pressure, leaving neopterin and ammonium formate (the latter sublimates out at high vacuum) behind. Stir the residue for 24 hours with EtOH, collect the solid and recrystallise it from H_2O . [Viscontini et al. *Helv Chim Acta* **53** 1202 1970, DOI: 10.1002/hlca.19700530537; cf. Wachter et al. Eds *Neopterin* W de Gruyter, Berlin 1992, ISBN 9783110117905, Beilstein **26** IV 4038.]

β -Nicotinamide adenine dinucleotide (diphosphopyridine nucleotide, NAD, DPN) [53-84-9] $C_{21}H_{27}N_7O_{14}P_2$, M 663.4, m 160°, $[\alpha]_D^{23}$ -34.8 (c 1, H_2O), pK_1 2.2 (PO_4H), pK_2 4.0 (adenine NH_2), pK_3 6.1 (PO_4^-). NAD is purified by paper chromatography or better on a Dowex-1 ion-exchange resin. The column is prepared by washing with 3M HCl until free of material absorbing at 260nm, then with water, 2M sodium formate until free of chloride ions and, finally, with water. NAD, as a 0.2% solution in water, is adjusted with NaOH to pH 8, and adsorbed onto the column, washed with water, and eluted with 0.1M formic acid. Fractions with strong absorption at 360nm are combined, acidified to pH 2.0 with 2M HCl, and cold acetone (ca 5L/g of NAD) is added slowly and with constant agitation. It is left overnight in the cold, then the precipitate is collected in a centrifuge, washed with pure acetone and dried under vacuum over $CaCl_2$ and paraffin wax shavings [Kornberg *Methods Enzymol* **3** 876 1957, DOI: 10.1016/S0076-6879(57)03468-0]. It has been purified by anion-exchange chromatography [Dalziel & Dickinson *Biochemical Preparations* **11** 84 1966.] The purity is checked by reduction to NADH (with EtOH and yeast alcohol dehydrogenase) which has ϵ_{340nm} 6220 $M^{-1}cm^{-1}$. [Todd et al. *J Chem Soc* 3727 1957, DOI: 10.1039/JR9570003727; 3733 1957, DOI: 10.1039/JR9570003733.] [For pK_a see Lamborg et al. *J Biol Chem* **231** 685 1958, PMID: 13539003.] The *free acid* crystallises from aqueous Me_2CO with $3H_2O$ and has m 140-142°. It is stable in cold neutral aqueous solutions in a desiccator ($CaCl_2$) at 25°, but decomposes at strong acid and alkaline pH. Its purity is checked by reduction with yeast alcohol dehydrogenase and EtOH to NADH and noting the OD at 340nm. Pure NADH (see below) has ϵ_{340} 6.2 x 10⁴ $M^{-1}cm^{-1}$, i.e. 0.1 μ mole of NADH in 3ml and in a 1cm path length cell has an OD at 340nm of 0.207. [Beilstein **26** IV 3644, **26** III/IV 3639.] NAD is a ubiquitous cofactor for oxido-reductase and other enzymes.

β -Nicotinamide adenine dinucleotide reduced di-Na salt trihydrate (reduced diphosphopyridine nucleotide sodium salt, NADH) [606-68-8] $C_{21}H_{27}N_7O_{14}P_2.2Na.xH_2O$, M 709.4 (anhydrous), pK_a as for NAD. This coenzyme is available in high purity, and it is advisable to buy a fresh preparation rather than to purify an old sample as purification will usually lead to a more impure sample contaminated with the oxidised form (NAD). It has UV with λ_{max} at 340nm (ϵ 6,200 $M^{-1}cm^{-1}$), at which wavelength the oxidised form NAD

has **no** absorption. At 340nm a 0.161mM solution in a 1cm (pathlength) cell has an absorbance of 1.0 unit. The purity is best checked by the ratio $A_{280\text{nm}}/A_{340\text{nm}} \sim 2.1$, a value which increases as oxidation proceeds. The dry powder is stable indefinitely at -20° . Solutions in aqueous buffers at pH ~ 7 are stable for extended periods at -20° and for at least 8 hours at 0° , but are oxidised more rapidly at 4° in a cold room (e.g. almost completely oxidised overnight at 4°). [For UV see Drabkin *J Biol Chem* **157** 563 1945, <http://www.jbc.org/content/157/2/563>; Fluorescence: Boyer & Theorell *Acta Chem Scand* **10** 447 1956, DOI: 10.3891/acta.chem.scand.10-0447; Redox: Rodkey *J Biol Chem* **234** 188 1959, PMID: 13610918; Schlenk in *The Enzymes* **2** 250, 268 1951, Kaplan in *The Enzymes* **3** 105, 112 1960.] Deuterated NADH, i.e. NADD, has been purified through the anion exchange resin AG-1 x 8 (100-200 mesh, formate form) then a Bio-Gel P-2 column. [Viola et al. *Anal Biochem* **96** 334 1979, DOI: 10.1016/0003-2697(79)90590-6.]

β -Nicotinamide adenine dinucleotide phosphate (NADP, TPN) [53-59-8] $\text{C}_{21}\text{H}_{29}\text{N}_7\text{O}_{17}\text{P}_3$, M 744.4, pK_1 1.1 (PO_4H_2), pK_2 4.0 (adenine NH_2), pK_3 6.1 (PO_4^-). Purify it by anion-exchange chromatography in much the same way as for NAD [Dalziel & Dickinson *Biochem J* **95** 311 1965, DOI: 10.1042/bj0950311; *Biochemical Preparations* **11** 87 1966]. Finally it is purified by dissolving in H_2O and precipitating with 4 volumes of Me_2CO and dried *in vacuo* over P_2O_5 . It is unchanged by storing *in vacuo* at 2° . [Hughes et al. *J Chem Soc* 3733 1957, DOI: 10.1039/JR9570003733; Shuster & Kaplan *J Biol Chem* **215** 183 1955, PMID: 14392153.] Deuterated NADPH, i.e. NADPD, has been purified through the anion exchange resin AG-1 x 8 (100-200 mesh, formate form) and through a Bio-Gel P-2 column. It has λ_{min} 259nm (ϵ 18.000) at pH 7.0. [Viola et al. *Anal Biochem* **96** 334 1979, DOI: 10.1016/0003-2697(79)90590-6; *Beilstein* **26** IV 3669, 3672.]

β -Nicotinamide adenine dinucleotide phosphate reduced tetrasodium salt (reduced diphosphopyridine nucleotide phosphate sodium salt, NADPH) [2646-71-1] $\text{C}_{21}\text{H}_{26}\text{N}_7\text{O}_{17}\text{P}_3 \cdot 4\text{Na} \cdot x\text{H}_2\text{O}$, M 833.4, pK_a as for NADP. Purification is mostly similar to that of NADH above. [*Beilstein* **26** III/IV 3671.]

β -Nicotinamide mononucleotide (NMN) [1094-61-7] $\text{C}_{11}\text{H}_{15}\text{N}_2\text{O}_8\text{P}$, M 334.2, m 90.3° , $[\alpha]_{\text{D}}^{23}$ -38.3 (c 1, H_2O), pK_{Est} ~ 6.1 (PO_4^-). Purify NMN by passage through a column of Dowex-1 (Cl $^-$ form) and washing with H_2O until no absorbance is observed at 260 nm. The tubes containing NMN are pooled, adjusted to pH 5.5-6 and evaporated *in vacuo* to a small volume. This is adjusted to pH 3 with dilute HNO_3 in an ice-bath and treated with 20 volumes of Me_2CO at $0-5^\circ$. The heavy white precipitate is collected by centrifugation at 0° . It is best stored wet and frozen or it can be dried to give a gummy residue. It has λ_{max} at 266nm (ϵ 4,600) and λ_{min} at 249nm (ϵ 3600) at pH 7.0 (i.e. no absorption at 340nm). It can be estimated by reaction with CN^- or hydrosulfite which form the 4-adducts (equivalent to NADH) which have UV with λ_{max} at 340nm (ϵ 6,200). Thus after reaction, an OD $_{340}$ of 'one' is obtained from a 0.1612mM solution in a 1cm path cuvette. [Plaut & Plaut *Biochemical Preparations* **5** 56 1957, Kaplan & Stolzenbach *Methods Enzymol* **3** 899 1957, DOI: 10.1016/S0076-6879(57)03473-4; Kaplan et al. *J Am Chem Soc* **77** 815 1955, DOI: 10.1021/ja01608a098; *Beilstein* **22/2** V 168.]

(-)-Nicotine [(2S)-1-methyl-2[3-pyridyl]-pyrrolidine] [54-11-5] $\text{C}_{10}\text{H}_{14}\text{N}_2$, M 162.2, m -79° , b $123-125^\circ/17\text{mm}$, $243-248^\circ/760\text{mm}$ (partial dec), d_4^{20} 1.097, n_{D}^{20} 1.5280, $[\alpha]_{\text{D}}^{20}$ -169 (c 1, Me_2CO), pK_1^{15} 6.16 (pyridine N^+), pK_2^{15} 10.96 (pyrrolidine N^+). (-)-Nicotine is a very pale yellow *hygroscopic* oil with a characteristic odour (tobacco extract) which turns brown in air on exposure to light. It is purified by fractional distillation under reduced pressure in an inert atmosphere. A freshly distilled sample should be stored in dark sealed containers under N_2 . It is a strong base; a 0.05 M aqueous solution has a pH of 10.2. It is very soluble in organic solvents. It is soluble in H_2O and readily forms salts. [For UV see Parvis *J Chem Soc* **97** 1035 1910, DOI: 10.1039/CT9109701035; Dobbie & Fox *J Chem Soc* **103** 1193 1913, DOI: 10.1039/CT9130301193.] The *hydrochlorides* (mono- and di-) form *deliquescent* crystals soluble in H_2O and EtOH but insoluble in Et_2O . It has also been purified *via* the ZnCl_2 double salt. [Ratz *Monatsh Chem* **26** 1241 1905, DOI: 10.1007/BF01526536; Biosynthesis: Nakane & Hutchinson *J Org Chem* **43** 3922 1978, DOI: 10.1021/jo00414a027.] The *picrate* has m 218° (from EtOH). [*Beilstein* **23/6** V 64.] Nicotine is the addictive principle in tobacco smoke with a biological half life of 1-2 hours. **POISONOUS**. (\pm)-Nicotine [22083-74-5] $\text{C}_{10}\text{H}_{14}\text{N}_2$, M 162.2, has b $113-115^\circ/10\text{mm}$, $242.3^\circ/\text{atm}$, d_4^{20} 1.082 (pK_a see above). It is purified by fractional distillation. Its solubility in EtOH is $\sim 5\%$. The *picrate* forms yellow needles from hot H_2O and has m 219° . The *methiodide* has m 219° (from MeOH). [Craig *J Am Chem Soc* **55** 2854 1933, DOI: 10.1021/

ja01334a036; Nakane & Hutchinson *J Org Chem* **43** 3922 1978, DOI: 10.1021/jo00414a027; *Beilstein* **23/6** V 64.] **POISONOUS.**

Nonactin [6833-84-7] $C_{40}H_{64}O_{12}$, **M 737.0**, **m 146°**, **147-148°**, $[\alpha]_D^{20}$ **0 ($\pm 2^\circ$)** (c **1.2**, $CHCl_3$). This cyclic macrotetrolide ionophore antibiotic crystallises from MeOH as colourless needles and is dried at 90°/20 hours/high vacuum. [Corbaz et al. *Helv Chim Acta* **38** 1445 1955, DOI: 10.1002/hlca.19550380617; for the crystal structure see Dobler *Helv Chim Acta* **55** 1371 1972, DOI: 10.1002/hlca.19720550504; Gombos et al. *Tetrahedron Lett* 3391 1975, DOI: 10.1016/S0040-4039(00)91406-0; *Beilstein* **19/12** V 751.]

L-Noradrenaline (Adrenor, R-2-amino-1-[3,4-dihydroxyphenyl]ethan-1-ol, L-) [51-41-2, 69815-49-2, 636-88-4 (bitartrate salt)] $C_8H_{11}NO_3$, **M 169.2**, **m 216.5-218°(dec)**, **~220-230°(dec)**, $[\alpha]_D^{20}$ **-45** (c **5**, N HCl), $[\alpha]_D^{25}$ **-37.3** (c **5**, 1 equivalent aqueous HCl), **pK₁²⁵ 5.58** (phenolic OH), **pK₂²⁵ 8.90** (phenolic OH), **pK₃²⁵ 9.78** (NH₂). Recrystallise adrenor from EtOH and store it in the dark under N₂. [For pKa see Lewis *Brit J Pharmacol Chemother* **9** 488 1954, PMID: 13219274; UV: Bergström & Hamberg *Acta Physiol Scand* **20** 101 1950, DOI: 10.1111/j.1748-1716.1950.tb00687.x; for fluorescence see Bowman et al. *Science NY* **122** 32 1955, DOI: 10.1126/science.129.3344.274; Tullar *J Am Chem Soc* **70** 2067 1948, DOI: 10.1021/ja01186a024.] The **L-tartrate salt monohydrate** has **m 102-104.5°**, and $[\alpha]_D^{25}$ **-11** (c **1.6**, H₂O), after recrystallisation from H₂O or EtOH. Store it at -20°. [*Beilstein* **13** III 2382.] A primary neurotransmitter.

L-Noradrenaline hydrochloride (Arterenol) [329-56-6] $C_8H_{11}NO_3 \cdot HCl$, **M 205.6**, has **m 145.2-146.4°**, **~150°(dec)**, $[\alpha]_D^{25}$ **-40** (c **6**, H₂O), **pKa** see above. Recrystallise arterenol from isoPrOH and store it in the dark as it is oxidised under light (see preceding entry). [Tullar *J Am Chem Soc* **70** 2067 1948, DOI: 10.1021/ja01186a024; *Beilstein* **13** III 2382.]

1R,2S-(-)-Norephedrine [L(-)-erythro-1R,2S-2-amino-1-phenyl-1-propanol] [492-41-1] $C_9H_{13}NO$, **M 151.2**, **m ~49-53°**, **50-52°**, $[\alpha]_D^{25}$ **-14.6** (c **3.4**, EtOH), $[\alpha]_D^{20}$ **-41** (c **7**, M HCl), **pK₂²⁵ 8.92**. It crystallises in plates from H₂O or Et₂O/petroleum ether. The **1R,2S-(-) hydrochloride** **M 187.7**, has **m 174-175°**, crystallises from iso-PrOH, and has $[\alpha]_D^{20}$ **-33.0** (c **5**, H₂O). [cf. Adkins & Cramer *J Am Chem Soc* **52** 4349 1930, DOI: 10.1021/ja01374a023; *Beilstein* **13** II 370, **13** III 1717, **13** IV 1875.]

1S,2R-(+)-Norephedrine [D(+)-erythro-1S,2R-2-amino-1-phenyl-1-propanol] [37577-28-9] $C_9H_{13}NO$, **M 151.2**, has **m ~49-53°**, **50-52°**, $[\alpha]_D^{27}$ **+14.8** (c **4**, EtOH), $[\alpha]_D^{20}$ **+40** (c **7**, M HCl), **pK₂²⁵ 8.92**. Purify it by recrystallisation from H₂O (plates), and it is soluble in Et₂O. Lewis [Brit *J Pharmacol Chemother* **9** 488 1954, PMID: 13219274] obtained **pK₂²⁰ 9.44** (H₂O). The **hydrochloride** [1S,2R-(+) 40626-28-7] **M 187.7**, has **m 172-175°** (plates from EtOH) and $[\alpha]_D^{20}$ **+33.4** (c **6**, H₂O), and the **sulfate** (plates from H₂O) has **m 285-286°** and $[\alpha]_D^{27}$ **+31.5** (H₂O). [cf. Adkins & Cramer *J Am Chem Soc* **52** 4349 1930, DOI: 10.1021/ja01374a023.]

DL-(±)-Norephedrine hydrochloride [Propadrin, (±)-erythro-1RS,2SR-2-amino-1-phenyl-1-propanol hydrochloride] [154-41-6] $C_9H_{13}NO \cdot HCl$, **M 187.7**, **m 194-196°** (also **194°** was reported), **pK 8.20**. It crystallises in plates from absolute EtOH or isoPrOH, and the (±)-oxalate has **m 245°(dec)** (plates from H₂O). The **2,4-dinitrobenzoate** has **m 86-88°** (from EtOH) and the **3,5-dinitrobenzoate** has **m 78-79°** (from EtOH). It is a **mixed anti-sympathomimetic** and used as a **nasal anticongestant**. [cf. Adkins & Cramer *J Am Chem Soc* **52** 4349 1930, DOI: 10.1021/ja01374a023; Krantz & Hartung *J Am Pharm Assoc* **19** 461 1930, DOI: 10.1002/jps.3080190507; Fischer & Plein *J Am Pharm Assoc* **44** 313 1955, DOI: 10.1002/jps.3030440517; *Beilstein* **13** I 252, **13** II 371, **13** III 1717, **13** IV 1875.]

(-)-1R,2R-Norpseudoephedrine [L(-)-threo-1R,2R-2-amino-1-phenyl-1-propanol] [37577-07-4] $C_9H_{13}NO$, **M 151.2**, **m ~50°**, **50-52°**, **77°**, **77.5-78°**, $[\alpha]_D^{20}$ **-34** (c **3.5**, EtOH), **pK₂²⁵ 8.92**. Purify (-)-nor-ψ-ephedrine by recrystallisation from H₂O, MeOH, EtOH, Et₂O/petroleum ether or *C₆H₆ (plates). The **mandelate salt** has **m 163.5°** (from EtOH/Et₂O) and $[\alpha]_D^{32}$ **-41.3** (c **0.8**, H₂O) [Jarowski & Hartung *J Org Chem* **8** 564 1943, DOI: 10.1021/jo01194a012]. The **hydrochloride** is purified by dissolving 1.44g in 96% EtOH (5ml), adding Et₂O (16ml) and cooling; it has **m 178-179°** (**m 180-181°** is also reported) and $[\alpha]_D^{30}$ **-42.9** (c **1.8**, H₂O) [Fles & Markovac-Prpic *Croat Chem Acta* **29** 186 1957]. [*Beilstein* **13** I 252, **13** II 370, **13** III 1716, **13** IV 1874.]

(+)-(1S,2S)-Norpseudoephedrine [(+)-Cathine, D(+)-threo-(1S,2S)-2-amino-1-phenyl-1-propan-1-ol] [492-39-7] $C_9H_{13}NO$, **M 151.2**, **m 77.5-78°**, $[\alpha]_D^{25}$ **+34.0** (c **4**, EtOH). Recrystallise it from *C₆H₆ and store away from CO₂ as it is a strong base readily forming a **hydrochloride** [1S,2S-(+) 2153-98-2] **M 187.7**, **m 180-**

181° (prisms from EtOH), and is soluble in H₂O with $[\alpha]_D^{20}$ +42.5. It is a *stimulant* and is an *anorexic* substance. [Nagai & Kanao *Justus Liebigs Ann Chem* **470** 157 1929, DOI: 10.1002/jlac.19294700110; Sicher & Pánková *Coll Czech Chem Commun* **20** 1409 1955, DOI: org/10.1135/cccc19551409; *Beilstein* **13** I 252, **13** II 370, **13** III 1716, **13** IV 1874.]

Novobiocin (7-[O³-carbamoyl-5-O⁴-dimethyl-β-L-lyso-6-desoxyhexahydropyranosyloxy]-4-hydroxy-3-[4-hydroxy-3-{3-methylbut-2-enyl}-benzylamino]-8-methylcoumarin) [303-81-1] C₃₁H₃₆N₂O₁₁, M 612.6, two forms m 152-156° and m 172-174°, 174-178°, λ_{max} at 330nm (acidic EtOH), 305nm (alkaline EtOH), $[\alpha]_D^{25}$ -63 (c 1, EtOH), pK₁ 4.03 (4.2), pK₂ 9.16. Crystallise novobiocin from EtOH and store it in the dark. It has also been recrystallised from Me₂CO/H₂O. [Hoeksema et al. *J Am Chem Soc* **77** 6710 1955, DOI: 10.1021/ja01629a129; Kaczka et al. *J Am Chem Soc* **77** 6404 1955, DOI: 10.1021/ja01628a121.] The *sodium salt* [1476-53-5] M 634.6, m 210-215°, 215-220°(dec), 222-229°, $[\alpha]_D^{25}$ -38 (c 1, H₂O) has been recrystallised from MeOH, then dried at 60°/0.5mm. [Sensi et al. *Anal Chem* **29** 1611 1957, DOI: 10.1021/ac60131a011; Kaczka et al. *J Am Chem Soc* **78** 4125 1956, DOI: 10.1021/ja01597a072; *Beilstein* **18/8** IV 8125.]

Nucleotide thiophosphate analogues. The preparation and purification of [³H]ATPyS, [³H]GTPyS, S⁶ITPyS (6-thioinosine), Cl⁶ITPyS (6-chloroinosine) and [³H]ATPyS were described, and general purification was achieved by chromatography of the nucleotide thiophosphates in the minimum volume of H₂O placed onto a DEAE-Sephadex A25 column and eluted with a linear gradient of triethylammonium bicarbonate (0.1 to 0.6M for G and I nucleotides and 0.2 to 0.5M for A nucleotides). [Goody et al. *Biochim Biophys Acta* **276** 155 1972, DOI: 10.1016/0005-2744(72)90016-2.]

Nystatin dihydrate (Mycostatin, Fungicidin) [1400-61-9] C₄₇H₇₉NO₁₉, M 962.1, m dec>160° (without melting by 250°), $[\alpha]_D^{25}$ -7 (0.1N HCl in MeOH), -10 (AcOH), +12 (Me₂NCHO), +21 (pyridine). Nystatin is a light yellow powder with the following solubilities at ~28°: MeOH (1.1%), ethylene glycol (0.9%), H₂O (0.4%), CCl₄ (0.12%), EtOH (0.12%), CHCl₃ (0.05%) and *C₆H₆ (0.03%). It has been precipitated from MeOH solution by addition of H₂O. Aqueous suspensions of this cyclic macrolide aminoglycoside antifungal antibiotic are stable at 100°/10 minutes at pH 7.0 but decompose rapidly at pH <2 and >9, and in the presence of light and O₂. [Birch et al. *Tetrahedron Lett* 1491 1964, DOI: 10.1016/S0040-4039(01)89518-6; 1485 1964, DOI: 10.1016/S0040-4039(01)89517-4; Weiss et al. *Antibiot Chemother* **7** 374 1957; Götzsche & Johansen *Cochrane Database Syst Rev* **9**: CD002033, 2014, DOI: 10.1002/14651858.CD002033.pub2. PMID: 25188770.] It may contain a mixture of components A₁, A₂ and A₃. [*Beilstein* **18** III/IV 7480.]

Omeprazole {5-methoxy-2-[(4-methoxy-3,5-dimethyl-2-pyridylmethylsulfinyl)]-1H-benzimidazole, Perizac, Losec Omeprazen, Pepticum) [73590-58-6] C₁₇H₁₉N₃O₃S, M 345.4, m 156°, pK₁²⁵ 3.98 (protonation of pyridine ring), pK₂²⁵ 8.70 (deprotonation of the imidazole NH). Clarke & Slemon have patented a novel synthesis which provides pure product free from coloured and other impurities and in which solids can be isolated [US Pat 5470983 A 1995 (lapsed)]. It involves linking a 4-methoxy-3,5-dimethylpyridin-2-yl moiety with a 5-methoxybenzimidazol-2-yl moiety via a bridge [—*CH(CONH₂)-S—] to form the thioether 4-OMe-3,5-MePy-2-CH(CONH₂)-S-2-(5-OMe)benzimidazole. This allowed the specific oxidation [e.g. with V(AcAc)₂/H₂O₂, at 0-5°, then 20-22°/1hr] of the sulfide group into a chiral sulfoxide (sulfinyl) group thus generating diastereoisomers (vicinal chiral centres at py-2-*C and the S(O)-benzimidazole). Mild treatment of 1g in H₂O (1ml) with 5.0g of sodium metabisulfite/75ml H₂O + 25ml MeOH (pH of mixture being at 7.2), followed by dropwise addition of glacial acetic acid (35 drops) to bring the pH to 4.8 resulted in vigorous evolution of CO₂ due to hydrolysis of the amide group and decarboxylation. The clear solution became turbid due to separation of *omeprazole* as an oil which, on addition of MeOH (2.0ml) and seeding with the drug, the oil solidified (30 minute). It was collected washed with H₂O and a little Me₂CO and dried to give 0.45g of substantially pure off-white *omeprazole* which can be recrystallised from MeCN. Allenmark et al. [*Anal Biochem* **136** 293 1984, DOI: 10.1016/0003-2697(84)90219-7] have succeeded in separating the enantiomers by HPLC on an affinity bovine serum albumin column (conditions: 0.08M phosphate pH 5.80, flow rate of 2.0ml/min, 10μl of 0.5mM solutions injected, using a 4.6x150mm Resolvosil 10μm analytical BSA-silica column, developed at Linköping University and available from Macherey-Nagel & Co. GmbH, Düren, GFR). *S-Omeprazole (Esomeprazole)* [11914-88-7] has been isolated as a syrup with $[\alpha]_D^{20}$ -155 (c 0.5, CHCl₃) and

elutes first from a chiral *trisphenylcarbamoylcellulose*-on 3-aminopropyl silica column. Erlandsson et al. [*J Chromatogr* **532** 305 1990, DOI: 10.1016/S0378-4347(00)83781-0] showed that the chiral drug is fairly optically stable since, in the minimum volume of 2-propanol diluted with 0.05M sodium phosphate buffer at pH 7.0, it **racemises** with half lives of 130 hours at 37° and 55 hours at 75°. They further studied the effect on acid formation in isolated gastric glands and found that the racemate, the (-)- and the (+)- enantiomer were about **equally effective**. The **magnesium salt of the S-enantiomer (Esomeprazole magnesium, Nexium)** [161973-10-0] ($C_{17}H_{19}N_3O_3S_2$)Mg, **M 713.1**, is a white powder with $[\alpha]_D^{20}$ -128.2 (c 1, MeOH) and similar **antiulcerative** activity.

Omeprazole is a **pro-drug** which has basic (pKa 3.98) and acidic (pKa 8.70) properties, and its charges are balanced, i.e. would have a pI of *ca* 6.3. It enters the gastric acid secreting parietal cells where it would be entirely in the protonated form which cannot get out of the cells. In these cells it undergoes an acid catalysed intramolecular rearrangement to a product which now has a free thiol group that forms an **irreversible** (almost) **disulfide bond** with the **ATPase proton pump**, and thus shuts down its activity (and acidity) [review: Lindberg et al. *Med Res Rev* **10** 1-54 1990, DOI: 10.1002/med.2610100102.] In time the pump is then reactivated by cytosolic glutathione to its normal state [Nagaya et al. *J Pharmacol Exp Ther* **252** 1289 1990, PMID: 2156997]. [For synthesis related compounds see Vidaillac et al. *Antimicrob Agents Chemother* **51** 831 2007, PMCID: 1803156; for reactions see Brändström et al. *Acta Chem Scand* **43** 536 1989, DOI: 10.3891/acta.chem.scand.43-0536; Brändström et al. *Acta Chem Scand* **43** 549 1989, DOI: 10.3891/acta.chem.scand.43-0549; Brändström et al. *Acta Chem Scand* **43** 569 1989, DOI: 10.3891/acta.chem.scand.43-0569; Brändström et al. *Acta Chem Scand* **43** 595 1989, DOI: 10.3891/acta.chem.scand.43-0595.] The **enantiomers** have also been separated using *Cellulase*(CBH I)*silica* as chiral stationary phase [Marle et al. *J Chromatogr* **586** 233 1991, DOI: 10.1016/0021-9673(91)85127-2; and Stenhoff et al. *J Chromatogr* **734** 191 1999, DOI: 10.1016/S0378-4347(99)00324-2].

Omeprazole is a H^+ , K^+ -ATPase inhibitor (IC_{50} = 5.8 μ M) that exhibits **antisecretory** and **antiulcer** activity. It inhibits gastric acid secretion (IC_{50} = 0.16 μ M for histamine-induced acid production), as well as being antibacterial towards *Helicobacter pylori* *in vitro*. It also blocks the swelling-dependent Cl^- channels (IC_{50} swell) in NIH3T3 fibroblasts. [Sato et al. *J Pharmacol Exp Ther* **248** 806 1989, PMID: 2537418; Schmarda et al. *Br J Pharmacol* **129** 598 2000, DOI: 10.1038/sj.bjp.0703070.]

Ondansteron (RS-9-methyl-(2-methyl-1H-imidazol-1-yl)-2,3-dihydro-1H-carbazol-4(9H)-one) [RS- 99614-02-5] **M 293.4, m 231-232°, 232-234°, pK²⁵ 7.40**. The synthesis of Ondansteron was described by Coates et al. [US Patent 4695578 1987, to Glaxo Group Ltd] with several variations. In one of these RS-3-[(dimethylamino)methyl]-2,3-dihydro-9-methyl-4(1H)-oxocarbazole (3.8g, see Evans *Aust J Chem* **26** 2555 1972, DOI: 10.1071/CH9732555) in MeI (100ml) was stirred while refluxing for 57 hours, the suspension was evaporated *in vacuo* to give the RS-3-methananium iodide (5.72g, m 192-195°). This iodide was stirred with excess of 2-methylimidazole [693-98-1] in dry DMF at 100° under N_2 for 17 hours, to give RS-2,3-dihydro-3-[(2-methylimidazol-1-yl)methyl]-4H-carbazol-4-one which was converted to the 9-sodio derivative with NaH in DMF and methylated at N-9 with Me_2SO_4 to provide **ondansteron** which was purified by recrystallisation from MeOH. The **citrate salt** was prepared by adding ondansteron (0.89g) to a hot citric acid (0.89g) solution in EtOH (20ml) and allowed to crystallise then recrystallising by dissolving in Me_2CO/H_2O (2:1, 2ml) and diluting with Me_2CO (20ml) to give pure **RS-citrate salt** (0.6g) **m 162°**. The **RS-maleate salt** formed white crystals **m 123.3°** after recrystallisation from hot EtOH. The **RS-monophosphate (1:1) salt, m 225°**, crystallised from hot H_2O .

RS-Ondansteron hydrochloride dihydrate (Zofran, Zofren) [99614-01-4] $C_{18}H_{19}N_3O \cdot HCl \cdot 2H_2O$, **M 365.9** was obtained by adding concentrated HCl (6.25ml) to a hot solution of ondansteron (18.3g) in *iso*-PrOH (98ml) and H_2O (90ml) which was filtered, the filtrate was diluted with *iso*-PrOH (90ml), stirred at ~25° for 17 hours, cooled to 2° and the salt (21.6g) was filtered off. A sample of the salt (6g) was recrystallised from a mixture of H_2O (6ml) and *iso*-PrOH (10ml) to provide **analytically pure white crystalline RS-hydrochloride dihydrate m 178.5-179.7°**. Its solubility in H_2O was 50mM, and its solubility in DMSO was 100mM.

Optical resolution of Ondansteron, which has an asymmetric carbon atom at C-3 of the dihydrocarbazolone ring, was achieved by mixing the free base (0.5g) in hot MeOH (30ml) with a hot solution of (-)-di-*p*-toluoyl-L-tartaric acid (0.7g) in MeOH (10ml) and cooled, whereby the (+)base-(-)acid diastereoisomeric salt (0.8g) separated. This salt was dissolved in hot DMF (20ml) diluted with hot H_2O (10ml) and allowed to crystallise for 3 days. The crystals were isolated, and dried *in vacuo* to give *ca* 95% enantiomerically **pure** (by 1H NMR) **(-)-di-*p*-toluoyl-L-tartaric salt** (0.26g) **m 190-192°**. This salt (0.2g) was mixed with 8% $NaHCO_3$ (25ml) and

extracted with CHCl_3 (2 x 25ml). Evaporation of the CHCl_3 extract gave pure **3-R-Ondansteron** that had $[\alpha]_{\text{D}}^{24} +16$ (c 0.34, MeOH) which was shown by $^1\text{H NMR}$ to have an enantiomer ratio of >95:5 (R:S). Similarly by using (+)-di-*p*-toluoyl-D-tartaric acid, the diastereomeric salt (-)base-(+)-acid was obtained. This salt gave the **free base, 3-S-Ondansteron**, which had $[\alpha]_{\text{D}}^{24} -14$ (c 0.19, MeOH) and shown by $^1\text{H NMR}$ to have an enantiomer ratio of 93:7 (S:R).

Ondansteron is a prescription drug, and is an *anti-emetic* for controlling post-operative nausea and vomiting, and is sometimes used in pregnancy. It is associated with a *long QT interval of the heart beat* which can lead to a fatal heart rhythm. [Review: Simpson & Hicks *J Pharm Pharmacol* **48** 774 1996, DOI: 10.1111/j.2042-7158.1996.tb03973.x.] **RS-Ondansteron** is an effective reversible *competitive antagonist of 5-HT₃ receptors* in rat vagus nerve and smooth muscle of guinea pig ileum; the **R-enantiomer is marginally more potent than the S-enantiomer** [Butler et al. *Brit J Pharmacol* **94** 397 1988, PMCID: PMC1854010]. [Cooke & Mehra in *Am J of Hospital Pharm* **51** 762 1994, PMID: 8010314; and Zoldan et al. *Neurology* **45** 1305 1995, PMID: 7617188.]

Oxcarbazepine (Trilapal, 10-oxo-10,11-dihydro-5H-dibenz[b,f]azepine-5-carboxamide) [28721-07-5] $\text{C}_{15}\text{H}_{12}\text{N}_2\text{O}_2$, M 252.3, m 215-216°. The synthesis of this drug has been described by Schindler [DE 2011087 1970 and US Patent 3642775 1972 to Ciba-Geigy]. Briefly it proceeds as follows: 10-methoxy-5H-dibenz[b,f]azepine [m 125-129°; 4698-11-7] + COCl_2 /toluene \rightarrow 10-methoxy-5H-dibenz[b,f]azepine-5-carbonyl chloride [77% yield, m 138° from EtOH; 28721-08-6] + EtOH/ NH_3 gas/4hrs \rightarrow 10-methoxy-5H-dibenz[b,f]azepine-5-carboxamide [93% yield, m 186-188° from EtOH; 28721-09-7] + 2N HCl/2hrs reflux, cool filter off crystals and recrystallise \rightarrow **Oxcarbazepine** [80% yield, m 215-216° from EtOH]. [see **Carbamazepine** [298-46-4] above and 5H-dibenz[b,f]azepine in 'Heterocyclic Compounds', Chapter 3.]

Oxcarbazepine is an *anticonvulsant* drug used for the evaluation and treatment of *epilepsy, psychosomatic disturbances* and of *trigeminal neuralgia*. [Dam et al. *Epilepsy Res* **3**(1) 70 1989, DOI: 10.1016/0920-1211(89)90070-3; Mazza et al. *Expert Opinion on Pharmacotherapy* **8**(5) 649 2007, PMID: 17376019.]

Oxacillin sodium salt (5-methyl-3-phenyl-4-isoxazolylpenicillin sodium salt) [1173-88-2, 7240-38-2 (H_2O)] $\text{C}_{19}\text{H}_{18}\text{N}_2\text{O}_3\text{S} \cdot \text{Na} \cdot \text{H}_2\text{O}$, M 423.4, m 188°(dec), $[\alpha]_{\text{D}}^{20} +29$ (c 1, H_2O), $\text{pK}_{\text{Est}} \sim 2.7$. This antibiotic, which is stable to penicillinase, is purified by recrystallisation from isoPrOH and dried *in vacuo*. Its solubility in H_2O at 25° is 5%. [Doyle et al. *Nature* **192** 1183 1961, DOI: 10.1038/1921183a0; Review: Chain et al. *Antibiotics* (Oxford University Press) **2** 1949.]

Oxolinic acid (5-ethyl-5,8-dihydro-8-oxo-1,3-dioxolo[4,5-g]quinoline-3-carboxylic acid) [14698-29-4] M 261.2, m 313-314°(dec), 314-316°(dec), $\text{pK}_{\text{Est}} \sim 2.3$. Purify the acid by recrystallisation from aqueous Me_2CO , 95% EtOH or dimethylformamide. It has UV with λ_{max} at 220, (255.5sh), 259.5, 268, (298sh, 311sh), 321 and 326nm [ϵ 14.8, (36.8sh), 38.4, 38.4, (6.4sh, 9.2sh), 10.8 and 11.2 x 10^3]. [Kaminsky & Meltzer *J Med Chem* **11** 160 1968, DOI: 10.1021/jm00307a041; *Beilstein* **17** III/IV 13, 17/1 V 11.] It is an antibacterial causing DNA cleavage *in vivo* in *E coli* [Snyder & Drlica *J Mol Biol* **131** 287 1979, DOI: 10.1016/0022-2836(79)90077-9].

Oxycodone [Endone, Roxicodon, (5R,9R,13S,14S)-4,5-epoxy-14-hydroxy-3-methoxy-17-methylmorphinan-6-one] [76-42-6] $\text{C}_{18}\text{H}_{21}\text{NO}_4$, M 315.4, m 218-220°, $\text{pK}_{\text{Est}} \sim 9.6$. Oxycodone is a *semi-synthetic opioid* prepared from thebaine, but also occurs naturally with other opium alkaloids. Purify it by recrystallisation from EtOH (colourless rods). It is almost insoluble in H_2O and Et_2O but quite soluble in EtOH and CHCl_3 . This base forms stable salts with acids e.g. tartaric, terephthalic acid as well as HCl, H_2SO_4 and H_3PO_4 . **The hydrochloride (Oxycon)** [124-90-3] crystallises in long rods from H_2O (solubility is 10w/v%/25°) with m 270-273°(dec) and $[\alpha]_{\text{D}}^{20} -125$ (c 2.5, H_2O). It is prepared by hydrolysis of thebaine with dilute H_2SO_4 to convert the dienol-ether into the enone (*codeinone*). The later can also be converted **4-OH-codeinone**, and reduction of this with hydrosulfite provides **oxycodone**. Oxycodone exhibits *keto-enol tautomerism* and apparently one form can crystallise out from EtOH and the more soluble isomer provides birefringent scales with m 219-220°. The two forms give the same salts [Freund & Speyer *J Prakt Chem* **94** 135 1916, DOI: 10.1002/prac.19160940112]. It is sold as an *analgesic*, alone or formulated with non-steroidal anti-inflammatory drugs. [Riley et al. *Curr Med Res Opin* **24** 175 2008, PMID: 18039433; Sunshine et al. *J Clin Pharmacol* **36** 595 1996, PMID: 8844441.]

Palmitoyl coenzyme A [1763-10-6] $\text{C}_{37}\text{H}_{66}\text{N}_7\text{O}_{17}\text{P}_3\text{S}$, M 1004.9. Possible impurities are palmitic acid,

S-palmitoyl thioglycolic acid and S-palmitoyl glutathione. These are removed by placing *ca* 200mg in a centrifuge tube and extracting with Me₂CO (20ml), followed by two successive extractions with Et₂O (15ml) to remove S-palmitoyl thioglycolic acid and palmitic acid. The residue is dissolved in H₂O (4 x 4 ml), adjusted to pH 5 and centrifuged to remove insoluble S-palmitoyl glutathione and other insoluble impurities. To the clear supernatant is added 5% HClO₄ (6ml) whereby S-palmitoyl CoA precipitates. The precipitate is washed with 0.8% HClO₄ (10ml) and finally with Me₂CO (3x 5ml) and dried *in vacuo*. It is stable for at least one year in dry form at 0° in a desiccator (dark). Solutions are stable for several months at -15°. Its solubility in H₂O is 4%. The adenine content is used as the basis of purity with λ_{\max} at 260 and 232nm (ϵ 6.4 x 10⁶ and 9.4 x 10⁶ cm²/mol, respectively). Higher absorption at 232nm would indicate other thio ester impurities, e.g. S-palmitoyl glutathione, which absorb highly at this wavelength. Also the phosphate content should be determined, and acid phosphate can be titrated potentiometrically. [Seubert *Biochemical Preparations* 7 80 1960, Srere et al. *Biochim Biophys Acta* 33 313 1959, DOI: 10.1016/0006-3002(59)90118-0; Kornberg & Pricer *J Biol Chem* 204 329 1953, PMID: 13084605; 345 1953, PMID: 13084606; Beilstein 26 III/IV 3665.] This thiol ester uses serine to biosynthesise sphingosine [Brady et al. *J Biol Chem* 244 491 1969, PMID: 4388074].

3-Palmitoyl-*sn*-glycerol (R-glycerol-1-palmitate, L- β -palmitin) [32899-41-5] C₁₉H₃₈O₄, M 330.5, d^{27.3} 0.9014, m 66.5° (α -form), 74° (β' -form) and 77° (β -form). The stable β -form is obtained by recrystallisation from EtOH or Skellysolve B, but recrystallisation from Et₂O provides the β' -form. The α -form is obtained on cooling the melt. Store it at -20°. [Malkin & el Shurbagy *J Chem Soc* 1628 1936, DOI: 10.1039/JR9360001628; Chapman *J Chem Soc* 55 1956, DOI: 10.1039/JR9560000055; Lutton & Jackson *J Am Chem Soc* 70 2445 1948, DOI: 10.1021/ja01187a043; Beilstein 2 III 966.]

D-Panthenol (Provitamin B, R-2,4-dihydroxy-3,3-dimethylbutyric acid 3-hydroxy-propylamide) [R- 81-13-0, RS- 16485-10-2] C₉H₁₉NO₄, M 205.3, 66-69°, b 118-120°/0.02mm, d₂₀²⁰ 1.2g/L, n_D²⁰ 1.4935, [α]_D²⁰ +30 (c 5, H₂O), pK²⁵ 13.03. Purify D-panthenol by distillation *in vacuo*. It is a slightly *hygroscopic* viscous oil and is soluble in H₂O and organic solvents. It is hydrolysed by alkali and strong acid. [Rubin *J Am Pharm Assoc (Sci Ed)* 37 502 1948, DOI: 10.1002/jps.3030371208; Bonati & Pitre *Farmaco Ed Scient* 14 43 1959, Beilstein 4 IV 1652.] Only the R(or D) enantiomer, *Dexpanthenol*, accelerates epidermal wound healing, but both enantiomers have moisturising properties for which they are used commercially. [Ebner et al. *Am J Clin Dermatol* 3 427 2002, DOI: 10.2165/00128071-200203060-00005, PMID:12113650.]

R-(+)-Pantothenic acid sodium salt (N-[2,4-dihydroxy-3,3-dimethylbutyryl] β -alanine Na salt) [867-81-2] C₉H₁₆NO₅. Na, M 241.2, [α]_D²⁵ +27.1 (c 2, H₂O), pK²⁵ 4.4 (for free acid). Recrystallise the salt from absolute EtOH. It is very *hygroscopic* (keep in sealed ampoules). The *free acid* [79-83-4] C₉H₁₇NO₅, M 219.2, is a viscous *hygroscopic* oil (m 179-179° ?) with [α]_D²⁵ +37.5 (c 5, H₂O), which is easily destroyed by acids and bases. Store it at -20°. See next entry. [Beilstein 4 IV 2569.]

R-(+)-Pantothenic acid Ca salt [(D(+)) 0.2Ca 137-08-6, 63409-48-3] (C₉H₁₆NO₅)₂. Ca, M 476.5, m 195-196°, 200-201°, [α]_D²⁰ +28.2 (c 5, H₂O). The salt crystallises as needles from MeOH, EtOH or isoPrOH (with 0.5mol of isoPrOH). Its solubility in H₂O is 50ml/ml at 25°, and is moderately *hygroscopic*. The *S-benzylisothiuronium salt* has m 151-152° (149° when crystallised from Me₂CO). [Kagan et al. *J Am Chem Soc* 79 3545 1957, DOI: 10.1021/ja01570a064; Wilson et al. *J Am Chem Soc* 76 5177 1954, DOI: 10.1021/ja01649a071; Stiller & Wiley *J Am Chem Soc* 63 1237 1941, DOI: 10.1021/ja01850a023; Beilstein 4 IV 2569.] It is a member of the *vitamin B complex* and essential for coenzyme A biosynthesis in mammalian cells.

Papaverine hydrochloride (6,7-dimethoxy-1-veratrylisoquinoline hydrochloride) [61-25-6] M 375.9, m 215-220°, 222.5-223.5°(dec), 231°, pK²⁵ 6.41. Recrystallise it from H₂O. It sublimes at 140°/0.1mm. Its solubility in H₂O is 5%. [For potentiometric titration of alkaloid salts see Saunders & Srivastava *J Pharm Pharmacol* 3 78 1951, DOI: 10.1111/j.2042-7158.1951.tb13047.x; and for pKa determination see Biggs *Trans Faraday Soc* 50 800 1954, DOI: 10.1039/TF9545000800.] The *free base* [58-74-2] has m 148-150°. The *picate* has m 186-189°(dec, 186-186.5° dec) [For regeneration of alkaloids from picrates see Bobbitt *J Org Chem* 22 1729 1957, DOI: 10.1021/jo01363a633]. [Beilstein 21 II 202, 21 III/IV 2788, 21/6 V 182.] The alkaloid is a cerebral vasodilator [Jayne et al. *J Clin Invest* 31 111 1952, DOI: 10.1172/JC1102568].

Paromomycin sulfate {amminosidin, *O*-2,6-diamino-2,6-dideoxy- β -L-idopyranosyl-(1 \rightarrow 3)-*O*- β -D-ribofuranosyl-(1 \rightarrow 5)-*O*-[2-amino-2-deoxy- α -D-glucopyranosyl-(1 \rightarrow 4)]-2-deoxystreptamine sulfate} [1263-89-4] $C_{33}H_{45}N_5O_{14} \cdot xH_2SO_4$, *M* 615.6 (free base), 713.7, amorphous, $[\alpha]_D^{25} +50$ (c 1.5, H_2O pH6), $pK_{est} \sim 9$. Purify this *antibiotic* by dissolving it in H_2O (0.5g/10ml) and adding excess EtOH, filter or collect and wash with EtOH, then Et₂O by centrifugation, and dry it *in vacuo*. An aqueous solution is stable at 37° for a week but longer at 0-5°. The *free base* [7542-37-2] is a white amorphous powder which stored under N₂ as it is strongly basic. It is soluble in MeOH (less so in EtOH) and has $[\alpha]_D^{25} +65$ (c 1.5, MeOH). It is an *antimicrobial* and is *antiamoebic*. It inhibits initiation and peptide elongation during protein synthesis. [Haskell et al. *J Am Chem Soc* **81** 3480 1959, DOI: 10.1021/ja01522a084; 3482 1959, DOI: 10.1021/ja01522a087; Hichens & Rinehart *J Am Chem Soc* **85** 1547 1963, DOI: 10.1021/ja00893a043; *Beilstein* **18** III/IV 7534.] It has *antiprotozoal* activity.

D-(-)-Penicillamine (*R*-3-mercapto-D-valine, 3,3-dimethyl-D-cysteine, from natural penicillin) [52-67-5] $C_5H_{11}NO_2S$, *M* 149.2, *m* 202-206°, 214-217°, $[\alpha]_D^{21} -63$ (c 1, N NaOH or pyridine), $pK_1^{25} 2.44$, $pK_2^{25} 7.97$ (7.88), $pK_3^{25} 10.43$ (10.46). The melting point of D-(-)-penicillamine depends on the rate of heating (*m* 202-206° is obtained by starting at 195° and heating at 2°/minute). It is soluble in H_2O and alcohols but insoluble in Et₂O, $CHCl_3$, CCl_4 and hydrocarbon solvents. Purify it by dissolving it in MeOH and adding Et₂O slowly. Dry it *in vacuo* and store under N₂. [Weigert et al. *Angew Chem Int Ed* **14** 330 1975, DOI: 10.1002/anie.197503301; Cornforth in *The Chemistry of Penicillin* (Clarke, Johnson and Robinson eds) Princeton Univ Press, 455 1949; Review: Chain et al. *Antibiotics* (Oxford University Press) **2** 1949; for polymorphism see Vidler *J Pharm Pharmacol* **28** 663 1976, PMID: 11326]. The *D-S-benzyl derivative* has *m* 197-198° (from H_2O), $[\alpha]_D^{17} -20$ (c 1, N NaOH), -70° (N HCl). [*Beilstein* **4** IV 3228.] It is a *penicillin metabolite* with *no* antibiotic activity. For pharmacokinetics dynamics see Joyce [*Pharmacol Therap* **42** 405 1989, DOI: 10.1016/0163-7258(89)90033-8]. *L-(-)-Penicillamine* [1113-41-3] $C_5H_{11}NO_2S$, *M* 149.2, has *m* 190-194°, 202-206°, 214-217°, $[\alpha]_D^{21} +63$ (c 1, N NaOH or pyridine), for *pKa* see *D-isomer*, with same chemistry as the *D-enantiomer* [*Beilstein* **4** IV 3228.], but is *toxic*, as it inhibits the action of pyridoxine.

D-Penicillamine disulfide hydrate (*S,S'*-di-[D-penicillamine] hydrate) [20902-45-8] $C_{10}H_{20}N_2O_4S_2$, *M* 296.4 + aqueous, *m* 203-204°(dec), 204-205°(dec), $[\alpha]_D^{23} +27$ (c 1.5, N HCl), -82 (c 0.8, N NaOH), $pK_{Est(1)} \sim 2.4$ (CO₂), $pK_{Est(2)} \sim 10.7$ (NH₂). Purify by recrystallisation from EtOH or aqueous EtOH. [Crooks in *The Chemistry of Penicillin* above, 469 1949; for use as *thiol reagent* for proteins see Garel et al. *Eur J Biochem* **123** 513 1982, DOI: 10.1111/j.1432-1033.1982.tb06561.x; Süss Justus Liebigs Ann Chem **561** 31 1948, DOI: 10.1002/jlac.19495610105; *Beilstein* **4** IV 3231.] For its use in *rheumatic disease* see Howard-Lock et al. [*Arthritis and Rheumatism* **15** 261 1986, DOI: 10.1016/0049-0172(86)90022-3].

Penicillic acid [5-hydroxy-5-isopropenyl-4-methoxy-2(5*H*)furanone (lactone), 3-methoxy-5-methyl-4-oxo-2,5-hexadienoic acid] [90-65-3] $C_8H_{10}O_4$, *M* 158.2, *m* 58-64°, 64-65° (monohydrate, acid), 83-84°, 87° (anhydrous, lactone), $pK^{25} 5.9$. The lactone (furanone, *anhydrous*) hydrolyses to the acid (3-methoxy-4-oxo-hexa-2,5-dienoic acid, *hydrate*). It crystallises from H_2O as the *monohydrate* (acid), or petroleum ether as the *anhydrous lactone*. The *free acid* and *lactone* are in equilibrium. Its UV has λ_{max} at 221nm (ϵ 12,500) in 0.02M KOH, and 228nm (ϵ 11,500) in 0.02M HCl [Raphael *J Chem Soc* 1508 1948, DOI: 10.1039/JR9480001508]. [*Beilstein* **3** II 519, **3** III 1467.] It is a possible *antineoplastic*.

(±)-Pentobarbital (5-ethyl-5-1'-methylbutyl barbituric acid, Nembutal is the Na salt) [76-74-4] $C_{11}H_{18}N_3O_2$, *M* 226.4, *m* ~127°(dec), $pK_{Est(1)} \sim 8.0$, $pK_{Est(2)} \sim 12.7$. A solution of the sodium salt in 10% HCl is prepared, and the acid is extracted with ether. Evaporation of the extract gives a solid which is then purified by repeated crystallisation from $CHCl_3$. It sublimates at 95-105°/10-12mm. [Bucket & Sandorfy *J Phys Chem* **88** 3274 1984, DOI: 10.1021/j150659a027.] The (+)- and (-)-*enantiomers* crystallise from 50% aqueous EtOH with *m* 120-121° and have $[\alpha]_D^{25} +4.73$ and -4.93 (EtOH) [Kleiderer & Shonle *J Am Chem Soc* **56** 1772 1934, DOI: 10.1021/ja01323a036]. [*Beilstein* **24** I 419, **24** II 287, **24** III/IV 1951.] It is a sedative and a hypnotic.

3-sn-Phosphatidylethanolamine (L- α -cephalin, from Soya bean) [39382-08-6] $C_{37}H_{74}NO_8P$, *M* 692.0, *M_r* ~600-800, amorphous, $pK_{Est(1)} \sim 5.8$ (PO₄⁻), $pK_{Est(2)} \sim 10.5$ (NH₂). Purify the cephalin by dissolving it in EtOH, adding Pb(OAc)₂·3H₂O (30g in 100ml H_2O) until presence of Pb²⁺. Filter off the solid, pass CO₂ until precipitation of PbCO₃ ceases. Filter the solid off and evaporate (while bubbling CO₂) *in vacuo*. H_2O (equal volume) is added to the residual oil and extracted with hexane. The extract is washed with H_2O until the aqueous phase is free from Pb [test with dithizone (2 mg in 100 ml CCl_4 ; Feigl *Spot Tests* Vol I, Elsevier p. 10

1954, ISBN-13: 978-0444409294, ISBN-10: 0444409297)]. The hexane is dried (Na_2SO_4), filtered and evaporated to give a yellow waxy solid which is *in vacuo*. It is insoluble in H_2O and Me_2CO , freely soluble in CHCl_3 (5%) and Et_2O , but slightly soluble in EtOH . [Schofield & Dutton *Biochemical Preparations* **5** 5 1957.]

O-Phosphocolamine (2-aminoethyl dihydrogen phosphate) [1071-23-4] $\text{C}_2\text{H}_8\text{NO}_4\text{P}$, **M 141.1**, **m 237-240°**, **242.3°**, **241.3-243°**, **234.5-244.5°**, **244-245°**(capillary), $\text{pK}_1^{20} < 1.5$ (PO_4H_2), $\text{pK}_2^{20} 5.77$ (PO_4H^-), $\text{pK}_3^{20} 10.26$ (NH^+). Purification by recrystallisation from aqueous EtOH gives a *hydrate* (**m 140-141°**). Its solubility in H_2O is 17% and 0.003% in MeOH or EtOH at 22°. [Fölsch & Österberg *J Biol Chem* **234** 2298 1959, PMID: 13823772; Baer & Stancer *Can J Chem* **34** 436 1956, DOI: 10.1139/v56-062; Christensen *J Biol Chem* **135** 399 1940, <http://www.jbc.org/content/135/2/399>.] A potent *inhibitor of ornithine decarboxylase* [Gilad & Gilad *Biochem Biophys Res Commun* **122** 277 1984, DOI: 10.1016/0006-291X(84)90471-6]. [Beilstein **4** IV 1415.]

Phosphoenolpyruvic acid monopotassium salt (KPEP) [4265-07-0] $\text{C}_3\text{H}_4\text{O}_6\text{P}$, **K**, **M 206.1**, $\text{pK}_1^{25} 3.4$ (CO_2), $\text{pK}_2^{25} 6.35$ (PO_4H^-) (for free acid). KPEP is purified *via* the cyclohexylamine salt (see below). The salt (534mg) in H_2O (10ml) is added to Dowex 50Wx4 H^+ form (200-400 mesh, 2ml, H_2O washed) and stirred gently for 30min then filtered. The resin is washed with H_2O (6ml), and combined solutions are adjusted to pH 7.4 with 3N KOH (~1.4ml) and volume adjusted to 18.4ml with H_2O , to give 0.1M KPEP that is lyophilised to a *pure powder* useful for enzyme work. Also recrystallise from $\text{MeOH}/\text{Et}_2\text{O}$. [Clark & Kirby *Biochemical Preparations* **11** 103 1966, Wold & Ballou *J Biol Chem* **227** 301 1957, PMID: 13449074; Cherbuliez & Rabinowitz *Helv Chim Acta* **39** 1461 1956, DOI: 10.1002/hlca.19560390536; Beilstein **3** IV 977.] **The triNa salt** [5541-93-5] **M 360.0**, is purified by dissolving 1g of salt in MeOH (40ml) and excess dry Et_2O is added. The white crystals are collected and dried ($\text{P}_2\text{O}_5/20^\circ$). [Cramer & Voges *Chem Ber* **92** 952 1959, DOI: 10.1002/cber.19590920429.]

Phosphoenolpyruvic acid tris(cyclohexylamine) salt (PRP-3CHA) [35556-70-8] $\text{C}_3\text{H}_5\text{O}_6\text{P}$, **3C}_6\text{H}_{13}\text{N}**, **M 465.6**, has **m 155-180°(dec)**, is recrystallised from aqueous Me_2CO and dried *in vacuo*. Store at -20° . It is stable at 4° for >2 years. Its IR has ν_{max} 1721cm^{-1} ($\text{C}=\text{O}$). [Wold & Ballou *J Biol Chem* **227** 301 1957, PMID: 13449074; see Clark & Kirby *Biochemical Preparations* **11** 103 1966 for the *monocyclohexylamine salt*.]

D(-)-3-Phosphoglyceric acid disodium salt (D-glycerate 3-phosphate di-Na salt) [80731-10-8] $\text{C}_3\text{H}_5\text{O}_7\text{P}$, **2Na**, **M 230.0**, $[\alpha]_{\text{D}}^{20} -13.9$ (c 5, H_2O), -745° (in aqueous NH_4^+ molybdate), $\text{pK}_{\text{Est}(1)} \sim 1.0$ (PO_4H_2), $\text{pK}_{\text{Est}(2)} \sim 6.66$ (PO_4H^-) (for free acid). It is best purified by conversion to the *Ba salt* by precipitation with BaCl_2 , which is recrystallised three times before conversion to the sodium salt. The *Ba salt* (9.5g) is shaken with 200ml of a 1:1 slurry of Dowex 50 (Na^+ form) for 2 hours. The mixture is filtered, and the resin is washed with H_2O (2 x 25ml). The combined filtrates (150ml) are adjusted to pH 7.0 and concentrated *in vacuo* to 30-40ml and filtered if not clear. Absolute EtOH is added to make 100ml, followed by *n*-hexane when a white solid and/or a second phase separates, which provides the complete solid *Na salt* on standing at $\sim 25^\circ$. The salt is removed by centrifugation, washed with Me_2CO , dried in air then in an oven at 55° to give a stable powder (4.5g), with no weight loss when kept over P_2O_5 at $78^\circ/8$ hours. The high optical rotation in the presence of $(\text{NH}_4)_6\text{Mo}_7\text{O}_{24}$ is not very sensitive to the concentration of molybdate or pH. Store it at -20° . It is **50% hydrolysed** in M HCl at $100^\circ/35.7\text{hrs}$, and at $125^\circ/3\text{hrs}$. [Towne et al. *J Biol Chem* **226** 777 1957, PMID: 13438863; Cowgill *Biochim Biophys Acta* **16** 613 1955, DOI: 10.1016/0006-3002(55)90299-7; Embdan et al. *Hoppe Seyler's Z Physiol Chem* **230** 20 1934, Beilstein **3** IV 1051.]

Phospholipids. For the removal of ionic contaminants from raw zwitterionic phospholipids, the lipids are purified twice by passage through mixed-bed ionic exchange resins (Amberlite AB-2) in methanolic solutions. (About 1g of lipid in 10ml of MeOH). In both runs the first 1ml of the eluate is discarded. The main fraction of the solution is evaporated at 40°C under dry N_2 and recrystallised (3x) from *n*-pentane. The resulting white powder is dried (4 hours at 50° *in vacuo*) and stored at 3° . Purify also by mixed-bed ion exchange resins suspended in the aqueous phase. [Kaatze et al. *J Phys Chem* **89** 2565 1985, DOI: 10.1021/j100258a028.]

O-Phospho-L-serine [407-41-0] $\text{C}_3\text{H}_5\text{O}_6\text{P}$, **M 185.1**, **m 175-176°**, **190°**, **228°**, $[\alpha]_{\text{D}}^{20} +4.3$ (c 3.2, H_2O), **+16.2** (c 3.2, 2N HCl), $\text{pK}_1^{25} < 1$ (PO_4H_2), $\text{pK}_2^{25} 2.08$ (CO_2H), $\text{pK}_3^{25} 5.65$ (PO_4H^-), $\text{pK}_4^{25} 9.74$ (NH_3^+). Recrystallise the phospho-serine by dissolving 10g in H_2O (150ml) at 25° , stirring for up to 20 minutes. Undissolved material

is filtered off (Büchner), and 95% EtOH (85ml) is added dropwise during 4 minutes, set aside at 25° for 3 hours, then at 3° overnight. The crystals are washed with 95% EtOH (100ml), then dry Et₂O (50ml), and dried in a vacuum (yield 6.5g). A further quantity (1.5g) can be obtained by keeping the mother liquors and washings at -10° for 1 week. The **DL-isomer** has **m 167-170°(dec)** after recrystallisation from H₂O/EtOH or MeOH. Store it at -20°. [Neuhaus & Korkes *Biochemical Preparations* **6** 75 1958, Neuhaus & Byrne *J Biol Chem* **234** 113 1959, PMID: 13610904; IR: Fölsch & Mellander *Acta Chem Scand* **11** 1232 1957, DOI: 10.3891/acta.chem.scand.11-1232; *Beilstein* **4** IV 3120.]

O-Phospho-L-threonine (L-threonine-O-phosphate) [1114-81-4] **C₄H₁₀NO₆P**, **M 199.1**, **m 194°(dec)**, **[α]_D²⁴ -7.4 (c 2.8, H₂O)** (**pK_a** as above). Dissolve the phosphate in the minimum volume of H₂O, add charcoal, stir for a few minutes, filter and apply onto a column of Dowex 50W (H⁺ form), then elute with 2N HCl. Evaporate the eluates under reduced pressure whereby the desired fraction produces crystals of the phosphate which can be recrystallised from H₂O/MeOH mixtures, and the crystals are then dried *in vacuo* over P₂O₅ at 80°. Store it at -20°. [de Verdier *Acta Chem Scand* **7** 196 1953, DOI: 10.3891/acta.chem.scand.07-0196; *Beilstein* **4** IV 3175.]

O-Phospho-L-tyrosine (L-tyrosine-O-phosphate) [21820-51-9] **C₉H₁₂NO₆P**, **M 261.2**, **m 225°, 227°, 253°**, **[α]_D²⁰ -5.5 (c 1, H₂O)**, **-9.2 (c 1, 2N HCl)**, **pK_{Est(1)} ~ 1.6 (PO₄H₂)**, **pK_{Est(2)} ~ 2.02 (CO₂H)**, **pK_{Est(3)} ~ 5.65 (PO₄H)**, **pK_{Est(4)} 9.2 (NH₃⁺)**. Purify it by recrystallisation from H₂O or H₂O/EtOH. [Levene & Schormüller *J Biol Chem* **100** 583 1933, <http://www.jbc.org/content/100/2/583>; Posternak & Graff *Helv Chim Acta* **28** 1258 1945, DOI: 10.1002/hlca.6602801179; *Beilstein* **14** III 1510.]

Phytol (d-2E-3,7R,11R,15-tetramethylhexadec-2-en-1-ol) [150-86-7, 7541-49-93] **C₂₀H₄₀O**, **M 296.5**, **b 145°/0.03mm**, **150-151°/0.06mm**, **203-204°/10mm**, **d₄²⁵ 0.8497**, **n_D²⁵ 1.437**, **[α]_D²² +0.06 (neat)**. This **acyclic triterpene** is commercially available as a mixture of isomers. Phytol is purified by distillation under high vacuum. It is almost insoluble in H₂O but soluble in most organic solvents. It has UV with λ_{max} at 212nm (log ε 3.04) in EtOH, and the IR has ν_{max} at 3300 and 1670cm⁻¹. [Demole & Lederer *Bull Soc Chim Fr* 1128 1958, Burrell *J Chem Soc (C)* 2144 1966, DOI: 10.1039/J39660002144; Bader *Helv Chim Acta* **34** 1632 1951, DOI: 10.1002/hlca.19510340546; *Beilstein* **1** IV 2208.]

Pirfenidone (5-methyl-1-phenylpyridin-2-one) [53179-13-8] **C₁₂H₁₁NO**, **M 185.2**, **m 103-105°, 107-111°**. It can be prepared by a general procedure where 2-cyano-*N*-(phenyl)acetamide (prepared by heating ethyl cyanoacetate with aniline under reflux for 2 hours, and crystallising from *C₆H₆-petroleum ether) is condensed with methylmalonaldehyde bis(dimethyl acetal) in boiling EtOH (3 hours) containing a catalytic amount of piperidine to give 3-cyano-5-methyl-1-phenylpyridin-2-one which is subsequently hydrolysed and decarboxylated to **Pirfenidone** [see Ismail & Noaman *Med Chem Res* **14** 382 2006, DOI: 10.1007/s00044-006-0146-2]. Purify it by recrystallisation from large volumes of H₂O (solubility 0.1w/v% at 20° and >1w/v% at 60°), dry it in air and sublime it *in vacuo*. Its solubility in DMSO > 2w/v% at 20°.

Pirfenidone is a **pulmonary antifibrotic drug** effective in models of pulmonary and lung fibrosis, is anti-inflammatory and is an analgesic.

[For novel Pirfenidone analogues see Ammar et al. *Archiv der Pharmazie* **339** 429 2006, DOI: 10.1002/ardp.200600017; for pharmacokinetics see Sun et al. *J Chromatogr B* **35** 981 2015, DOI: 10.1016/j.jchromb.2014.12.027; Kehrer & Margolin *Toxicol Lett* **90** 125 1997, DOI: 10.1016/S0378-4274(96)03845-3; Richeldi et al. *Nature Reviews Drug Discovery* **10**, 489 2011, DOI: 10.1038/nrd3495; Kaneko et al. *Clin Exp Immunol* **113** 72 1998, DOI: 10.1046/j.1365-2249.1998.00618.x; PMID: PMC1905007.]

Podophylotoxin [Wartec, Podofilox, (10R,11R,15R,16R)-16-hydroxy-10-(3,4,5-trimethoxyphenyl)-4,6,13-trioxatetracyclo-[7.7.0.0^{3,7}.0^{11,15}]hexadeca-1,3(7),8-trien-12-one] [518-28-5] **C₂₂H₂₂O₈**, **M 414.4**, **m 181-181°, 183-184°, 188-189°**, **[α]_D²⁰ -132 (c 1, CHCl₃)**. This **toxin** recrystallises from *C₆H₆ (with 0.5C₆H₆), EtOH/*C₆H₆, aqueous EtOH (with 1-1.5H₂O, **m 114-115°**) and CH₂Cl₂/pentane. When dried at 100°/10mm it has **m 183-184°**. Its biological half life is 1 to 4.5 hours. [UV: Stoll et al. *Helv Chim Acta* **37** 1747 1954, DOI: 10.1002/hlca.19540370620; IR: Schrecker et al. *J Org Chem* **21** 288 1956, DOI: 10.1021/jo01109a008.] It is an **inhibitor of microtubule assembly** [Prasad et al. *Biochemistry* **25** 739 1986, DOI: 10.1021/bi00351a035]. [*Beilstein* **19/10** V 666.] In the form of a cream, it is used for the topical treatment of **genital and other warts**.

Polyethylene glycol [25322-68-3] $C_{2n}H_{4n+2}O_{n+1}$, M_r various, from PEG ~200 to ~35,000, m 182-287°. PEG is available commercially as a powder or as a solution in various degrees of polymerisation depending on the average molecular weight, e.g. PEG 400 and PEG 800 have average molecular weights of 400 and 800, respectively. They may be contaminated with aldehydes and peroxides. Solutions *deteriorate* in the presence of air due to the formation of these contaminants. Methods available for purification are as follows:

Procedure A: A 40% aqueous solution of PEG 400 (2L, average molecular weight 400) is de-aerated under vacuum and made 10mM in sodium thiosulfate. After standing for 1 hour at 25°, the solution is passed through a column (2.5x20cm) of mixed-bed R-208 resin which has a 5cm layer of Dowex 50-H⁺ at the bottom of the column. The column is previously flushed with 30% aqueous MeOH, then thoroughly with H₂O. A flow rate of 1ml/minute is maintained by adjusting the fluid head. The first 200ml are discarded, and the effluent is then collected at an increased flow rate. The concentration of PEG solution is checked by density measurement, and it is stored (preferably anaerobically) at 15°.

Procedure B: A solution of PEG 800 (500g in 805ml H₂O) is made 1mM in H₂SO₄ and stirred overnight at 25° with 10g of treated Dowex 50-H⁺ (8% crosslinked, 20-50 mesh). The resin, after settling, is filtered off on a sintered glass funnel. The filtrate is treated at 25° with 1.5g of NaBH₄ (added over a period of 1 minute) in a beaker with tight but removable lid through which a propeller-type mechanical stirrer is inserted and continuously flushed with N₂. After 15 minutes, 15g of fresh Dowex 50-H⁺ are added, and the rate of stirring is adjusted to maintain the resin suspended. The addition of an equal quantity of Dowex 50-H⁺ is repeated and the reaction times are 30 and 40 minutes. The pH of a 1 to 10 dilution of the reaction mixture should remain above pH 8 throughout. If it does not, more NaBH₄ is added or the addition of Dowex 50-H⁺ is curtailed. (Some samples of PEG can be sufficiently acidic, at least after the hydrolysis treatment, to produce a pH that is too low for efficient reduction when the above ratio of NaBH₄ to Dowex 50-H⁺ is used.) About 30 minutes after the last addition of NaBH₄, small amounts of Dowex 50-H⁺ (~0.2g) are added at 15 minute intervals until the pH of a 1 to 10 dilution of the solution is less than 8. After stirring for an additional 15 minutes the resin is allowed to settle, and the solution is transferred to a vacuum flask for brief de-gassing under a vacuum. The de-gassed solution is passed through a column of mixed-bed resin as in procedure A. The final PEG concentration would be about 40% w/v. Assays for aldehydes by the purpuril method and of peroxides are in the reference below.

Treatment of Dowex 50-H⁺ (8% crosslinked, 20-50 mesh): The Dowex (500g) is suspended in excess 2N NaOH, and 3ml of liquid Br₂ is stirred into the solution. After the Br₂ has dissolved, the treatment is repeated twice, and then the resin is washed with 1N NaOH on a sintered glass funnel until the filtrate is colourless. The resin is then converted to the acid form (with dilute HCl, H₂SO₄ or AcOH as required) and washed thoroughly with H₂O and sucked dry on the funnel. The treated resin can be converted to the Na salt and stored. [Ray & Puvathingal *Anal Biochem* 146 307 1985, DOI: 10.1016/0003-2697(85)90544-5.]

Porphobilinogen (PBG, 5-amino-4-carboxymethyl-1H-pyrrole-3-propionic acid) [487-90-1] $C_{10}H_{14}N_2O_4$, M 226.2, m 172-175°(dec), 175-180°(dec, darkening at 120-130°), pK_1 3.70 (4-CH₂CO₂H), pK_2 4.95 (3-CH₂CH₂CO₂H), pK_3 10.1 (NH⁺). Porphobilinogen recrystallises as the *monohydrate* (pink crystals) from dilute NH₄OAc solutions of pH 4, and is dried *in vacuo*. The *hydrochloride monohydrate* has m 165-170°(dec) (from dilute HCl). [Jackson & MacDonald *Can J Chem* 35 715 1957, DOI: 10.1139/v57-100; Westall *Nature* 170 614 1952, DOI: 10.1038/170614a0; Granick & Bogorad *J Am Chem Soc* 75 3610 1953, DOI: 10.1021/ja01110a526; Beilstein 22/14 V 210.] PBG levels increase in *acute intermittent porphyria* [Aarsand et al. *Clin Chem* 52 (4) 650 2006, DOI: 10.1373/clinchem.2005.060772, PMID: 16595824].

Porphyrin A {from ox heart, coproporphyrin, 3-[18-(2-carboxyethyl)-7-ethenyl-17-formyl-12-[(4E,8E)-1-hydroxy-5,9,13-trimethyltetradeca-4,8,12-trienyl]-3,8,13-trimethyl-22,24-dihydroporphyrin-2-yl]propanoic acid} [5162-02-1] $C_{49}H_{58}N_4O_4$, M 799.0, m dec on heating. It is purified on a cellulose powder column followed by extraction with 17% HCl and fractionated with HCl. [Morell et al. *Biochem J* 78 793 1961, DOI: 10.1042/bj0780793.] It recrystallises from CHCl₃/petroleum ether or Et₂O/*C₆H₆ [detailed UV-VIS and NMR data: Caughey et al. *J Biol Chem* 250 7602 1975, PMID: 170266; Lemberg *Adv Enzymol* 23 265 1961, DOI: 10.1002/9780470122686.ch6].

Prazosin hydrochloride (2[4-((2-furoyl)piperazin-1-yl)4-amino-6,7-dimethoxyquinazoline hydrochloride, Minipress, Vasoflex] [19237-84-4, 19216-56-9 free base] $C_{19}H_{21}N_5O_4$. HCl, M 419.9, 383.4 (free base), m 278-280°, 280-282° (free base), pK^{25} 6.5. The salt is recrystallised by dissolving it in hot MeOH, adding a small

volume of MeOH/HCl (dry MeOH saturated with dry HCl gas) followed by dry Et₂O until crystallisation is complete. Dry it *in vacuo* over solid KOH till free of HCl. Also recrystallise from hot H₂O, and remove any H₂O azeotropically with CH₂Cl₂, then dried *in vacuo*. The solubility at ~25° in H₂O is ~0.5mg/ml; in MeOH it is 6mg/ml, but is poorly aqueous mineral acids. [NMR and IR: Honkanen et al. *J Heterocycl Chem* **17** 797 1980, DOI: 10.1002/jhet.5570170436; cf. Armarego & Reece *Aust J Chem* **34** 1561 1981, DOI: 10.1071/CH9811561.] Several syntheses of **Prazosin** and related compounds have been devised, but most of them run along similar lines. They start with 4,5-dimethoxyanthranilic acid that is converted **6,7-dimethoxyquinazolin-2,4-dione** (**m 323-325°**, platelets from aqueous AcOH), and provides **2,4-dichloro-6,7-dimethoxyquinazoline** (**m 158°**, colourless needles from light petroleum b 80-100°) on treatment with POCl₃/Me₂NPh/4.5hrs [Curd et al. *J Chem Soc* 1759 1948, DOI: 10.1039/JR9480001759]. Dry NH₃ is bubbled through a solution of this dichloroquinazoline in dry THF at 0°/30min and set aside to give **4-amino-2-chloro-6,7-dimethoxyquinazoline** (**m 302° dec**, from MeOH). When the latter is mixed with a slight excess of 1-(2-furoyl)piperazine [Althuis & Hess *J Med Chem* **20** 146 1977, DOI: 10.1021/jm00211a031] in *iso*-pentanol and refluxed for 2.5 hours (clear after 1.5hrs) and cooled, **Prazosin hydrochloride** is formed (>73% yield, **m 280-282°**, on crystallisation by dissolving in MeOH adding a slight excess of ~8N MeOH/HCl then by Et₂O until crystal separation is complete). [See Hess US3511836 A 12 May 1970 to Pfizer & Co; also Baker & Davies EP0520722 A1 30 Dec 1992, to Zeneca Ltd.] Many studies have been made to identify and quantify **Prazosin** and its **metabolites** in human blood and urine from users of this **antihypertensive** drug. Althuis and Hess [reference above] prepared the following: **2,4-di-amino-6,7-dimethoxyquinazoline** **m 244-246°** (from H₂O) R_F 0.41; **2-(1-piperazinyl)-4-amino-6,7-dimethoxyquinazoline HCl** **m 285-287°** (from 1.0 N HCl) R_F 0.24; **6-demethyl-prazosin trifluoroacetate hydrate** **m 220°** (from CF₃CO₂H-Et₂O) R_F 0.47; **7-demethyl-prazosin trifluoroacetate hydrate** **m 230°** (from CF₃CO₂H-Et₂O) R_F 0.43; and using TLC silica gel plates, eluting with EtOAc-MeOH-Et₂NH (70:20:5) where **Prazosin** has R_F 0.70. Armarego and Reece [reference above] prepared **8-chloroprazosin hydrochloride** (**m 207-208° dec**) and as a **non-interfering internal standard in the HPLC** for quantitative estimation of **Prazosin** and its metabolites in human plasma. Armarego et al. have also prepared a water soluble **Prazosinamine hydrochloride** [1-(4'-amino-6',7'-dimethoxyquinazolin-2'yl)-4-(6''-aminohexanoyl)piperazine hydrochloride, **m >210°** (with effervescence on slow heating, from MeOH/~7N MeOH-HCl followed by Et₂O; also TLC, IR and ¹H NMR data provided] which **reversibly inhibits the calcium-mobilizing action** of α₁-adrenergic agonists in perfused rat liver [*Biochem Pharmacol* **36** 1583 1987, DOI: 10.1016/0006-2952(87)90040-2].

Prazosin is an **antihypertensive drug** and is an α₁- and α_{2B}-adrenergic antagonist [Brosman et al. *Proc Natl Acad Sci USA* **82** 5915 1985, PMID: 2994053]. It is also a potent **antagonist** at the **melatonin MT₃ receptor** (K_i = 10.2nM) [Pickering & Niles *Eur J Pharmacol* **175** 71 1990, DOI: 10.1016/0014-2999(90)90154-X], and **inhibits MK-801** (a psychotomimetic, non-competitive NMDA receptor antagonist)—**induced hyperlocomotion and dopamine release** in the **nucleus accumbens** [Mathé et al. *Eur J Pharmacol* **309** 1 1996; DOI: 10.1016/0014-2999(96)00315-9].

Procaine hydrochloride (Novocain, **2-diethylaminoethyl-4-aminobenzoate HCl**) [51-05-8] C₁₃H₂₀N₂O₂. HCl, **M 272.8, 236.3**(free base), **m 153-156°, 154-156°, 156°, pK_{Est(1)}~ 2.52 (NH₂⁺) pK₂₀²⁰ 9.0 (Et₂N⁺)**. Novocain is recrystallised from aqueous EtOH. It is soluble at 25° in H₂O (86.3%), EtOH (2.6%) and Me₂CO (1%), it is slightly soluble in CHCl₃, but is almost insoluble in Et₂O. The anhydrous **free base** [59-46-1] is recrystallised from ligroin or Et₂O and has **m 61°**. Its biological half life is 40-84 seconds. [Einhorn *Justus Liebigs Ann Chem* **371** 125 1909, DOI: 10.1002/jlac.19093710202; IR: Szymanski & Panzica *J Am Pharm Assoc* **47** 443 1958, DOI: 10.1002/jps.3030470618; *Beilstein* **14** IV 1138.]

PropERICIAZINE [Pericyazine, **10-{3-(4-hydroxy-1-piperidiny)-propyl}-10H-phenothiazine-2-carbonitrile**] [2622-26-6] C₂₁H₂₃N₃OS, **M 365.4, m 116-117°**. It crystallises from a saturated solution in cyclohexane. It is an antipsychotic drug, and is a sensitive reagent for Pd, Ru, Rh and Au. [Gowda et al. *Anal Chem* **55** 1816 1983, DOI: 10.1021/ac00261a041; Gowda et al. *Anal Chim Acta* **154** 347 1983, DOI:10.1016/0003-2670(83)80039-7.]

R(+)-Propranalol hydrochloride (**R-1-isopropylamino-3-(1-naphthyloxy)-2-propanol HCl**) [13071-11-9] C₁₆H₂₁NO₂. HCl, **M 295.8, 259.3**(free base), **m 192°, 193-195°, [α]_D²⁰ +25 (c 1, EtOH), pK₂₀²⁰ 9.5 (for free base)**. The hydrochloride is recrystallised from *n*-PrOH or Me₂CO. It is soluble in H₂O and EtOH but is insoluble in Et₂O, *C₆H₆ or EtOAc. The **racemate salt** [318-98-9, 3506-09-0 100mM in H₂O] has **m 163-164°**,

and is *less active* than an equal mixture of *R*- and *S*- isomers. The *free base* recrystallises from cyclohexane with **m 96°**. [Howe & Shanks *Nature* **210** 1336 1966, DOI: 10.1038/2101336a0.] The *S*(-)-*isomer* (below) is physiologically the *more active* isomer. [For *RS*- and *R*- isomer: Xe et al. *Br J Pharmacol* **123** 599 1998, DOI: 10.1038/sj.bjp.0701630; Litwin et al. *Br J Pharmacol* **127** 1671 1999, DOI: 10.1038/sj.bjp.0702701.]

***S*(-)-Propranolol hydrochloride (S-1-isopropylamino-3-(1-naphthoxy)-2-propanol HCl) [4199-10-4] C₁₆H₂₁NO₂. HCl, M 295.8, has m 192°, 193-195°, [α]_D²⁰ -25.5 (c 1, EtOH) pK_a²⁰ 9.5.** See preceding entry for physical properties and purification. The (-)-salt is the more active isomer which *blocks isoprenaline tachycardia* and is a β-adrenergic blocker and *antiarrhythmic*. It is also a local *anaesthetic*. [Leclerc et al. *Trends Pharmacol Sci* **2** 18 1981, DOI: 10.1016/0165-6147(81)90248-0; Howe & Shanks *Nature* **210** 1336 1966, DOI: 10.1038/2101336a0.] The *S*(-)-isomer is the physiologically more active (by 2 orders of magnitude), but the *R*-isomer is not devoid of activity as it exhibits significant prolongation of the PR interval of the electrocardiogram. The enantiomers have been separated using *Cellulase*(CBH I)*silica* as chiral stationary phase [Marle et al. *J Chromatogr* **586** 233 1991, DOI: 10.1016/0021-9673(91)85127-2; Bond et al. [*Nature* **213** 721 1967, DOI: 10.1038/213721a0; PMID: 6031788; Barrett et al. [*Br J Pharmacol* **34** 43 1968, PMID: 19108278].

6-Propyl-2-thiouracil (propacil, propycil) [51-52-5] M 170.2, m 218-220°, pK_a²¹ -6.54 (aqueous H₂SO₄), pK_a²¹ 8.25 (4% aq EtOH). Purify by recrystallisation from H₂O (solubility: 900 parts at 20°, and 100 parts at 100°). UV, MeOH has λ_{max} 277nm. [Anderson et al. *J Am Chem Soc* **67** 2197 1945, DOI: 10.1021/ja01228a042; Vanderhaegue *Bull Soc Chim Belg* **59** 689 1950, DOI: 10.1002/bscb.19500590911; Beilstein **24** III/IV 1333.] It is an antihyperthyroid drug and promotes fattening.

Protoporphyrin IX (3,18-divinyl-2,7,13,17-tetramethylporphyrin-8,12-dipropionic acid, ooporphyrin) [553-12-8] C₃₄H₃₄N₄O₄, M 562.7, pK_{Est} ~ 4.8. Purify it by dissolving (4g) in 98-100% HCOOH (85ml), diluting with dry Et₂O (700ml) and keeping at 0° overnight. The precipitate is collected and washed with Et₂O, then H₂O, and dried in a vacuum at 50° over P₂O₅. It crystallises from aqueous pyridine or Et₂O in monoclinic, brownish-yellow prisms. The UV (25% HCl) has λ_{max} at 557.2, 582.2 and 602.4nm. It is freely soluble in ethanolic HCl, AcOH, CHCl₃, and Et₂O containing AcOH. It forms sparingly soluble diNa and diK salts. [Ramsey *Biochemical Preparations* **3** 39 1953; Drabkin *Haematin Enzymes* pp 142-172 1961, DOI:10.1016/B978-1-4831-9646-6.50016-8; Granick *J Biol Chem* **175** 333 1948, PMID: 18873307; see Falk 'Porphyrins and Metalloporphyrins' Vol 2 1964 Elsevier, NY, Lib Cong No: 62-19821; Beilstein **26** IV 3042.]

Protoporphyrin IX dimethylester [5522-66-7] C₃₆H₃₈N₄O₄, M 590.7, has m 225-228°, 228-230°. The crude dimethyl ester (1g) in CHCl₃ (200 ml) is mixed with petroleum ether (b 70-90°, 600ml), and any porphyrin (m > 260°) which is insoluble in this mixture is filtered off. The filtrate is passed through a column of CaCO₃ [from CaCO₃ (130g) which is kept overnight in a mixture of CHCl₃/petroleum ether (b 70-90°, 1:3), and the slurry is poured into a glass tube (2.5 x 26cm) to form the column]. After all the filtrate is applied, the column is eluted with a solution of CHCl₃/petroleum ether (b 70-90°, 1:3). All the coloured eluates are collected, evaporated at room temperature in a vacuum to give a residue (0.8g), m 208-211°. The residue (0.8g) in CHCl₃ (66ml) is heated briefly to its boiling point, then boiling MeOH (198ml) is added immediately to it. The mixture is allowed to cool to room temperature, refrigerated for 2 days and the solid is filtered off. The solid is washed on the filter funnel with CHCl₃/MeOH (1:9, 50ml) and dried at 50°/vacuum (yield 0.62-0.66g). It can also be recrystallised by dissolving in as little hot dry *C₆H₆ as possible and left overnight at 20°, or with CHCl₃/MeOH (1:9). Its UV has λ_{max} at 631, 576, 541, 506 and 407nm in CHCl₃ and 601, 556 and 406nm in 25% HCl. [see Ramsey above.]

Pterin-6-carboxylic acid (2-amino-4-oxo-3,4-dihydropteridine-6-carboxylic acid) [948-60-7] C₇H₅N₅O₃, M 207.2, m >360°, pK_a²⁰ 1.43, pK_a²⁰ 2.88, pK_a²⁰ 7.72. The acid gives yellow crystals by repeated dissolution in aqueous NaOH and adding aqueous HCl. It has UV with λ_{max} at 235, 260 and 265nm (ε 11,000, 10,500 and 9,000) in 0.1N HCl, and 263 and 365nm (ε 20,500 and 9,000) in 0.1N NaOH. [For UV see Pfeleiderer et al. *Justus Liebigs Ann Chem* **741** 64 1970, DOI: 10.1002/jlac.19707410108; Stokstad et al. *J Am Chem Soc* **70** 5 1948, DOI: 10.1021/ja01181a003; and for fluorescence see Kavanagh & Goodwin *Arch Biochem* **20** 315 1949, PMID: 18108925; Beilstein **26** III/IV 4053.]

Purine-9-β-ribofuranoside (Nebularin) [550-33-4] C₁₀H₁₂N₄O₄, M 252.2, m 178-180°, 181-182°, [α]_D²⁵ -48.6 (c 1, H₂O), -22 (c 0.8, 0.1N HCl) and -61 (c 0.8, 0.1N NaOH), pK_a²⁵ 2.05. Nebularin is recrystallised from

butanone/Mebutanone/MeOH or EtOH and forms a **MeOH photo-adduct**. It is a strong **inhibitor of adenosine deaminase** [EC 3.5.4.4]. [Nair & Wiechert *Bioorg Chem* **9** 423 1980, DOI: 10.1016/0045-2068(80)90002-4; Löfgren et al. *Acta Chem Scand* **7** 225 1953, DOI: 10.3891/acta.chem.scand.07-0225; UV: Brown & Weliky *J Biol Chem* **204** 1019 1953, PMID: 13117878; *Beilstein* **26** III/IV 1740.]

Puromycin dihydrochloride (*O*-methyl-L-tyrosine[*N*⁶,*N*⁶-dimethylaminoadenosin-3'-yl-amide]) [58-58-2] **C₂₂H₂₈N₇O₅·2HCl**, **M 616.5**, **m 174°**, **178-180°**, [α]_D²⁵ **-11** (free base in EtOH), **pK₁ 6.8**, **pK₂ 7.2**. Puromycin dihydrochloride is purified by recrystallisation from H₂O (solubility 50mg/ml). It has some solubility in EtOH (~1mg/ml), DMSO (~13mg/ml) and DMF (~14mg/ml). The **free base** [58-60-6] **M 544.4**, has **m 175.5-177°** (**172-173°**) (from H₂O). The **sulfate** has **m 180-187° dec** (from H₂O), and the **picrate monohydrate** has **m 146-149°** (from H₂O). [Baker et al. *J Am Chem Soc* **77** 1 1955, DOI: 10.1021/ja01606a001; Fryth et al. *J Am Chem Soc* **80** 2736 1958, DOI: 10.1021/ja01544a039.] An **aminopeptidase inhibitor** that terminates protein synthesis [Reboud et al. *Biochemistry* **20** 5281 1981, DOI: 10.1021/bi00521a029; *Beilstein* **26** III/IV 3704.]

Pyridoxal hydrochloride [65-22-5] **C₈H₉NO₃·HCl**, **M 203.6**, **m 173°(dec)**, **176-180°(dec)**, **pK₁²⁰ 4.23** (**3-OH**), **pK₂²⁰ 8.7** (**Pyridinium⁺**), **pK₃²⁰ 13.04** (**CH₂OH?**). Dissolve it in water and adjust the pH to 6 with NaOH. Set aside overnight to crystallise. The crystals are washed with cold water, dried in a vacuum desiccator over P₂O₅, and stored in a brown bottle at room temperature. The **free base** [66-72-8] **C₈H₉NO₃**, **M 167.2**, **m 165° dec**, is then converted to the hydrochloride with one equivalent of HCl. [Fleck & Alberty *J Phys Chem* **66** 1678 1962, DOI: 10.1021/j100815a028; *Beilstein* **21/13** V 44.]

Pyridoxal-5'-phosphate monohydrate (**PLP**, **codecarboxylase**) [54-47-7, 41468-25-1] **C₈H₁₀NO₆P·H₂O**, **M 265.2**, **m 140-143°(dec)**, **pK₁²⁵ <2.5** (**PO₄**), **pK₂²⁵ 4.14** (**3-OH**), **pK₃²⁵ 6.20** (**PO₄**), **pK₄²⁵ 8.69** (**pyridinium⁺**). PLP has been purified by dissolving 2g in H₂O (10-15ml, in a dialysis bag a third full) and dialysing with gentle stirring against 1L of H₂O (+ two drops of toluene) for 15 hours in a cold room. The dialysate is evaporated to 80-100ml, then lyophilised. Lemon yellow microscopic needles of the **monohydrate** remain when all the ice crystals have been removed. Check purity by paper chromatography (in EtOH or *n*-PrOH/NH₃), and visualise spot(s) under UV light after spraying with *p*-phenylene diamine, NH₃ and molybdate. Solutions in a freezer are 2-3% hydrolysed in 3 weeks. Only 4-6% hydrolysis occurs in N NaOH or HCl at 25°, and 2% at 37° in 1 day — but is complete at 100° in 4 hours. Store as a dry solid at -20°. In aqueous acid the solution is colourless but is yellow in alkaline solutions. It has UV with λ_{\max} at 305nm (ϵ 1100) and 380nm (ϵ 6550) in 0.1 N NaOH; 330nm (ϵ 2450) and 388nm (ϵ 4,900) in 0.05M phosphate buffer pH 7.0, and 295nm (ϵ 6700) in 0.1N HCl. [Peterson et al. *Biochemical Preparations* **3** 34, 119 1953.] The **oxime** decomposes at **229-230°** and is insoluble in H₂O, EtOH and Et₂O. The ***O*-methyloxime** decomposes at **212-213°**. [Heyl et al. *J Am Chem Soc* **73** 3430 1951, DOI: 10.1021/ja01151a126.] Also purify by chromatography through a column of Amberlite IRC-50 (H⁺) [Peterson & Sober *J Am Chem Soc* **76** 169 1954, DOI: 10.1021/ja01630a045]. [*Beilstein* **21/13** V 46.]

Pyridoxamine hydrochloride [5103-96-8, 524-36-7 (*free base*)] **C₈H₁₂N₂O₂·2HCl**, **M 241.2**, **m 224-226°(dec)**, **226-227°(dec)**, **pK₁²⁵ 3.54** (**3-OH**), **pK₂²⁵ 8.21** (**ring N⁺**), **pK₃²⁵ 10.63** (**NH₂**). The amine salt is recrystallised from hot MeOH. The **free base** [85-87-0] **C₈H₁₂N₂O₂**, **M 168.2**, crystallises from EtOH with **m 193-193.5°** [Harris et al. *J Biol Chem* **154** 315 1944, <http://www.jbc.org/content/154/1/315>; *J Am Chem Soc* **66** 2088 1944, DOI: 10.1021/ja01240a025]. [*Beilstein* **22** IV 6064, **22/12** V 324.]

(±)-Quinacrine [Atebrine, Mepacrine, **3-chloro-9(4-diethylamino-1-methyl)butylamino-7-methoxy)-acridine**] dihydrochloride [69-05-6] **C₂₃H₃₀ClN₃O·2HCl**, **M 472.9**, **m 248-250°(dec)**, **pK₁³⁰ -6.49** (aq **H₂SO₄**), **pK₂³⁰ 7.73** (**ring NH⁺**), **pK₃³⁰ 10.18** (**Et₂N**). It crystallises from H₂O (soly 2.8w/v% at ~25°) as yellow crystals. Poorly soluble in MeOH and EtOH. The **free base** has **m 86-88°** (from Me₂CO or petroleum ether) or **m 85-87.5°** (aqueous EtOH). The **bismethiodide** has **m 224°** (from MeOH/EtOAc/Et₃N), and **picrate** has **m 207-208° dec** (from Me₂CO/EtOH). It is an antimalarial, antiprotozoal and intercalates DNA. [Wolfe *Antibiot* **3** (Springer-Verlag) 203 1975, *Beilstein* **22** III/IV 6247, **22/12** V 235.]

Quizalofop ethyl {[ethyl 2-[4-(6-chloro-2-quinoxalinyloxy)phenoxy]propionate} [(±) 76578-14-8; *R*-100646-51-3, 100646-52-4, 100760-08-5, 100760-10-9] **C₁₉H₁₇ClN₂O₄**, **M 372.8**, **m 91.7-92.1°**, **92-93°**, **b 220**

/0.2mm. This (\pm)-**herbicide** forms white crystals from Me₂CO/EtOH and sublimes *in vacuo*. Large quantities can be distilled at high vacuum. Its solubilities at 20° in g/10ml are 0.09 (EtOH), 1.1 (Me₂CO), 1.2 (xylene), 2.9 (*C₆H₆), and is nearly insoluble in H₂O (0.3mg/l). The (\pm)-**acid** has [95977-28-9]. The **R-ester enantiomer** has **m 76-77°** (pale brown crystals from EtOH), [α]_D²⁰ **+35.9 (EtOH)**. It is the **more active fatty acid synthase inhibitor** (designated DPX-Y6202) and is used to control grassy weeds in broadleaf crops. The **R-acid** has [94051-08-8]. [Shiu et al. [Rev Environ Contam Toxicol **116** 15 1990, PMID: 2205893]. **POISONOUS**

Racetams are pyrrolidin-2-ones with **nootropic properties**, and of importance for the treatment of Alzheimer. The effects of **racetams** potentiation already present neurotransmission, and appear to modulated ion flux. They have low toxicity and lack serious side effects. For a review on Piracetam and related nootropics' see Gouilaev & Senning [Brain Research Rev **19** 180 1994, DOI: 10.1016/0165-0173(94)90011-6 and Waegemans et al. Dementia and geriatric cognitive disorders **13**(4) 217 2002, DOI:10.1159/000057700, PMID: 12006732]. Some of the compounds studied are listed below.

PIRACETAMIDE (2-oxo-1-pyrrolidine acetamide) [7491-74-9] C₆H₁₀N₂O₂, **M 142.2, m 151.5-152.5°**. This typical nootropic (Alzheimer) drug modulates Na flux in AMPA receptors and is purified by recrystallisation from isoPrOH. **ANIRACETAM (1-[4-methoxybenzyl]-2-pyrrolidinone)** [72432-10-1] C₁₂H₁₃NO₃, **M 219.1, m 121-122°**. Purify aniracetam by recrystallisation from EtOH. [Hoffmann-La Roche & Co AG Jpn Kokai Tokkyo Koho JP 54,117,468 1979; Chem Abstr **92** 41755t.] **OXIRACETAM (4-hydroxy-2-oxo-1-pyrrolidine acetamide)** [62613-82-5 unspecified, 68567-97-5 racemate] C₆H₁₀N₂O₃, **M 158.2, m 160-162°, 165-168°**. This nootropic (Alzheimer, anti-amnesic, memory enhancing) drug is purified by recrystallisation from MeOH or aqueous Me₂CO and dried *in vacuo*. [NMR, IR: Pifferi & Pinza Farmaco Ed Sci **32** 602 1977, Banfi et al. Farmaco Ed Sci **39** 16 1984.] **R-(+)-OXIRACETAM (4-hydroxy-2-oxo-1-pyrrolidine acetamide)** [68252-28-8] C₆H₁₀N₂O₃, **M 158.2, m 135-136°, [α]_D²⁰ +36.2 (c 1, H₂O)**, and **S-(-)-oxiracetam** [88929-35-5] **m 135-136°, [α]_D²⁰ +36.5 (c 1, H₂O)**, have been recrystallised from aqueous Me₂CO. There are differences in activity between the enantiomers in which it appears that the **S-enantiomers** exhibit **significantly higher activities**. [Gouilaev & Senning Brain Research Rev **19** 180 1994, DOI: 10.1016/0165-0173(94)90011-6] **PRAMIRACETAM [amacetam, N-(2-(diidopropylamine)ethyl-2-oxo-1-pyrrolidineacetamide** [68497-62-1, 72869-16-0 sulfate] C₁₄H₂₇N₃O₇, **M 269.4**, has **m 47-48° (monohydrate), b 164°/0.15mm**. [Preparation: L'Italien & Nordin (Parke, Davis & Co) Ger Offen DE 2,808,067 1978; Chem Abstr **90** 22798b; US Pat US 4,145,347 1979; Chem Abstr **91** 39332p.] **ETIRACETAM (α -ethyl-2-oxo-1-pyrrolidineacetamide)** [(\pm)- 103833-73-4, (R)-103765-01-1, (S)-102767-28-2] C₈H₁₄N₂O₂, **M 170.1**, has **m 122°**, [Gobert et al. (UCB S.A.) for S-isomer Eur Pat Appl, EP 162,036 1985; CA **105** 18467d; Gobert et al. (UCB S.A.) for R-isomer see Eur Pat Appl, EP 165,919 1985; Chem Abstr **105** 97305a.] **NEFIRACETAM (2-oxo-pyrrolidineacetic acid 2,6-dimethylanilide)** [77191-36-7] C₁₄H₁₈N₂O₂, **M246.3**, [Betzing et al. (GmBH) Ger Offen DE 2,924,011 1980; Chem Abstr **94** 156740t.] **ROLZIRACETAM (tetrahydropyrrolizine-3,5-dione)** [18356-28-0] C₇H₉NO₂, **M 139.2**, has **m 176-177° (from EtOH), 181° (from *C₆H₆), 181.5-182° (from H₂O) and b 173°/5mm; 192°/0.5mm)**. [Leonard et al. J Am Chem Soc **69** 690 1947, DOI: 10.1021/ja01195a067.]

Ramipril {Prilace, Ramace, Altace, (2S,3aS,6aS)-1-[(2S)-2-[(1S)-1-ethoxycarbonyl-3-phenylpropyl]amino]-1-oxopropyl]-octahydrocyclopenta[b]pyrrole-2-carboxylic acid, (1S,5S,7S)-8-[(2S)-2-[(1S)-1-ethoxycarbonyl-3-phenylpropyl]amino]propanoyl]-8-azabicyclo[3.3.0]octane-7-carboxylic acid} [87333-19-5, mixt diastereomers 87269-97-4] C₂₃H₃₂N₂O₅, **M 416.5, m 109°, d²⁰ 1.20g/cm³, [α]_D²⁴ +33.2 (c 0.1N HCl in EtOH), pK_a, 3.17**. Prepared in various ways by Gold et al. [US 4587258 1986], e.g. condensing benzyl *cis,endo*-octahydrocyclopenta[b]pyrrole-2-carboxylate hydrochloride with N[1(S)-carboethoxy-3-phenylpropyl]-(S)-alanine using a carbodiimide method then de-benzylating (with 10% Pd/C) and recrystallising of the resulting **1-[N-(1(S)-carboethoxy-3-phenylpropyl)-(S)-alanyl]-cis,endo-octahydrocyclopenta[b]pyrrole-2(S)-carboxylic acid** (**m 110-112° dec**) from Et₂O. It provides a **hydrochloride** when a 1.3M HCl in Et₂O is added to it. See also the patent of Stepankova et al. [WO2005121084, 2005]. This drug is an **ACE** (angiotensin converting enzyme, see 'Proteins, Enzymes...' in this Chapter) **inhibitor**, causing vasodilation by blocking the conversion of angiotensin I to angiotensin II which lowers blood. [See Ruggerenti et al. Lancet **352** 1252 1998, PMID: 9788454; and Bosch et al. Br Med J **324** 699 2002, PMCID: PMC99052.]

Captopril {Acepress, Capoten, 1-[(2S)-3-mercapto-2-methyl-1-oxopropyl]-L-proline} [62571-86-2] C₉H₁₅-

NO₃S, **M 217.3**, **m 103-104°** (polymorphic unstable form **m 86°**, melts at **87-88°** solidifies and then melts again at **104-105°**), $[\alpha]_D^{25}$ **-131.0** (c 1.7, EtOH), **pK₁²⁵ 3.7** (COOH), **pK₂²⁵ 9.8** (SH). It is the first **antihypertensive ACE** (Angiotensin-Converting Enzyme) **inhibitor** from the work of Nobel Laureate John Vane and developed by Squibb & Sons Pharmaceuticals. The synthesis involved reaction of L-proline with (2S)-acetylthiopropionyl chloride in basic conditions (NaOH) followed by deacylation (NH₃) [Shimazaki et al. *Chem Pharm Bull Jpn* **30** 3139 1982, DOI:10.1248/cpb.30.3139], and later improved by Nam et al. [*J Pharm Sci* **73** 1843 1984, DOI:10.1002/jps.2600731251. PMID: 6396401]. Purify it by recrystallisation from EtOAc/hexane. It is purified also by dissolving in the minimum volume of EtOAc and fractionating through a column of Wakogel C200 with a linear gradient (0 to 100) of MeOH and EtOAc, and the fractions which gave a positive test with sodium nitroprusside (test for SH) are combined, evaporated and the residue recrystallised from EtOAc/hexane (1:1) to give the white crystalline drug [Nam et al. *J Pharm Sci* **73** 1843 1984, DOI:10.1002/jps.2600731251. PMID: 6396401]. Alternatively, dissolve it in H₂O, purify through AG-50Wx2 (BioRad)/eluting with H₂O. The free acid is converted to the **dicyclohexylamine salt** in MeCN by addition of the amine until the pH is 8-9. The salt free acid is obtained by shaking with EtOAc/10% aqueous KHSO₄ or passage through AG50Wx2. The EtOAc solution is dried (MgSO₄), evaporated and the residue is recrystallised from EtOAc/hexane [Cushman et al. *Biochemistry* **16** 5484 1977, DOI: 10.1021/bi00644a014]. Of the N-proline derivatives that Cushman et al. studied for competitive ACE inhibition **Captopril** was the **most potent** with a K_i of 0.0017 μM (in 0.1M phosphate, 0.3M NaCl, pH 8.3 with Hip-His-Leu (5mM) as substrate at 37°).

Resazurin sodium salt (Alamar blue, 7-hydroxy-3H-phenoxazin-3-one-10-oxide Na salt) [62758-13-8] **C₁₂H₆NO₄. Na**, **M 251.2**, **decomposes on heating**. It is a bluish powder that can be dissolved in hot MeOH and precipitated out on adding Et₂O. Alternatively, dissolve it in conc H₂SO₄ and dilute the solution (**care**, use ice cooling). Filter off the orange solid, wash well with hot H₂O and dry *in vacuo*. The **Na salt** can be obtained by careful titration with NaOH or Na₂CO₃. It is soluble in H₂O and MeOH, less in higher alcohols, and has λ_{\max} (MeOH) at 604nm with ϵ_{\max} 60,000cm⁻¹M⁻¹. It is an **indicator**: pH 3.8 (orange) —6.5 (blue). It is a versatile **biological stain**. [For preparation and properties see Sabnis in *Handbook of Biological Dyes and Stains* Wiley, Hoboken pp 406-407 2010, onlinelibrary.wiley.com/doi/10.1002/9780470586242.fmatter/pdf; for oxidation-reduction of Resazurin and Resorufin (below) see Twigg *Nature* **155** 401 1945, DOI: 10.1038/155401a0.]

Rescinamine (Anaprel, Apoterin, a β-carboline alkaloid) [24815-24-5] **C₂₅H₄₂N₂O₉**, **M 634.7**, **m 238-239°**(vacuum), **240°**, $[\alpha]_D^{20}$ **-97** (c 1, CHCl₃), **pK_{Est(1)} ~<0** (carbazole N), **pK_{Est(2)} ~7.0** (quinolizidine N), **pK₂²⁵ 6.4** (75% aqueous HCONMe₂). Crystallise it from *benzene, MeOH or aqueous Me₂CO. The **hydrochloride** has **m 232° dec** (from MeOH) and $[\alpha]_D^{20}$ **-74** (MeOH). It is an antihypertensive. [Klohs et al. *J Am Chem Soc* **77** 2241 1955, DOI: 10.1021/ja01613a065; *Beilstein* **25** III/IV 1323.]

Resorufin (7-hydroxy-3H-phenoxazin-3-one Na salt) [34994-50-8] **C₁₂H₆NO₃. Na**, **M 235.2**, **decomposes on heating >300°**, **pK₂²⁵ 5.8** (H₂O), **6.10** (50% aqueous MeOH). The **free acid** [635-78-9] **C₁₂H₇NO₃**, **M 213.2**, is purified by dissolving the salt in ice cold concentrated H₂SO₄ which on dilution precipitates resorufin (pink) and is filtered off, washed with hot H₂O and dried. [Eichler *J Prakt Chem* **139** 113 1934, DOI: 10.1002/prac.19341390305.] Detailed UV and IR studies were reported by Musso & Matthies *Chem Ber* **90** 1814 1957, DOI: 10.1002/cber.19570900919]. For pK_a see Musso & Rathjen [*Chem Ber* **92** 751 1959, DOI: 10.1002/cber.19590920331; *Beilstein* **27** IV 2263.] See also entry [635-78-9] in 'Organo-metallic Compounds' in Chapter 4. The **O⁷-monoacetate** [1152-14-3] **C₁₄H₉NO₄**, **M 255.2**, **m 223°** is a **fluorogenic substrate** used for hydrolytic enzymes; its fluorescence has λ_{em} 593nm (λ_{ex} 500nm, in 0.1M phosphate pH 8.0 with lipase) [Kitson & Kitson *Biochem J* **322** 701 1997, DOI: 10.1042/bj3220701]. [*Beilstein* **27** IV 2263.]

The **O⁷-methylresorufin (7-methoxy-3H-phenoxazin-3-one)** [5735-89-3] **C₁₃H₉NO₃**, **M 227.2**, **m >220°** has fluorescence at λ_{em} 585nm (λ_{ext} 571 nm in deacylase solution) and is used to **differentiate isoenzymes of cytochrome P-450**. It is insoluble in H₂O and dilute alkali, but is soluble in EtOH and CHCl₃ to give an orange yellow colour. [Kehrmann *Justus Liebigs Ann Chem* **372** 287 1910, DOI: 10.1002/jlac.19103720303; *Beilstein* **27** II 108.] Several **O⁷-substituted** derivatives are commercially available. They are **biological stains and dyes**.

Rifampicin (Rifampin, Rifadin) [13292-46-1] **C₄₃H₅₈N₄O₁₂**, **M 823.0**, **m 183-185°**, **pK₁ 1.7**, **pK₂ 7.9**. This **macrolide antibiotic** crystallises from Me₂CO in red-orange plates. It has UV with λ_{max} at 237, 255, 334, and 475nm (ϵ 33,200, 32,100, 27,000 and 15,400) at pH 7.38. It is stable in Me₂SO and H₂O, and is freely soluble in

most organic solvents, but slightly soluble in H₂O at pH <6. [Binda et al. *Arzneim.-Forsch* **21** 1907 1971.] It *inhibits cellular RNA synthesis* without affecting DNA [Calvori et al. *Nature* **207** 417 1965, DOI: 10.1038/207418a0].

Rifamycin B [13929-35-6] C₃₉H₄₉NO₁₄, M 755.8, m 300° (darkening at 160-164°), [α]_D²⁰ -11° (MeOH), pK₁ 2.60, pK₂ 7.76. Rifamycin B forms yellow needles from *C₆H₆. Its solubilities are: H₂O (0.027%), MeOH (2.62%) and EtOH (0.44%), also soluble in DMSO and EtOAc. It has UV with λ_{max} at 223, 304 and 245nm (A_{1cm}^{1%} 555, 275 and 220). [Oppolzer & Prelog *Helv Chim Acta* **56** 2287 1973, DOI: 10.1002/hlca.19730560717; Oppolzer et al. *Experientia* **20** 336 1964, PMID: 5855864; for X-ray structure see Brufani et al. *Experientia* **20** 339 1964, PMID: 5855865.] It is a macrolide antibacterial agent.

Rifamycin SV [6998-60-3] C₃₇H₄₆NO₁₂·Na, M 697.8, m 300° (darkening >140°), [α]_D²⁰ -4° (MeOH), pK_{Est} ~7.8. Rifamycin SV gives yellow-orange crystals from Et₂O/petroleum ether or aqueous EtOH, is very soluble in MeOH, EtOH, Me₂CO and EtOAc, and is less soluble in Et₂O and HCO₃⁻, but slightly soluble in H₂O and petroleum ether. Its UV has λ_{max} at 223, 314 and 445nm (A_{1cm}^{1%} 586, 322 and 204) in phosphate buffer pH 7. [For NMR see Bergamini & Fowst *Arzneim.-Forsch* **15** 951 1965.] It is a macrolide antibacterial agent.

(-)-Scopolamine hydrobromide 3H₂O (6β,7β-epoxy-3α-tropanyl S(-)-tropate HBr, hyoscine HBr) [114-49-8, 6533-68-2, 51-34-3 free base] C₁₇H₂₁NO₄·HBr·3H₂O, M 438.3, m 193-194°, 195°, 195-199°, [α]_D²⁵ -25(c 5, H₂O), pK₂₀ 8.15. The *hydrobromide* is crystallised from Me₂CO, H₂O or EtOH/Et₂O, and dried *in vacuo*. It is soluble in H₂O (60%) and EtOH (5%) but insoluble in Et₂O and slightly in CHCl₃. The *hydrochloride* has m 300° (from Me₂CO). The *free base* is a viscous liquid which forms a crystalline *hydrate* with m 59° and [α]_D²⁰ -28 (c 2.7, H₂O). It hydrolyses in dilute acid or base. [Meinwald *J Chem Soc* 712 1953, DOI: 10.1039/JR9530000712; Fodor *Tetrahedron* **1** 86 1957, DOI: 10.1016/0040-4020(57)85013-3; Beilstein **6** III 4185.] A drug for treatment of motion sickness (Kwells) and post-operative nausea and vomiting.

Serotonin hydrochloride (5-HT, 3-[2-aminoethyl]-5-hydroxyindole HCl) [153-98-0] C₁₀H₁₂N₂O·HCl, M 212.7, m 149-154°, 167-168°, 178-180°, pK₁²⁵ 4.9, pK₂²⁵ 9.8 (10.0, NH₂), pK₃²⁵ 11.1 (5-OH), pK₄²⁵ 18.25 (acidic indole NH). 5-HT is purified by recrystallisation from EtOH/Et₂O or Et₂O to give the *hygroscopic* salt. Store it in the dark as it is light sensitive. The *free base* has m 84-86° (from Et₂O). The *5-benzyloxy* derivative has m 84-86° (from Et₂O). [Ek & Witkop *J Am Chem Soc* **76** 5579 1954, DOI: 10.1021/ja01651a001; Hamlin & Fischer *J Am Chem Soc* **73** 5007 1951, DOI: 10.1021/ja01154a551.] The *picrate* 1H₂O has m 196-197.5° (dec with sintering at 160-165°) after crystallisation from Et₂O. Serotonin is a natural neurotransmitter [Roth & Chuang *Life Sci* **41** 1051 1987, DOI: 10.1016/0024-3205(87)90621-7]. [Beilstein **22/12** V 16.]

Spectinomycin dihydrochloride pentahydrate (Actinospectacin) [22189-32-8, 21736-83-4] C₁₄H₂₄N₂O₇·2HCl·5H₂O, M 495.3, m 205-207° (dec), [α]_D²⁰ +14.8° (c 0.4, H₂O), pK₁ 6.95, pK₂ 8.70. The salt is purified by recrystallisation from aqueous Me₂CO and is soluble in H₂O, MeOH and dilute acid and base, but only slightly soluble in Me₂CO, EtOH, CHCl₃ and *C₆H₆. The *free base* is an amorphous solid, m 184-194° with [α]_D²⁰ -20 (H₂O). [Wiley et al. *J Am Chem Soc* **93** 2652 1963, X-ray: Cochran et al. *JCS Chem Commun* 494 1972, DOI: 10.1039/C39720000494.] It is an *aminoglycoside antibiotic* which interacts with 16S ribosomal RNA [Moazed & Noller *Nature* **327** 389 1987, DOI: 10.1038/nature01493, PMID: 2953976] and is used for the *treatment of gonorrhea* [Rinehart *J Infect Dis* **119** 345 1969, DOI: 10.1093/infdis/119.4.5.345].

D-Sphingosine (2S,3S-D-erythro-2-aminooctadec-4t-ene-1,3-diol from bovine brain) [123-78-4] C₁₈H₃₇NO₂, M 299.5, m 79-82°, 82° 82.5° (softens at ~70°), [α]_D²² -3.4 (c 2, CHCl₃), pK_{Est} ~8.8. D-Sphingosine is purified by re-crystallisation from EtOAc, Et₂O or petroleum ether (60-80°). It is insoluble in H₂O but is soluble in Me₂CO, EtOH and MeOH and CHCl₃ (20mg/ml). It has IR bands at 1590 and 875 cm⁻¹, and is characterised as the *tribenzoate* m 122-123° (from 95% EtOH). Store it at -20°. [Tipton *Biochemical Preparations* **9** 127 1962; for 'Biochemistry of the sphingolipides. III. Structure of sphingosine' see Carter et al. *J Biol Chem* **170** 285 1947, <http://www.jbc.org/content/170/1/285>; and Carter et al. *J Biol Chem* **169** 77 1947, PMID: 20240540.] Important lipid for membranes, and in cellular signaling and other processes. For 'Killing tumours by ceramide-induced apoptosis: a critique of available drugs' see Radin [*Biochem J* **371** (Pt 2) 243

2003, DOI: 10.1042/BJ20021878, PMID: 12558497].

Statins. These are a group of lipid lowering drugs that act by strongly inhibiting 3-hydroxy-3-methylglutamylCoA reductase (HMG-CoA reductase K_i is ~in the nM range) which is the first enzyme of the *de novo* biosynthesis of cholesterol (e.g. in mammalian liver) and have the effect of increasing LDL (low density lipoproteins) uptake by hepatocytes, decrease HDL-C (high density lipoprotein cholesterol) and lower triglyceride levels. The pyrrol-type statins related to Lipitor have been mentioned above, and here are included four 1,2,6,7,8,8a-hexahydronaphthalene-type statins (denoted *S*) with a *S-I*-(*R*-3,4-dihydroxyheptanoic acid, or its lactone group), an *S-2*-methyl group, a 6-unsubstituted or a *S-6*-(methyl or hydroxyl) group, *S-8*-hydroxy group (esterified), and an *R-8a* hydrogen at the bridgehead. The lactones are *pro-drugs* which are hydrolysed to the corresponding hydroxyl-heptanoic acids *in vivo*. These acids can be converted into their lactones by azeotropic removal of H₂O in boiling toluene.

Pravastatin Na salt {Pravachol, (β *R*, δ *R*-1*S*,2*S*,6*S*,8*S*,8*aR*)-1,2,6,7,8,8a-hexahydro- β , δ ,6-trihydroxy-2-methyl-1-oxo-2-methyl-8-[*S*-2-methyl-1-oxobutoxy]-1-naphthaleneheptanoic acid monoNa salt) [81131-70-6, free acid 81093-37-0] C₂₃H₃₅O₇Na, M 446.5, m 171.2-173° (for free acid), m 138-142° (for δ -lactone), $[\alpha]_D^{25}$ +194 (c 0.5, MeOH for δ -lactone), pK₁²⁵ 4.21 (CO₂H), pK₂²⁵ -2.7 (basic). Crystallise from H₂O (solubility 0.45g/L) by addition of Me₂CO, or MeOH/Me₂CO then wash with Et₂O and dry *in vacuo* to give a pale white powder or fine crystals. It is soluble in MeOH but insoluble in Me₂CO, MeCN and CHCl₃. The UV (MeOH) has λ_{max} at 230, 237 and 245nm, as for the δ -lactone. Its K_i (*in vivo*) for HMG-CoA reductase is ~1nM, and displays cardioprotective properties. [Hamelin & Turgeon *Trends Pharmacol Sci* **19** 26 1998, PMID: 9509899.]

Lovastatin {Mevacore, Mevinolin, (2*S*)-(1*S*,3*R*,7*S*,8*S*,8*aR*)-1,2,3,7,8,8a-hexahydro-3,7-dimethyl-8-[2-[(2*R*,4*R*)tetrahydro-4-hydroxy-6-oxo-2*H*-pyran-2-yl]ethyl]-1-naphthalenyl-methyl butanoate} [75330-75-5] C₂₄H₃₆O₅, M404.5, m 174.5° (under N₂), $[\alpha]_D^{25}$ +323 (c 0.5, MeCN), pK₁²⁵ 14.9 (v weak acid), pK₂²⁵ -2.8 (v weakly basic). This fungal metabolite was isolated from *Aspergillus Terreus*. Recrystallise it from a combination of solvents which do not allow hydrolysis of the lactone ring. It crystallises from EtOH in thick colourless needles, and crystallises also from MeCN then EtOAc and dried at 40°/16hrs. Its solubility at ~25° (g/100ml) is: CHCl₃ (35), DMF (9.0), Me₂CO (4.7), DMSO (4.0), MeCN (2.8), MeOH (2.8), *iso*PrOH (2.0), EtOH (1.6), *iso*BuOH (1.4), *n*-PrOH (1.1), *n*-BuOH (0.7), *n*-octanol (0.2), H₂O (almost insol, 0.4 x 10⁻⁵). The UV (MeOH) has λ_{max} at 231 (ϵ 21,490), 238 (ϵ 25,090) and 247 (ϵ 16,890 M⁻¹cm⁻¹) nm, and ¹H NMR and MS are consistent with its structure and similar to those reported below for Mevastatin. [Alberts et al. *Proc Natl Acad Sci USA* **77** 3957 1980; PMID: 6933445; Hirama & Iwashita *Tetrahedron Lett* **24** 1811 1983, DOI: 10.1016/S0040-4039(00)81777-3]. Its K_i (*in vivo* and *in vitro*) for HMG-CoA reductase is ~0.6nM, decreases CDK2, 4, 6, and cyclin E levels, and induces G₁ arrest and apoptosis in tumour cell lines *in vitro* [Alberts *Am J Cardiol* **62** 10J 1988, PMID: 3055919; Park et al. *Anticancer Res* **19** 3133 1999, PMID: 10652602.]

Mevastatin {Compactin (2*S*)-2-methyl-(1*S*,7*S*,8*S*,8*aR*)-1,2,3,7,8,8a-hexahydro-7-methyl-8-[2-[(2*R*,4*R*)tetrahydro-4-hydroxy-6-oxo-2*H*-pyran-2-yl]ethyl]-1-naphthalenyl-methyl butanoate} [73573-88-3] C₂₃H₃₄O₅, M 390.5, m 152°, $[\alpha]_D^{22}$ +283 (c 0.84, Me₂CO), pK₁²⁵ 4.21 (CO₂H), pK₂²⁵ -2.7 (basic) of hydrolysed lactone. It has been isolated from *Penicillium brevicompactum* and purified by recrystallisation from aqueous EtOH. Its UV (EtOH) has λ_{max} at 230 (log ϵ 4.28), 237 (log ϵ 4.30) and 246 (log ϵ 4.11) nm; the IR (KBr) has ν_{max} at 3520, 1750 (lactone CO) and 1710 (ester CO) cm⁻¹; the IR (CHCl₃) has ν_{max} at 3510 and 1724 (lactone and ester CO) cm⁻¹; the ¹H NMR (CDCl₃) has δ at 5.59 (1H, d, *J*=10Hz, H-4), 5.71 (1H, dd, *J*=10 and 5Hz, H-3'), 5.44 br (1H, s, H-5'), 5.33br (1H, s, H-8'), 4.62 (1H, m, H-3), 4.35 (1H, m, H-5), 3.28br (1H, s, OH), 2.64 (2H, d, *J*=4Hz, H-6), 2.37 (2H, m, H-2' and H-2''), 2.2–1.3 (14H, m, CH₂ and CH), 1.13 (3H, d, *J*=7Hz, 2''-Me), 0.90 (3H, d, *J*=7Hz, 2'-Me) and 0.88 (3H, t, *J*=7.5Hz, CH₃-CH₂); MS has *M*⁺ 390.2403, C₂₃H₃₄O₅ requires *M* 390.2406, and *m/e* 390(4%), 372(3), 288(4), 273(4), 270(12), 210(14), 186(12), 185(42), 184(57), 183(24), 169(11), 159(34), 158(56), 155(30), 145(100), 144(40), 143(81), 129(28), 91(15) and 57(31). The *benzoate* has m 88-89° after recrystallisation from aqueous EtOH and MS has *M*⁺ 494.2667, C₃₀H₃₈O₆ requires *M* 494.2668. A fragment was cut from a large crystal and subjected to X-ray crystallographic studies which showed that the structure and relative configurations of the chiral centres were as expected for this lactone. [Brown et al. *JCS Perkin Trans 1* 1165 1976, DOI: 10.1039/P19760001165.] [For the total chiral synthesis of Compactin see Hirama & Uei *J Am Chem Soc* **104** 4251 1982, DOI: 10.1021/ja00379a037.]

Simvastatin {Zocor, Sivastin (1*S*,3*R*,7*S*,8*S*,8*aR*)-1,2,3,7,8,8a-hexahydro-3,7-dimethyl-8-[2-[(2*R*,4*R*)-tetrahydro-4-hydroxy-6-oxo-2*H*-pyran-2-yl]ethyl]-1-naphthalenyl-2,2-dimethylbutanoate} [79902-63-9]

C₂₈H₃₈O₅, M 418.6, m 135-138°, [α]_D²⁵ +292 (c 0.5, MeCN), pK₁²⁵ 4.21 (CO₂H), pK₂²⁵ -2.7 (basic) of **hydrolysed lactone**. By altering the acid which esterifies the OH group at S-1 from 2-methylbutanoic acid (in Mevinolin, Lovastatin above) to 2,2-dimethylbutanoic acid (in Simvastatin, Zocor) Hoffman et al. [*J Med Chem* **29** 849 1986, DOI: 10.1021/jm00155a040] have succeeded in increasing the intrinsic inhibition of HMG-CoA reductase by ~2.5 times. Prepared by adding 2,2-dimethylbutyryl chloride (4.6mmol, b 130-134°/atm) over 2 minutes to a stirred solution of the **S-I-OH** compound (0.5g, 1.15mmol) and DMAP (50mg) in dry pyridine (5ml) at 0°/N₂, and stirring at 0°/1hr then at ~25°/18hrs. Et₂O (100ml) is added to the mixture which is washed with 2% aqueous HCl (3 x 25ml), saturated NaHCO₃ solution (25ml), and brine (2 x 25ml), the organic layer is dried (MgSO₄), filtered and evaporated to dryness. The residual oil is chromatographed through a silica gel (230-400 mesh) column, eluted with CHCl₃/Me₂CO (32:1 v/v, 200ml), discarded this fraction, then continue with the same eluant (200ml) to give the **desired ester**. This is then recrystallised from *n*-C₄H₉Cl/hexane. It is soluble (in g/100ml at ~25°) in *n*-hexane (0.015), propylene glycol (3), polyethylene glycol-400 (7), EtOH (16), MeOH (20), DMSO (54), CHCl₃ (61), 0.1M HCl (0.006) and 0.1N NaOH (7, most probably the lactone has hydrolysed to give the Na salt of the hydroxyl acid formed). [For the preparation by fermentation of *Penicillium citrinum* and further reactions, see patents to Merck and Co 1980, by Hoffman et al. US 444.4784, and Willard et al. EP 33538.] Apart from inhibiting HMG-Co reductase and decreasing levels of LDL [White *J Clin Pharmacol* **39** 111 1999, DOI: 10.1177/00912709922007642], it has many other biological effects such as stimulating bone formation (*in vitro* and *in vivo*) [Garrett et al. *Curr Pharmaceut Des* **7** 715 2001, DOI: 10.2174/1381612013397762], inhibition of smooth muscle cell proliferation and migration [Reinso et al. *Methods Find Exp Clin Pharmacol* **24** 593 2002, PMID: 12616706], and anticancer and anti-inflammatory activity [Kaushal et al. *Endothelium* **10** 49 2003, PMID: 12699077].

Sterigmatocystin (3a,12c-dihydro-8-hydroxy-6-methoxy-3H-furo[3',2':4,5]furo[2,3-c]xanthen-7-one) [10048-13-2] **C₁₈H₁₂O₆**, M 324.3, m 246°, 247-248°, [α]_D²⁰ -398 (c 0.1, CHCl₃), pK_{Est} ~ 8.0. It crystallises from amyl acetate, Me₂CO or EtOH and sublimes *in vacuo*. It has UV with λ_{\max} at 208, 235, 249 and 329nm (log ϵ 4.28, 4.39, 4.44 and 4.12). [For UV see Bullock et al. *J Chem Soc* 4179, 1962, DOI: 10.1039/JR9620004179; for UV and IR see Holker & Mulheirn *JCS Chem Commun* 1576 1968, DOI: 10.1039/C19680001576; Birkinshaw & Hammady *Biochem J* **65** 162 1957, DOI: 10.1042/bj0650162.] This **mycotoxin** induces **bone marrow changes** in mice [Curry et al. *Mutation Res* **137** 111 1984]. [Beilstein **19/10** V 575.]

Stigmatellin A (2-[4,6-dimethoxy-3,5,11-trimethyltridecatri-7*t*,9*t*,11*t*-enyl]-8-hydroxy-5,7-dimethoxy-3-methyl-4H-1-benzopyran-4-one) [91682-96-1, 94234-27-2] **C₃₀H₄₂O₇**, M 514.6, m 128-130°, [α]_D²⁰ +38.5 (c 2.3, MeOH), pK_{Est} ~7 (phenolic OH). Stigmatellin A is stable in aqueous solution at neutral pH but decomposes at pH <5. It is purified by recrystallisation from toluene/hexane. It has UV with λ_{\max} nm (ϵ) at 248sh (4,100), 258 (59,500), 267 (65,500), 279 (1,400) and 335 (5,200) in MeOH; 249sh (45,600), 258 (60,000), 268 (72,700), 277 (54,100), 320 (2,500) and 370 (3,000) in MeOH + 1 drop of N KOH; 243sh (29,300), 264 (63,200), 274 (64,100), 283sh (45,800), 329 (4,800) and 420 (21,000) in MeOH + 6N HCl; and IR (CHCl₃) with ν_{\max} at 3550m, 1645chs, 1635ss, 1620ss, 1590s, 1510m and 905m cm⁻¹. It gives colour reactions at 110° with vanillin/H₂SO₄ (grey), Ce(IV)/(NH₄)₂SO₄ (yellow) and phosphomolybdate (blue-grey). [Höfle et al. *Justus Liebigs Ann Chem* 1883 1984, DOI: 10.1002/jlac.198419841202.] It **inhibits electron transport** by interfering with the quinol oxidation (Q_o) site of cytochrome Bc1 [von Jagow & Link *Methods Enzymol* **126** 253 1986, DOI: 10.1016/S0076-6879(86)26026-7; Robertson et al. *Biochemistry* **32** 1310 1993, DOI: 10.1021/bi00056a016], and has antibiotic properties [Kunze et al. *J Antibiot* **37** 454 1984]. The **7*t*,9*t*,11*c*-isomer** is **Stigmatellin B**.

Streptomycin sulfate [3810-74-0] **C₂₁H₃₉N₇O₁₂ · 1.5H₂SO₄**, M 7268.7, 1457.4 (for 2 base. 3 H₂SO₄), [α]_D²⁰ -84.3 (c 3, H₂O), pK_{Est(1)} ~ 9.5 (MeNH), pK_{Est(2,3)} ~ 13.4 (guanidino). The sulfate is recrystallised from H₂O/EtOH, washed with a little EtOH, Et₂O and dried in a vacuum. [For UV and IR see Grove & Randall *Antibiotics Monographs* **2** 163 1955, Heuser et al. *J Am Chem Soc* **75** 4013 1953, DOI: 10.1021/ja01112a041; Kuehl et al. *J Am Chem Soc* **68** 1460 1946, DOI: 10.1021/ja01212a019; Regna et al. *J Biol Chem* **165** 631 1946, PMID: 20276129.] It **inhibits initiation and causes misreading of mRNA** during protein synthesis [Zierhut et al. *Eur J Biochem* **98** 577 1979, DOI: 10.1111/j.1432-1033.1979.tb13219.x; Chandra & Gray *Methods Enzymol* **184** 70 1990, DOI: 10.1016/0076-6879(90)84261-E]. [Beilstein **18/11** V 82.]

Streptonigrin (nigrin, 5-amino-6-[7-amino-5,8-dihydro-6-methoxy-5,8-dioxo-2-quinolinyl]-4-[2-hydroxy-3,4-dimethoxyphenyl]-3-methyl-2-pyridinecarboxylic acid) [3930-19-6] $C_{25}H_{22}N_4O_8$, **M 506.5, m 262-263°**, **275°(dec)**, **pK²⁵ 6.3 (1:1 aqueous dioxane)**. Streptonigrin is purified by TLC on pH 7-buffered silica gel plates and eluted with 5% MeOH/CHCl₃. The extracted band crystallises from Me₂CO or dioxane (black plates or needles) and is soluble in pyridine, Me₂NCHO, aq NaHCO₃ (some dec), and poorly soluble in MeOH, EtOH, EtOAc and H₂O. It has UV with λ_{\max} at 248, 375-380nm (ϵ 38,400 and 17,400). [Weinreb et al. *J Am Chem Soc* **104** 536 1982, DOI: 10.1021/ja00366a028; Rao et al. *J Am Chem Soc* **85** 2532 1963, DOI: 10.1021/ja00899a051.] It is an *anti-neoplastic antibiotic* [Wilson et al. *Antibiot Chemother* **11** 147 1961].

Succinyl coenzyme A trisodium salt [108347-97-3] $C_{25}H_{40}N_7O_{19}P_3S$, **M 933.5, 867.7 (free acid)**. If impure dissolve it in H₂O (0.05g/ml) adjusted to pH 1 with 2M H₂SO₄ and extract many times with Et₂O. Remove excess Et₂O by bubbling N₂ through it and store frozen at pH 1. When required, adjust the pH to 7 with dilute NaOH and use within 2 weeks (freeze samples). It is estimated by the hydroxamic acid method [Hersh & Jencks *J Biol Chem* **242** 3468 1967, <http://www.jbc.org/content/242/15/3468>; Hersh & Jencks *J Biol Chem* **242** 3481 1967, <http://www.jbc.org/content/242/15/3481>]. More stable in acidic than neutral aqueous solutions, but neutral solutions are stable at -15°. [Jordan & Laghai-Newton *Methods Enzymol* **123** 435 1986, DOI: 10.1016/S0076-6879(86)23054-2; Beilstein **26** III/IV 3666.] An intermediate in the citric acid cycle.

Terramycin (oxytetracycline) [79-57-2; 6153-64-6 (2H₂O)] $C_{22}H_{24}N_2O_9$, **M 460.4 (anhydrous), 496.5 (2H₂O), sinters at 182°, melts at 184-185°(dec)**, **[α]_D²⁰ -196.6 (equilibrium in 0.1M HCl), -2.1°(equilibrium in 0.1M NaOH)**. It crystallises (as *dihydrate*) from H₂O (0.25mg/ml at 25°) or aqueous EtOH and is soluble in MeOH and Me₂CO but insoluble in Et₂O and petroleum ether. It is amphoteric; aqueous solutions have pH 2.0-5.0. It has UV with λ_{\max} at 247, 275 and 353nm in 0.1 M phosphate (pH 4.5). [Finlay et al. *Science* **111** 85 1950, DOI: 10.1126/science.111.2874.85.] The *hydrochloride*, [2058-46-0] **M 496.9 [Beilstein 14 IV 2630]**, crystallises from MeOH in needles and from H₂O at 50° it forms plates. Terramycin has also been purified *via* the *hydrochloride* by dissolving it in H₂O, adjusting to pH 6, and the solid is filtered off after 1 hour. The crystals of the *dihydrate* are dried to constant weight in vacuum/CaCl₂/25°. Drying at 60° *in vacuo* gives the *anhydrous base m 184.5-185.5°* (sintering at 180°). The *dihydrate* has **m 181-182°**, and its optical rotation in MeOH decreases from **[α]_D²⁵ +26 (c 0.5%)** to **[α]_D²⁵ +11.3** after standing for 16 hours. Its biological half life is 6-8 hours. It forms a *sodium salt* and a *CaCl₂ complex*. [Regna et al. *J Am Chem Soc* **73** 4211 1951, DOI: 10.1021/ja01153a050; Beilstein **14** IV 2633.] A broad spectrum tetracycline antibiotic.

Tetracycline [60-54-8] $C_{22}H_{24}N_2O_8$, **M 444.4, m 172-174°(dec)**, **[α]_D²⁰ -291 (c 1, MeOH), pK₁²⁵ 3.30, pK₂²⁵ 7.68, pK₃²⁵ 9.69**. Recrystallise the antibiotic from toluene or aqueous MeOH as the *trihydrate*. Its biological half life is 8-11hrs (normally), but 57-108hrs (in kidney impairment). Store it at -20°. [Stephens et al. *J Am Chem Soc* **76** 3568 1954, DOI: 10.1021/ja01642a064; Beilstein **14** IV 2625.] **Tetracycline hydrochloride** [64-75-5] $C_{22}H_{24}N_2O_8 \cdot HCl$, **M 480.9, has m 214°(dec), 215-220°, [α]_D²⁵ -258 (c 0.5, 0.1N HCl), [α]_D²⁰ -245 (c 1, MeOH), [α]_D²⁵ -257.9 (c 0.5, aqueous HCl), pK₁ 1.4 (enolic OH), pK₂ 3.30, pK₃ 7.68 (phenolic OH), pK₄ 9.27 (Me₂N)**. The hydrochloride is recrystallised from MeOH/*n*-BuOH or *n*-BuOH/HCl. It is insoluble in Et₂O and petroleum ether. It has UV with λ_{\max} at 270 and 366nm in MeOH. [Gottstein et al. *J Am Chem Soc* **81** 1198 1959, DOI: 10.1021/ja01514a046; Conover et al. *J Am Chem Soc* **84** 3222 1962, DOI: 10.1021/ja00875a063; Stephens et al. *J Am Chem Soc* **78** 4155 1956, DOI: 10.1021/ja01597a081; Beilstein **14** IV 2627.]

Terreic acid (2-hydroxyl-3methyl-1,4-benzoquinone-5,6-epoxide) [121-40-4] $C_7H_6O_4$, **M 154.1, m 127-127.5°, [α]_D²² +74 (pH 4, phosphate buffer), pK²⁵ 4.5**. Recrystallise the acid from *C₆H₆, *C₆H₆/petroleum ether or hexane, and sublime at 80-100°/1mm. It is an inhibitor of Burton's tyrosine kinase. [Kawakami et al. *PNAS* **26** 2227 1999, PMID: 10051623; Beilstein **17** IV 6698.]

6R-Tetrahydro-erythro-biopterin dihydrochloride (BH₄·2HCl, 6R-2-amino-4-hydroxy-6-[(1R,2S)-1,2-dihydroxypropyl]-5,6,7,8-tetrahydropteridine 2HCl) [69056-38-8] $C_9H_{15}N_5O_3 \cdot 2HCl$, **M 314.2, m 245-246°(dec)**, **[α]_D²⁵ -6.8 (c 0.67, 0.1N HCl), pK₁ 1.37 (pyrimidine⁺), pK₂ 5.6 (5-NH⁺), pK₃ 10.6 (acidic, 3NH)**. Recrystallisation of BH₄·2HCl from HCl enriches BH₄ in the natural 6R isomer. Dissolve the salt (~6g) in concentrated HCl (15ml) under gentle warming, then add EtOH (30ml) dropwise, chill and collect the colourless needles (67%, up to 99% if the mother liquors are concentrated), and dry it *in vacuo* immediately over P₂O₅ and

KOH. It is stable indefinitely at -20° in a dry atmosphere. Better store it in sealed ampoules under dry N_2 . It can be recrystallised from 6N aqueous HCl. It has UV with λ_{\max} (in 2N HCl) at 264nm (ϵ 16,770; pH 3.5 phosphate buffer) 265nm (ϵ 13,900); (in pH 7.6) at 297nm (ϵ 9,500) and 260nm sh (ϵ 4,690). It has been separated from the 6S-isomer by HPLC on a Partisil-10SCX column using 30mM ammonium phosphate buffer (pH 3.0) containing 3mM $NaHSO_3$ (2ml/minute flow rate; 275nm detector) with retention times of 5.87 minutes (6R) and 8.45 minutes (6S). It is stable in acidic solutions and can be stored for extended periods at -20° in 0.04M HCl. Above pH 7 the neutral species are obtained, and these are readily oxidised by the oxygen in the solvent to the *quinonoid species*, and then further oxidation and degradation occurs at room temperature. These changes are slower at 0° . The *sulfate salt* can be obtained by recrystallisation from 2M H_2SO_4 and is less soluble in water than the hydrochloride salt. The **6R-2,5,1',2'-tetraacetylbiopterin derivative** has **m 292^o(dec)** after recrystallisation from MeOH (100 parts) and $[\alpha]_{589}^{20} -144$ (c 0.5, $CHCl_3$), $[\alpha]_{589}^{20} +12.8$ (c 0.39, Me_2SO). [For NMR and UV see Matsuura et al. *Heterocycles* **23** 3115 1985, DOI: 10.3987/R-1985-12-3115; Furrer et al. *Helv Chim Acta* **62** 2577 1979, DOI: 10.1002/hlca.19790620811; Armarego et al. *Aust J Chem* **37** 355 1984, DOI: 10.1071/CH9840355; *Beilstein* **26** III/IV 4032.] It is a natural cofactor for several oxygenases.

Tetrahydrofolic acid dihydrochloride $2H_2O$ (THFA, 6S- or 6RS- 5,6,7,8-tetrahydrofolic acid $2HCl$ $2H_2O$, 5,6,7,8-tetrahydropteroyl-L-glutamic acid $2HCl$ $2H_2O$) [135-16-0] $C_{19}H_{23}N_7O_6 \cdot 2HCl \cdot 2H_2O$, **M 544.4, m >200^o(dec)**, $[\alpha]_D^{27} +16.9$ (H_2O pH 7.0 + 2-mercaptoethanol), **pK₁ 1.7** (pyrimidine N^+), **pK₂ 2.4** ($10N^+$), **pK₃ 3.5** (α - CO_2H), **pK₄ 4.9** (γ - CO_2H), **pK₅ 5.6** ($5-NH^+$), **pK₆ 10.4** (acidic, $3NH$). Very high quality material is now available commercially, and should be a white powder. Dry it over P_2O_5 *in vacuo* and store in weighed aliquots in sealed ampoules. It is stable at room temperature in this way for many months and much longer at -10° . When moist, it is extremely sensitive to air whereby it oxidises to the yellow 7,8-dihydro derivative. In solution it turns yellow in colour as it oxidises, and then particularly in the presence of acids it turns dark reddish brown in colour. Aqueous solutions should be frozen immediately when not in use. It is advisable to add 2-mercaptoethanol (if it does not interfere with further procedures) which depletes the solution of O_2 . The *sulfate salt* is more stable but is much less soluble. The best way to prepare standard solutions of this acid is to dissolve it in the desired buffer and estimate the concentration by UV absorption in pH 7 buffer at 297nm (ϵ 22,000 $M^{-1}cm^{-1}$). If a sample is suspect, it is not advisable to purify it because it is likely to deteriorate further as 'dry box' conditions are necessary. Either a new sample is purchased or one is freshly prepared from folic acid. [Hafeti et al. *Biochemical Preparations* **7** 89 1960, UV: Mathews & Huennekens *J Biol Chem* **235** 3304 1960, PMID: 13768089; Osborn & Huennekens *J Biol Chem* **233** 969 1958, PMID: 13587525; O'Dell et al. *J Am Chem Soc* **69** 250 1947, DOI: 10.1021/ja01194a023; Blakley *Biochem J* **65** 331 1957, DOI: 10.1042/bj0650331; *Beilstein* **26** III/IV 3879.] A natural cofactor and vitamin.

5,6,7,8-Tetrahydropterin sulfate (2-amino-5,6,7,8-tetrahydropteridin-4-one H_2SO_4) [20350-44-1] $C_6H_9N_5O \cdot H_2SO_4$, **M 265.2, m >200^o(dec)**, **pK₁²⁵ 1.3** (pyrimidine⁺), **pK₂²⁵ 5.6** ($5-NH^+$), **pK₃²⁵ 10.6** (acidic, $3NH$). If its colour is strongly violet, then it needs to be reduced again. It is best to check the UV in N HCl where the peak at ~265nm drops sharply to zero at ~340nm. UV absorption at 340nm indicates oxidation to quinonoid or 7,8-dihydropterin. If absorption is weak, then dissolve it in the minimum volume of dry trifluoroacetic acid (fume hood), add charcoal, filter, add two drops of N H_2SO_4 then dry Et_2O at 0° , allow the white tetrahydro salt to settle, collect it, and wash it with dry Et_2O , by centrifugation. Dry the residue *in vacuo* over P_2O_5 and KOH. Store in aliquots in the dark at $<0^{\circ}$. For the UV spectra of the monocation, dication, neutral species and anion see references. [Blakley *Biochem J* **72** 707 1959, DOI: 10.1042/bj0720707; Pfeleiderer in *Pterins and Folate* (Benkovic and Blakley Eds) J Wiley **Vol 2** p 97 1985, Blakley *The Biochemistry of Folic Acid and Related Pteridines*, North-Holland Publ Co., Amsterdam, pp 157-163 1969, ISSN 0071-965X.]

Theophylline (1,3-dimethyl-2,6-dioxopurine, Theocin) [58-55-9] $C_7H_8N_4O_2$, **M 180.2, m 272-274^o, pK₁⁴⁰ -0.24, pK₂⁴⁰ 8.79, pK₃ 11.5** (acidic). It crystallises from H_2O as the *monohydrate* which dehydrates $>100^{\circ}$. It complexes with heavy metals, is a diuretic, vasodilator and cardiac stimulant. [Lister *Purines Part II. Fused Pyrimidines* Brown Ed. p 253 1971, J.Wiley & Sons NY, ISBN 047138205-1; *Beilstein* **26** III/IV 2331.]

Thiamine monophosphate chloride $2H_2O$ (Aneurine monophosphate chloride) [532-40-1, 273724-21-3] $C_{12}H_{18}ClN_4O_4PS \cdot 2H_2O$, **M 416.8, m 193^o(dec), 200^o(dec), 200-203^o(dec)**, **pK₁ 2.40, pK₂ 4.80, pK₃ 6.27, pK₄ 9.65, pK₅ 10.20**. Purify the salt by recrystallisation from aqueous HCl, EtOH slightly acidified with HCl,

EtOH/Me₂CO, H₂O, or H₂O/EtOH/Et₂O. Dissolve it in a small volume of H₂O and mix it with EtOH/Me₂CO (1:1) to give the HCl.H₂O as crystals. Filter it off, wash it with Et₂O and dry it in a vacuum. The **chloride hydrochloride**, **m 215-217°(dec)** is obtained when it is crystallised from aqueous HCl. [Wenz et al. *Justus Liebigs Ann Chem* **618** 210 1958, DOI: 10.1002/jlac.19586180124; Viscontini et al. *Helv Chim Acta* **34** 1388 1951, DOI: 10.1002/hlca.19510340521; Leichszenring & Schmidt *Chem Ber* **95** 767 1962, DOI: 10.1002/cber.19620950330; Matsukawa et al. *Methods Enzymol* **18A** 141, 147 1970, DOI: 10.1016/0076-6879(71)18293-6; *Beilstein* **27** III/IV 1766.] It is a vitamin.

Thiamphenicol (Thycocymetin, 1R,2R-2-[2,2-dichloroacetyl-amino]-1-[4-methanesulfonyl-phenyl]-propan-1,3-diol) [15318-45-3 (D-threo), 90-91-5] C₁₂H₁₅Cl₂NO₅S, **M 356.2, m 163-166°, 165.2-165.6°, 165-166°, [α]_D²⁵ +15.6 (c 2, EtOH), pK_a²⁵ 7.2.** Recrystallise thiamphenicol from H₂O or CHCl₃. The UV has λ_{max} at 224, 266 and 274nm (ε 13,700, 800 and 700) in 95% EtOH. The **1S,2S-isomer** [14786-51-7] has **m 164.3-166.3°** (from H₂O/EtOAc/petroleum ether) and [α]_D²⁵ -12.6 (c 1, EtOH); and the **racemate 1RS,2RS-Racefenical** [847-25-6] has **m 181-183° (dec)** from CHCl₃/EtOAc/petroleum ether. [Cutler et al. *J Am Chem Soc* **74** 5475 1952, DOI: 10.1021/ja01141a074; For UV see Nachod & Cutler *J Am Chem Soc* **74** 6291 1952, DOI: 10.1021/ja01144a517; Suter et al. *J Am Chem Soc* **75** 4330 1953, DOI: 10.1021/ja01113a057; Cutler et al. *J Am Pharm Assoc* **43** 697 1954, DOI: 10.1002/jps.3030431121; *Beilstein* **13** IV 2957.] It is an antibiotic and has biological half life of 0.5 hours. For the 'Role of thiamphenicol in the treatment of community-acquired lung infections' see Raymond et al. [*Med Trop (Mars)* **64** 33 2004, PMID: 15224555].

ε-[2-(4-Thiazolidinone)]hexanoic acid (Mycobacidin, Acidomycin, 6[4-oxothiazolidin-2-yl]hexanoic acid) [539-35-5] C₉H₁₅NO₃S, **M 215.3, m 140°, pK_a²⁵ 5.1.** The **dl-form is dimorphic**; it crystallises from CHCl₃ with **m 116-117°**, and from H₂O with **m 123°**. The **l(-)-enantiomer** (from Actinomyces) crystallises from H₂O, MeOH (**m 139-140°**), aqueous EtOH (**m 140-141°**) or EtOAc, and has [α]_D²⁰ -54 (c 1, MeOH). The **l(-)-methyl ester** C₁₀H₁₇NO₃S, **M 231.3**, has **m 53-54°** (needles from Et₂O/hexane), [α]_D²⁰ -50.9 (MeOH). The **d(+)-enantiomer** (from optical resolution of the brucine salt) has **m 138-139°** (from H₂O) and [α]_D²⁵ +57 (c 1, MeOH). The optically active acids **racemise** in hot alkali. [McLamore et al. *J Am Chem Soc* **75** 105 1953, DOI: 10.1021/ja01097a030; isolation: Tejera et al. *Antibiot Chemother* **2** 333 1952; *Beilstein* **27** III/IV 4281.]

6-Thioguanosine (2-amino-6-mercapto-9-β-D-ribofuranosylpurine) [85-31-4, 345909-25-3] C₁₀H₁₃N₅O₄S, **M 299.3, m 224-227°(dec), 230-231°(dec), [α]_D²⁰ -64 (c 1.3, 0.1N NaOH), pK_a²⁵ 8.33.** Thioguanosine is crystallised (as **hemihydrate**) from hot H₂O (charcoal) and cooled slowly to give tapered prisms. It also crystallises by dissolving in dilute NH₃ and acidifying with acetic acid, and then recrystallising from H₂O. Its UV (pH 4-6) has λ_{max} at 257nm (ε 8,820) and 342nm (ε 24,800), and at pH 10.4-12.0 it has λ_{max} at 252nm (ε 14,700) and 319.5nm (ε 21,000). [Fox et al. *J Am Chem Soc* **80** 1669 1958, DOI: 10.1021/ja01540a041; *Beilstein* **26** III/IV 3927.]

dl-α-Tocopherol (see **vitamin E**) [10191-41-0] C₂₉H₅₀O₂, **M 430.7, A_{1cm}^{1%} 74.2 at 292 nm in MeOH.** Dissolve dl-α-tocopherol in anhydrous MeOH (15ml/g) cool to -6° for 1 hour, then chill in a Dry-ice/acetone bath; crystallisation is induced by scratching with a glass rod. The **dl-α-acetate** [52225-20-4, 7695-91-2] (see **DL-vitamin E acetate** below) is a viscous yellow liquid with **m -7°, b 184°/0.01mm, 224°/0.3mm, d₄²⁰ 0.953, n_D²⁰ 1.496.** It is used as a standard for Vitamin E activity where the unit of activity is attained with 1mg of pure dl-α-acetate. [Friedrich 'Vitamins' Water de Guyter Publ, Berlin 1988, *Beilstein* **17/4** V 168.]

γ-Tocopherol (3,4-dihydro-2,7,8-trimethyl-2-(4,8,12-trimethyltridecyl)-2H-benzopyran-6-ol) [54-28-4] C₂₈H₄₈O₂, **M 416.7, m -30°, b 200-210°/0.1mm, d₄²⁰ 0.951, n_D²⁰ 1.505, [α]_D²⁰ -2.4 (EtOH).** γ-Tocopherol is purified by distillation at high vacuum and stored in dark ampoules under N₂. Its UV has λ_{max} at 298nm (A_{1cm}^{1%} 92.8). It is insoluble in H₂O but soluble in organic solvents. The **allophanate** (used for separating it from its isomers) has **m 136-138°, [α]_D¹⁸ +3.4 (CHCl₃).** [Baxter et al. *J Am Chem Soc* **65** 918 1943, DOI: 10.1021/ja01245a041; Emerson et al. *Science* **83** 421 1936, DOI: 10.1126/science.83.2157.421; Evans et al. *J Biol Chem* **113** 319 1936, <http://www.jbc.org/content/113/1/319>; *Beilstein* **17/4** V 158.] It is a food additive.

Tomatidine (5α,20β,22α,25β,27-azaspirostan-3β-ol) [77-59-8] C₂₇H₄₅NO₂, **M 415.7, m 202-206°, [α]_D²⁰ +5.9 (c 1, MeOH), [α]_D²⁰ +8 (CHCl₃).** Tomatidine forms plates from EtOAc. It is also purified by dissolving

80mg in $^*\text{C}_6\text{H}_6$ and applying to an Al_2O_3 column (3.0g) and eluting with $^*\text{C}_6\text{H}_6$, evaporating and recrystallising the residue three times from EtOAc. The **hydrochloride** [6192-62-7] $\text{C}_{27}\text{H}_{45}\text{NO}_2 \cdot \text{HCl}$, M 452.1, has **m 265-270° (281-284° dec)** from EtOH and $[\alpha]_{\text{D}}^{25} -5$ (MeOH). Store it at -20° . [For IR see Uhle *J Am Chem Soc* **83** 1460 1961, DOI: 10.1021/ja01467a045; Kessar et al. *Tetrahedron* **27** 2869 1971, DOI: 10.1016/S0040-4020(01)98078-0; Schreiber & Adams *Experientia* **17** 13 1961, PMID: 13748592; Beilstein **27** III/IV 1950.] Dyle et al. proposed that it is can inhibit muscle atrophy [*J Biol Chem* **289** 14913 2014, DOI: 10.1074/jbc.M114.556241, PMID: 24719321; for ‘Hedgehog signalling within airway epithelial progenitors and in small-cell lung cancer’ see Watkins, et al. *Nature* **422** 313 2003, DOI: 10.1038/nature01493; PMID: 12629553.

Tomatine (22S,25S-3 β , β -lycotetraosyloxy-5 α -spirosolan) [17406-45-0] $\text{C}_{50}\text{H}_{83}\text{NO}_{21}$, M 1034.2, **m 263-268°(dec), 290-291°(evacuated capillary), 283.5-287°(dec), 272-277°(dec), 300-305°(dec), $[\alpha]_{\text{D}}^{20} -18$ to -34 (c 0.55, pyridine).** Tomatine is recrystallised from MeOH, EtOH, aqueous EtOH or dioxane/ NH_3 . It is almost insoluble in petroleum ether, Et_2O or H_2O . [Reichstein *Angew Chem* **74** 887 1962, DOI: 10.1002/ange.19620742202; Beilstein **27** III/IV 1954.] It is an antimicrobial.

Tubercidin (7-deazaadenosine) [69-33-0] $\text{C}_{11}\text{H}_{14}\text{N}_4\text{O}_4$, M 266.3, **m 247-248°, $[\alpha]_{\text{D}}^{17} -67$ (50% aqueous AcOH), $\text{pK}^{10} 5.2-5.3$.** 7-Deazaadenosine forms needles from hot H_2O . It is soluble in H_2O (0.33%), MeOH (0.5%) and EtOH (0.05%). It has UV with λ_{max} at 270nm (ϵ 12,100) in 0.001N NaOH. The **picrate** has **m 229-231°(dec)**. [Tolman et al. *J Am Chem Soc* **91** 2102 1969, DOI: 10.1021/ja01036a040; Mizuno et al. *J Org Chem* **28** 3329 1963, DOI: 10.1021/jo01047a012; Beilstein **26** IV 1117.] Antifungal that blocks purine biosynthesis.

Tunicamycin [11089-65-9] $\text{C}_{39}\text{H}_{64}\text{N}_4\text{O}_{16}$, M ~780, **m 234-235°(dec), $[\alpha]_{\text{D}}^{20} +52$ (c 0.5, pyridine), $\text{pK}_{\text{Est}} \sim 9.4$.** The components of this homologous nucleoside antibiotic from *Streptomyces* sp. are purified by recrystallising 3 times from hot glass-distilled MeOH, and the white crystals are dissolved in 25% aqueous MeOH and separated on a Partisil ODS-10 μ column (9.4 x 25 cm) [Magnum-9 Whatman] using a 260nm detector. The column is eluted with a MeOH/ H_2O mixture adjusted to 1:4 (v/v) then to 2:4 (v/v). The individual components are recovered and lyophilised. Ten components have been isolated, and all were active (to varying extents) depending on the lengths of the aliphatic side-chains. The mixture has UV with λ_{max} at 205 and 260nm ($A_{1\text{cm}}^{1\%}$ 230 and 110). It is stable in H_2O at neutral pH but unstable in acidic solution. It inhibits protein glycosylation. [Mahoney & Duksin *J Biol Chem* **254** 6572 1979, PMID: 447736; Elbein *Trends Biochem Sci* **6** 219 1981, DOI: 10.1016/0968-0004(81)90080-3; for **teichoic acid synthesis inhibition** see Hashizume et al. *J Antibiot* **68** 373 2015, DOI:10.1038/ja.2014.169.]

Uracil, uridine and uridine nucleotides. These are resolved by ion-exchange chromatography with AG1 (Cl⁻ form). [Lindsay et al. *Anal Biochem* **24** 506 1968, DOI: 10.1016/0003-2697(68)90158-9.]

Uridine 5'-(1-thio) monophosphate [15548-52-4, 18875-72-4 (Absolute Stereochemistry specified)] and **Uridine 5'-(α -thio) diphosphate** [*RS*(α -P) 27988-67-6; *R*(α -P) 72120-52-6], **$\text{pK}_{\text{Est}(1)} \sim 6.4$, $\text{pK}_{\text{Est}(2)} \sim 9.5$** The Et_3N salts are purified by dissolving ~4g in 500ml of H_2O (adding a drop or two of Et_3N if they do not dissolve) and chromatograph by applying to a column (3 x 30cm) of DEAE-Sephadex A-25 and eluting with 1.4L of a linear gradient of $\text{Et}_3\text{NH} \cdot \text{HCO}_3$ from 0.05 to 0.55M, pH 7.8 and 4° . The product elutes between 0.2-0.3M $\text{Et}_3\text{NH} \cdot \text{HCO}_3$. The pooled fractions are evaporated, and the residue is twice taken up in EtOH and evaporated to dryness to remove the last traces of $\text{Et}_3\text{NH} \cdot \text{HCO}_3$. ^{31}P NMR: P_α is a doublet at -40.81 and -40.33, and P_β at 7.02ppm, $J_{\alpha,\beta}$ 32.96Hz. [Sheu et al. *Biochemistry* **18** 5548 1979, DOI: 10.1021/bi00592a004.]

Uridylic acid (di-Na salt) [27821-45-0, 3387-36-8, 58-97-9 free acid] $\text{C}_9\text{H}_{11}\text{N}_2\text{O}_9\text{P} \cdot 2\text{Na}$ M 368.2, **m 198.5°, 202°(dec), $\text{pK}_1^{25} 1.0$, $\text{pK}_2^{25} 6.63$, $\text{pK}_3^{25} 9.71$.** Crystallise it from MeOH. It may contain some Ba salt(s); hence stir it with Amberlite IR-120 cation exchanger (25ml, wet resin, 15-50 mesh in H^+ form) in H_2O (50ml) until the nucleotide dissolves. Filter, wash resin with H_2O until the eluate is neutral. Combine the filtrate and washings, and adjust the pH to 8.0 with 2.0 M aqueous NaOH. Concentrate it *in vacuo* to ~ 20ml, and add Me_2CO dropwise until crystallisation begins. Cool to 0° , and the shiny plates of di **Na uridine-5'-phosphate dihydrate** are filtered off and dried over P_2O_5 at $25^\circ/0.1\text{mm}$ for 24 hours (>76% recovery). Its solubility in H_2O at 20° is

40g/100ml. The UV has λ_{\max} at 262nm (ϵ 10,000 M⁻¹cm⁻¹) in 0.1 M HCl. [Brown et al. *J Chem Soc* 408 1950, DOI: 10.1039/JR9500000403; Smith *Biochemical Preparations* 8 130 1960.] [Beilstein 24 IV 1214.]

(+)-Usnic acid (2,6-diacetyl-7,9-dihydroxy-8,9b-dimethyldibenzofuran-1,3(2H,9bH)-dione) [7562-61-0, 125-46-2] C₁₈H₁₆O₇, M 344.3, m 201-204°, 203-206°, [α]_D²⁰ +630 (c 0.7, CHCl₃), [α]_D²⁰ -488 (c 0.7, CHCl₃), pK₁ 4.4, pK₂ 8.8, pK₃ 10.7. This very weak acid is the natural form which is recrystallised from Me₂CO, MeOH or *C₆H₆. At 25° it is soluble in H₂O (<0.01%), Me₂CO (0.77%), EtOAc (0.88%), MeOCH₂CH₂OH (0.22%), and furfural (7.32%). [Curd & Robertson *J Chem Soc* 894 1937, DOI: 10.1039/JR9370000894; Barton & Bruun *J Chem Soc* 603 1953, DOI: 10.1039/JR9530000603; resolution: Dean et al. *J Chem Soc* 1250 1953, DOI: 10.1039/JR9530001250; for synthesis see Barton et al. *J Chem Soc* 530 1956, DOI: 10.1039/JR9560000530; Beilstein 18/5 V 586.] It is a phytotoxic lichen metabolite. Optical and racemic isomers occur in lichens, and used in medicine, perfumery and cosmetics. The sodium salt (Lipokinetix) is claimed to induce weight loss and increase metabolic rate. For 'A review on usnic acid, an interesting natural compound' see Cocchietto et al. [*Naturwissenschaften* 89(4) 137 2002, DOI: 10.1007/s00114-002-0305-3, ISSN 0028-1042].

(-)-Usnic acid (2,6-diacetyl-7,9-dihydroxy-8,9b-dimethyldibenzofuran-1,3(2H,9bH)-dione) [6159-66-6, 7562-61-0] C₁₈H₁₆O₇, M 344.3, m 201-204°, 204°, [α]_D²⁰ -495 (c 0.9, CHCl₃). Its properties are similar to those of the acid in the preceding entry. [Beilstein 18 II 241, 18 III/IV 3522.] **Racemic (±)-usnic acid** [125-46-2] has m 200°, 201-203°, 204°.

Valinomycin (Potassium ionophore I) [2001-95-8] C₅₄H₉₀N₆O₁₈, M 1111.3, m 172°, 186-187°, 190°, [α]_D²⁰ +31.0 (c 1.6, *C₆H₆). Recrystallise the cyclic dodecadepsipeptide valinomycin from dibutyl ether or Et₂O. It is **dimorphic**: modification A crystallises from *n*-octane, and modification B crystallises from EtOH/H₂O. It is soluble in petroleum ether, CHCl₃, AcOH, BuOAc, Me₂CO and Me₂SO (~10mg/ml) but is insoluble in H₂O. [Smith et al. *J Am Chem Soc* 97 7242 1975, DOI: 10.1021/ja00858a008; for UV, IR and NMR see Brockmann & Schmidt-Kastner *Chem Ber* 88 57 1955, DOI: 10.1002/cber.19550880111; Beilstein 27 I 9728. 17 IV 9728.] For 'Interaction of valinomycin and monovalent cations, e.g. Li⁺, Na⁺, K⁺, Cs⁺, Rb⁺, with the (Ca²⁺, Mg²⁺)-ATPase of skeletal muscle sarcoplasmic reticulum' see Davidson & Berman *J Biol Chem* 260 7325 1985, PMID: 3158656; and for 'The effects of the ionophore valinomycin on biomimetic solid supported lipid DPPTE/EPC membranes' see Rose & Jenkins [*Bioelectrochem* 70 387 2007, DOI: 10.1016/j.bioelechem.2006.05.009].

(±)-Verapamil hydrochloride (5-[N-{3,4-dimethoxyphenylethyl}methylamino]-2-[3,4-dimethoxyphenyl]-2-isopropylvaleronitrile HCl) [152-11-4, 23313-68-0] C₂₇H₃₈N₂O₄. HCl, M 491.1, m 138.5-140.5°, 142°(dec), pK_{Est} ~ 10.6. The salt is purified by dissolving it in EtOH, filtering (if insoluble particles are present), and adding Et₂O, filtering the salt, washing it with Et₂O and drying it *in vacuo*. It has the following solubilities: hexane (0.001%), CH₂Cl₂ (~10%), MeOH (~10%), EtOH (20%) and H₂O (8.3%). It has UV with λ_{\max} at 232 and 278nm. The **free base** [52-53-9] C₂₇H₃₈N₂O₄, M 454.6, is a viscous yellow oil **b** 243-246°/0.01mm (n_D²⁵ 1.5448) and is almost insoluble in H₂O but soluble in organic solvents. Its biological half life is 2.8-7.4 hours. It is a Ca channel antagonist and is a coronary vasodilator; and is used for treating hypertension in cases of atrial fibrillation or other types of arrhythmia. Also used for controlling high blood pressure and angina. [Ramuz *Helv Chim Acta* 58 2050 1975, DOI: 10.1002/hlca.19750580720; Harvey et al. *Biochem J* 257 95 1989, DOI: 10.1042/bj2570095.]

Veratridine (3-veratroylveracevine) [71-62-5] C₃₆H₅₁NO₁₁, M 673.8, m 160-180°, ~180° (after drying at 130°, pK 9.54 (quinolizidine N), [α]_D²⁰ +8.0 (c ~5, EtOH). It is an alkaloid neurotoxin which prevents inactivation of Na⁺ channels. Its solubility in EtOH is ~5%; and it separates as a pale yellow powder from an ethanolic solution on addition of Et₂O. It forms nitrate, sulfate and perchlorate salts. [McKinney et al. *Anal Biochem* 153 33 1986, DOI: 10.1016/0003-2697(86)90056-4; Beilstein 21 V/13 709.] It binds to activated Na⁺ leading to nerve excitability.

Vinblastine sulfate (VLB, vincaleucoblastine sulfate) [143-67-9] C₄₆H₅₈N₄O₉. H₂SO₄, M 909.5, m 284-285°(dec), 267°(dec), [α]_D²⁶ -28 (MeOH), pK₁ 5.5, pK₂ 7.4. Crystallise the sulfate from MeOH or EtOH and

dry it *in vacuo* over conc H_2SO_4 . The **free base** crystallises from EtOH or MeOH **m 211-216°** (with 2MeOH and 1 H_2O) and forms a stable **etherate** from Et_2O with **m 201-211°**, and $[\alpha]_{\text{D}}^{25} +42$ (CHCl_3); and its UV has λ_{max} at 214 and 259nm (log ϵ 4.73 and 4.21). The **dihydrochloride** has **m 244-246° dec** (MeOH). Store it at -20°. It is a **monoamine oxidase B inhibitor** and **induces microtubule aggregation**, and is an **antineoplastic** drug for Hodgkin's lymphoma. [Neuss et al. *J Am Chem Soc* **81** 4754 1959, DOI: 10.1021/ja01526a086; Son et al. *J Med Chem* **33** 1845 1990, DOI: 10.1021/jm00169a002; Warfield & Bouck *Science* **186** 1219 1974, DOI: 10.1126/science.186.4170.1219; *Beilstein* **26** III/IV 3167.]

Vincristine sulfate (22-oxovincalocoblastine sulfate) [2068-78-2] $\text{C}_{46}\text{H}_{56}\text{N}_4\text{O}_{10} \cdot \text{H}_2\text{SO}_4$, **M 925.1, m 218-220°**, $[\alpha]_{\text{D}}^{25} +26.2$ (CH_2Cl_2), **pK₁ 5.0, pK₂ 7.4** (in 33% $\text{Me}_2\text{NCHO}/\text{H}_2\text{O}$). The salt is recrystallised from MeOH. It has UV with λ_{max} at 220, 255 and 296nm (log ϵ 4.65, 4.21 and 4.18). Store it at -20°. It is a **monoamine oxidase inhibitor** and is used in cancer research [Son et al. *J Med Chem* **33** 1845 1990, DOI: 10.1021/jm00169a002; Horio et al. *Proc Natl Acad Sci USA* **85** 3580 1988, PMID: 3368466]. For 'Comparison of the effects of vinblastine, vincristine, vindesine, and vinepidine on microtubule dynamics and cell proliferation *in vitro*' see Jordan et al. [*Cancer Res* **45** 2741 1985, PMID: 3986806].

Viomycin sulfate (Viocin, Tuberactinomycin B) [37883-00-4] $\text{C}_{23}\text{H}_{46}\text{N}_6\text{O}_{13} \cdot 2.5\text{H}_2\text{SO}_4$, **M 614.7, 685.7, m 266°(dec)**, $[\alpha]_{\text{D}}^{17} -29.5$ (c 1, H_2O), **pK₁ 7.2 (8.2), pK₂ 10.3**. The aminoglycoside antibiotic viocin crystallises from $\text{H}_2\text{O}/\text{EtOH}$ and is dried in a vacuum. The dry material is **hygroscopic** and should be stored dry. The UV has λ_{max} at 268 and 285nm (log ϵ 4.4 and 4.2) in H_2O . [Kitigawa et al. *Chem Pharm Bull Jpn* **20** 2176 1972, DOI: org/10.1248/cpb.20.2176.] The **hydrochloride** forms **hygroscopic** plates with **m 270° dec**, and $[\alpha]_{\text{D}}^{18} -16.6$ (c 1, H_2O), with λ_{max} at 268nm (log ϵ 4.5) in H_2O ; 268nm (log ϵ 4.4) in 0.1N HCl and at 285nm (log ϵ 4.3) in 0.1N NaOH. [*Beilstein* **26** III/IV 4245.] Antibiotic used in topical medications like ointments, creams and eyedrops.

Vitamin A acetate (retinyl acetate) [127-47-9] $\text{C}_{22}\text{H}_{32}\text{O}_2$, **M 328.5, m 57°, 57-58°**. The acetate is separated from retinol by column chromatography, then crystallised from MeOH. [Kofler and Rubin *Vitamins and Hormones* (NY) **18** 315 1960, DOI: 10.1016/S0083-6729(08)60867-5; for purification methods]. Store it in the dark, under N_2 or argon, at 0°. [*Beilstein* **6** IV 4135.]

Vitamin A acid [Retinoic acid, 3,7-dimethyl-9-(2,6,6-trimethyl-1-cyclohexenyl)-2,4,6,8-nonatetraen-1-oic acid, (2E,4E,6E,8E)-3,7-dimethyl-9-(2,6,6-trimethylcyclohexen-1-yl)nona-2,4,6,8-tetraenoic acid] [302-79-4] $\text{C}_{20}\text{H}_{28}\text{O}_2$, **M 300.4, m 180-181°, 180-182°, pK_{Est} ~ 4.2**. Purify the acid by chromatography on silicic acid columns, and eluting it with a small amount of EtOH in hexane. Also dissolve it in Et_2O , wash it with H_2O , dry (Na_2SO_4), evaporate and the solid residue is recrystallised from MeOH (0.53g /3.5ml MeOH to give 0.14g) or EtOH. It also recrystallises from *i*-PrOH, or as the **methyl ester** from MeOH. Its UV(MeOH) has λ_{max} at 351nm (ϵ 45,000). **9-Cis-acid** forms yellow needles from EtOH, with **m 189-190°**, and its UV(MeOH) has λ_{max} at 343nm (ϵ 36,500); the **13-cis-acid** forms red-orange plates from *i*-PrOH with **m 174-175°**, and its UV has λ_{max} at 345nm (ϵ 39,800). Store it in the dark, in an inert atmosphere, at 0° [Robeson et al. *J Am Chem Soc* **77** 4111 1955, DOI: 10.1021/ja01620a043]. [*Beilstein* **9** IV 2387.]

Vitamin A alcohol (retinol) [68-26-8] $\text{C}_{20}\text{H}_{30}\text{O}$, **M 286.5, m 62-64°, A_{1cm}^{1%} (max) (all-trans) 1, 832 (325 nm), (13-cis) 1686 (328nm), (11-cis) 1230 319 nm), (9-cis) 1480 (323 nm), (9,13-di-cis) 1379 (324 nm), (11,13-di-cis) 908 (311 nm) in EtOH**. Purify retinol by chromatography on columns of water-deactivated alumina and elute with 3-5% acetone in hexane. Separate the isomers by TLC plates on silica gel G, developed with petroleum ether (low boiling)/methyl heptanone (11:2). Store it in the dark, under N_2 , at 0°, or in Et_2O , Me_2CO or EtOAc. [See Ganguly et al. *Arch Biochem Biophys* **38** 275 1952, DOI: 10.1016/0003-9861(52)90032-5; *Beilstein* **6** IV 4133.]

Vitamin A aldehyde [all-trans-retinal; 3,7-dimethyl-9-(2,6,6-trimethyl-1-cyclohexenyl)-2,4,6,8-nonatetraen-1-al] [116-31-4] $\text{C}_{20}\text{H}_{28}\text{O}$, **M 284.4, m 61-64°**. The aldehyde is separated from retinol by column chromatography on water-deactivated alumina. Elute with 1-2% acetone in hexane, or on TLC plates of silica gel G and using the same eluting solvent. It crystallises from petroleum ether or *n*-hexane as yellow-orange crystals, and the UV in hexane has λ_{max} at 373nm ($A_{1\text{cm}}^{1\%}$ 1,548) and 368nm (ϵ 48,000). It is an **irritant** and is light sensitive. Store it in sealed ampoules under N_2 . The **semicarbazone** forms yellow crystals from

$\text{CHCl}_3/\text{Et}_2\text{O}$ or EtOH , **m 199-201° dec.** The **9-cis-isomer** [514-85-2] and the **13-cis-isomer** [472-86-6] [λ_{max} at 375nm (ϵ 1,250) in EtOH] are also available commercially. [Beilstein 7 III 1742.]

Vitamin A palmitate (retinyl palmitate) [79-81-2] $\text{C}_{36}\text{H}_{60}\text{O}_2$, **M 524.9**, **m 28-29°**, $\epsilon_{1\text{cm}}^{1\%}$ (all-trans) **1000 (325 nm) in EtOH**. The palmitate is separated from retinol by column chromatography on water-deactivated alumina with hexane containing a very small percentage of acetone. It is also chromatographed on TLC silica gel G, and developed with pet. ether/isopropyl ether/acetic acid/water (180:20:2:5) or petroleum ether/acetonitrile/acetic acid/water (190:10:1:15). Then recrystallize it from propylene at below -47°. [Beilstein 6 IV 4135.]

Vitamin B₁ Hydrochloride [Aneurine hydrochloride, Thiamine hydrochloride, 3{(4-amino-2-methyl-5-pyrimidinyl)methyl}-4-methylthiazolium chloride monohydrochloride] [67-03-8, 59-43-8 chloride] $\text{C}_{12}\text{H}_{17}\text{ClN}_4\text{OS} \cdot \text{HCl}$, **M 337.3**, **m 248°(dec)**, **249-250°**, monohydrate **m 135°(dec)**, **pK₁²⁵ 4.8**, **pK₂²⁵ 9.2**. The hydrochloride crystallises from 95% EtOH (solubility is ca 1%). The *monohydrate* is dehydrated at 100° *in vacuo* over H_2SO_4 , but is *hygroscopic* and picking up one H_2O readily. It can be sterilised at 100° if the pH is below 5.5. The *nitrate* has **m 196-200° (dec)** and is more stable than the hydrochloride. The *picrolonate* crystallises from H_2O and is *dimorphic*, **m 164-165°** and **228-229°(dec)**. [Todd & Bergel *J Chem Soc* 364, 367 1937, DOI: 10.1039/JR9370000364; Williams et al. *J Am Chem Soc* 58 1063 1936, DOI: 10.1021/ja01297a515; 1504 1936, DOI: 10.1021/ja01299a505; 59 526 1937, DOI: 10.1021/ja01282a028; Beilstein 27 IV 1766.]

Vitamin B₂ [Riboflavin, Lactoflavin, 6,7-dimethyl-9-(D-1'-ribityl)isoalloxazine] [83-88-5] $\text{C}_{17}\text{H}_{20}\text{N}_4\text{O}_6$, **M 376.4**, **m 278-282°(dec with darkening at 240°)**, **281-282°**, $[\alpha]_{\text{D}}^{25}$ -9.8 (H_2O), $[\alpha]_{\text{D}}^{25}$ -112 to -125 (c 2.5, 0.02N NaOH), $[\alpha]_{\text{D}}^{20}$ -59 (c 0.23, AcOH), **pK₁ 1.7**, **pK₂ 9.69 (10.2, acidic NH)**. It crystallises from H_2O as a yellow-orange powder in three different forms with differing amounts of H_2O . It melts if placed in an oil bath at 250°, but decomposes at 280° if heated at a rate of 5°/minute. It is also purified by crystallisation from 2M acetic acid, then extracted with CHCl_3 to remove lumichrome impurity. [Smith & Metzler *J Am Chem Soc* 85 3285 1963, DOI: 10.1021/ja00903a051.] Its solubility in H_2O is 1g in 3-15L depending on the crystal structure. Its solubility in EtOH at 25° is 4.5mg in 100ml. Store it in the dark because it is decomposed to lumichrome by UV light. [Pearson *The Vitamins* vol V pp1-96 1967, vol VII pp 1-96 1972, Academic Press, Beilstein 26 IV 2542.]

Vitamin B₃ (Nicotinamide, Niacin, vitamin PP) [98-92-0] $\text{C}_6\text{H}_6\text{N}_2\text{O}$, **M 122.1**, **m 128-131°**, **b ~150-160°/high vac**, **pK₁²⁵ 0.5**, **pK₂²⁵ 3.33**. Crystallise niacin from *benzene. Its solubility in g/ml is H_2O (1), EtOH (0.7) and glycerol (0.1). [Methods in Enzymology 66 23 1980, UV: Armarego *Physical Methods in Heterocyclic Chemistry* (Ed Katritzky, Academic Press) Vol III 83 1971, Beilstein 22 III/IV 389, 22/2 V 80.]

Vitamin B₆ hydrochloride (adermine, pyridoxine HCl, 3-hydroxy-4,5-bis[hydroxymethyl]-2-methylpyridine HCl) [58-56-0] $\text{C}_8\text{H}_{11}\text{NO}_3 \cdot \text{HCl}$, **M 205.6**, **m 208-208.5°**, **208-209°(dec)**, **209-210°(dec)**, **205-212° (sublimes)**, **214-215°**, **pK₁²⁵ 5.0 (3-OH)**, **pK₂²⁵ 8.96 (pyridinium⁺)**. Purify the vitamin by recrystallisation from $\text{EtOH}/\text{Me}_2\text{CO}$, *n*-BuOH or $\text{MeOH}/\text{Et}_2\text{O}$. Its solubility in H_2O is 22%, and in EtOH it is 1.1%. It is insoluble in Et_2O and CHCl_3 . Acidic aqueous solutions are stable at 120°/30minute. The *free base* [65-23-6] $\text{C}_8\text{H}_{11}\text{NO}_3$, **M 168.2**, has **m 159-160°** after recrystallisation from Me_2CO and sublimation at 140-145°/0.0001mm. It has UV with λ_{max} at 290nm (ϵ 84,000), 253 and 325nm (ϵ 3,700 and 7,100) in 0.1N aqueous HCl. [Khun & Wendt *Chem Ber* 71 780 1938, DOI: 10.1002/cber.19380710415; 72 311 1939, DOI: 10.1002/cber.19390720214; Harris et al. *J Am Chem Soc* 61 1242 1939, DOI: 10.1021/ja01874a068; 62 3198 1940, DOI: 10.1021/ja01868a086; Beilstein 21/5 V 492.] See also **Pyridoxal-5'-phosphate H_2O** above.

Vitamin B₇ [Levocarnitine, carnitine, α -hydroxy- β -N,N,N-trimethylaminopropionic acid] [*R*(+) 541-14-0, *S*(-) 541-15-1, *RS* 461-06-3] $\text{C}_7\text{H}_{15}\text{NO}_3$, **M 161.2**, **m *R* or *S* isomer 197-198°(dec)**, **210-212°(dec)**, ***RS* isomer 195-197°**, $[\alpha]_{\text{D}}^{20}$ (+) and (-) 36 (c 10, H_2O), **pK²⁵ 3.6**. The *S*(L) isomer is *levocarnitine*, the vitamin. The *R* or *S* isomers crystallise from $\text{EtOH}/\text{Me}_2\text{CO}$ (*hygroscopic*). The *R* or *S* *hydrochlorides* crystallise from hot EtOH or $\text{EtOH}/\text{Et}_2\text{O}$ and have **m 142°(dec)**. The *racemate* crystallises from hot EtOH (*hygroscopic*), and its *hydrochloride* crystallises in needles from hot EtOH and has **m 196°(dec)**. [For racemate see Mazzetti & Lemmon *J Org Chem* 22 228 1957, DOI: 10.1021/jo01353a606; Beilstein 4 H 513, 4 I 548, 4 II 937-8, 4 III 1632-5, 4 IV 3185.] It is a nutrient involved in lipid metabolism [see Steiber et al. 'carnitine perspectives' *Mol Aspects Med* 25(5-6) 455 2004, DOI: 10.1016/j.mam.2004.06.006, PMID: 15363636].

Vitamin B₁₂ (cyanocobalamine, α -[5,6-dimethylbenzimidazolyl]cobamidcyanide) [68-19-9] **C₆₃H₈₈CoN₁₄O₁₄P**, **M 1355.4**, **m** darkens at 210-220° and does not melt below 300°, [α]_D²³₆₅₆ -59 (H₂O). Vitamin B₁₂ crystallises from de-ionised H₂O, with a solubility in H₂O of 1g/80g, and is dried under vacuum over Mg(ClO₄)₂. The dry red crystals are *hygroscopic* and can absorb ~12% of H₂O. A solution at pH 4.5-5 can be autoclaved for 20 minutes at 120° without decomposition. Aqueous solutions are stabilised by addition of (NH₄)₂SO₄. [Golding *Comprehensive Organic Chem* Vol 5 (Ed. Haslam; Pergamon Press, NY, 1979) pp 549-584.]

Alternatively, an aqueous solution of the coenzyme can be concentrated, if necessary in a vacuum at 25° or less, until the concentration is 0.005 to 0.01M (as estimated by the OD at 522nm of an aliquot diluted with 0.01M K-phosphate buffer pH 7.0). If crystals begin to form on the walls of the container, they should be re-dissolved with a little H₂O. The concentrated solution is placed in a glass stoppered flask and diluted with 5 volumes of Me₂CO. After 2-3 hours at 3° it is centrifuged (10,000xg/10minutes) in Me₂CO-insoluble plastic tubes to remove some amorphous precipitate. The clear supernatant is inoculated with a small crystal of the vitamin and allowed to crystallise overnight at 3°. Crystals are formed on the walls and the bottom of the container. A further 2 volumes of Me₂CO are added and set aside at 3° to further crystallise. Crystallisation is followed by observing the OD₅₂₂ of the supernatant. When the OD falls to 0.27, then *ca* 94% of the crystals have separated. The supernatant is decanted (saved for obtaining a second crop), and the crystals are washed with a little cold 90% aqueous Me₂CO (2x), 100% Me₂CO (2x), Et₂O (2x) at which time the crystals separate from the glass walls. Allow them to settle and remove residual Et₂O with a stream of dry N₂. The process can be repeated if necessary. The crystals can be dried in air or in a vacuum for 2 hours over silica gel at 100° with an 8-9% weight loss. [Barker et al. *Biochemical Preparations* 10 33 1963.] This material gives a single spot on paper chromatography [see Weissbach et al. *J Biol Chem* 235 1462 1960, PMID: 13843764.] The vitamin is soluble in H₂O (16.4mM at 24°, 6.4mM at 1°), in EtOH and PhOH but insoluble in Me₂CO, Et₂O, CH₂Cl₂ and dioxane. Its UV has λ_{\max} at 260, 375 and 522nm (ϵ 34.7 x 10⁶, 10.9 x 10⁶ and 8.0 x 10⁶ /mole) in H₂O. The dry crystals are stable for months in the dark, but aqueous solutions decompose on exposure to VIS or UV light or alkaline CN⁻, but are stable in the dark at pH 6-7. The vitamin is inactivated by strong acids or alkalis. It is an essential cofactor for various enzymes, and deficiency leads to *pernicious anaemia*, hence its use as a food supplement. Its biological half life is ~6 days (400 days in the liver) and it is excreted *via* the kidneys. [Barker et al. *J Biol Chem* 235 480 1960, PMID: 13796809; see also *Vitamin B₁₂* (Zagalak & Friedrich Eds) Walter de Gruyter, Berlin 1979, ISBN 10: 3110076683, ISBN 13: 9783110076684; for the publication series ‘Corrin Synthesis—Parts I to IV’ which include the A. Eschenmoser and R.B. Woodward synthesis of *Vitamin B₁₂* see Eschenmoser et al. *Helv Chim Acta* 98 1475 2015, DOI: 10.1002/hlca.201400399, and subsequent papers; *Beilstein* 26 IV 3117.]

Vitamin C See ascorbic acid entry in ‘Carbohydrates’ in this Chapter.

Vitamin D₂ (ergocalciferol) [50-14-6] **C₂₈H₄₄O**, **M 396.7**, **m** 114-116°, 114-118°, [α]_D²⁰₅₄₆ +122 (c 4, EtOH) It is converted into its 3,5-dinitrobenzoyl ester and crystallised repeatedly from acetone. The ester is then saponified and the free vitamin is isolated. [Laughland & Phillips *Anal Chem* 28 817 1956, DOI: 10.1021/ac60113a012; *Beilstein* 6 IV 4404.] A secosteroid which enhances intestinal absorption of Ca, Fe, Mg, Zn and phosphate in humans.

Vitamin D₃ (cholecalciferol, Calcitol, activated 7-dehydrocholesterol, (+)-vitamin D₃) [67-97-0] **C₂₇H₄₄O**, **M 384.6**, **m** 83-85°, 83-86°, [α]_D²⁰₅₄₆ +126 (c 2, EtOH). It is converted into its 3,5-dinitrobenzoyl ester and crystallised repeatedly from acetone. The ester is then saponified and the free vitamin is isolated. Store it in sealed ampoules under argon below 8°. It acts through a receptor which modulates differentiation and proliferation of normal and neoplastic cells. [Laughland & Phillips *Anal Chem* 28 817 1956, DOI: 10.1021/ac60113a012; DeLuca & Schnoes *Ann Rev Biochem* 52 411 1983, DOI: 10.1146/annurev.bi.52.070183.002211; *Beilstein* 6 III 2811, 6 IV 4149.]

Vitamin E (2*R*,4*R*,8*R*- α -tocopherol, natural active isomer) [59-02-9] **C₂₉H₅₀O₂**, **M 430.7**, **m** 2.5-3.5°, **b** 200-220°/0.1mm, 200°/0.005mm, **d**₄²⁵ 0.950, **n**_D²⁵ 1.5045, [α]_D²⁵ +3.58 (c 1, *C₆H₆). Vitamin E is a viscous yellow oil which is distilled at high vacuum. It has λ_{\max} at 294nm ($E_{1\text{cm}}^{1\%}$ 71). It is oxygen and light sensitive and is best stored as its stable *D*- α -acetate [58-95-7] which is purified by evaporative distillation at **b** 180-200°

(bath temperature)/0.7mm, and has $[\alpha]_D^{25} +3.3$ (c 5.1, EtOH). It forms needles at -30° and has **m** 26.5-27.5°, and $[\alpha]_D^{25} +0.25$ (c 10, CHCl₃). [For NMR see Cohen et al. *Helv Chim Acta* **64** 1158 1981, DOI: 10.1002/hlca.19810640422; Burton & Ingold *Acc Chem Res* **19** 194 1986, DOI: 10.1021/ar00127a001; Karrer et al. *Helv Chim Acta* **21** 520 1938 DOI: 10.1002/hlca.19380210173; Robeson *J Am Chem Soc*, **64** 1487 1942, DOI: 10.1021/ja01258a507; **65** 1660 1943, DOI: 10.1021/ja01248a510.] Of the **eight isomers** the D- α -isomer is the most active. [See W. Friedrich 'Vitamins' Walter de Gruyter Publ, Berlin 1988.] [*Beilstein* **17/4** V 168.] (\pm)- **α -Tocopherol** [10191-41-0], which is a mixture of **four racemates**, is a thick oil with **b** 200-220°/0.1mm, **d** $^{25}_4$ **0.950**, **n** $^{25}_D$ **1.5045**, and λ_{\max} at 294nm ($E_{1\text{cm}}^{1\%}$ 71) see above. Store in the dark below 10°. Tocopherols generally are freely soluble in most organic solvents, insoluble in H₂O; and in the absence of oxygen are stable to heat alkali and acids (<100°), but are sensitive to light and traces of metal salts (e.g. Fe, Ag) in air.

DL-Vitamin E acetate (DL- α -tocopheryl acetate) [7695-91-2] **C₃₁H₅₂O₃**, **M 472.8**, **m** -27.5°, **b** 194-196°/0.01mm, 222-224°/0.3mm, **d** $^{20}_4$ **0.958g/ml**, **n** $^{20}_D$ **1.4958**. It is a viscous liquid which is purified by distillation under high vacuum under N₂ or argon, and stored in sealed ampoules in the dark. It is considerably more stable to light and air than the parent unacetylated vitamin. It is insoluble in H₂O but freely soluble in organic solvents. All **eight stereoisomers** have been synthesised. The commercially pure **d- α -tocopheryl acetate (2R,4'R,8'R)** has **b** 180-200°/0.7mm and $[\alpha]_D^{20} +3.9$ (c 5, EtOH); see above. [Cohen et al. *Helv Chim Acta* **64** 1158 1981, DOI: 10.1002/hlca.19810640422; *Beilstein* **17/4** V 169.]

Vitamin K₁ (2-methyl-3-phytyl-1,4-naphthoquinone) [84-80-0] **C₃₁H₄₆O₂**, **M 450.7**, **m** -20°, **b** 141-140/0.001mm, **b** 140-145°/10⁻³ mm, **d** $^{25}_{25}$ **0.967**, **n** $^{25}_D$ **1.527**, $[\alpha]_D^{20}$ **-0.4** (c 57.5, *C₆H₆). Vitamin K₁ is a yellow viscous oil, which can be distilled at high vacuum practically unchanged. It is insoluble in H₂O, but soluble in common organic solvents. Store it in the dark under N₂ as it is oxygen sensitive. It has $A_{1\text{cm}}^{1\%}$ 328 at 248nm. [Fieser et al. *J Am Chem Soc* **61** 2559 1939, DOI: 10.1021/ja01878a514; DOI: 10.1021/ja01878a515; Hirschmann et al. *J Am Chem Soc* **76** 4592 1954, DOI: 10.1021/ja01647a026; Isler & Doebl *Helv Chim Acta* **37** 225 1954, DOI: 10.1002/hlca.19540370128; *Beilstein* **7** IV 2496.]

Vitamin K₃ (2-methyl-1,4-naphthoquinone, Menadione, Menaphthone) [58-27-5] **C₁₁H₈O₂**, **M 172.2**, **m** 105-106°, 105-107°. Recrystallise it from 95% EtOH, or MeOH after filtration. It forms bright yellow crystals which are decomposed by light. Its solubility in EtOH is 1.7% and in *C₆H₆ it is 10%. It **IRRITATES** mucous membranes and skin. [Fieser *J Biol Chem* **133** 391 1940, <http://www.jbc.org/content/133/2/391>; *Beilstein* **7** IV 2430.]

Vitamin K₅ (4-Amino-2-methyl-1-naphthol hydrochloride) [130-24-5] **C₁₁H₁₂NO**. HCl, **M 209.6**, **m** 283°(dec), **pK_{Est(1)} ~5.6** (NH₂), **pK_{Est(2)} ~10.4** (OH). Crystallise it from dilute HCl. [Sah *Recl Trav Chim Pays-Bas* **59** 454 1940, DOI: 10.1002/recl.19400590505; Sah *Recl Trav Chim Pays-Bas* **60** 373 1941, DOI: 10.1002/recl.19410600509; Veldstra & Wiardi *Recl Trav Chim Pays-Bas* **61** 547 1942, Veldstra & Wiardi *Recl Trav Chim Pays-Bas* **62** 75 1943, DOI: 10.1002/recl.19430620203; *Beilstein* **13** III 1921.]

Vitamin P [(+)-Quinacr, quercetin-3-rubinoside, 2-(3,4-dihydroxyphenyl)-5,7-dihydroxy-3-[α -L-rhamno-pyranosyl-(1 \rightarrow 6)- β -D-glucopyranosyloxy]-4H-chromen-4-one, (+)-Rutin] [153-18-4] **C₂₇H₃₀O₁₆**, **M 610.5**, **664.5** (trihydrate), **m** 188-189°, 195-196°, 242° (anhydrous?) $[\alpha]_{546}^{20} +13$ (c 5, EtOH), $[\alpha]_D^{24} -37.6$ (c 1.24, pyridine), (polyphenolic flavone **pKs 7—10**). The vitamin crystallises from MeOH or water/EtOH, dry it in air, then dry it further for several hours at 110° or in high vacuum at 120°. It forms yellow crystals from EtOH/Me₂CO/H₂O (2:1:1). Its solubility in H₂O is 13mg/100ml (25°). It has also been purified by passing (0.5g) through a Kieselgel column (30 x 5cm) with EtOAc/MeOH/H₂O (100:20:15), and after 750ml have passed through, the yellow fraction of 250ml gives the glycoside (0.3g) on evaporation. [Hörhammer et al. *Chem Ber* **101** 1183 1968, DOI: 10.1002/cber.19681010407; Marini-Bettolo *Gazz Chim Ital* **80** 631 1950, *Beilstein* **18/5** V 519.] Rutin has antioxidant and antibacterial properties.

Vitamin U (S-Methyl-L-methionine chloride) [S- 1115-84-0, S-ion- 6708-35-6 and 4727-40-6, RS- 44901-24-8] **C₆H₁₅NO₂S**. Cl, **M 199.5**, **164.3** (ion), **m** 135° (dec), 139° (dec), $[\alpha]_D^{23} +33$ (0.2M HCl), **pK₁ 1.9**, **pK₂ 7.9**. Likely impurities are methionine, methionine sulfoxide and methionine sulfone. Crystallise Vitamin U from water by adding a large excess of EtOH. It is **hygroscopic**. It should be stored in a cool, dry place and protected

from light. It has also been purified on a column of Dowex 50 H⁺ form, washed with H₂O and eluted with 6N ammonia, lyophilised, the residue is dissolved in dilute HCl, lyophilised again and then it is obtained as colourless prisms by vapour diffusion of Et₂O in a 1:1 (v/v) mixture of MeOH/EtOH in which the **hydrochloride** is dissolved [Del Re *Acta Cryst B* **33** 3289 1977, DOI: 10.1107/S0567740877010887]. The **iodide salt**, **m** ~150°(dec), is obtained by dissolving it in 50% aqueous EtOH and adding ~3.5 volumes of absolute EtOH. The **bromide salt** has **m** 140°(dec) [Toennies & Kolb *J Am Chem Soc* **67** 849 1945, DOI: 10.1021/ja01221a046]. It is a **biological methylating agent**. For ‘Anti-peptic ulcer dietary factor (vitamin ‘U’) in the treatment of peptic ulcer’ see Cheney [*J Am Diet Assoc* **26**(9) 668 1950, PMID: 15436263].

dl-Warfarin (Coumafene, 4-hydroxy-3-(3-oxo-1-phenylbutyl)-2H-1-benzopyran-2-one) [81-81-2] C₁₉H₁₆O₄, **M** 308.3, **m** 162-164°, **pK**²⁰ 4.85 (the 4-mercapto derivative has **pK**²⁰ 6.60). *dl*-Warfarin crystallises from EtOH or MeOH. Its UV has λ_{max} at 308nm (ε 13,610) in H₂O. The **acetate** has **m** 117-118°, the **O-triflate** has **m** 90-91°, and the **2,4-dinitrophenylhydrazone** has **m** 215-216°. [West et al. *J Am Chem Soc* **83** 2676 1961, DOI: 10.1021/ja01473a020; HPLC: Banfield & Rowland *J Pharm Sci* **72** 921 1983, DOI: 10.1002/jps.2600720820; *Beilstein* **17** III/IV 6794.]

R-(+)-Warfarin has **m** 170-171°, [α]_D²³ +149 (c 1.2, in 0.5N NaOH), +25.5 (c 2, AcOH), +15.5 (c 2, MeCN) and -19 (c 1.1, propan-2-ol), and **S-(-)-Warfarin** (*more active enantiomer*) has **m** 172-173°, C₁₉H₁₆O₄, **M** 308.3, [α]_D²³ -148 (c 1.2, in 0.5N NaOH), -25.5 (c 2, AcOH), -15.5 (c 2, MeCN) and +19 (c 1.1, propan-2-ol). *dl*-Warfarin is **optically resolved** via recrystallisation of the **quinidine salt**, and the **free acids** are recrystallised (70g) from 600ml of 80% aqueous Me₂CO. Large prismatic crystals of the pure enantiomers are obtained by slow crystallisation from Me₂CO or AcOH. The solubilities of the pure enantiomers at 25° are 11.2% in Me₂CO and 2.6% in AcOH, whereas the racemate has solubilities of 6.5% in Me₂CO and 2% in AcOH. The IR spectra are the same with ν_{max} (CHCl₃) at 2.78 (w), 5.88, 6.16 and 6.38μ. [West et al. *J Am Chem Soc* **83** 2676 1961, DOI: 10.1021/ja01473a020; Cbz-proline diastereoisomeric esters were used for HPLC analysis: Banfield & Rowland *J Pharm Sci* **72** 921 1983, DOI: 10.1002/jps.2600720820.] Although both enantiomers produce anticoagulant results, the **S-enantiomer** is 2-5 times **more potent** than the **R-enantiomer**, but they are cleared differently because the metabolite of the former is the 7-OH derivative, whereas the 10-OH metabolite is cleared from *R-warfarin*. For ‘Foundation guide to warfarin therapy’ see Hirsch et al. *J Am Coll Cardiol* **41**(9) 1633 2003, DOI: 10.1016/S0735-1097(03)00416-9, PMID: 12742309; and for metabolites see Naveed et al. *J Chromatogr B* **1008**(1) 164–173 2016, DOI: 10.1016/j.jchromb.2015.11.036. They are effective **anticoagulants** and **rodenticides**.

Xanthopterin monohydrate (2-amino-4,6-dihydroxypteridine, 2-aminopteridin-4,6(1*H*,5*H*)-dione) [5979-01-1 (H₂O), 119-48-8 (anhydrous)] C₆H₅N₅O₂·H₂O, **M** 197.2, **m** <300°, **pK**₁ 1.6 (basic), **pK**₂ 6.59 (acidic), **pK**₃ 9.31 (acidic)(anhydrous species), and **pK**₁ 1.6 (basic), **pK**₂ 8.65 (acidic), **pK**₃ 9.99 (acidic)(7,8-hydrated species). Purification is as for isoxanthopterin (see [529-69-1] ‘Heterocyclic Compounds’, Chapter 3). It is crystallised by acidifying a hot ammoniacal solution with formic acid, and collecting the crystals by centrifugation followed by washing with EtOH, ether and drying at 100° *in vacuo*. Its R_F values on paper chromatography are 0.15 (*n*-PrOH, 1% aqueous NH₃, 2:1), 0.36 (*n*-BuOH/AcOH/H₂O, 4:1:1) and 0.47 (3% aqueous NH₃). [Inoue & Perrin *J Chem Soc* 2600 1962, DOI: 10.1039/JR9620002600; Inoue *Tetrahedron* **20** 243 1964, DOI: 10.1016/S0040-4020(01)93212-0; see also Blakley *Biochemistry of Folic Acid and Related Pteridines* North Holland Publ Co, Amsterdam 1969, ISSN 0071-965X; *Beilstein* **26** II 313, **26** III/IV 4000.] It is a urinary metabolite as a marker of malignant disease [for ‘Altered urinary excretion of pteridines in neoplastic disease. Determination of biopterin, neopterin, xanthopterin, and pterin’ see Rokos et al. *J Clin Chim Acta* **105** 275 1980, DOI: 10.1016/0009-8981(80)90470-2].

Xanthotoxin (Methoxalen, 9-methoxypsoralen, Ammoidin, 9-methoxyfuro[3,2-*g*][1]benzopyran-7-one) [298-81-7] C₁₂H₈O₄, **M** 216.2, **m** 146-148°, 148°, 148-149°, 148-150°. Purify xanthotoxin by recrystallisation from *C₆H₆/petroleum ether (b 60-80°) to give silky needles, or from EtOH/Et₂O to give rhombic prisms or from hot H₂O to give needles. It is soluble in aqueous alkali due to ring opening of the cyclic lactone but recyclises upon acidification. It has UV with λ_{max} (EtOH) at 219, 249 and 300nm (log ε 4.32, 4.35 and 4.06) and ¹H NMR in CDCl₃ with δ at 7.76 (d, 1H, *J* = 10Hz), 7.71 (d, 1H, *J* = 2.5Hz), 7.38 (s, 1H), 6.84 (d, 1H, *J* = 2.5Hz), 6.39 (d, 1H, *J* = 10Hz) and 4.28 (s, 3H). [Nore & Honkanen *J Heterocycl Chem* **17** 985 1980, DOI:

10.1002/jhet.5570170527.] It is a **DNA intercalator**, is used in the treatment of dermal diseases, and is a **human carcinogen** [Tessman et al. *Biochemistry* **24** 1669 1985, DOI: 10.1021/bi00328a015.] [Beilstein **19** I 711, **19/6** V 15.] The drug has been used for the treatment of eczema, psoriasis, vitiligo and cutaneous lymphomas.

Zeatin (*trans-N⁶-[4-hydroxy-3-methylbut-2-en-1-yl]adenine*) [1637-39-4] $C_{10}H_{13}N_5O$, **M 219.3**, **m 207-208°**, **209-209.5°**, **208-210°**, **pK₁ 4.4 (basic)**, **pK₂ 9.8 (acidic)**. Purify zeatin by recrystallisation from EtOH or H₂O. Its UV has λ_{max} at 207 and 275nm (ϵ 1,400 and 14,650) in 0.1N aqueous HCl; 212 and 270nm (ϵ 17,050 and 16,150) in aqueous buffer pH 7.2; and 220 and 276nm (ϵ 15,900 and 14,650) in 0.1N aqueous NaOH. The **picrate** has **m 192-194°** (from H₂O) from which zeatin can be recovered by treatment with Dowex-1 x 8 (200-400 mesh, OH⁻ form). Zeatin is isolated from sweet corn kernels (1mg from 70g of corn). It was synthesised from 6-mercaptopurine [see 154-42-7] and *trans*-4-hydroxy-3-methyl-2-buten-1-ylamine at 134° overnight followed by isolation as the **picrate m 180-190°**, and liberating the **free base Zeatin** by passing an aqueous solution through an analytical Dowex 1x8 (200-400 mesh) column in the OH⁻ form. The *trans*-buten-1-ylamine was prepared from reaction of methyl γ -bromotiglate [Ratney & English *J Org Chem* **25** 2213 1960, DOI: 10.1021/jo01082a600; Inhoffen et al. *Justus Liebigs Ann Chem* **580** 7 1953, DOI: 10.1002/jlac.19535800103] and NaN₃ in MeCN followed by reduction of the resulting methyl γ -azidotiglate (b 52-54°/0.2mm, ν 2100cm⁻¹ for -N₃) with LiAlH₄. With acetic anhydride in dry pyridine (shaking for 30 minutes to give a clear solution then overnight at ~25°), and the isolated **mono-O-acetylzeatin** has **m 168-169°** (needles) after recrystallisation from H₂O. It has strong IR bands at ν 1740 and 1240cm⁻¹. [For isolation and characterisation see: Letham et al. *Aust J Chem* **22** 205 1969, DOI: 10.1071/CH9690205; Letham et al. *Proc Chem Soc* p.230 (in **notes** starting on p.201) 1964, DOI: 10.1039/PS9640000201; for synthesis see Shaw & Wilson *Proc Chem Soc* p.231 (in **notes** starting on p.201) 1964, DOI: 10.1039/PS9640000201, and Shaw et al. *J Chem Soc (C)* 921 1966, DOI: 10.1039/J39660000921.]

It is a cell division factor (**plant growth regulator**) [Letham & Palni *Ann Rev Plant Physiol* **34** 163 1983, DOI: 10.1146/annurev.pp.34.060183.001115] and **inhibits mitochondrial function** [Miller *Plant Physiol* **69** 1274 1982, DOI: 10.1104/pp.69.6.1274]. **Trans-Zeatin hydrochloride** [6025-81-6] $C_{10}H_{13}N_5O \cdot HCl$, **M 255.7**, is commercially available as is the **trans-9-riboside derivative**, [6025-53-2] $C_{15}H_{21}N_5O_5$, **M 351.4**, and is a cytokine which separates from aqueous AcOH with **m 177-179°**. It solubility in AcOH is ~5%. [McDonald & Morris *Methods Enzymol* **110** 347 1985, DOI: 10.1016/S0076-6879(85)10093-5]. For 'Gerontomodulatory and youth-preserving effects of zeatin on fibroblasts of human skin' see Rattan & Sodagam [*Rejuvenation Res* **8**(1) 46 2005, DOI: 10.1089/rej.2005.8.46, PMID: 15798374].

Zoxazolamine (Flexin, 5-chloro-2-benzoxazolamine) [61-80-3] $C_7H_5ClN_2O$, **M 168.6**, **m 184-185°**, **185-185.5°**, **189-190°**. Purify flexin by recrystallisation from *C₆H₆ or 50% aqueous EtOH. Its solubility at ~25° in DMSO is 34mg/ml, in EtOH it is 34mg/ml and in H₂O it is <1mg/ml. It sublimes at high vacuum. In MeOH it has UV λ_{max} at 244 and 285nm. The synthesis of this oxazoline is briefly described by Nagano et al. [Nagano et al. *J Am Chem Soc* **75** 2770 1953, DOI: 10.1021/ja01107a511; English Patent 240,969 1925], but the synthesis of the 4,6d₂ derivative used in pharmacological studies described by Tanabe et al. [*J Med Chem* **13** 30 1970, DOI: 10.1021/jm00295a008] is reported here because more details about the reaction are provided by the latter authors. Thus, a mixture of 2-amino-4-chlorophenol-3,5d₂ (2.351g) and CNBr (3.4g) in H₂O (225ml) are heated in a steam bath for 15 minutes (**CARE as CNBr is highly toxic**). The mixture is cooled to ~25° and neutralised with 15M NH₄OH to pH 8 with cooling, and the solid product is filtered off. After drying *in vacuo*, it is sublimed at 160°/0.020mm to provide **5-chloro-2-benzoxazolamine-4,6d₂** (1.926g) **m 182-183°** with the correct deuterium content.

The known (quantified) **muscle relaxant** (paralytic) potency of **Zoxazolamine** (the so-called '**Zoxazolamine Effect**') is employed as a benchmark standard when new compounds are tested as muscle relaxant inhibitors [see Decken & Gossage *J Inorg Biochem* **99**(2) 664 2005, PMID: 15621301]. For 'A quantitative assessment of motor function in cerebral palsy and evaluation of zoxazolamine (flexin) as a new muscular relaxant drug' see Hadra & Millichap [*Neurology* **6**(12) 843 1956, PMID: 13378587]. Zoxazolamine, as a centrally-acting muscle relaxant, reportedly acts by decreasing CNS interneuronal activity, by decreasing dopaminergic turnover and by inducing a pacemaker-like discharge pattern in dopaminergic neurons without being anxiolytic [McMillen et al. *J Neural Transm Gen Sect* **89** 11 1992, PMID: 1329854]. In addition to its muscle relaxant activity, Zoxazolamine is also a useful tool for assessing hepatic cytochrome P-450 activity [Sternson & Gammans *J Med Chem*, 19 174 1976, DOI: 10.1021/jm00223a034].

CHAPTER 7

NANOMATERIALS AND NANOTECHNOLOGY

INTRODUCTION

Nanomaterials are substances that are, or have been, reduced in size to the range from 1nm to ~100nm (i.e. 1 to ~100 nanometers, or 1 to ~100 x 10⁻⁹ meters). **Nanotechnology** is the science and applications of nanomaterials, and is growing at an ever increasing pace. At this particle size the properties of materials can be altered dramatically. Properties such as solubility, reactivity, spectroscopy, electrical and magnetic, transport through membranes etc. are generally different from those of the same materials with large particle size. The applications of materials of nano size have escalated in the last fifteen or so years and are currently gaining momentum. The technology has broad applications in performance materials, health, consumer products, water, information technology and energy. The discovery of **fullerenes** (1996 Nobel Prize for Chemistry to H.W. Kroto, R.E. Smalley and R.F. Curl) and of **Graphene** (2010 Nobel Prize for Physics to A. Geim and K. Novoselov) are very important factors in the development of nanotechnology. Not very many nanomaterials have so far been included in chemical catalogues of commercially available substances, e.g. as compared with catalysts (although some catalysts are now available as nanoparticles), hence the relatively smaller size of this chapter. Fullerenes and related materials such as carbon nanotubes have found applications in medicine for the transport of drugs and biological materials to targeted sites and some readily available derivatives are described in this chapter, together with other commercially available nanoparticulate substances, e.g. catalysts, magnetic and electrically conducting thin films, and quantum dots. General safety aspects of handling nanomaterials have been briefly addressed. Nanoscience and nanotechnology is developing explosively as the awareness and usefulness of materials at the nano scale is becoming evident. In the past few years this has resulted in the appearance of an enormous number of journals and periodicals in which scientific, technological and medical discoveries are readily published. Many of these publications are listed at the end of this chapter and can be accessed *via* their International Standard Series Numbers (ISSNs). A large number of books have also been published on these subjects and can be accessed *via* their International Standard Book Numbers (ISBNs).

IDENTIFICATION AND MEASUREMENT OF NANOMATERIALS

The 1 to ~ 100nm (10 to ~1000Å) particles are smaller than the wavelength of UV light, and thus the use of appropriate small particles e.g. electrons, or radiation e.g. X-rays is necessary to identify and measure these particles. Instrumentation for these purposes has been developed and the resolutions are continuously being improved. These instruments can measure very accurately, and depending on the physical principles of the machines, they measure different parameters of the nanoparticles, e.g. diameter (if spherical), length (if rod like), width (if bulky) etc, and not necessarily all parameters at the same time with one instrument. The commonly used instruments include the **Transmission Electron Microscope** (TEM) in which the electron beam runs past the particles placed on a transparent surface, and then through the surface giving a 2D image and magnifying one million times. In the **Scanning Electron Microscope** (SEM) and **Scanning Tunneling Microscope** (STM, or *Spectroscopy* STS) the electron beam scans the surface of the particles to provide a 3D picture, e.g. of hairs on a fly. **Dynamic Light Scattering** (DLS) uses light scattering through solutions of nanoparticles. Another technique, **Scattering Probe Microscopy** (SPM), allows scanning with a probe over a surface and includes **Atomic Force Microscopy** (AFM, or *non-contact atomic force microscopy* NC-AFM) where a ceramic or semiconductor tip scans the surface (much like a phonograph needle scans a record), and measures the atomic forces beneath it. The NC-AFM tip is attached to a very small tuning fork that oscillates at amplitudes of 0.2 to 0.5Å, smaller than the diameter of an atom. When the tip is brought very close to a molecule on a surface, this frequency alters slightly thus **detuning** due to the atomic forces between the tip and the molecule to give a complete picture of the molecule on the surface. By placing a CO (carbon monoxide)

molecule at the tip, and keeping it there throughout the imaging process, very high resolution can be obtained. Gross and coworkers [Gross et al. *Science* **325** 1110 2009, DOI: 10.1126/science.1176210] obtained the complete image of pentacene on a copper surface in this way, and succeeded in imaging the amount of charge on a single atom [Gross et al. *Science* **324** 1428 2009, DOI: 10.1126/science.1172273]. The sensitivity of **Magnetic Resonance Imaging** (MRI) has been considerably increased (with a resolution of ~4nm), by Rugar and coworkers [Degen et al. *Proc Natl Acad Sci* **106** 1313 2009, DOI: 10.1073/pnas.0812068106; cf H. Birch *MRI on the Nanoscale*, RSC Chemistry World **6**(2) 28 2009, J. M. Crow *Picture perfect pentacene*, RSC Chemistry World **6**(1) 462011] who have developed the **Magnetic Resonance Force Microscopy** (MRFM) technique. Here the sample is placed at the end of a tiny silicon cantilever to which is applied, via a magnetic tip, a radio frequency magnetic field. The spin of the nuclei alternate (*up* and *down*) when the frequency reaches their ‘wobbling’ frequencies making the cantilever vibrate. Photon density 3D images of the specimen (e.g. tobacco mosaic virus, *ca* 18 nanometers across) can be obtained by using a laser to detect the cantilever vibrations. The technique does not affect the sample and gives superior images to those from STM and AFM. [A.I. Kirkland and J. I. Hutchinson (eds) *Nanocharacterisation* RSC Publishing UK, 2007, ISBN 9781847557926, 1847557929; A. Turley *Sizing it up*, RSC Chemistry World **8**(3) 50 2011; M. Thompson RSC *The Characterisation of Nanoparticles; Analytical Methods Committee Technical Briefs* 48, December 2010; ISSN 1757-5958, <http://bit.ly/AMCparticle>). Other techniques include **Near-Infrared Spectroscopy** (NIR) and solution phase NIR [Itkis et al. *Nano Lett* **3** 309 2003, DOI: 10.1021/nl025926e], general **Electron Microscopy** (EM), **Field Emission Microscopy** (FEM), **FT-Raman Scattering Spectroscopy**, **Nuclear (proton) Magnetic Resonance Spectroscopy** (NMR for derivatised carbon nanotubes, CNTs), **Thermo Gravimetric Analysis** (TGA), **Energy Dispersive X-ray Spectroscopy** (EDX), as well as other optical techniques (see Bibliography at the end of this chapter).

Preparation of the specimen is very important since the atoms or/and molecules have to be spread over the surface in as close to one atom or molecule thick. Where thicknesses (*t*) of 7 to ~13nm need to be measured **Grazing Incident X-Ray Reflectometry** (GICRR/XRR) provides an initial measure of particle lengths (*l*) of 1 to 35nm. **Spectroscopic Ellipsometry** (SE, *t* = ~ 10.5 to 12.3nm, *l* = 1 to 10.5nm), **Scanning Ion Mass Spectrometry** (SIMS, *t* ~ 10.4 to 10.8nm, *l* = ~12 to 13 nm), **Transmission Electron Microscopy** (TEM, *t* ~ 7.5 to 12.5nm, *l* = 12.5 to 18.5 nm), **X-Ray Fluorescence Spectroscopy** (XFS, *t* ~ 7.5 to 9.3nm, *l* = 16.0 to 19.5 nm), **X-Ray Photoelectron Spectroscopy** (XPS, *t* ~ 9.2 to 11.7, *l* = 18.5 to 25.5 nm) and higher resolution **Grazing Incident X-Ray Reflectometry** (GIXRR/XRR, *t* ~ 9.2 to 11.7, *l* = 24.0 to 36.0 nm) provide the approximate *t* and *l* values stated in the brackets. It must be noted that these values are approximate as they do vary depending on the nature of the materials that are measured.

Average particle size: The accepted method for determining the primary particle size of nanomaterials is by TEM analysis. However, this is expensive and time-consuming. Studies have shown that an average primary particle size for spherical or cubic shaped dry nanoparticles of roughly uniform shape and size can be calculated from the equation:

$$\text{Average particle size (nanometers)} = (6/\text{BET surface area}^* \times \text{density}^{**}) \times 1000$$

Where surface area* is in sq. m/g, and density** is in g/cc. **Micromeritics Analytical Services** (MAS) will measure **BET surface area** by a gas absorption procedure and the real density by gas pycnometry. This gives a good estimate of average primary particle size. For further information see www.micromeritics.com, Micromeritics Analytical Services Downloads.

FULLERENES AND RELATED SUBSTANCES

Fullerenes and related substances are made by heating carbon (e.g. soot, graphite) to form nano-sized carbon ‘balls’. These nanoparticles look like ‘footballs’ in being constructed of five (pentagon) and/or six (hexagon) carbon rings. Their carbon ring sizes make for stable ‘ball-like’ molecules of 60 or more carbon atoms and have some solubility in organic solvents. These carbon **allotropes** have a variety of exotic properties including electrochemical [Echegoyen & Echegoyen *Acc Chem Res* **31** 593 1998, DOI: 10.1021/ar970138v], ferromagnetism [Allemand et al. *Science* **253** 301 1991, DOI: 10.1126/science.253.5017.301], superconductivity [Hebard et al. *Nature* **350** 600 1991, DOI: 10.1038/350600a0], photophysical [Guldi *Chem Commun* 321 2000, DOI: 10.1039/A907807J], anti-HIV bioactivity [Friedman et al. *J Am Chem Soc* **115** 6506 1993, DOI: 10.1021/ja00068a005], and their ‘aromatic’ nature allows them to react accordingly and be functionalised (see below) for further chemical reactions [Kordatos et al. *J Org Chem* **66** 4915 2001, DOI:

10.1021/jo015608k]. New and useful applications are continually being found in nano science, medicine, pharmaceuticals, and nanotechnology [see references below].

FULLERENES

Fullerene C₆₀ (Buckminsterfullerene C₆₀, Footballene, Buckyball 60) [99685-96-8] M 720.66 and Fullerene C₇₀ [115383-22-7] M 840.77. *Purification procedures:* (a) These were purified from the soluble toluene extract (400mg) of the soot (Fullerite), formed from resistive heating of graphite, by adsorption on neutral alumina (100g, Brockmann I, 60 x 8cm). Elution with toluene/hexane (5:95 v/v) gives *ca* 250mg of quite pure C₆₀. It has characteristic spectral properties (see below). Further elution with toluene/hexane (20:80 v/v, i.e. increased polarity of solvent) provides 50mg of 'pure' C₇₀ [Allemand et al. *J Am Chem Soc* **113 1050 1991, DOI: 10.1021/ja00003a053].**

Chromatography on alumina can be improved by using conditions that favour adsorption rather than crystallisation. Thus the residue from toluene extraction (1g) in CS₂ (*ca* 300ml) is adsorbed on alumina (375g, standard grade, neutral *ca* 150 mesh, Brockmann I) and loaded as a slurry in toluene/hexanes (5:95 v/v) to a 50 x 8cm column of alumina (1.5Kg) in the same solvent. To avoid crystallisation of the fullerenes, 10% of toluene in hexane is added quickly followed by 5% of toluene in hexane after the fullerenes had left the loading fraction (2-3 hours). With a flow rate of 15ml/minute the purple C₆₀ fraction is eluted during a 3-4 hour period. Evaporation of the eluates gives 550-630mg of product which, after recrystallisation from CS₂/cyclohexane yields 520-600mg of C₆₀ which contains adsorbed solvent. On drying at 275°/10⁻³mm for 48 hours a 2% weight loss is observed although the C₆₀ still contains traces of solvent. Further elution of the column with 20% of toluene in hexane provides 130mg of C₇₀ containing 10-14% of C₆₀ (by ¹³C NMR). This was re-chromatographed as above using a half scale column and adsorbing the 130mg in CS₂ (20ml) on alumina (24g) which gave 105mg of recrystallised C₇₀ (containing 2% of C₆₀). The purity of C₆₀ can be improved further by washing the crystalline product with Et₂O and Me₂CO followed by recrystallisation from *C₆H₆ and vacuum drying at high temperatures.

(b) Carbon soot from resistive heating of a carbon rod in a partial helium atmosphere (0.3bar) under specified conditions is extracted with boiling *C₆H₆ or toluene, filtered and the red-brown solution is evaporated to give crystalline material in 14% yield which is mainly a mixture of fullerenes C₆₀ and C₇₀. Chromatographic filtration of the 'crude' mixture with *C₆H₆ allows no separation of components, but some separation was observed on silica gel TLC with *n*-hexane or *n*-pentane, but not cyclohexane as eluants. Analytical HPLC with hexanes (5µm Econosphere silica) gave satisfactory separation of C₆₀ and C₇₀ (retention times of 6.64 and 6.93 minutes respectively) at a flow rate of 0.5ml/minute and using a detector at 256nm. HPLC indicated the presence of minor (<1.5% of total mass) unidentified C_n species with retention times of 5.86 and 8.31 minutes. Column chromatography on flash silica gel with hexane gives a few fractions of C₆₀ with ≥95% purity, but later fractions contain mixtures of C₆₀ and C₇₀. These can be obtained in 99.85 and >99% purity, respectively, by column chromatography on neutral alumina. [Ajie et al. *J Phys Chem* **94** 8630 1990, DOI: 10.1021/j100387a004.]

(c) Separation of C₆₀ and C₇₀ can be achieved by HPLC on a dinitroanilinopropyl (DNAP) silica (5µm pore size, 300Å pore diameter) column with a gradient from *n*-hexane to 50% CH₂Cl₂ using a diode array detector at wavelengths 330nm (for C₆₀) and 384nm (for C₇₀). [Cox et al. *J Am Chem Soc* **113**, 2940, 1991, DOI: 10.1021/ja00008a023.] By using Soxhlet extraction of 'soot' and recrystallisation from toluene, and using the greater affinity of 95.% pure C₆₀

Soxhlet extraction of the 'soot' with toluene is a good preliminary procedure, and as C₇₀ has greater affinity for the solvent and recrystallisation from toluene gave 99.5% of pure C₆₀. [See *Purification of Laboratory Chemicals* 7th edition and Coustel et al. *JCS Chem Commun* 1402 1992, DOI: 10.1039/C39920001402.]

(d) Purification of C₆₀ from a C₆₀/C₇₀ mixture was also achieved by dissolving it in an aqueous solution of γ (but not β) cyclodextrin (0.02M) with refluxing and recovery of C₆₀ of >99 purity [See *Purification of Laboratory Chemicals* 7th edition, and Andersson et al. *JCS Chem Commun* 604 1992, DOI: 10.1039/C39920000604.]

(e) Similarly C₆₀ and C₇₀ from fresh carbon-arc soot can also be readily purified by using inclusion complexes with *p*-tert-butylcalix[6] *p*-tert-butylcalix[8]arenes. [See *Purification of Laboratory Chemicals* 7th edition, 2013, and Atwood et al. *Nature* **368** 229 1994, DOI: 10.1038/368229a0.]

(f) **Rapid and economical purification of C₆₀ fullerene** [22685-96-8, 99685-96-8] **M 720.6, m >280°, flash point 94°** from crude soot enriched in fullerenes with ~5% of soluble fullerenes available from suppliers such as

Texas Fullerene Corporation (8926 Kirby Drive, Houston, USA) and Polygon Enterprises (P.O.Box 5536, Waco) was used [See *Purification of Laboratory Chemicals* 7th edition, 2013].

(g) A higher purity C₆₀ can be obtained by using a glass filter funnel with a D4 frit (diameter 6.5cm) on a suction flask filled with a slurry of silica gel (~84g, 230-400 mesh) and Darco G60 charcoal (42g) in toluene to a plug height of 9cm; and covered with sea sand (1cm) [See *Purification of Laboratory Chemicals* 7th edition 2013, and for HPLC and NMR see Isaacs et al. *Helv Chim Acta* **76** 1231 1993, DOI: 10.1002/hlca.19930760310].

(h) Tour and coworkers [Scrivens et al. *J Am Chem Soc* **114** 7917 1992, DOI: 10.1021/ja00046a051] described another **gram-quantity purification procedure for C₆₀ fullerene** where a slurry of alkaline decolorising carbon Norit-A (36g) and silica gel (72g, flash chromatography grade 60 of 230-240 mesh ASTM, 0.040-0.063mm particle size) in toluene (200ml) was used. [See *Purification of Laboratory Chemicals* 7th edition, 2013.] Repeated chromatography on neutral alumina yields minor quantities of solid samples of C₇₆, C₈₄, C₉₀ and C₉₄ believed to be higher fullerenes have been separated by repeated flash chromatography on alumina with gradient elution using hexane/toluene mixtures. A stable oxide C₇₀O has been identified [Diederich et al. *Science* **252** 548 1991, DOI: 10.1126/science.252.5005.548, see also *Purification of Laboratory Chemicals* 7th edition, 2013].

Physical properties of Fullerene C₆₀: [135105-52-1] C₆₀ fullerene, M 720.64, does not melt below 360°, and starts to sublime at 300° in vacuo, and is now available commercially in a high state of purity. It is a mustard-coloured solid that appears brown or black with increasing film thickness. It is soluble in common organic solvents, particularly aromatic hydrocarbons which give a beautiful magenta colour. Toluene solutions are purple in colour. It is soluble in *C₆H₆ (5mg/ml), but dissolves slowly. C₆₀ crystallises in needles or plates. [Taylor *JCS Chem Commun* 1423 1990, DOI: 10.1039/C39900001423.]

UV-Vis in hexanes: λ_{max}nm(log ε) 211(5.17), 227sh(4.91), 256(5.24), 328(4.71), 357sh(4.08), 368sh(3.91), 376sh(3.75), 390(3.52), 395sh(3.30), 403(3.48), 407(3.78), 492sh(2.72), 540(2.85), 568(2.78), 590(2.86), 598(2.87) and 620(2.60). IR (KBr): ν_{max} 1429m, 1182m, 724m, 576m and 527s cm⁻¹. ¹³C NMR: one signal with δ at 142.68. C₆₀ fullerene is a nano-channel organic semiconductor [Newman et al. *Chem Mater* **16** 4436 2004, DOI: 10.1021/cm049391x].

Purification of [5,6] C₇₀ fullerene for derivatisation [115383-22-7] C₇₀ fullerene, M 840.8, m >280°. By using a toluene extract (845mg) of Norit A (Aldrich) and a single-column purification using activated charcoal and derivatised polystyrene as stationary phases with toluene/1,2-dichlorobenzene then pure 1,2-dichlorobenzene as eluants, the six fractions obtained (confirmed by HPLC) were: (i) 205mg of pure C₆₀, (ii) 231mg of 38:62 C₆₀/C₇₀, (iii) 101mg of 88% pure C₇₀, (iv) 66mg of 92% pure C₇₀, (v) 131mg of 96% pure C₇₀ and (vi) 57mg of 94% pure C₇₀. Fractions with >90% purity were satisfactory for the preparation of fullerene C₇₀ derivatives. [Herrmann et al. *Helv Chim Acta* **78** 1673 1995, DOI: 10.1002/hlca.19950780705; for column used see Tour above and below.] Fullerene C₇₀ forms a **nano adduct** with 4,5-dimethoxy-1,2-quinonedimethane [Smith et al. *J Am Chem Soc* **117** 9359 1995, DOI: 10.1021/ja00141a031; Tour *J Org Chem Perspective* **72** 7477 2007, DOI: 10.1021/jo070543s].

Tour and coworkers [Scrivens et al. *J Am Chem Soc* **116** 6939 1994, DOI: 10.1021/ja00094a060] described a **gram-quantity purification procedure for C₇₀ fullerene** and separation of C₆₀/C₇₀ by HPLC [Scrivens & Tour *J Org Chem* **57** 6932 1992, DOI: 10.1021/jo00051a047] [For details of purification and ¹³C NMR in *C₆H₆ see *Purification of Laboratory Chemicals* 7th edition, 2013].

Physical properties of Fullerene C₇₀: C₇₀ (5,6)-fullerene [115383-22-7] M 840.78, does not melt below 360°, and starts to sublime at 300° in vacuo, and is now available commercially in a high state of purity. It is a reddish-brown solid but greenish black in thicker films. Solutions are port-wine red in colour. Mixtures of C₆₀ and C₇₀ are red due to C₇₀ being more intensely coloured. It is less soluble than C₆₀ in *C₆H₆ and also dissolves slowly. C₇₀ gives orange-coloured solutions in toluene. Drying at 200-250° is not sufficient to remove all the solvent. Samples need to be sublimed to be free from solvent. [Taylor *JCS Chem Commun* 1423 1990, DOI: 10.1039/C39900001423.] UV-Vis in hexanes: λ_{max} nm(log ε) 214(5.05), 235(5.06), 249sh(4.95), 268sh(4.78), 313(4.23), 330(4.38), 359(4.29), 377(4.45), 468(4.16), 542(3.78), 590sh(3.47), 599sh(3.38), 609(3.32), 623sh(3.09), 635sh(3.13) and 646sh(2.80). IR (KBr): ν_{max} 1430 m, 1428m, 1420m, 1413m, 1133mw, 1087w, 795s, 674ms, 642ms, 5778s, 566m, 535ms and 458m cm⁻¹. ¹³C NMR [run in the presence of Cr (pentan-2,4-dione)₃ which induces a ca 0.12ppm shift in the spectrum]: Five signals with δ at 150.07, 147.52, 146.82, 144.77 and 130.28, which are unaffected by proton decoupling. C₇₀ fullerene is a nano-channel organic semiconductor [Newman et al. *Chem Mater* **16** 4436 2004, DOI: 10.1021/cm049391x].

For *Isomers of C₇₀ dimers* see Zerbetto and Fowler [*J Phys Chem* **105** 1140 2001, DOI: 10.1021/jp0036036].

C₇₈ (C_{2v})-fullerene [136316-32-0] **M 936.98, melts above 350°**. It is now available commercially. Pure material is obtained as in the previous purification and elutes after C₇₆ fullerene, followed by **C₇₈ (D_{3h})-fullerene**. The identities are confirmed by an HPLC/GPC system with Waters 600E UV/VIS detection, mass and NMR spectroscopy. [Seleque et al. in Kadish and Ruoff (Eds) '*Fullerenes: Recent Advances in the Chemistry and Physics of Fullerenes and Related Materials*' The Electrochemical Soc. Inc, Pennington, NJ, 1994 ISBN 1566770823, Diederich & Whetten *Acc Chem Res* **25** 119 1992, DOI: 10.1021/ar00015a004; Diederich et al. *Science* **254** 1768 1991, DOI: 10.1126/science.254.5039.1768, MS and NMR: Taylor et al. *JCS Chem Commun* 1043 1992, DOI: 10.1039/C39920001043.] For '*Experimental and Theoretical Study of the Infrared, Raman, and Electronic Spectra of Two Isomers of C₇₈ of C_{2v} Symmetry*' see Benz et al. [*J Phys Chem* **100** 13399 1996, DOI: 10.1021/jp9602282].

C₈₄ fullerene [135113-16-5] **M 1008.94, melts above 350°**. It is now available commercially. Pure material is obtained as in the previous purification and elutes after C₇₈ (D_{3h})-fullerene. It consists of at least two isomers. Common impurities are C₈₂ and C₈₆ fullerenes. The identities are confirmed by an HPLC/GPC system with Waters 600E UV/VIS detection, mass and NMR spectroscopy. [Seleque et al. In Kadish and Ruoff (Eds) '*Fullerenes: Recent Advances in the Chemistry and Physics of Fullerenes and Related Materials*' The Electrochemical Soc. Inc, Pennington, NJ, 1994 ISBN 1566770823, Diederich & Whetten *Acc Chem Res* **25** 119 1992, DOI: 10.1021/ar00015a004; Diederich et al. *Science* **254** 1768 1991, DOI: 10.1126/science.254.5039.1768; for '*IR fingerprints of nine C₈₂ fullerene isomers: a semi-empirical prediction*' see Orlandi, Zerbetto and Fowler [*J Phys Chem* **97** 13575 1993, DOI: 10.1021/j100153a026].

Higher Fullerenes, e.g. **C₃₉₆ fullerene** [175833-78-0] have also been isolated [*Chem Abstr* **124** 299339 1996.], and for '*Competition between Even and Odd Fullerenes: C₁₁₈, C₁₁₉ and C₁₂₀*', see P.W. Fowler, T. Heine and F. Zerbetto [*J Phys Chem* **104** 9625 2000, DOI: 10.1021/jp0019815].

For '*Structural Motifs and the Stability of Fullerenes*' see Austin and Fowler et al. [*J Phys Chem* **99** 8076 1995, DOI: 10.1021/j100020a035].

FUNCTIONALISED FULLERENES.

A large number of functionalised fullerenes have been reported. They are synthesised by chemical reactions at the double bonds of the 'ball' to form anchors for further condensation with small or large molecules such as proteins, lipids, DNA etc. They find applications in a variety of industries including drugs and pharmaceutical industries and in medicine. They are of nano size, can circulate in the animal's body and can be used to deliver substances to targeted tissues. A few of these are commercially available in chemical catalogues and a few are described here to show how fullerenes can be made to react with reagents.

1,4-Bis(pentafluorobenzyl)[C₆₀]fullerene {7,8-dihydro-7,8-bis[(2,3,4,5,6-pentafluorophenyl)methyl]-[5,6]-fullerene-C₆₀-I_h} [1260376-31-5] **C₆₀(C₆F₅CH₂)₂, M 914.6**. This bifunctional [C₆₀]fullerene was prepared in an argon atmosphere by adding potassium metal (124mg, 3.19mmol) in one portion to a freeze-thawed degassed mixture of fullerene (1.0g, 1.93mmol) and 1-methylnaphthalene (5.93g, 4.17 mmol, 30 equivalents, see [90-12-0]) in THF (150ml). This produced a dark red solution after stirring at ~25° for 3 hours under argon. Pentafluorobenzyl bromide (3.63g, 13.9mmol, tenfold excess) is then added, stirring is continued for 8 hours, the reaction is stopped by addition of aqueous NH₄Cl (0.5ml), the mixture is concentrated *in vacuo* down to ~10ml and the crude desired product is precipitated by addition of MeOH. It is purified by chromatography through silica gel and eluting first with CS₂/hexane (1:1) then CS₂ to provide the **bis-perfluorobenzyl fullerene** (830mg, 55%) which is fully characterised by its spectroscopic properties. It is stable in air and in solution (e.g. CHCl₃). For further purification preparative HPLC or GPC may be necessary. It is identified by investigations of dynamic light scattering (DLS), X-ray diffraction (XRD), differential scanning calorimetry (DSC), thermogravimetric analysis (TG-DTA), and of its electrochemical (OVP) properties. The data showed that the perfluoroaromatic rings interact in a face-to-face π - π manner with the [C₆₀]fullerene surfaces and exhibit unique aggregation-deaggregation behaviour. These properties allow this fullerene to be used for **high-performance organic photovoltaic devices**. [see Y. Matsuo and coworkers in Li et al. *Chem Commun* **46** 8582 2010, DOI: 10.1039/C0CC03028G; Matsuo et al. *Synfacts* **2** 148 2011, DOI: 10.1055/s-0030-1259252, Darwish *Annu Rep Prog Chem, Sect A: Inorg Chem* **107** 473 2011, DOI: 10.1039/C1IC90014E.] **Note** that the name within the chain brackets above is the name given by the *Chemical Abstract Service*.

tert-Butyl (1,2-methanofullerene C₆₀)-61-carboxylate [6,6-closed 1,2-dihydro-1,2-methanofullerene C₆₀]-61-carboxylic acid tert-butyl ester [150493-29-1] C₆₀[=CHCO₂C(CH₃)₃]₁, M 834.8. The crude *tert*-butyl ester was prepared as for the ethyl ester below and obtained in 25% yield as a 1:1:3 mixture of **6,6-closed**, **trans-5,6-open** and **cis-5,6-open esters**. As for the diethyl ester below, the mixture (20mg) was equilibrated in boiling toluene to give pure **6,6-closed-tert-butyl ester** (15mg 75%) which held toluene very strongly. It has UV/VIS with λ_{max} nm(ε) in toluene at 334 (40250), 406 (*sh* 3850), 429 (1800), 503 (1150), 593 (*br* 750); the ¹H NMR (500 MHz, in *C₆D₆) has δ at 4.38 (s 1 H), 1.51 (s, 9 H); and for IR-DRIFT, ¹³C NMR and FAB-MS see references. [Isaacs et al. *Helv Chim Acta* **76** 1231 1993 DOI: 10.1002/hlca.19930760310; Isaacs & Diederich *Helv Chim Acta* **76** 2454 1993, DOI: 10.1002/hlca.19930760704.]

Diethyl (1,2-methanofullerene C₆₀)-61,61-carboxylic acid [6,6-closed-1,2-dihydro-1,2-methanofullerene C₆₀]-61,61-carboxylic acid diethyl ester [155679-98-4] C₆₀{=C=[CO₂C(CH₃)₃]₂}₂, M 878.8, m >270°. If the ¹H NMR spectrum indicated the presence of isomeric esters then dissolve it in chlorobenzene (0.2g in 100ml) and heat it under reflux for 24 hours to convert the isomers to the more stable **[6,6-closed]-ester**. Column chromatography on silica and elution first with toluene/hexane (1:1) then toluene followed by recrystallisation from CHCl₃/MeOH and drying (60°/0.1 Torr, 6 hours) provided the desired diester as a dark solid [R_F: SiO₂/toluene is 0.50]. Store it at 2-8°. The ¹H NMR (200MHz; CDCl₃) has δ_H at 4.75 (q, *J* = 7Hz) and 1.53 (t, 6H, *J* = 7Hz); and for IR and ¹³C NMR see references. The **di-tert-butyl ester** was similarly purified (see above). [Bingel *Chem Ber* **126** 1957 1993, DOI: 10.1002/cber.19931260829; Diederich et al. *JCS Perkin Trans II* 391 1994, DOI: 10.1039/P29940000391.]

Diethyl (1,2-methanofullerene C₇₀)-71,71-carboxylate [6,6-closed-1,2-dihydro-1,2-methanofullerene C₇₀]-71,71-carboxylic acid diethyl ester [153218-95-2] C₇₀{=C=(CO₂CH₂CH₃)₂}₂, M 998.9. It was prepared by reaction of diethyl bromomalonate with fullerene C₇₀ in the presence of DBU in toluene (4 hours, ~25°), filtered and purified by chromatography through Kieselgel (0.063–0.2mm) by eluting with a gradient of toluene/*iso*-hexane (1/1 to 4/1). This gave the **diester** in 60% yield [R_F: SiO₂/toluene is 0.50], with a 40% recovery of the fullerene. Store it at 2-8°. The ¹H NMR (360MHz; CDCl₃) has δ_H at 4.75 (q, *J* = 7.1Hz) and 1.46 (t, 6H, *J* = 7.1Hz); and for ¹³C NMR see references. [Bingel *Chem Ber* **126** 1957 1993, DOI: 10.1002/cber.19931260829.]

1,2-Dihydro-[1,2]fullereneC₆₀[3,4]pyrrolidin-1-ylethoxyethoxyethylammonium bis-trifluoroacetate {2-[2-[2-(2'*H*-[5,6]fullereno-C₆₀-I_h-[1,9-*c*]pyrrol-1'(5'*H*)-yl)ethoxy]ethoxy]ethanamine bis-2,2,2-trifluoroacetate} [C₆₀=CH]₂NH⁺-CH₂CH₂OCH₂CH₂OCH₂CH₂NH₃⁺ · 2CF₃COO⁻, C₇₀H₂₀O₆N₂F₆] [356066-52-9 for mono(trifluoroacetate)] M 1123.0. *N*-Boc-aminoethoxyethoxyethylaminoacetic acid (423mg, 1.38mmol, see below in SWCNT[=CH]₂N-CH₂CH₂O-CH₂CH₂O-CH₂CH₂NH₃⁺ · HCl]_n), paraldehyde (456mg, 3.45mmol) and fullerene C₆₀ (500mg, 0.69mmol) in toluene (300ml) are refluxed for 1 hour. After cooling to ~25° the mixture is applied onto a column of silica gel (NM Kieselgel 60, 70-320 mesh) and eluted with 9:1 toluene/EtOAc to give the **analytically pure** (elemental C, H, N) C₆₀=*N*-Boc-pyrrolidinyl derivative (212mg, 0.29mmol, 31%). The UV-VIS (cyclohexane) has λ_{max} at 254, 309, 429, 466sh, 546sh, 703 nm; the ¹H NMR (200 MHz, CDCl₃, TMS) has δ at 5.08 (bs, 1H), 4.50 (s, 4H), 4.05 (t, *J* = 5.5Hz, 2H), 3.77 (m, 4H), 3.60 (t, *J* = 5.3Hz, 2H), 3.36 (m, 4H), 1.44 (s, 9H); and for IR-DRIFT, ¹³C NMR and ES-MS see references.

The *N*-Boc group is removed by treating the pure C₆₀=*N*-Boc-pyrrolidinyl derivative (150mg, 0.14mmol) in CH₂Cl₂ (3ml) with CF₃COOH (3ml), and stirring for 3 hours. The solvent is removed *in vacuo*, the residue is washed with toluene and dried *in vacuo* to give the **analytically pure** (elemental C, H, N) functionalised title **fullerene C₆₀=*N*-pyrrolidinyl bis-trifluoroacetate salt** (168mg, 0.15mmol, ~99%). The UV-VIS (cyclohexane) has λ_{max} at 331, 430, 484sh, 684 254, 309, 429, 466sh, 546sh, 703 nm; the ¹H NMR (200 MHz, DMSO-*d*₆, TMS) has δ at 7.81 (brs, 3H), 4.58 (s, 4H), 3.93 (t, *J* = 5.5Hz, 2H), 3.71-3.43 (m, 6H), 3.32 (m, 2H), 2.95 (m, 2H); and for IR, ¹³C NMR and ES-MS see references. Related compounds with a two carbon atoms and a thirteen carbon atoms chain between the pyrrolidine nitrogen atom and the terminal amino nitrogen atom have also been prepared. The ω-primary amino groups at the end of the chains were derivatised (e.g. with 12-acetylsulfanildodecanoic acid in the presence of EDCI and HOBt for **self-assembled monolayers (SAMs) purposes**, or various **fluorescent** indole-2-carboxylic acids) by standard procedures. Useful applications for nano materials science and nano medicinal chemistry (e.g. **by linking to DNAs or other biological materials**) have been made possible with such fullerene precursors by M. Prato and coworkers. [Kordatos et al. *J Org Chem* **66** 4915 2001, DOI: 10.1021/jo015608k.]

Ethyl (1,2-methanofullerene C₆₀)-61-carboxylate [6,6-closed 1,2-dihydro-1,2-methanofullerene C₆₀]-61-carboxylic acid ethyl ester [50493-27-9] C₆₀(=CHCO₂CH₂CH₃), **M 806.8**. Fullerene C₆₀ reacts with an equimolar amount of ethyl diazoacetate in boiling toluene for 7 hours to provide a 30% yield of a 1:1:3 mixture of **6,6-closed: trans-5,6-open : cis-5,6-open tert-butyl** (1,2-methanofullerene C₆₀)-61-carboxylate. This mixture was isolated by applying the residual solid from the reaction onto a silica gel column which was first eluted with hexane to remove fullerene C₆₀ then with toluene-hexane (1:1) to provide the mixture at R_F ca 0.43. The mixture was isomerised almost entirely to the **more stable 6,6-close ester** by refluxing it (35mg) in toluene (105ml) for 24 hours when the initial purple solution turned pink in colour. Chromatography on silica gel as before provided a pink-red product eluting at the same R_F ca 0.43 which was then evaporated to dryness. The black microcrystalline residue was washed with Et₂O and dried at 25°/0.1 Torr (31mg, 89%). It holds residual toluene tenaciously. It has UV/VIS with λ_{max} nm(ε) in toluene at 331 (40250), 395 (sh 5250), 404 (sh 3300), 417 (1900), 429 (2250), 495 (br 1250); the ¹H NMR (400 MHz, in *C₆D₆) has δ at 4.31 (s 1 H), 4.10 (q, J = 7.1Hz, 2 H), 1.06 (t, J = 7.1Hz, 3 H); the elemental analysis is consistent with C₆₂H₆O₂, and for IR, ¹³C NMR and FAB-MS see references. [Isaacs et al. *Helv Chim Acta* **76** 1231 1993, DOI: 10.1002/hlca.19930760310; Isaacs & Diederich *Helv Chim Acta* **76** 2454 1993, DOI: 10.1002/hlca.19930760705.]

Fullerene C₆₀-γ-lactone esters. G-W. Wang and coworkers [Li et al. *Org Lett* **12** 4896 2010, DOI: 10.1021/ol102056k] have improved the condensation yields of diethyl malonate esters with [C₆₀] fullerene to form the disubstituted fused γ-lactone esters by using Fe(ClO₄)₃ instead of Mn(OAc)₃ for assistance (catalysis). Although the yields are still low, they are however, considerably higher when mediated by the iron perchlorate, and the products are cleaner. The solvent is *o*-dichlorobenzene and the ratios of reagents are 1:2:2:20 for [C₆₀]fullerene: EtOCOCHRCOOEt: Fe(ClO₄)₃:Ac₂O. The following results were obtained: **R** ester (reaction temperature, reaction time, **yield** of γ-lactone ester, consumed fullerene): **Me** (80°, 20 minutes, **34%**, 55%); **Et** (80°, 20 minutes, **37%**, 71%); **PhCH₂** (80°, 30 minutes, **27%**, 61%); **Ph*** (0°, 180 minutes, **12%**, 67%); **Br** (80°, 30 minutes, **22%**, 56%); and **EtOCO** (110°, 20 minutes, **16%**, 53%); * reactant ratio was 1:2:2:50. The structures of the lactones are consistent with their HRMS, ¹H NMRs, ¹³C NMRs, FT-IRs and UVs. It should be possible to convert these lactones into hydrofullerenes, fullerene hemiacetals, fullerene hemiketals and fullerenols. [Li et al. *Synfacts* **1** 38 2011, DOI: 10.1055/s-0030-1259158.]

Fullerene C₆₀/Poly(bisphenol A)carbonate. C₆₀ is a known '**radical sponge**' [Morton et al *J Am Chem Soc* **114** 5454 1992, DOI: 10.1021/ja00039a083; McEwen et al. *J Am Chem Soc* **114** 4412 1992, DOI: 10.1021/ja00037a064]. It reacts readily with free radicals and undergoes photolysis to generate C₆₀ radical ions. Thus irradiating a mixture of C₆₀ and poly(bisphenol A)carbonate polymer (PC, [25037-45-0]) at room temperature with a conventional UV lamp, or warming the mixture at 60° with AIBN (a radical initiator, see [78-61-1]) results in fullerenated-PC. By controlling the ratio of reagents (i.e. 5.4mg C₆₀/500mg PC in 1,1,2,2-tetrachloroethane), a **C₆₀/PC** can be obtained with a C₆₀ content as high as 6.3% (i.e. >2 C₆₀ per PC chain) in ~99% yield. Multi-additions of PC did not lead to heavy cross-linking. The fullerenated polymer is a brown powder which is soluble in common organic solvents such as THF and CHCl₃. When the polymer is dissolved in THF (a solvent that does not dissolve C₆₀) and the solution is precipitated into hexane through a filter, no particles are left on the filter, and the hexane supernatant is colourless. This shows that all the C₆₀ is incorporated into the polymer. The precipitate is then collected and dried under high vacuum to constant weight. The C₆₀ content is 1.16 wt%, has M_n ~27,000; the UV (0.3mg/ml in THF) has λ_{max} at 238.9, 259.4, 262.0, 287.2, 329.5sh nm, and for IR see references. This fullerenation provides a versatile synthetic tool for making processable fullerene polymers. [Tang et al. *Macromolecules* **30** 2848 1997, DOI: 10.1021/ma961731f.]

(1,2-Methanofullerene C₆₀)-61-carboxylic acid [6,6-closed-1,2-dihydro-1,2-methanofullerene C₆₀]-61-carboxylic acid [155116-19-1] C₆₀(=CHCO₂H), **M 778.7**. Attempted hydrolysis of the ethyl ester (see [155679-98-4] above) was unsuccessful, however, treatment of the **6,6-closed-ethoxycarbonylmethyl ester** with BBr₃ in *C₆H₆ under N₂ and stirring for 9 hours gave an 82% yield of the acid. Similarly the **tert-butyl ester** (see [150493-29-1] above), but in refluxing toluene for 8 hours, gave a 77% yield of the acid. After hydrolysis, the acid was purified by dissolving it in CHCl₃/Me₂SO, re-precipitating with hexane and drying overnight at 25°/0.1 Torr, or at 60°/0.1 Torr for 12 hours. The brown or black solid **6,6-closed-acid** was mostly insoluble in common solvents and slightly soluble in bromobenzene and 1,2-dichlorobenzene. This 6,6-closed acid (i.e. cyclopropane acid) has IR (KBr) with strong bands ν_{max} at 525, 699, 810, 847, 1013, 1426, 1785, 1794 cm⁻¹ and twenty other medium to weak bands. The ¹H NMR (300 MHz, in CHCl₃/Me₂SO, 1:1) has one signal for

the cyclopropane H at 5.13 ppm; the ^{13}C NMR consists of 32 signals between δ 136 and 167, and signals at δ 40.34 and 71.66. Elemental analysis is consistent with the formula $\text{C}_{62}\text{H}_2\text{O}_2 \cdot 0.5\text{Me}_2\text{SO}$.

With alcohols in the presence of dicyclohexylcarbodiimide, 1-*H*-benzotriazol-1-ol and a base [e.g. 4-(Me_2N) $\text{C}_6\text{H}_4\text{N}$ or Et_3N] in bromobenzene, the corresponding esters were obtained. Similarly peptides with **methyl glycinate** and **methyl-L-phenylalaninate** were obtained by using the corresponding amino acid esters. [Isaacs & Diederich *Helv Chim Acta* **76** 2454 1993, DOI: 10.1002/hlca.19930760705; Diederich et al. *Chem Soc Rev* **23** 243 1994, DOI: 10.1039/CS9942300243.]

6,9,12,15,18-Pentamethyl-1,6,9,12,15,18-hexahydro($\text{C}_{60}\text{-I}_h$)fullerene [244229-54-7] $\text{C}_{60}[\text{H}, (\text{CH}_3)_5]$, **M 796.7**. Preparation of these substituted fullerenes should be carried out in Schlenk-type equipment under an inert atmosphere (N_2 or argon), in dry solvents and reagents (at least in the early stages of the reactions), and any oxygen or air in solvents should be flushed out by bubbling dry inert gas through them. Microcrystalline [C_{60}] fullerene (2.0g, 2.78mmol, [99685-96-8]) in 1,2-dichlorobenzene (90ml), cooled in an ice-water bath, stirred under reduced pressure (1mm) to remove dissolved O_2 , and warmed to $\sim 25^\circ$, is added during 15 minutes to a stirred mixture of $\text{CuBr} \cdot \text{SMe}_2$ (6.84g, 33.3mmol, use an efficient fume cupboard) in THF (47ml) at $\sim 25^\circ$ to which is previously added MeMgBr in THF (3 M, 11.1ml, 33.3mmol) followed by DMI (3.62ml, 33.3mmol, see [80-73-9], 1,3-dimethyl-2-imidazolone) and rapidly warmed to $\sim 35^\circ$ within 5 minutes. The colour of the white suspension soon turned to dark brown. After stirring at 35° for 40 minutes under a flow of N_2 (**note** that some liberation of ethane may occur), a degassed saturated aqueous NH_4Cl solution (3.0ml) is added *via* a syringe, the colour of the solution changes from dark brown to red-brown and the mixture is stirred under a vacuum (*ca* 1mm) at $\sim 25^\circ$ to remove THF and Me_2S and reduce it to half its volume. This is diluted with degassed toluene (200ml) and subjected to silica gel flash chromatography (45 x 200mm size, using 90g of silica gel 230-450 mesh) with toluene as eluant (total volume 100ml). It should be done as rapidly as possible, as care should be taken to avoid oxygen and formation of $\text{C}_{60}\text{Me}_5\text{O}_n\text{H}$ ($n = 2-3$). The vermilion eluate is evaporated at $\sim 40^\circ/10\text{mm}$ then at 80° (to remove 1,2-dichlorobenzene) until solid begins to appear on the sides of the flask. N_2 is allowed to enter the evacuated flask and degassed MeOH is added along the inside wall of the flask, whereby the mixture becomes cloudy, and MeOH ($\sim 400\text{ml}$ total) is added until precipitation is complete. The solid is filtered off, washed with MeOH (5 x 10ml) and hexane (3 x 10ml) and dried *in vacuo* (1mm) to give red microcrystals of pentamethylfullerene (2.08g, 94%) of $\sim 91\%$ purity. Purity and/or purification is carried out using a Buckyprep or ODS column (4.6 x 150mm, flow rate 1ml/minute) and eluting with toluene/*i*-PrOH (7:3) or (3:7) respectively. The solid should be stored in an inert atmosphere, as on storage in air it deteriorates slowly over several months, and in solution in the presence of air its purity decreases by 80% in 24 hours. It has ^1H NMR (500 MHz, $\text{CS}_2/(\text{CD}_3)_2\text{CO}$ 5%) has δ at 2.34 (s, 6H), 2.35 (s, 6H), 2.46 (s, 3H), 4.50 (s, 1H, $\text{C}_{60}\text{-H}$); and for IR and ^{13}C NMR see references. [Matsuo, Mueamatsu, Tahara, Koide and Nakamura *Org Synth* **83** 80 2006, DOI: 10.15227/orgsyn.083.0080; Matsuo et al. *Functional Organic Materials* 58-80 2007, ISBN: 978-3-527-31302-0.]

The procedure described above is unique as all five substituents are symmetrically placed on the bridgehead carbons of five six-membered rings surrounded by a fused five membered cyclopentadiene ring (X-ray evidence). Similar reactions also produce very high yields of fullerenes with *five* C_6H_5 , *five* $4\text{-CF}_3\text{C}_6\text{H}_5$, *five* $4\text{-MeOC}_6\text{H}_4$, *five* $4\text{-ClC}_6\text{H}_5$, *five* $4\text{-PhC}_6\text{H}_5$, *five* (*E*)-1-propenyl, *five* (*Z*)-1-propenyl and *five* (*E*)-2-phenylethenyl groups all symmetrically placed as in the pentamethyl derivative above. [see Matsuo, Mueamatsu, Tahara, Koide and Nakamura *Org Synth* **83** 80 2006, DOI: 10.15227/orgsyn.083.0080; and references therein.] The hydrogen atom of the central cyclopentadiene ring can be displaced by potassium with *t*-BuOK to form the **$K(\text{C}_{60}\text{Me}_5)$ complex** in which the potassium atom can then be displaced to form iridium complexes such as **$\text{Ir}(\eta^5\text{-C}_{60}\text{Me}_5)(\text{CO})_2$** [Matsuo, Iwashita and Nakamura *Organometallics* **24** 89 2005, DOI: 10.1021/om0493331], and with rhodium to form **$\text{Rh}(\eta^5\text{-C}_{60}\text{Me}_5)(\text{CO})_2$** [Sawamura, Kuninobu and Nakamura *J Am Chem Soc* **122** 12407 2000, DOI: 10.1021/ja005564h], structures which are supported by X-ray structure analyses. The same cyclopentadiene can complex with Fe and cyclopentadiene (Cp) to form a hybrid of '**buckyferrocene**' **$\text{Fe}(\text{C}_{60}\text{Me}_5)\text{Cp}$** [Sawamura et al. *J Am Chem Soc* **124** 9354 2002, DOI: 10.1021/ja026069j; Nakamura *Pure Appl Chem* **75** 427 2003].

***N*-Tosyl[1,2]-aziridino[C_{60}]fullerene** {2a[(4-methylphenyl)sulfonyl]-2a-aza-1,2(2a)-homo[5,6]fullerene- $\text{C}_{60}\text{-I}_h$ [226909-63-3] $\text{C}_{60}[\text{=N-(p-C}_6\text{H}_4\text{CH}_3)]$, **M 889.0**. This useful fullerene precursor is readily prepared. In a dry Schlenk flask with flushing argon is added CuCl (1.2mg, 12 μmol), *o*-dichlorobenzene (40ml) and 2,6-toluidine (2.0 μL , 24 μmol), then stirring at $\sim 25^\circ$ for 30 minutes, followed by adding C_{60} fullerene (432mg,

600 μ mol) and ***TsN=IPh** (224mg, 600 μ mol, see synthesis below*). The mixture is stirred $\sim 25^\circ$ for 12 hours and purified by flash chromatography through silica gel (toluene/hexane 1.2 \sim 1.1) to provide the pure aziridinofullerene (228mg, 43%) as a dark brown solid [and recovered 185mg, 43%, of unreacted C₆₀ fullerene]. It has ¹H NMR (400 MHz, CDCl₃/CS₂ = 1:1, TMS) with δ at 2.56 (s, 3H), 7.50 (d, *J* = 8.3Hz, 2H), 8.19 (d, *J* = 8.3Hz, 2H); for the ¹³C NMR and the HRMS (ESI-TOF; negative) see references. **Note** that other Cu catalysts and other pyridine bases resulted in lower yields of this reaction. Itami and coworkers [Nambo et al. *J Am Chem Soc* **133** 2402 2011, DOI: 10.1021/ja111213k] developed the use of this aziridinofullerene as a versatile platform for preparing a variety of functionalised fullerenes. Thus it reacts with aryl and heteroaryl compounds (5% TfOH, 1,2-Cl₂C₆H₄, 100°, 12 hours) to replace the 1,2-aziridine by two aryl or heteroaryl groups to form 1,4-disubstituted fullerenes in over 80% yields; it reacts with bifunctional nucleophiles of the type R-Z(**H**)=Y(X-**H**)-R' (10% TfOH, 1,2-Cl₂C₆H₄, 100°, 12 hours) to form 1,2-disubstituted fullerenes (1,2-disubstituted fullerenes fused with a 5-membered ring where the **H**'s are replaced by bonds with C1 and C2 of the fullerene), such as **2,3-furano**, **2,3-pyrrolo**, **2,3-thiazolo**-, **1,2-cyclic[1,3,2]-dioxaborolan-** fullerenes in over 60% yields; it reacts with **2,2'-bithiophenes** in various ways, and undergoes formal [2+2] cycloaddition reactions with **1,2-[bis(*p*-methoxyphenyl)]acetylene** (10% TfOH, 1,2-Cl₂C₆H₄, 100°, 12 hours) to form a **3,4-adduct (cyclobutene)** which retains the **1,2-aziridino** moiety in 81% yields. Some of the structures were supported by X-ray crystal structure analysis.

***[N-(*p*-Toluenesulfonyl)imino]phenyliodinane (TsN=IPh)** [55962-05-5] C₇₀(C₁₃H₁₂INO₂S, **M 373.2, m 102-104° (dec)**). Toluene-*p*-sulfonamide (3.42g, 20mmol) and diacetoxyiodobenzene (6.4g, 20mmol, see [2340-34-4]) are added to a solution of KOH (2.8g, 50mmol) dissolved in MeOH (80ml, or better use CH₂Cl₂) at 0°, and then stirred at 25° for 3 hours. The mixture is poured into distilled water at 0°, kept at 4° overnight, the solid is filtered off, drained and recrystallised from hot MeOH (20ml) to give TsN=IPh (2.6g, 35%). **Note** the reagent reacts slowly with DMSO-*d*₆ and MeOH, so allow minimum contact time with these solvents (work rapidly). It has ¹H NMR (200 MHz, DMSO-*d*₆, TMS) with δ at 7.80-7.73 (m, 2H), 7.56-7.48 (m, 3H), 7.40-7.31 (m, 2H), 7.15-7.10 (m, 2H), 2.34 (s, 3H). [Heuss et al. *Inorg Chim Acta* **342** 301 2003, DOI: 10.1016/S0020-1693(02)01145-3; Besenyi et al. *Tetrahedron Lett* **34** 6105 1993, DOI: 10.1016/S0040-4039(00)61740-9; Evans & Barnes *e-EROS Encyclopedia of Reagents for Organic Synthesis* Published Online 15 APR 2001, DOI: 10.1002/047084289X.rt139].

2,5,10-Triphenyl-2,5,6,10-tetrahydro(C₇₀-I_h)fullerene [244237-40-9] C₇₀(C₆H₄)₃H, **M 1073.0**. When a procedure similar to the preparation of C₆₀Me₅H above is applied to fullerene-C₇₀ only **three substituents are inserted**. A suspension of CuBr.SMe₂ (374mg, 1.82mmol, 30 equivalents) in THF (23.0ml) is treated with PhMgBr (0.98M, 1.86ml, 1.82mmol, 30 equivalents) and stirred at 28° for 20 minutes. To the resulting yellow suspension is added a degassed solution of C₇₀ (49.6mg, 59 μ mol) in 1,2-dichlorobenzene (25ml) and stirring is continued for 24 hours. The reaction is treated with 5% aqueous HCl, the organic layer is washed with H₂O, brine, dried (MgSO₄), evaporated to a small volume and diluted with MeOH to give a dark brown precipitate which is washed thoroughly with MeOH, Et₂O, H₂O again and dried *in vacuo* to give **C₇₀Ph₃H** (61.2mg, 95%, 96% purity, *cf* C₆₀Me₅H above). The ¹H NMR (400 MHz, CDCl₃) has δ at 7.8-7.78 (m, 2H), 7.74-7.70 (m, 2H), 7.61-7.57 (m, 3H), 7.39-7.23 (m, 8H), 4.43 (s, 1H); and for the ¹³C NMR and the APCI-MS see references. Similarly prepared are **C₇₀Me₃H**, **C₇₀(4-PhC₆H₄)₃H**, **C₇₀(4-ClC₆H₄)₃H**, and **C₇₀(1-naphthyl)₃H**. [Sawamura et al. *J Mater Chem* **12** 2109 2002, DOI: 10.1039/B202130G; *cf* Matsuo, Mueamatsu, Tahara, Koide and Nakamura *Org Synth* **83** 80 2006, DOI: 10.15227/orgsyn.083.0080; and references therein.] **C₇₀(4-CF₃C₆H₄)₃H** is also prepared similarly and is believed to proceed by formation of [(4-CF₃C₆H₄)₂Cu⁻] from the reaction between the arylMgBr and CuBr.SMe₂ which attacks successively three six-membered rings around a cyclopentadiene ring to give the **intermediate C₇₀(4-CF₃C₆H₄)₃Cu-Ar** which decomposes on addition of acid or NH₄Cl to provide the desired **C₇₀(4-CF₃C₆H₄)₃H**. When the latter, in THF-*d*₈, is treated with 1 equivalent of *t*-BuOK or TIOEt at 25°, the colour changes from reddish-brown to dark red to provide **K[η^5 -C₇₀(4-CF₃C₆H₄)₃]** {whose ¹H NMR (400 MHz, THF-*d*₈) has δ at 8.07-8.05 (m, Ar-H, 6H), 7.57-7.54 (m, ArH, 6H); and for the ¹³C NMR see references. For **Tl[η^5 -C₇₀(4-CF₃C₆H₄)₃]** {whose ¹H NMR (400 MHz, THF-*d*₈) has δ at 8.09-8.07 (m, Ar-H, 6H), 7.59-7.53 (m, ArH, 6H)} also see Sawamura et al. [*J Am Chem Soc* **120** 8285 1998, DOI: 10.1021/ja981256w].

1,4,11,15,30-Pentakis(4-hydroxyphenyldimethylsilylmethyl)-2H-1,2,4,11,15,30-hexahydro-(60)fullerene {[4-HOC₆H₄Si(Me)₂CH₂]₅-2H-1,2,4,11,15,30-hexahydro-(C₆₀-I_h)fullerene, 1,7,8,11,24,27-hexahydro-1,7,11,24,27-pentakis(dimethyl 4-hydroxyphenylsilyl)methyl-[5,6]fullerene-C₆₀-I_h} [658080-04-7] [4-HOC₆H₄Si(Me)₂CH₂]₅- (6H)- C₆₀, M 1549.1. This functionalised fullerene can be used as an example of a molecule to which *long chains can be coupled* for specialised purposes. Typically, using Schlenk equipment under N₂ or argon in strict absence of H₂O, the protected Grignard compound Me₂(4-tetrahydro-pyranyloC₆H₄)SiCH₂MgCl (0.656M in THF, 13.0ml, 9.84mmol) is added to a purple suspension of a white powder of CuBr.SMe₂ (3.00g, 14.6mmol) and microcrystalline [C₆₀]fullerene (400mg, 0.555mmol) in 1,2-dichlorobenzene (80ml), and stirred at 25° for 1 hour. The mixture is treated with saturated aqueous NH₄Cl (0.5ml), concentrated *in vacuo* to ca 60ml and diluted with toluene. The brown mixture is applied onto a silica column and washed with toluene (~100ml, i.e. until the effluent becomes yellow), followed by EtOAc/toluene (3:97) and the effluent is collected until its colour is pale yellow. This is evaporated *in vacuo* to a small volume and MeOH is added rapidly to precipitate a brown powder which is collected under N₂, dissolved in MeOH/toluene (1:1, 46ml) and TsOH.H₂O (92mg, 0.48mmol) is added to it. After stirring for 1 hour at 25°, the mixture is neutralised with NaHCO₃, the solvent is evaporated, the residue is dissolved in MeOH/toluene (1:9, ~30ml), filtered from some insoluble material and evaporated *in vacuo*. The crude residue is purified by preparative HPLC (Nomura Chemical RPFullerene, with toluene/MeOH 1:9 as eluant) to give on evaporation the *desired fullerene* as an orange powder (99.5mg, 0.064mmol, 12% yield). Its ¹H NMR (400 MHz, acetone-*d*₆, TMS) has δ at 7.32-7.28 (m, 8H), 7.23 (d, *J* = 8.0Hz, 2H), 6.69-6.64 (m, 10H), 4.55 (s, 1H), 2.38 (s, 2H), 2.31 (d, *J* = 14.4Hz, 2H), 2.23 (d, *J* = 14.4Hz, 2H), 2.11 (d, *J* = 14.8Hz, 2H), 2.05 (d, *J* = 14.8Hz, 2H), 0.36 (s, 6H), 0.35 (s, 6H), 0.34 (s, 6H), 0.33 (s, 6H), 0.27 (s, 6H); and for ¹³C NMR and HRMS (APCI+) see references. This fullerene now has *five phenolic groups for condensation* with lipid molecules. When a mixture of it (9.40mg, 6.07μmol) and 4-(dodecan-1-yloxy)benzoyl chloride (13.0mg, 9.81μmol) in THF (3.0ml) is treated with Et₃N (5.8 μL, 42mmol) and 4-dimethylaminopyridine (3.5mg, 29μmol), stirred at 25° for 3 hours, evaporated *in vacuo*, and the residue in toluene is filtered through a silica gel pad, and then purified by GPC (JAIGEL 3H, eluting with toluene), precipitated with MeOH/toluene as above and dried *in vacuo* (at 25° for 12 hours), it provided **1,4,11,15,30-pentakis(4-[4-(dodecan-1-yloxy)benzoyloxy]phenyldimethylsilylmethyl)-2H-1,2,4,11,15,30-hexahydro-(60)fullerene** (12.9mg, 71% yield). Similarly the [4-tetradecan-1-yloxy] (75% yield), [4-hexadecan-1-yloxy] (72% yield), [4-octadecan-1-yloxy] (61% yield) and [3,4-di(octadecan-1-yloxy)] (37% yield) *derivatives* were obtained in high purity and characterised by ¹H NMR, ¹³C NMR and APCI MS analysis. These form beautifully coloured liquid crystals and X-ray diffraction studies showed interesting stacking of these fullerene molecules. [Matsuo, Muramatsu, Hamasaki, Mizoshita, Kato and Nakamura *J Am Chem Soc* **126** 432 2004, DOI: 10.1021/ja038816y.]

[Further reading: H.W. Kroto et al. *Chem Rev* **91** 1213 1991, DOI: 10.1021/cr00006a005; H. Kroto, Fischer and Cox *Fullerenes*, Pergamon Press, Oxford, 1993 ISBN 0080421520; Kadish and Ruoff (Eds) *Fullerenes: Recent Advances in the Chemistry and Physics of Fullerenes and Related Materials*, The Electrochemical Soc. Inc, Pennington, NJ, 1994, ISBN 1566770823; *Self-assembly of fullerenes*: Smalley *Acc Chem Res* **25** 98 1992, DOI: 10.1021/ar00015a001; and following papers; Hammond & Kuck *Fullerenes: Synthesis, Properties and Chemistry of Large Carbon Clusters*, American Chemical Society, Washington, 1992, ISBN 0-841221820; K. Jinno *Separation of Fullerenes by LC*, RSC Publ., 1999, ISBN 9780854045204; S. Nagase & T. Akasaki *Endofullerenes: a new family of carbon clusters*, Springer, 2003, ISBN 9781402009822; F.J.M. Reitmeijer *Natural Fullerenes and related structures of elemental carbon*, NetLibrary, 2006, eBook ID 190026, eISBN 9781402041358; P.W. Fowler & D.E. Manolopoulos *An Atlas of Fullerenes*, Dover Publications Inc, 2007, ISBN-10: 0486453626, ISBN-13: 9780486453620; P. O'Brien, H. Craighead, H.W. Kroto, F. Langa and J-F. Nierengarten *Fullerenes: principles and applications* RSC Publ., 2007, ISBN 9780854045518; *Fullerenes, Nanotubes and Carbon Nanostructures*, Marcel Dekker Inc, New York, on line series on *World Wide Web.online*; N. Chaniotakis *Fullerenes-bifunctionalisation (nanostructured for biosensing)* in 'Nanomaterials for Biosensors', C. Kumar ed., Wiley-VCH, 2007, ISBN 9783527313884; C.N. Kramer *Fullerene research advances*, Nova Science Publishers Inc, 2007, ISBN 9781600218248; M. Lang *Progress in fullerene research*, Nova Science Publishers Inc, 2007, ISBN 9781600218415; S. Margadonna *Fullerene-related materials: recent advances in their chemistry and physics*, Springer 2007, ISBN 9781402044588.] See Bibliography for further reading.

CARBON NANOTUBES (CNTs) AND RELATED MATERIALS

The development of carbon nanotubes (CNTs) since their discovery by Sumio Iijima [*Nature* **354** 56 1991, DOI: 10.1038/354056a0] has been explosive, not only because of their mechanical, thermal and electronic properties but because they can be modified physically and chemically in a variety of ways to suit a plethora of applications. Structurally, they are made of rolled up sheets of carbon atoms forming fused hexagonal carbon rings just like a monolayer of graphite. The sides of the sheet are fused to form complete tubes. When one or both ends of a tube come together, like the bottom of a test tube, the round portion is more like *half a 'fullerene ball'*. The way that the carbon atoms in the half fullerene ball '*finger tip*' structure are stable is by forming both hexagonal and pentagonal rings (see Marchand et al. *Nano Lett* **9** 2961 2009, DOI: 10.1021/nl901380u; FEM imaging below).

Many carbon nanotubes are now available commercially in chemical catalogues, and a selected number are entered here. Their chemical and biochemical applications include diagnostics, novel devices for rapid DNA sequencing, human personalised medicine, bio-labelling for cancer research, novel biomaterials for human implants, and for tissue engineering implants. The non-chemical applications have also been extended to attachments to surfaces with patterned and unpatterned textures to form conductors, and exciting applications as for nano-filters, in LEDs, nanostructured optical fibres, lasers, and solar cells. Carbon nanotubes are very strong materials and could be ~200 stronger than steel. When mixed with building materials, such as cement or aluminium composites, they not only increase their strength but make them much lighter. Carbon sheets such as graphene are conducting surfaces and have found extensive use in touch screens, i-Pods, mobile phones, bicycle frames, fishing rods, air-craft materials to name a few. [Haddon *Carbon Nanotubes (editorial)*, *Acc Chem Res* **35** 997 2002, DOI: 10.1021/ar020259h; see also P. Calvert *A recipe for strength*, *Nature* **399** 210 1999, DOI: 10.1038/20326].

PREPARATION OF CNTs (NTs)

MWCNTs (multi walled CNTs) were first prepared by the *arc discharged evaporation method* (as for fullerene synthesis) where the needles (nanotubes) grew at the negative end of the carbon electrode in an argon filled vessel (at 100Torr). TEM micrographs showed tubes of 5 concentric graphic sheets (6.7nm diameter), 2 concentric graphic sheets (5.5nm diameter, DWCNT, double walled CNTs), and 7 concentric sheets (2.2nm diameter MWCNT). Electron diffraction patterns show that the tubes were formed from rolled up graphite sheets of carbon hexagons into helical structures. [Iijima *Nature* **354** 56 1991, DOI: 10.1038/354056a.] In chemical arc deposition, acetylene (at partial pressure ~10⁻²mbar, 0.0075mm Hg at 0°) is often used as the hydrocarbon gas. J-M Bonard and coworkers prepared CNTs by growing them on 0.3mm diameter wires of Kanthal (Fe-Al-Cr alloy) in the centre of the anode at the rather low temperatures of 700-900°, hence the name **Cold Atmosphere Chemical Vapour Deposition CVD** (CACVD, compare below). The technique produces homogeneous films of well-graphitised NTs allowing control of their length and density. [Crocì et al. *Chem Vap Deposition* **8** 89 2002, DOI: 10.1002/1521-3862(20020503)8; Bonard et al. *Appl Phys Lett* **81** 2836 2002, <http://dx.doi.org/10.1063/1.1511539>]. MWCNTs are generally formed to a smaller or larger extent in the procedures used, together with small quantities of **buckyballs** and **amorphous carbon** which can be invariably sublimed out, annealed out, or separated by chromatography. Annealing at high temperatures (~2500-2700°) removes volatile carbon, metals (generally bound to the open end of the tubes) and improves the structural order of the tubes.

SWCNTs (single walled CNTs): Among the techniques that have been developed for making SWNTs are **EA (electric arc) methods** which use a carbon source and catalytic amounts (~ 1 to 6 atom%) of transition metals, commonly Ni/Co, Ni/Y, Rh/Pt or Fe that are allowed to supersaturate the carbon [Kokai et al. *J Phys Chem B* **103** 4346 1999, DOI: 10.1021/jp990065s; Sen et al. *Chem Phys Lett* **349** 383 2001, DOI: 10.1016/S0009-2614(01)01208-8]. The methods have been studied in detail in order to optimise the sizes and yields of CNTs. The technique developed by R.C Haddon and co-workers [Itkis et al. *J Phys Chem B* **108** 12770 2004, DOI: 10.1021/jp0487307], and the purity estimated by solution phase NIR [Itkis et al. *Nano Lett* **3** 309 2003, DOI: 10.1021/nl025926e] adopts the EA method where an electric arc discharge is applied between a graphite cathode and composite anode containing catalytic amounts of metal(s), e.g. 4:1 atom% of Ni/Y, under He buffer gas at ~680Torr, and arc current 90A with a 10mm arc gap. These conditions produce high yields of AP-SWCNT (high purity, As-Prepared nanotubes). The purity can be assessed by Near-Infrared (NIR) spectroscopy.

Large scale production of SWCNTs by the EA technique is achieved by using an arc generated between a graphite rod (16mm diameter, 40mm long) cathode and an anode of graphite (6mm diameter, 100mm long) in which a hole 3.5mm diameter, 40mm deep, is drilled and filled with a mixture of a metallic catalyst (Ni 4.2 atom%/Y1 atom% and graphite) in a He atmosphere (660mbar) formed by a current of 100A/30V and keeping a constant distance (~ 3 mm) between the electrodes. Within ~ 2 minutes, the total carbon mass was in ~ 2 g quantities consisting of $\sim 70\%$ SWCNTs in highly crystalline bundles. The nature of the product(s) was assessed by SEM, high resolution TEM, Raman spectroscopy and X-ray diffraction. [Journet et al. *Nature* (letter) **388** 756 1997.]

Another technique developed by R.E. Smalley and coworkers is the **laser ablation** of carbon targets (**Carbon Vapour Deposition, CVD**), e.g. laser-vaporised carbon-Ni/Co mixture (e.g. 1:1 1.2 atom% in C) at high temperatures (e.g. 1200°C) and vacuum (e.g. 500Torr) and an argon stream. [Thess et al. *Science* **273** 483 1996, DOI: 10.1126/science.273.5274.483, Sen et al. *Chem Phys Lett* **332** 467 2000, DOI: 10.1016/S0009-2614(00)01320-8; for the *Role of temporal delay in dual-laser ablated plumes* see Witanachchi & Mukherjee *J Vac Sci Technol A* **13** 1171 1995, <http://dx.doi.org/10.1116/1.579856>]. This CVD produces high yields ($\sim 70\%$) of SWCNTs as assessed by X-ray diffraction and electron microscopy (EM) methods at a rate of ~ 80 mg/day. The production of SWCNTs can be increased to ~ 1 g/day by using a 2" tube and dual laser pulses. However, by modifying the apparatus, using up and down scanning of the laser pulses onto a rotating carbon-Ni/Co target in a 4" diameter tube, a lower temperature (1100°C) can be used to generate 20g of 40-50vol% SWCNT material in 48 hours of continuous (largely unattended) operation [Rinzler et al. *Appl Phys A* **67** 29 1998, DOI: 10.1007/s003390050734]. The high purity was checked by EM, XRD, Raman spectroscopy and TGA. **Note** that higher temperature operations produce tubes of larger diameters, albeit well within the nanometer scale.

In a more recent, essentially CVD procedure, S.T. Purcell and co-workers [Marchand et al. *Nano Lett* **9** 2961 2009, DOI: 10.1021/nl901380u; which provide an animated supplement, and summarised by H. Birch 'Nanotube growth on camera', *RSC Chemistry World* **6**(9) 26 2009] have demonstrated by *Field Emission Microscopy* (FEM) how the growth of SWCNTs takes place. They used a W-tip connected to the electrodes (at 1200°C) in a chamber at ultra high vacuum (5×10^{-10} Torr base pressure). The W-tip (tip radius ~ 60 nm) is first covered with a graphite diffusion barrier by heating in acetylene (at $P_{C_2H_2} \sim 1 \times 10^{-4}$ Torr), and Ni nanoparticles are induced to grow directly onto the tip [by dewetting, from a circular Ni wire in the chamber using CVD in acetylene ($\sim 1-2 \times 10^{-7}$ Torr during growth) at 800°C] while FEM imaging is performed. The images are collected on camera, and a frame-by-frame analysis of the video showed that the SWCNT grows at one end of the carbon tube at a time. The tube rotates in discrete steps, ~ 24 per rotation (half of the number of carbon atoms on the circumference of a common SWCNT) with the CNT turning ~ 180 times during its 11 minutes growth (as in the '**screw-dislocation-like**' model). Growth starts at the Ni atom (of the catalyst), the carbon attaches itself to the metal then it forms carbon rings (as hexagons, because these are the more stable configuration), and the molecule rotates as the tube is being formed. In the end, the tube is sealed by forming a semi-fullerene ball '**finger tip**' structure, and produces a '**test-tube like**' nano carbon structure with the metal atom at its mouth. The metal can then be removed by heating the SWCNT strongly (annealing).

The **HiPco process** for preparing SWCNTs developed by R.E. Smalley and co-workers [Nikolaev et al. *Chem Phys Lett* **313** 91 1999, DOI: 10.1016/S0009-2614(99)01029-5] involves formation of the catalysts by *in situ thermal decomposition of Fe(CO)₅* in a heated flow of CO in the gas phase at 1-10atm and 800-1200°C. By adjusting the processing parameters (best at 1-10atm and 800-1200°C), SWCNTs were produced in high yields (79 mole%, i.e. 44 wt%) with narrowest tubes (as small as 0.7nm in diameter, same as in C₆₀ fullerene) at the rate of 1.24mg/hour. The structures were confirmed by TEM, SEM, EDX and TGA. An important advantage of this procedure is that it is a **continuous-flow process** that can be used for bulk production of narrow SWCNTs. See below for purification of HiPco tubes.

PURIFICATION OF CNTs (NTs)

MWCNTs: Raw tubes, as prepared by the EA method, contain about one third of other nanomaterial. This material is best removed by oxidation. Pure tubes can thus be prepared, although the procedure is wasteful. The purest tubes are obtained when 99% is oxidised; but when 95% is oxidised only 10-20% of product contains pure nanotubes. A ground raw sample is placed in an oven and the temperature is raised to 750° in air or oxygen for ~ 30 minutes until $\sim 1\%$ is left. This consists mostly of pure tubes with length/diameter ratios that exceed 100, and traces of open cylinders with the ratio of 20. [Ebbesen et al. *Nature* **367** 519 1994, DOI: 10.1038/367519a0; see also Chen et al. *Adv Mater* **8** 1012 1996, DOI: 10.1002/adma.19960081216.]

MWCNTs, prepared by the CVD procedure using ferrocene and xylene (0.75% Fe/C) as catalyst and precursor with argon/H₂ at ~700° [Andrews et al. *Chem Phys Lett* **303** 467 1999, DOI: 10.1016/S0009-2614(99)00282-1; (a step closer to commercialisation), Rao et al. *Appl Phys Lett* **76** 3813 2000, <http://dx.doi.org/10.1063/1.126790>], or acetylene as precursor and 2.5wt% Co-2.5wt% Fe/NaY zeolite as catalyst with argon/H₂ at ~700° [Bulusheva et al. *J Phys Chem B* **105** 4853 2001, DOI: 10.1021/jp010056v], were purified by heating the raw tubes (200mg) in 2.6M aqueous HNO₃ (40ml) for 48 hours. The cooled solution was centrifuged (use PTFE tubes, ~20,000g/30 minutes), the sediment was washed/re-centrifuged with de-ionised H₂O until the supernatant was barely acidic and the MWCNT sediment was dried *in vacuo*. This material was **satisfactory for functionalising** [Lin et al. *J Phys Chem B* **106** 1294 2002, DOI: 10.1021/jp013501v]. Alternatively, to a dispersed suspension of MWCNTs (100mg), formed by sonication (1 minute), is added an acidic solution of KMnO₄ (4g in 120ml of 1N H₂SO₄) dropwise and refluxed for 12 hours, cooled, and centrifuged. The sediment is treated with concentrated HCl (20ml) and refluxed for 24 hours to dissolve the MnO₂, cooled, centrifuged and the sediment is refluxed with 68% (azeotropic) HNO₃ (40ml) for 24 hours, cooled, and centrifuged again. The solids are washed repeatedly (using several centrifugation/washing cycles) with de-ionised H₂O until almost neutral then dried in a vacuum oven. This **procedure exposes the CO₂H groups** on the CNTs and can be **used for conversion to COCl groups**, and **further reactions** to make soluble MWCNTs. [Fu et al. *Nano Lett* **1** 439 2001, DOI: 10.1021/nl010040g.]

SWCNTs: (i) Mechanically ground cloth-like raw SWCNTs from the electric arc process (EA) can be purified in reproducibly high yields (optimally 25-30wt% and containing <1wt% of transition metal) in two steps. The **first** involves thermal annealing in air by rotating the powders at 470°C for 50 minutes in a quartz tube at 30rpm which burns the carbonaceous particles out, and the **second** is acid treatment which etches out the metals. The powder is immersed in aqueous 6M HCl for 24 hours, filtered (or centrifuged) several times until the colour of the acid is unaltered, and then washed with de-ionised H₂O. The SWCNTs are then **'unbundled'** by boiling in 30% aqueous HNO₃ for 4-6 hours, and the suspension is filtered with a polytetrafluoroethylene (PTFE) membrane in deionised H₂O, rinsed and dried to give a greyish black, thin mat of SWCNTs. The purity at each stage can be observed by SEM, TEM, FT-Raman spectroscopy and TGA which usually show ~96% purity after HCl treatment. The HNO₃ treatment, however, appears to break down the CNTs into small pieces and sometimes forms MWCNTs. [Moon et al. *J Phys Chem B* **105** 5677 2001, DOI: 10.1021/jp0102365.]

(ii) A gas-phase purification procedure developed by Margrave and co-workers [Zimmerman et al. *Chem Mater* **12** 1361 2000, DOI: 10.1021/cm990693m] involved purging the CNT sample [prepared by the pulsed laser method (Rinzler et al. *Appl Phys A* **67** 29 1998, DOI: 10.1007/s003390050734), from Graphite (1-2mm), or from EA] in a quartz tube with a mixture of Cl₂ (7.2ml/min), H₂ (2.7ml/min) and argon (3.0ml/min) for 1 hour at ~25°, which is then lowered into a furnace at 500° while the evolving gases (CO, COCl₂, CCl₄ and CO₂) are monitored by IR. Purging is continued until the CO partial pressure is <0.5Torr. The temperature of the sample is lowered to ~25°, sonicated in DMF/0.6M HCl (1:1) to remove metals, then pure DMF, filtering, and washing with MeOH to form an **SWCNT 'paper'**. (Alternatively, the metals, if present, can be removed by sublimation in HCl gas at higher temperature.) The SWCNT 'paper' is dried at 160° and the purity assessed by SEM, TGA and UV-VIS. Any fullerenes present may be removed by refluxing in toluene or by sonication. The yield is ~15wt% to give CNTs that are generally **more reactive due to their larger curvature**.

(iii) Raw HiPco SWCNTs can be purified in a multi-step process. Low density raw HiPco (~100mg, d ~0.01g/cc) is compressed onto dry filter paper (vacuum), placed in a ceramic boat and inserted into a quartz tube furnace while a gas mixture of 20% O₂ (or air) in argon (wetted by bubbling through H₂O) is allowed to flow over the sample in the furnace at 100cc/sec. The nanotubes are heated at 225° for 18 hours, sonicated for ~15 minutes (or stirring overnight) in concentrated HCl (yellow colour due to Fe³⁺ ions). The suspension is filtered onto a Teflon membrane (47mm, 1.0µm pore size), washed several times with de-ionised H₂O, MeOH, and dried in a vacuum oven at 100° for 2 hours (weight loss 33.7%; 0.67% residual metal). This oxidation with wet O₂ (or air) in argon, and extraction cycle is repeated at 325° for 1.5 hours (further weight loss 8.3%, 0.05% residual metal), and again at 425° for 1 hour (further weight loss 22.9%, 0.09% residual metal). After drying *in vacuo*, the HiPco tubes are annealed at 800° in argon for 1 hour (further weight loss 4.2%, 0.03% residual metal). Assessment of purity is by SEM, TEM, TGA, Raman and UV-VIS-NIR spectroscopy. The onset of oxidation of the smaller diameter HiPco SWCNTs is *ca* 100° lower than for the purification of large diameter tubes obtained in the laser-oven process, and is consistent with the greater steric strain present in small diameter SWCNTs. [Chiang et al. *J Phys Chem B* **105** 8297 2001, DOI: 10.1021/jp0114891.]

(iv) A **general purification procedure** for raw SWCNTs obtained by the EA process (reaching 2-3g, containing ~70% SWCNTs) [Ebbesen & Ajayan *Nature* **358** 220 1992, DOI: 10.1038/358220a0; Journet et al. (letter) *Nature* **388** 756 1997], or the laser ablation method (reaching ~20g, containing up to 50% SWCNTs) was developed by A. Hirsch and coworkers [Holzinger et al. *Appl Physics A* **70** 599 2000,]. The process involves three steps. **(a)** Raw material (100mg) was refluxed with 65% HNO₃ (150ml) for 3 hours, and centrifuged. The sediment was washed with H₂O and centrifuged repeatedly until the supernatant (first colourless then darkens, still containing some SWCNTs and was kept for further use) has pH ~1. The sediment was suspended in distilled H₂O, subjected to two and three 0.5 seconds of ultrasonic pulses when the liquid became weakly acidic (pH 3-4), almost free from metal catalyst, and contained ~1mg/ml H₂O of only SWCNTs and nanoparticles. For storage, the pH is adjusted to 8-9 with K₂CO₃ which increased the repulsive interactions between the carboxylate groups of the SWCNTs. **(b)** The suspension is sonicated for 1 minute which reduces the nanoparticles to smaller pieces. The tubes and bundles were also smashed and the difference in size between the SWCNTs and degraded particles increased (as observed by AFM). **(c)** This most important step involved elution of the suspension through a column of potassium polyacrylate which was swollen (the cross-linked polymer particle size swells from <1µm to ~300µm) creating pores large enough to entrap nanoparticles as well as SWCNTs and bundles of comparable size. [Optimum: 15ml of swollen polymer per 1ml of a 1g/L aqueous SWCNTs, and choosing a diameter to gave a column height of 6-7cm held by a glass filter pore size 2]. A vacuum was applied at the bottom of the column which squeezes the swollen polymer particles. The SWCNTs were so large that they cannot now be entrapped and eluted freely to give large quantities (> 40% of the total mass) in the first fraction. The remaining material contained a large portion of SWCNT fraction of lesser quality which can be recycled. Most of the degraded material and smaller nanoparticles remain in the cavities. The purification can be followed by AFM and the purity can be assessed by Raman spectroscopy. In the Raman spectrum, **pristine** SWCNTs have a doublet at 1546 and 1575 cm⁻¹ (of about equal intensity — **G**-lines) and a less intense band (**D**-line, *ca* 1/4.75th of the **G**-lines) at ~1342 cm⁻¹. The relative intensity of the G-lines increase as the purity increases. The broadness of the D-band may originate from disordered or carboxylated carbon formed by oxidation.

PREPARATION AND PURIFICATION OF FULLERENE PIPES (TUBES)

These are **short SWCNTs** of 100 to 300nm lengths and ~1.2 to 1.4nm in diameter which are most useful for preparing functionalised SWCNTs. They are distinct from the above **fullerenes** which are ‘bucky balls’ and are made up of tubes consisting of a network of hexagonal carbon rings like in CNTs. As prepared, the CNTs are long almost endless single walled highly convoluted, entangled, bundled tubes. R.E. Smalley and coworkers [Rinzler et al. *Appl Phys A* **67** 29 1998, DOI: 10.1007/s003390050734; Liu et al. *Science* **280** 1253 1998, DOI: 10.1126/science.280.5367.1253] have purified, and cut such material from large batches into small lengths of open-ended tubes, which form stable colloidal suspensions in the presence of detergents. These can be made to react like ordinary organic molecules to form any variety of derivatives. In a typical preparation, the raw tubes (10g) in 2.6M HNO₃ were refluxed for 45 hours, cooled, and the black solution was centrifuged in PTFE (Teflon) tubes (20,000g, 20 minutes, Sorvall R5C centrifuge). The supernatant was decanted off, the sediment was resuspended in de-ionised H₂O by vigorous shaking and the centrifuging/washing cycle repeated 3 to 4 times until the supernatant was almost neutral, although black in colour. The acid treatment not only assists in separating the bundles, but also oxidises the material, thus producing carboxylic acid groups at the ends as well as at the sides of the tubes. This makes ‘**holes**’ in the sides of the tubes and causes them to be ‘**cut**’ [cf K. Kinoshita *Carbon Electrochemical and Physicochemical Properties* Wiley, New York 1988, ISBN 978047184802-8]. The sediment as examined by SEM and TEM contains carboxylated carbon tubes and other species such as fullerenes, polycyclic aromatic sheets and cross-linked sheets edged with CO₂H groups that, by virtue of de-protonation allow mutual repulsion between the particles. The sediment (using 10g) was dispersed in aqueous NaOH (1.8L, pH 10) containing 0.5vol% of the non-ionic surfactant Triton X 100 by sonication for ~1 hour, and purified by **cross-flow-filtration** (CFF, cellulose ester M22M 600 01N, mini-Kros® Spectrum cartridge: 0.6mm diameter, 200nm pores and 0.56m² of surface area) against a buffer of aqueous NaOH (40L, pH 10) containing 0.2vol% of Triton X 100. At a head pressure of 5-6psi and flow rate of ~70ml/min the process was complete in ~10 hours. Salt was removed by washing with de-ionised H₂O (10L). Clogging occasionally occurred and was overcome by momentarily reversing the flow before continuing. Clamping off the buffer line and opening a vent allowed concentration of the SWCNTs down to ~200ml of solution. When a portion of this solution was filtered through a PTFE membrane (Millipore LS 5µ pore) using a vacuum, it deposited a thick pad of SWCNTs on the membrane. After washing with MeOH, the pad can be easily peeled

off to give a freestanding mat which was called '**Bucky Paper**' and was estimated as typically 10 to 20% yield depending on the initial raw material. TEM images of the SWCNT '**Bucky paper**' showed that the pipes still contained impurities which were removed by two further oxidising acid treatment which '**cut**' (by reacting at the sides of the tubes) and '**polished**' the tubes.

In a typical **cutting**, the purified SWCNT '**Bucky Paper**' (10mg) was suspended in a 3:1 mixture of concentrated $\text{H}_2\text{SO}_4/\text{HNO}_3$ (40ml), sonicated (55kHz) at 35° to 40° for 24 hours. The suspension was diluted with H_2O (200ml), larger tubes were collected onto a 100nm pore filter membrane (Millipore, VCTP type), and washed with 10mM NaOH solution. **Polishing** was performed by suspension in a 4:1 mixture of concentrated $\text{H}_2\text{SO}_4/30\% \text{H}_2\text{O}_2$, stirring at 70° for 30 minutes, filtering and washing again on a 100nm filter, and re-suspending at 0.1mg/ml of 0.5wt% Triton X100 in H_2O to avoid flocculation. On a larger scale **CFF purification** was carried out after each oxidation step. Finally, to obtain **best purified 'Bucky Paper'**, annealing *in vacuo* at 1200° was necessary. **Note** that in 3:1 $\text{H}_2\text{SO}_4/\text{HNO}_3$ at 70° the average cutting was at a rate of ~130nm/hour, and in 4:1 $\text{H}_2\text{SO}_4/30\% \text{H}_2\text{O}_2$ the shortening rate was at ~200nm/hour.

Characterisation of these fullerene tubes was done by **electro-deposition** [developed by Smalley and coworkers, see above references] which drives the suspended tubes onto the surface of highly oriented pyrolytic graphite surface and are scanned by AFM. The molecular nature of the tubes was demonstrated by converting their CO_2H groups to COCl , followed by reaction with $\text{H}_2\text{N}(\text{CH}_2)_{11}\text{SH}$ and exposure to 10nm gold particles. Gold (which reacts with SH) was only found on the carboxylated fullerene tubes.

SOLUBILISATION OF CNTs

CNTs, whether MWCNTs or SWCNTs, readily form bundles, and attempts to separate the bundles by means such as sonication or use of detergents have been uniformly unattractive. The carbon material is 'non-wettable' and quite insoluble in polar and non-polar liquids. Considerable effort has been expended to find means of separating the bundles and much success has been achieved by functionalising the carbon matrices, which destabilise the interaction between the fibres, but also introduce bound groups which afford solubility in organic solvents as well as aqueous solutions. These provide opportunity for purification because the partly purified CNTs are not homogeneous. The non-homogeneity is not only in the length and/or diameter, but also in their nature since during the formation of the tubes, the row of hexagonal rings do not only align to form straight tubes (with parallel hexagonal rows), but can also form spiral tubes in which the row can be out of line by one and/or more than one hexagon while forming the tube. [For the *Dislocation theory of chirality-controlled nanotube growth* see B.I. Yakobsen and coworkers *Proc Natl Acad Sci USA* **106**(8) 2506 2009, and for a review see H. Birch *RSC Chemistry World* **6**(3) 25 2009.] This provides the opportunity for a variety of **chiral CNTs**. In his respect M Zheng and coworkers [Zheng & Semke *J Am Chem Soc* **129** 6084 2007, DOI: 10.1021/ja071577k; Tu et al. *Nature* **460** 250 2009, DOI: 10.1038/nature08116; see also P. Broadwith *RSC Chemistry World* **6**(8) 28 2009] have selected short strands of DNA [8 to 12 mers, with repeating nucleotides e.g. $(\text{GTC})_2$, $(\text{GTC})_2\text{GT}$, $(\text{GTT})_3\text{G}$, $(\text{TAT})_4$ etc] which entwine around the SWCNTs (e.g. produced from HiPco, see above) of *ca* 1nm diameter with different selectivities depending on the '*n,m* chirality' of the nanotubes [e.g. (8,6), (9,4), (7,6), (6,5) etc respectively] to form barrels around the tubes. The mixture is dispersed in appropriate aqueous solutions (e.g. 0.1M NaCl, 0.1M NaCl/10% glycerol) by sonication (~60 minutes), incubated for a period of time, centrifuged (16,000g/90 minutes) and the supernatant is fractionated by Size Exclusion Chromatography (SEC) through an IEX resin (CNT-NS 1500, Biochrom, Terre Haute, IN, USA), and the various DNA-wrapped-SWCNTs are eluted with $2\times(0.3\text{M NaCl}, 0.03\text{M sodium citrate})/0.5\text{mM EDTA/pH } 7.0$ and a 0 to 1M sodium benzoate gradient. The fractions are monitored at 300nm and 425nm. The DNA interaction is electrostatic and is easily removed by thermal treatment.

Strong acids solubilise SWCNT and MWCNT bundles by protonating them causing the strands to separate, but chlorosulfonic acid is by far (~1,000 times) the best acid as shown by M. Pasquali and the late R.E. Smalley [Davis et al. *Nature Nanotechnology* **4** 830 2009, DOI: 10.1038/nnano.2009.302; summarised by P. Broadwith *RSC Chemistry World* **6**(12) 32 2009]. SWCNTs spontaneously dissolve in ClSO_3H at a concentration of as low as 0.5wt%. The Phase Diagram in strong H_2SO_4 exhibited true thermodynamic behaviour forming isotropic (I), liquid crystal (LC), crystal solvate (CS) and solid (S) phases at different concentrations of SWCNTs, ClSO_3H and H_2SO_4 . When an 8.5vol% of SWCNTs was coagulated in 96% aqueous H_2SO_4 by extrusion through an orifice while co-flowing with CHCl_3 or CH_2Cl_2 , the resulting nanotube bundles (10-100nm) combined into larger bundles to form macroscopic fibres which were easily **spun into tens of meters** of continuous fibres in a matter of minutes. **These fibres can yield a strength in excess of 320 MPa.** By

sandwiching the dope between glass plates, long ribbons of material were produced, and possibly any shapes (e.g. coils) can be made. The technological applications are enormous.

For examples of some currently commercial CNTs and functionalised CNTs see the following paragraphs.

CARBON NANOTUBES (CNTs)

Carbon annealed nanodiscs/nanocones [7440-44-0, EC Number 231-153-3] **C, M 12.0, m 3652-3697^o(sublimes), d²⁰ 1.8-2.1g/cm³**. These are available as a black powder containing a mixture of 20% cones, 70% discs and 10% of carbon black impurities by weight. The carbon cones have a nearly perfect geometry. The five theoretically possible cones with apical angles of 19.2°, 38.9°, 60°, 84.6° and 112.9° are all present in the mixture with cone lengths of 0.3-0.8μ, maximum base diameter of 1-2μ and wall thickness of 20-50 nanometers. The diameter and thickness of the discs are 0.88-3.5μ and 20-50 nanometers respectively. A product of almost 100% carbon is obtained by annealing at 2500-2700° which increases the structural order and reduces the concentration of impurities particularly of metals (from catalyst). They are similar to multi-walled carbon nanotubes (MWCNTs).

Carbon multi-walled nanotubes (MWCNTs or MWNTs) [1333-86-4, EC Number 215-609-9] **C, M 12.0, m 3652-3697^o(sublimes), bulk density 0.346g/cm³**. These are produced as a black powder by chemical vapour deposition (CVD) and contain >90% of nanotubes which are almost free from metal impurities. The tubes are ~7μ in length and ~140 nanometers in diameter. Multi-walled smaller nanotubes of ~1.5μ in length and ~20-25 nanometers in diameters are similarly prepared. When the multi-walled carbon nanotubes are '**arc-produced**' they are even smaller in size with ~2μ in length and ~2-50 nanometers in diameter [*double-walled 308068-56-6*]. This '**arc process**' produces a mixture of 55-65wt% of straight multi-walled nanotubes and 35-45wt% of graphite nanoparticles. They contain almost 100% carbon. By growing the tubes at 3000-4000° using this process they contain less defects than by other procedures, and are purified in this way. They are stable in air up to 700°.

MWCNT arrays for *nanoelectronics* with nickel-carbide particle tips with same diameters as above are also available. Vertically aligned **MWCNTs on a silicon wafer substrate** with low resistivity (1-30 ohm-cm), as well as on a **copper wafer substrate** (1cm x 1cm x 0.05cm high conductivity low-oxygen copper) have been prepared. The latter are produced by **plasma-enhanced chemical vapour deposition (PECVD)**. MWCNTs have also been made in 3-19, 7-13 and 5-15 graphene layers wall thicknesses. See below for breaking up the bundles and graphene.

Carbon double-walled nanotubes (DWCNTs or DWNTs) [308068-56-6] MDL number MFCD06411993. These have been prepared by the *Carbon Vapour Deposition* (CVD) method with >90% carbon content and 50-80% DWCNT. They can be prepared with approximately the following dimensions 5nm OD, 1.3-2.0nm ID and 50μm length; and with 0.12-0.14g/ml bulk density and >600m²/g surface area. The rest (10-40%) consists of amorphous carbon and residual metal catalysts that are deliberately placed in the original carbon in order to synthesise the carbon tubes. [Double-walled Carbon Nanotubes by Pfeiffer et al. *Topics in Appl Phys* **111** 495 2008, DOI: 10.1007/978-3-540-72865-8_16.]

Carbon single-walled nanotubes (SWCNTs or SWNTs) [308068-56-6] **M 3652-3607, d²⁵ 1.7-1.9g/ml**. The SWCNTs are of particular use for derivatisation and the applications stated above. They can be prepared in various bundle dimensions, viz: $d = 1.2-1.5\text{nm}$, $l = 2-5\mu\text{m}$ (40-60% SWCNT, arc method); $d = 1.1-0.5\text{nm}$, $l = 100\mu\text{m}$ (>50% SWCNT, Carbon Vapour Deposition, CVD, method); $d = 2-10\text{nm}$, $l = 1-5\mu\text{m}$ (50-70% SWCNT, arc method); individual short tubes $d = 1-2\text{nm}$, $l = 0.5-2\mu\text{m}$ (90% SWCNT, electric arc, EA, method), and individual long tubes $d = 1.3-1.5\text{nm}$, $l = 1-5\mu\text{m}$ (40-60% SWCNT, arc method, EA) and the HiPco process (see above). [See 'Single-walled Carbon Nanotubes' by Nanot et al. in *Springer Handbook of Nanomaterials (Part A)* Springer Verlag, pp 105-146 2013, DOI: 10.1007/978-3-642-20595-8_4, ISBN: 978-3-642-20594-1 print, 978-3-642-20595-8 on line.]

FUNCTIONALISED CNTs.

MWCNTs and SWCNTs have been functionalised successfully, and the products have found numerous applications. They are conducive to further chemical reactions as they are readily soluble, thus amenable to solution chemistry [see R.C. Haddon and coworkers in *Chemistry of Single-Walled Carbon Nanotubes*, Niyogi et al. *Acc Chem Res* **35** 1105 2002, DOI: 10.1021/ar010155r; and Sun and coworkers in *Functionalized Carbon*

Nanotubes: Properties and Chemistry, Sun et al. *Acc Chem Res* **35** 1096 2002, DOI: 10.1021/ar010160v]. When purified by the acid and oxidising conditions the tubes invariably are oxidised to form CO₂H or quinone groups, particularly at the ends of the tubes. Depending on the conditions, ‘*cutting*’ may occur leaving open-ended tubes (see fullerene tubes above). Annealing at high temperatures *in vacuo* causes some decarboxylation and closing up of the ends that form half **fullerene tips** (like the end of a glass test tube, see above). Because of the larger curvature at the tips than on the sides of the tubes, the tips are more readily functionalised (due to the easier conversion of sp² to sp³ carbon atoms); but then again they would lose the added function more readily on annealing. The tips are also more easily cleaved by oxidants (KMnO₄, OsO₄ and RuO₄) at 100° under acidic conditions [Hwang *JCS Chem Commun* 173 1995, DOI: 10.1039/C39950000173]. Functionalised SWCNTs dissolve in organic solvents (e.g. THF, CH₂Cl₂) and can be examined by optical spectroscopy. Tubes with CO₂H groups on the sides are commonly formed, are readily converted to COCl groups, and can react with a variety of reagents possessing the appropriate functional group. Also by virtue of the double bonds in the hexagonal rings, the tubes condense with nitrenes (to form aziridino compounds with UV light), react with carbenes (to form cyclopropane derivatives), are reduced (by Birch reduction), form metal derivatives which can be reacted further, react with aryl diazonium compounds (arylation), undergo 1,3-dipolar cycloaddition reactions (e.g. to form pyrrolidine derivatives), undergo nucleophilic reactions, radical reactions, and halogenation. J.L. Musgrave and coworkers [Khabashesku et al. *Acc Chem Res* **35** 1087 2002, DOI: 10.1021/ar020146y] have prepared ‘**fluoronanotubes**’ by direct fluorination of SWCNTs, and the fluoronanotubes dispersed in THF reacted with alkyl-Li reagents, Grignard reagents, alkoxides, hydrazine and ω-diamines to form a variety of useful functionalised tubes. Sun and coworkers [Fu et al. *Nano Lett* **1** 439 2001, DOI: 10.1021/nl010040g] functionalised SWCNTs with **lipophilic and hydrophilic dendra** and showed that they can be defunctionalised in homogeneous solutions by base- and acid- catalysed hydrolysis demonstrating the existence of ester linkages. Absorption in the UV/VIS spectra of these **functionalised tubes obey Beer’s law** demonstrating no aggregation effects, and SEM analysis was not successful due to the extent of substitution, whereas the de-functionalised SWCNTs and MWCNTs gave successful SEM imaging. SWCNTs attached to a silicon substrate can be used as very small highly sensitive **chemical sensors or memristors** for gases. They also adsorb chemicals such as alcohols, aromatics, amines and phosphonates, some of which do not readily desorb and slow the process down unless they are removed by other means such as high temperatures which tend to be time consuming and degrade the sensor. Masel and coworkers [Salehi-Khojin et al. *Science* **329** 1327 2010, DOI: 10.1126/science.1194210, reviewed by H. Birch, ‘**Electric shock resets nanotube sensor**’, *RSC Chemistry World* **7** (10) 2010] showed that current induced voltage above the Poole-Frenkel conduction threshold (>12V) desorbed most molecules effectively after 3 hours and the sensors returned rapidly to baseline. The carbon of SWCNTs can be functionalised when coated with SDS in H₂O by reaction with *p*-substituted phenyldiazonium tetrafluoroborate to form **functionalised SWCNTs**. These have 1 to 9 carbon atoms along their backbones which will be attached to the organic moiety, and they remain unbundled throughout their entire lengths. Typically the SDS coated CNTs (10ml, 2.08μM, 0.02mmol) and the aryldiazonium salt (0.32nmol, 16 equiv/mol of carbon; with *p*-Cl, *p*-Br, *p*-*t*-Bu, *p*-CO₂Me, *p*-NO₂, *p*-C=C-Ph) are stirred in a flask for 10min. Completion of reaction can be ascertained from the UV spectrum of an aliquot, then the mixture is diluted with Me₂CO, filtered through a PTFE (0.2μm) membrane. The filtrate is washed with H₂O (100ml) the Me₂CO (100ml x 3) to remove the SDS and unreacted diazonium salt completely. The functionalised solid nanotubes are collected and dried overnight at 65° *in vacuo* and characterised. AFM (atomic force microscopy) analysis of the original nanotubes and the functionalised tubes showed that the latter are either incapable of bundling or easily exfoliated in organic solvents. Some solutions of them can remain without precipitation or flocculation for months. [Dyke & Tour *Nano Lett* **3** 1215 2003, DOI: 10.1021/nl034537x.] Although functionalisation affects the electronic properties of the nanotubes, their use for rheological modification of blended materials can be extensive. For the application of this procedure to carbon or graphite coated nanometals see for example carbon coated cobalt nanoparticles below.

Rapid determination of the carbonaceous content of bulk SWCNT soot uses a solution-phase near-IR procedure. Thus the AP-SWCNT (10g, AP = as prepared) is homogenised mechanically to a fine powder, then a small portion (50mg) is dispersed in DMF (100ml) by ultrasonication and mechanical stirring (5min) to a slurry. A few drops of the slurry are further dissolved in DMF (10ml) to give a faintly coloured non-scattering dispersion after further ultrasonication (2min). Keep the sonication to a minimum to avoid damaging the sample. The concentration of the dispersion should be adjusted to obtain an optical density of 0.15–0.2 at 12,000 cm⁻¹. This concentration gives the minimum signal/noise ratio for recording spectra and the range

corresponds to a concentration of *ca* 0.01mg/ml. It should be noted that at higher concentrations dispersions were less stable. A reference sample (from highest purity material) is used to evaluate the purity and utilising the region of the second interband transition (S_{22}) for semiconducting SWCNTs. [Itkis et al. *Nano Lett* **3** 309 2003, DOI: 10.1021/nl025926e].

SWCNT—(CO₂H)_n, 89-90% SWCNT (carbonaceous purity), bundle dimension: diam 4-5nm x length 0.5-1.5μm. The SWCNTs were produced by EA discharge [Journet et al. *Nature* (letter) **388** 756 1997, Itkis et al. *J Phys Chem B* **108** 12770 2004, DOI: 10.1021/jp0487307] or the HiPco procedure (see above). SWCNT-CO₂H is prepared as follows: HiPco SWCNTs (0.2g) are sonicated in a 3:1 mixture of concentrated H₂SO₄/concentrated HNO₃ (80ml) for 4 hours, diluted with de-ionised H₂O and filtered through a 0.2μm pore acid-resistant membrane. The solid is dried at ~25°, then sonicated in a solution of 4:1 concentrated H₂SO₄/30% H₂O₂ for 15 minutes, diluted with excess de-ionised H₂O, filtered again and dried *in vacuo* at 25° to give 79w/w% of ‘shortened’ product (85% have length < 600nm, i.e. ~0.6μm). [Zhao et al. *Adv Funct Mater* **14** 71 2004I, DOI: 10.1002/adfm.200304440.] Solution phase NIR spectroscopy against a standard [see Itkis et al. *Nano Lett* **3** 309 2003, DOI: 10.1021/nl025926e], gave a carbonaceous purity of 80-90% and 3-6 atom% carboxylic acid. Its solubility is 1mg/ml in DMF, and 0.1mg/ml in H₂O. Metal content is 5-10%. This is suitable for further reactions. [see 652490 ALDRICH.]

SWCNT—(COCl)_n. The preceding shortened SWCNT-acid (100mg) [see also Liu et al. *Science* **280** 1253 1998, DOI: 10.1126/science.280.5367.1253] is stirred in SOCl₂ (20ml) containing DMF (1ml) at 70° for 2 hours, centrifuged, the brown coloured supernatant is decanted, the sediment is washed with anhydrous THF, centrifuged again; the pale yellow supernatant is decanted and the sedimented SWCNT-(COCl)_n is dried at ~25° *in vacuo*. This product is reactive and should be used immediately. [Chen et al. *Science* **282** 95 1998, DOI: 10.1126/science.282.5386.95]. Alternatively, the preceding acid (12mg) in dry DMF (20ml) is sonicated for 30 minutes to give a homogeneous suspension, to which is added redistilled oxalyl chloride (0.4ml) dropwise at 0° under N₂. The mixture is stirred at 0° for 2 hours, then stirred at 70° overnight to remove excess oxalyl chloride, and dried *in vacuo*. This reactive product must be used immediately. [See Itkis et al. *Nano Lett* above.]

SWCNT—(CONH₂)_n, 89-90% SWCNT (carbonaceous purity), bundle dimension: diam 4-6nm x length 0.7-1.0μm, average diameter of individual SWCNT being 1.4nm ±0.1nm [MFCD09753863]. The preceding cooled SWCNT-(COCl)_n in dry DMF (~0°) is treated with liquid NH₃ (~0.5ml) and stirred while the temperature rose to ~25°, and kept there with stirring for 24 hours. Dry N₂ is bubbled through the mixture to remove excess of NH₃, excess of dry THF is added and the mixture is centrifuged, the supernatant is decanted off, the sediment is washed with dry THF, centrifuged again and the SWCNT-NH₂ residue is dried *in vacuo* at ~25°. The extent of labelling is the same as in the original acid, i.e. 3-6 atom % (amide groups) and metals ~6-8%. Their solubilities are 0.5 to 1.0mg/ml, in each of the alcohols, acetone and DMF; and can be functionalised with any reagent that will react with amide groups.

SWCNT-(ODA)_n {SWCNT-[CONHCH₂(CH₂)₁₆CH₃]_n, octadecylamide functionalised}, 89-90% SWCNT (carbonaceous purity), bundle dimension: diam 2-10nm x length 0.5-2.0μm, average diameter of individual SWCNT being 1.4nm ±0.1nm, [MFCD07370655]. It is prepared from the above SWCNT-(COCl)_n (obtained from 100mg of the acid) by mixing with octadecylamine (ODA, 2g, m 55-57°) and heating at 90-100° for 96 hours under dry N₂, and cooled. Excess of ODA is removed by washing with EtOH four times (5-10 minutes sonication at 40 KHz) by centrifugation/decantation, the sediment is dissolved in CH₂Cl₂, filtered, the black coloured filtrate is evaporated to dryness in a rotavap and the residue is dried *in vacuo* at ~25° to give >60% yield based on shortened SWCNTs. **Note** that the reaction of SWCNTS-(COCl)_n and excess of ODA in toluene at ~25° for several days gives only traces of product and the success of the former procedure was attributed to expansion and defoliation of the SWCNTs bundles to give the more reactive individual nanotubes [Chen et al. *Science* **282** 95 1998, DOI: 10.1126/science.282.5386.95]. The extent of labeling in the commercial sample is 30-40wt% (ODA) and contains 4% of metals. The amide is soluble in CHCl₃, CH₂Cl₂, *C₆H₆, toluene, and the solubility in CS₂ or THF is >1mg/ml. The solubility of the ODA derivative made it amenable to purification, and R.C. Haddon and coworkers purified it by gel permeation chromatography (SEC) using Styragel HMW7 [Niyogi et al. *J Am Chem Soc* **123** 733 2001, DOI: 10.1021/ja0024439] and PLgel MIXED-A [Zhao et al. *J Am Chem Soc* **123** 11673 2001, DOI: 10.1021/ja010488j], both being polystyrene divinylbenzene resins. The latter proved to be a superior gel (300 x 7.5mm column), and when using THF as eluent three bands

were separated at a flow rate of 0.5ml/minute. The first band contained 74% of SWCNTs-ODA (as detected by AFM, UV and NIR) [MW range 2000 to 4×10^7 , particle size 15-20 μ m, retention time 8 minutes], the second band with retention time of 9 minutes contains mostly nanoparticles with traces of SWCNTs, and the third band with retention time of 19 minutes contained amorphous carbon. IR, Raman and UV spectra confirmed the ODA component, and ^1H NMR (200MHz, CDCl_3) demonstrated bands characteristic of CH_2 groups and the terminal CH_3 group. The nature (e.g. such as broadness) of the bands was indicative that both ionic (charge transfer) and covalent interactions had occurred.

SWCNT- $\{\text{CONH-}p\text{-}[\text{C}_6\text{H}_3(m\text{-SO}_3\text{H})\text{-}p\text{-NHC}_6\text{H}_3(m\text{-SO}_3\text{H})\text{-}]_n\}$ [SWCNT- n -poly- p -aminobenzene- m -sulfonic acid, SWCNT-(CO-PABS) $_n$], 75-85% SWCNT (carbonaceous purity), average size of individual SWCNT is L 1.1nm x 0.5-1.0 μ m, average PABS Mw ~400-600g/mol, [MFCD05865412]. The PABS polymer was prepared by mixing m -aminobenzenesulfonic acid (ABS, 0.865g) and aniline (15-20mol% of ABS, as inhibitor of polymerisation) and 1M HCl with ammonium persulfate as oxidant, stirred at 0° for 6 hours, concentrated at $\sim 25^\circ$ (*in vacuo*), filtered and the solid was washed with Me_2CO . This was dissolved in H_2O , and the aqueous solution was slowly added to a large excess of Me_2CO . The black solid was filtered off and dried at $\sim 25^\circ$ (*in vacuo*) to give PABS (340mg, 40%, Av $M_w \sim 400\text{-}600$ g/mol), which was identified by its UV spectrum that has λ_{max} at 290 and 510nm in aqueous 1N NaOH [Roy et al. *Synth Met* **100** 233 1999, DOI: 10.1016/S0379-6779(98)01505-7]. **SWCNT-(CO-PABS) $_n$** was obtained from SWCNT-(COCl) $_n$ [prepared as above by the oxalyl method from HiPco SWCNTs (12mg, Carbon Nanotechnologies Inc.)] in dry DMF ($\sim 20\text{ml}$) by mixing with PABS (120mg) in DMF ($\sim 50\text{ml}$) and stirring at 100° for 5 days. After cooling to 25° the solid was filtered through a 0.2mm pore-size membrane and washed thoroughly with DMF and EtOH, and dried *in vacuo*. The black **SWCNT-(CO-PABS) $_n$** (57mg) on the membrane was collected and dried *in vacuo* overnight. Its solubilities are 0.05mg/ml in EtOH, 0.1mg/ml in DMF and 5.0mg/ml in H_2O . The commercial product has 65% (PABS) and 4% metals. The water soluble graft polymer had a much higher conductivity ($5.6 \times 10^{-3} \text{ Scm}^{-1}$) than PABS ($5.4 \times 10^{-7} \text{ Scm}^{-1}$), and IR spectrum was consistent with an amide bond being formed; the ^1H NMR (300MHz, D_2O) exhibited a very weak broad spectrum compared with the sharp signals of PABS itself, characteristic of the effect of ring currents in the nanotubes. The UV/VIS/MIR spectrum showed the presence of the interband transitions of the semiconducting SWCNTs and an absorption at $17,750 \text{ cm}^{-1}$ due to the PABS moiety. [Zhao et al. *Adv Funct Mater* **14** 71 2004, DOI: 10.1002/adfm.200304440.] Solution phase NIR spectroscopy against a standard, provided a carbonaceous purity of 80-90% — a procedure which has an accuracy of $\sim 3\%$ [see Itkis et al. *Nano Lett* **3** 309 2003, DOI: 10.1021/nl025926e].

SWCNT-(PEG) $_n$ {SWCNT-[COO(CH $_2$ CH $_2$ -O-) $_m$ -H] $_n$, polyethylene glycol functionalised}, 80-90% SWCNT (carbonaceous purity), bundle dimension: diam 4-5nm x length 0.5-0.6 μ m, average diameter of individual SWCNT being 1.4nm \pm 0.1nm, PEG Mw ~600g/mol, [MFCD07370656]. The above SWCNT-(COCl) $_n$ prepared from 30mg of SWCNT-CO $_2\text{H}$ is mixed with PEG (250mg, m 20-25 $^\circ$) and heated under N_2 at 75° with vigorous stirring for 48 hours. The mixture is cooled to 25° extracted with CHCl_3 several times, filtered, and the dark coloured solution is repeatedly precipitated with EtOH to give SWCNT-(PEG) $_n$ which is collected (filtration or centrifugation) and dried *in vacuo*. Its solubility in H_2O is high (5.0mg/ml) and has $\sim 30\text{wt}\%$ (PEG) and $\sim 6\%$ trace metals. The ester function can be identified by IR ($\nu_{\text{max}} \sim 1700 \text{ cm}^{-1}$), but the ^1H NMR signals are weak and broad (see above). The ester-free SWCNTs can be recovered by acid- and base-catalysed hydrolysis [see above Fu et al. *Nano Lett* **1** 439 2001, DOI: 10.1021/nl010040g]. Solution phase NIR spectroscopy against a standard, provided a carbonaceous purity of 80-90% — a procedure which has an accuracy of $\sim 3\%$ [see Itkis et al. *Nano Lett* **3** 309 2003, DOI: 10.1021/nl025926e].

SWCNT-($p\text{-C}_6\text{H}_4\text{-R}$; R = F, Cl, Br, I, SO_3H , CO_2H , NO_2 , n -butyl, $t\text{-Bu}$ or CO_2Me) $_n$. These were prepared by ‘on water’ functionalisation of bundled SWCNTs. The term ‘on water’ refers to the water-based reactions of water-insoluble organic substrates [see K.B. Sharpless and coworkers in Narayan et al. *Angew Chem Int Ed* **44** 3275 2005, DOI: 10.1002/anie.200462883; and Klijn & Engberts *Nature* **435** 746 2005, DOI: 10.1038/435746a]. This technically involves the reaction of p -substituted benzene diazonium compounds with the hexagonal rings on the walls of SWCNTs in aqueous medium and represents a ‘green’ or ‘environmentally friendly’ process. The following are optimal conditions for functionalisation. HiPco SWCNTs (10mg) and de-ionised H_2O (30ml) were homogenised in a flask (100ml) by stirring at medium setting for 30 minutes, then heating at 80° with the substituted aniline (4 equivalents/SWCNT) and isoamyl nitrite (2 equivalents/SWCNT) with vigorous stirring (stirrer bar) under a reflux condenser overnight. The mixture is then cooled, filtered

through a 0.45µm Teflon filter, the filter cake is washed with de-ionised H₂O, and Me₂CO until the filtrate is clear. The cake is collected, sonicated in DMF (25ml) to remove any remaining organic compounds, collected by filtration (0.45µm Teflon filter) and rinsed with Me₂CO to give the desired functionalised SWCNTs as evidenced by 20-30% weight increases. TGA, Raman [elevated D (diamondoid, ν_{\max} at 1290 cm⁻¹)/G (graphic, ν_{\max} 1590 cm⁻¹) band ratios] and UV-VIS-NIR spectroscopy, XPS, AFM and TEM confirmed the structures. It is interesting that TEM images (on a carbon grid) showed that the SWCNT bundles have smooth edges, whereas functionalised SWCNT bundles have ‘*bumps*’ all along the edges. These functionalised nanotubes should be useful for further functionalisation reactions [Price & Tour *J Am Chem Soc* **128** 12899 2006, DOI: 10.1021/ja063609u]. By using the above procedure **SWCNT-(*p*-C₆H₄-CH₂NH₂)** and **MWCNT-(*p*-C₆H₄-CH₂NH₂)** were prepared and successfully coupled, *via* their terminal amino group to the carboxy group of *N*(1)-carboxymethyl-thymine, to form thymine ends. These functionalised CNTs readily form stable *double* hydrogen bonds with other thymine groups of these CNTs. This induces controlled non-covalent self-assembled supramolecular aggregation of the CNTs in solvents that do not break hydrogen bonds such as CH₂Cl₂, and can form good dispersions in polar aprotic solvents such as DMF. All characterisations were performed using spectroscopic, analytical and microscopic techniques. [Quintana & Prato *Chem Commun* 6005 2009, DOI: 10.1039/B915126E; reviewed by K. Davies *RSC Chemistry World* **6**(11) 6005 2009.] These properties can be of importance in *nanoelectronics*, or in *biological applications* such as making patterned active substrates for neuronal growth [Cellot et al. *Nat Nanotechnol* **4** 126 2009, DOI: 10.1038/nnano.2008.374].

I-SWCNT, I = {-COOCH₂-[3,5-di(hexadecyloxy)phenyl]}_n and **I_{PEG}-SWCNT, I_{PEG} = {-COOCH₂-[3,5-di(methyltriglycoloxy)phenyl]}_n**. These are respectively hydrophobic and hydrophilic SWCNTs which were prepared as described by Y.-P. Sun and coworkers [Fu et al. *Nano Lett* **1** 439 2001, DOI: 10.1021/nl010040g] by stirring vigorously SWCNT-(COCl)_n (30mg, see above) and carefully dried **I** (250mg) under N₂ at 75°/48 hours. The cooled mixture is extracted several times with CHCl₃, the combined dark-coloured extracts were repeatedly precipitated with EtOH to give **I-SWCNT** with ¹H NMR (500MHz, CDCl₃) which had δ at 0.88 (t, *J* = 6.5Hz) and typically weak broad bands at 1.1-1.5, 1.6-1.8, 3.5-3.9, 4.0-4.2 and 6.0-6.5 caused by the effect of the SWCNT. **I_{PEG}-SWCNT** was prepared in a similar manner except that its solubility in H₂O allowed further purification by dialysis for several days against de-ionised H₂O (dialysis tubing with Mr ~100,000 cut off to remove PEG). **Dendron I** was prepared by reaction of methyl 3,5-dihydroxybenzoate and hexabromodecane followed by reduction with LAH, and **Dendron I_{PEG}** was obtained from the same benzoate and triethylene glycol monoethyl ether in the presence of Ph₃P and diethyl azidodicarboxylate in THF, followed LAH reduction. The corresponding hydrophobic **I-MWCNT** and hydrophilic **I_{PEG}-MWCNT dendra** were similarly prepared by reacting SWCNT-(COCl)_n with the benzylic OH group in the **dendra**, and all were characterised by ¹H NMR, UV-Vis spectroscopy, TGA and SEM as well by de-functionalisation of homogeneous solutions under acid- and base- catalysed reaction conditions.

MWCNT-(COOH)_n are commercially available [755125 *ALDRICH*], with average diameter, 9.5nm length x 1.5µm and >8% -COOH functionalised. For the preparation of the polyurethane (**PU**) carbon nanotube composite **MWCNT-(COEDA-PU-PTMO)_n**, the MWCNT-(COOH)_n was converted into MWCNT-(COEDA)_n in two steps (SOCl₂ then H₂NCH₂-CH₂NH₂, EDA), and the grafted amide groups reacted readily with toluene-2,4-diisocyanate (TDI) to form MWCNT-(COEDA-TDI)_n which were condensed with PTMO {polyoxytrimethylene glycol, HO[-(CH₂)₄-O-(CH₂)₄]_n-OH, *M_n* = 1000} to form the hard segment of the polyurethane, **PU**. The microstructure of this composite containing 2 wt% of MWCNT was investigated by FESEM (field emission scanning electron microscopy) and TEM (transmission electron microscopy); and the IR of a wafer in KBr confirmed that condensation had occurred. The thermal and mechanical properties (tensile strength) of the polyurethane/multi-walled carbon nanotube composite were found to be superior in comparison with pure polyurethane composite [Xiong J, et al. *Carbon* **44** 2701 2006, DOI: 10.1016/j.carbon.2006.04.005].

MWCNT-(PPEI-ED)_n. These functionalised MWCNTs were prepared in two ways. *Firstly*, the MWCNTs (20mg) were refluxed with SOCl₂ (5ml) for 24 hours and evaporated *in vacuo*, then the co-polymer PPEI-EI [200mg, poly(propionylethylenimine-co-ethylenimine) [as prepared by Y.-P. Sun and coworkers *Macromolecules* **32** 8747 1999, DOI: 10.1021/ma9906736; *Photochem Photobiol* **66** 301 1997, DOI: 10.1111/j.1751-1097.1997.tb03152.x; *Chem Commun* 2699 1996, DOI: 10.1039/CC9960002699] was added and heated at 160-180° for 12 hours under N₂, cooled, repeatedly extracted with CHCl₃ and the dark combined

extracts were precipitated with hexane. The isolated solid was dissolved in de-ionised H₂O and dialysed against fresh H₂O (dialysis tubing with Mr ~100,000 cut off to remove PPEI-EI) for 3 days. Further purification was by re-precipitation from CHCl₃ solution into hexane, and drying the dark **MWCNT-(PPEI-EI)_n** solid which had ¹H NMR (500MHz, CDCl₃) with broad weak bands at δ 0.8-1.4, 1.9-2.6, 2.7-2.9, 3.1-4.3. **Secondly**, (without forming the COCl derivative) by directly heating the MWCNTs (20mg) with PPEI-EI (200mg) at 160-180° for 12 hours under N₂ and worked up as above provided **MWCNT-(PPEI-EI)_n** which had ¹H NMR (500MHz, CDCl₃) with broad weak bands at δ 0.8-1.4, 1.9-2.6, 2.7-2.9, 3.1-4.3. Raman, UV-VIS spectroscopy, SEM, TEM and TGA showed that both procedures were effective in producing functionalised MWCNTs which were readily soluble in common organic solvents and in H₂O. The latter method may have caused amidation as well as ionic interaction between the amino-polymer and the MWCNT. [Lin et al. *J Phys Chem B* **106** 1294 2002, DOI: 10.1021/jp013501v.]

SWCNT[=CH]₂N-CH₂CH₂OCH₂CH₂OCH₂CH₂NH₃⁺. HCl]_n and **MWCNT[=CH]₂N-CH₂CH₂OCH₂CH₂-OCH₂CH₂NH₃⁺. HCl]_n. These salts are very soluble in H₂O and solutions are stable for more than a month at concentrations of 20mg and 12mg per ml respectively. The free amino terminal group is a useful handle for attachment to physiologically active molecules (see below). They are formed in a 1,3-dipolar cycloaddition reaction between the C=C bonds of the CNTs with the N-glycine N atom and formaldehyde. The glycine in this case is BocNHCH₂CH₂O-CH₂CH₂OCH₂CH₂NHCH₂CO₂H and is prepared as follows: BocNHCH₂CH₂O-CH₂CH₂OCH₂CH₂NH₂ (30mmol) in dioxane (20ml) at 0° is treated dropwise with a solution of benzyl bromoacetate (2.3g, 10mmol) in dioxane (30ml) during 1 hour, and the mixture is stirred overnight. The solvent is evaporated off *in vacuo*, the residue is dissolved in H₂O (70ml) and extracted with EtOAc (3 x 50ml). The combined organic phases are dried (Na₂SO₄), evaporated *in vacuo*, and the residue is purified by chromatography on Silica gel [NM Kieselgel 60 (70-230 mesh)] and eluted with 1:1 EtOAc/petroleum ether then pure EtOAc to give **N-Boc-aminoethoxyethoxyethylaminoacetic** as an oil. To a solution of this oil (5.05mmol) in MeOH (50ml) is added 10% Pd/C (50mg), the mixture is stirred under H₂ for 24 hours, the catalyst is filtered off (through Celite), the solvent is evaporated and the residue is triturated with dry Et₂O to give **aminoethoxyethoxyethylaminoacetic acid** as a pure white solid (1.6g, 99%, 5.05mmol), **m 105-106°** with the expected elemental (C, H and N) analyses. The acid has IR-DRIFT (KBr) with ν_{max} at 3250, 2970, 1706, 1620, 1540, 1365, 1115, 686, 590, 480 cm⁻¹; the ¹H NMR (200MHz, CDCl₃, TMS) has δ at 1.40 (s, 9H), 3.22 (m, 2H), 3.49 (t, *J* = 5.1 Hz, 2H), 3.64-3.53 (m, 8H), 3.79 (bt, 2H), 5.54 (bt, 1H), 6.23 (bs, 1H), 8.21 (bs, 1H); the ¹³C NMR (50MHz, CDCl₃, TMS) has δ at 170.5, 156.2, 79.1, 70.4, 70.3, 70.1, 66.6, 49.8, 46.8, 40.4, 28.6; and EI-MS found *m/z* 306 (M⁺). [Kordatos et al. *J Org Chem* **66** 4915 2001, DOI: 10.1021/jo015608k.]**

For **functionalisation**, a suspension of full length SWCNTs or MWCNTs [diameter 20-30nm, from Carbon Nanotechnologies, Inc USA (www.cnanotech.com), and Nanostructured & Amorphous Materials Inc USA, (www.nanoamor.com) respectively] in DMF is treated with the preceding **N-Boc-aminoethoxyethoxyethylaminoacetic** (or and paraformaldehyde, and the mixture is heated at 130° for 96 hours. Unreacted material is removed by filtration, the filtrate is evaporated, the residue is dissolved in CHCl₃, washed with H₂O, dried, evaporated, redissolved in CHCl₃, precipitated with Et₂O, collected (on a 0.45mm Teflon filter) and washed several times with Et₂O to give the functionalised CNTs in ~10% yields based on the amount of starting CNTs. They are soluble in solvents such as CH₂Cl₂, CHCl₃, toluene and Me₂CO. Removal of the *N*-Boc group is achieved by dissolving the previous functionalised SWCNTs or MWCNTs in CH₂Cl₂, dry HCl gas is bubbled through the solution whereby the **CNT-chloride hydrochloride salts** (desired materials) separate out. They are collected (or the solvent is evaporated), dissolved in MeOH and precipitated with dry Et₂O. *Alternatively*, the *N*-Boc group is removed by treatment with TFA (~25°/3hrs, evaporate, wash residue with toluene and dry *in vacuo*) in which case the **CNT-trifluoroacetate salts** are obtained. A variety of such functionalised CNTs were prepared and some were methylated (at the pyrrolidine that was produced) to form the quaternary bases. The desired products, as identified by TEM showed that the functionalised SWCNTs and MWCNTs have 10-50nm and 20-30nm mean diameters respectively, their ¹H NMR spectra are similar with the signals from the oligoethylene glycol chains appearing as broad peaks at ~3.6 ppm, and absence of the Boc methyl groups which would have been at 1.2 ppm from TMS. [Georgakilas et al. *Chem Commun* 3050 2002, DOI: 10.1039/B209843A.] These functionalised CNTs were successfully coupled *via* their terminal amino group to the carboxy group of *N*(1)-carboxymethyl-thymine to produce thymine ends. The thymine functionalised CNTs readily form stable double hydrogen bonds with other thymine groups of these CNTs. This induces controlled non-covalent self-assembled supramolecular aggregation on the CNTs in solvents that do not break hydrogen bonds such as CH₂Cl₂, and can form good dispersions in polar aprotic solvents such as DMF. All characteris-

ations were performed using spectroscopic, analytical and microscopic techniques. [Quintana & Prato *Chem Commun* 6005 2009, DOI: 10.1039/B915126E; reviewed by K. Davies RSC *Chemistry World* 6(11) 6005 2009.] These properties can be of importance in *nanoelectronics* or in *biological applications* such as making patterned active substrates for neuronal growth [Cellot et al. *Nat Nanotechnol* 4 126 2009, DOI: 10.1038/nnano.2008.374.]

SWCNT[=(CHR')(CH₂N-R'')], where R' = H, 4-MeOC₆H₄- or 2-pyrenyl-, and R'' = -(CH₂CH₂O)₃CH₃ or -(CH₂)₇CH₃]_n and MWCNT[=(CHR')(CH₂N-R'')], where R' = H, 4-MeOC₆H₄- or 4-pyrenyl-, and R'' = -(CH₂CH₂O)₃CH₃ or -(CH₂)₇CH₃]_n were prepared by 1,3-dipolar cycloaddition reactions with short oxidised SWCNTs or purified MWCNTs as in the preceding entry and using paraformaldehyde, 4-methoxybenzaldehyde or 4-formylpyrene to form the substituted pyrrolidines with the double bond(s) of the CNTs. They were characterised by UV-VIS, Raman and ¹H NMR spectroscopy and by TEM. The fluorescence of the pyrene derivatives were of interest, and all provide means for the preparation of nanocomposites. [Georgakilas et al. *J Am Chem Soc* 124 760 2002, DOI: 10.1021/ja016954m; Calvert *Nature* 399 210 1999, DOI: 10.1038/20326.]

GRAPHENE MATERIALS

Graphene is the name given to a flat monolayer of carbon atoms packed in a two-dimensional (2D) honeycombe lattice of six-membered carbon rings. It is the unit block (2D sheet) of graphitic materials which stack into multilayers to form graphite (3D), rolls to form nanotubes (1D) or wraps to form buckyballs (0D)[see above]. 'Graphenes' are defined as single-, double-, and few (3 to <10) carbon layers (2D), with characteristic electrical properties not found in graphite. *Graphene* is the accepted name for a single, or at most a double carbon thick layer which is ≈ 5 Å thick. Single layer (SLG) 2D crystals have been prepared (see below) that have high crystal quality where charge carriers can travel thousands of atomic distances without scattering. Its electrical behaviour includes a pronounced ambipolar electric field effect in which charged carriers could be tuned continuously between electrons and holes at concentrations *n* as high as 10¹³ cm⁻², with mobilities *μ* greater than 15,000 cm² V⁻¹ s⁻¹ under laboratory conditions. Films of 5 layers would be considered as '*bulk material*', and films with more than 100 layers are considered as thin films of 3D graphite (see graphite [7782-42-5] below). [Geim & Novosolov *The Rise of Graphene: Nature Materials* 6 183 2007, DOI: 10.1038/nmat1849.]

Graphene films were obtained by mechanical exfoliation (repeated peeling) of highly oriented pyrolytic graphite. The 2D crystallites were collected from a fresh surface of a layered crystal using *few-layered graphene*, *FLG*, by rubbing against another surface (any solid is suitable, but an oxidised silicon wafer, i.e. SiO₂ would be also useful later for identification). This allows flakes of graphene to be attached to the surface for characterisation. This approach was very reproducible, has also been used for preparing thicker films (≥ 3nm) that were up to 100 μm across, as well as FLGs of up to 10 μm in size and visible to the naked eye.

Preliminary identification of single layer films among the resulting flakes was done with an optical microscope. These became visible when they were on top of an oxidised silicon wafer (see above) because even a monomolecular layer adds enough to the optical path of reflected light that the interference colour changes reveal the presence and contours of the film. The process is simple and the 2D crystallites can be identified in about half an hour. Further identification of selected single layers is then done by AFM. It was pointed out that only a small number of single films among the many thicker films were produced by this method, they could not be identified by TEM, but they were transparent to visible light on surfaces other than SiO₂ wafers, and thus required AFM and high resolution AFM imaging for definitive identification, and are difficult to find by scanning surfaces at random. These atomically thin sheets, particularly very large 2D molecules, unprotected from the environment are stable in air under laboratory conditions, have high crystal quality and maintain macroscopic continuity. [Novoselov, Geim and coworkers, '*Electric field effects in atomically thin carbon films*' *Science* 306 666 2004, DOI: 10.1126/science.1102896, '*Two dimensional atomic crystals*' *Proc Natl Acad Sci USA* 102 10451 2005, DOI: 10.1073/pnas.0502848102.]

The sp² character of all the C-C bonds of graphene confer remarkable electric conductivity to it due to the delocalisation of electrons, and it is difficult to stop the flowing current at will. Geim and Novoselov and their coworkers [Elias et al. '*Control of Graphene's properties by reversible hydrogenation: Evidence for graphite*' *Science* 323 610 2009, DOI: 10.1126/science.1167130] have succeeded in hydrogenating the double bonds of graphene to make *graphane* which now has C-C bonds with sp³ character. The reduction transforms graphene from the highly conductive zero-overlap semimetal into an *insulator*. Reduction was achieved with a hydrogen-

helium mixture (10% H₂) at low pressure (0.075mm Hg) and the direct current plasma ignited between two aluminium electrodes with the samples 30cm away from the discharge zone to avoid damage by energetic ions. Graphene and graphane have distinctly different Raman properties. The reaction is reversed by annealing graphane at 450° (higher temperatures damage graphene) in an argon atmosphere for 24 hours. Thus the original metallic state, lattice spacing, Raman and TEM properties, and even the quantum '*Hall effect*' are restored. **Note** that when the thin film is on a substrate, reduction takes place on the upper face of the film, and for more effective reduction they used free-standing graphene membranes. If one can imagine a magic pencil which can remove hydrogen from graphane and drawing channels of graphene into it, better control of current flow can be achieved [see review by J. Urquart RSC *Chemistry World* **6** (3) 23 2009].

Cutting of single-layer graphene (SLG) by anisotropic etching using thermally activated Ni nanoparticles has been developed by Jarillo-Herrero and coworkers [Campos et al. *Nano Lett* **9** 2600 2009, DOI: 10.1021/nl900811r; reviewed by L. Brindley, '*Cutting Graphene to ribbons*' RSC *Chemistry World* **9** (8) 2009]. Unlike previous methods for making graphene nanoribbons and the like, their procedure makes SLG nanoribbons and other SLG structures (e.g. triangles, rectangles) in which all cuts (sub-10nm width) are oriented with the same edge-chirality, i.e. aligned along a single crystallographic directions with no hanging edges. Thus the cuts are straight, and always at an angle of 60° and/or 120°. With graphite and FLGs, etching occurs with chirality changing angles of 30°, 90°, and 150°, as well as 60° and 120°. Their method is described here in some detail as it involves ***purification procedures useful for SLG preparation***. To ensure reliable cuts in nanoparticle assisted etching, the SiO₂/Si substrates (wafers, see above) have to be cleaned with Me₂CO and *iso*-PrOH, and submitted to UV illumination for 5 minutes to remove organic material and for making the SiO₂/Si surface more hydrophilic to ensure proper wetting during the spin-coating deposition of Ni from solution. Graphene is exfoliated on this clean surface using semiconductor grade tape. The SLG on the clean substrate is identified by optical spectroscopy (see above), AFM (typically SLGs are at a height of 0.8nm) and Raman spectroscopy (graphene's G' 2D peak, see above). The tape residue is removed by heating the SLGs in a quartz tube at 500° for 15 minutes under an argon:H₂ flow (850:150 sccm). After this heat 'cleaning', an aqueous solution of NiCl₂ (2.4mg/ml) is centrifuged onto this surface at 1800rpm for 60 seconds, then heated at 90° for 10 minutes on a hot plate to eliminate H₂O. The NiCl₂ treated sample in an argon: H₂ flow (850:150 ratio) is firstly annealed at 500° for 20 minutes to convert the salt into Ni nanoparticles, and then the etching is carried out at 1000° for 25 minutes in the same gas flow. The best etching conditions require a slow heating/cooling rate of ~50°/minute. During the etching the Ni nanoparticle (with H₂ adsorbed) furrows through the SLG by catalytically hydrogenating the carbon atoms in its path and converting each atom into a molecule of methane. With SLGs, but not with FLGs or graphite, the cutting paths never cross each other; as soon as the Ni nanoparticle comes within a 10nm distance of a preformed path it changes direction at a 60° or 120° angle. The purification procedure and the Ni concentration stated are very critical if the formation of CNTs are to be avoided. This opens many future directions for studying SLG nanoribbons and other shapes, constrictions and quantum dots with crystallographic edges. H-H. Ahn and B.H. Hong and coworkers [Bae, et al, *Nature Nanotechnology* **5** 574 2010, DOI: 10.1038/nnano.2010.132] have pressed rectangular graphene films (76cm diagonal) with a roller against an adhesive polymer support and the copper etched away. The graphene is pressed with rollers onto a polyethylene terephthalate substrate and the polymer adhesive is released on heating. In this way further layers of graphene can be added. When treated with HNO₃, the graphene sheet acts as a transparent electrode for touchscreen devices. It is tougher and has better transparency than indium-tin oxide (ITO) electrodes [see below, and review by S. Hadlington RSC *Chemistry World* **7** (8) 22 2010].

Stacked Graphene Platelet Nanochips (stacked graphene nanofilms, SGNF heat treated) and Platelet Nanofibres (SGNF acid washed) are commercially available as black powders with 40-50nm mean width, 0.1-10μ length, 0.3g/cm³ density, and 120m² of surface area. The electrical resistivities are 55μWcm and 120μWcm for the chips and fibres respectively. These SGNFs are used in nanotechnology research. [US Patents 6,995,115 and 7,001,586.]

Graphite [7782-42-5] **m 3652-3697°, d²⁵ 2.09, 2.23.** It is available commercially in various forms such as **flakes** (mesh 75+), **rods** (6mm x 150mm, low density and high density, 3mm x 150mm low density), **powder** (≤20μm, ≤45μm, and 150≤μm), **nanofibres** (I.D. 0.5-10nm x O.D 80-200nm x L. 0.5-20μm, bulk density 0.06-0.08g/ml), **graphitised nanopowder** (particle size <200nm with large mesopores, 10% graphite lattice content, surface area 70m²/g, average pore ~137Å and d 1.818g/cm³) and **platelet nanofibres** (prepared by catalytic CVD then de-mineralised by HCl, with surface area 80m²/g, particle width ~100nm and particle length of

~2.5µm, TEM). [For purification by acid treatment see entry [7782-42-5] in Chapter 4, 'Inorganic Compounds'.] The applications of graphite include the manufacture of 'lead' pencils, lubricants, polishing, pigments, explosives, commutator brushes, carbon anodes, and arc-lamp carbon among many others. **Avoid breathing it** as it can cause coughing, decreased lung function and fibrosis, **use a mouth mask**. [See Holliday et al. in '*Comprehensive Inorganic Chemistry*' vol 1, Bailar Jr et al. Eds, Pergamon Press, Oxford, pp1250-1294 1973; ISBN 9780080172750.]

SELF ASSEMBLED MONOLAYERS (SAMs).

SAMs have been of considerable interest for some time [see Chechik, Crooks and Stirling, '*Reactions and reactivity of self-assembled nanolayers*' *Adv Mater* **12** 1161 2000, DOI: 10.1002/1521-4095(200008)12: 16<1161] and several useful ones have been prepared using a variety of materials and methods. Only a recent report will be described here, and it is of 2D carbon nanolayers prepared by Mo(IV) and Cu catalysed crosslinking of 1,4-bispropyne- (**1**), 1,4-bisacetylene- (**2**), and 1,4-bis(2,3,4-triacetylenephényl)propyne- (**3**) benzenes with each benzene ring further substituted with a 3-(triethoxysilyl)propylaminocarbonyloxymethyl—[(EtO)₃SiCH₂CH₂CH₂-NH-COOCH₂—] group which acts as a linker in the polymerisation process. These three organic compounds are connected by vacuum-driven Mo(IV) catalysed alkyne metathesis [cf Zhang & Moore *Adv Synth Catal* **349** 93 2007, DOI: 10.1002/adsc.200600476], or oxidative Cu catalysed Hay-type coupling [cf Siemsen et al. *Angew Chem Int Ed* **39** 2632 2000, DOI: 10.1002/1521-3773(20000804)39:15<2632] which polymerise them to form self-assembled monolayers on SiO₂-, Si₃N₄-coated substrates (wafers) or quartz/glass slides when immersed in monomer solutions.

Preparation and purification of substrates: The freshly cleaned and dried SiO₂-, Si₃N₄-coated substrates or quartz/glass slides are placed in a reaction tube containing a 15mM solution of (**1**) or (**2**), or (**3**) in toluene and Me₃N (10mM) under N₂, sealed and heated at 95-100° for 24 hours. The liquid is removed, the substrate is rinsed with toluene (1x), CH₂Cl₂ (2x), and sonicated for 5 minutes in toluene. The rinse is repeated, the substrate is sonicated in MeOH for 5 minutes, the rinse is repeated again and the substrate is blown dried under a stream of N₂ ready for monolayer formation.

Preparation and purification of polymerised coated substrate linked monolayers by Cu Coupling: CuCl (20mg) and degassed Me₂CO (3ml) in a reaction vial are treated with TMEDA (61µl), stirred under N₂ at 20° for 30 minutes and the preceding coated substrates are added. The mixture is purged with O₂ from a balloon and stirred under O₂ for 15 hours. The polymerised coated substrate is collected, rinsed with DMF (1x), with a solution of 0.1M sodium diethyldithiocarbamate in DMF (1x), with toluene (1x), with CH₂Cl₂ (2x), sonicated in toluene for 5 minutes, the rinse is repeated, the substrate is finally sonicated for 5 minutes in MeOH, and the *polymerised linked substrate* blown dry with a stream of N₂.

Preparation and purification of polymerised coated substrate linked monolayers by alkyne metathesis: A mixture of trisamidomolybdenum(IV) propylidyne (5.0mg) and *p*-nitrophenol (3.3mg) dissolved in trichlorobenzene (3.3ml, 1,2,4-) are placed in a vial, the coated substrate is added, the flask is sealed under a vacuum (5 Torr) for 22 hours. The substrate is collected. Rinsed with DMF (1x), 0.1M sodium diethyldithiocarbamate in DMF (1x), toluene (1x), CH₂Cl₂ (2x), then sonicated in toluene for 5 minutes. The rinse is repeated, the substrate is sonicated in MeOH for 5 minutes, the rinse repeated again and finally the *polymerised linked substrate* is blown dry in a stream of N₂.

These monolayers (SAMs) are characterised by UV-Vis, fluorescence and Raman spectroscopy. This is made easy because of the presence of benzene cores within the acetylene and linker chains, these being made of poly(1,4-phenylene-1,3-butadiynylene) polymers. AFM then provides the topology, i.e. shapes (contours), thicknesses and aspect ratios (length and width), of the 2D SAMs. This is done at all stages including those described below.

Lift-off and transfer of the SAMs: The '*lifting off*' is carried out by simple oxygen-reactive ion etching through a layer of 'photoresist' (e.g. 5µm x 5µm 'aspect ratio') patterned by photolithography. The 'photoresist' is placed on top of the monolayer which lies on top of the silicon substrates (support, i.e. the films of SiO₂, or Si₃N₄, or glass). Selective etching from the support is done with concentrated HF which dissolves the silicon, leaving photoresist/SAM or photoresist/monolayer membrane hybrid which can be transferred onto other substrates and then freed from the 'photoresist' layer by sonication in Me₂CO. Characterisation of the monolayers during these stages showed that very little, if any, changes in structure has occurred. The atomic force micrographs (AFMs) showed that the SAMs derived from (**1**) are monolayers with thicknesses of ≈ 1.5nm (average roughness of 0.18-0.27nm), and those from (**2**) and (**3**), unlike from (**1**), formed multilayers of 2.8nm

and 4.7nm respectively (average roughness of 0.5nm and 0.3nm respectively). These SAMs, with aspect ratios being the same as the ‘photoresist’ can be transferred by contact onto a variety of surfaces such as films, pleated sheets, spheres, tubes, cones and any other shapes (followed by removal of the ‘photoresist’ by sonication in Me₂CO). With a free-standing shape, a free standing silicon support can be used which is then removed by etching to give a **free standing SAM**. The monolayers are quite robust and can be stretched over ~440nm holes without tearing. This approach and these procedures that were developed by J.S. Moore, J.A. Rogers and coworkers [Schultz et al. *Proc Natl Acad Sci USA* **105** 7353 2008, DOI: 10.1073/pnas.0710081105] have considerable merit for **synthesising monolayer carbon networks** of varying shapes which can be accessible for analysis and device applications.

DIAMOND NANOMATERIALS

Nanodiamond (ND) [7782-40-3] **M 12.01** is commercially available in powder form with particle spherical size <10nm and ≥ 97% trace metal basis, bulk density 0.2-0.7g/ml with BET surface area of 200-450m²/g, and in powder form with particle spherical size <10nm and ≥ 95% trace metal, bulk density 0.17g/ml with BET surface area of 278-335m²/g. The diamond powder is made by **explosive detonation procedures** that are performed in stainless steel hermetic tanks (e.g. of 0.17m). Carbon condensed products [the HE (High Explosive) composite explosives] such as cast trotyl/cyclotrimethylene-trinitramine (TNT/RDX) in various proportions from 50/50 and increasing RDX, with various initial inert gas (N₂ or Ar) pressures were used to determine the best conditions to produce diamonds. Thirty minutes after the explosion, the pressure in the tank is slowly released, and the carbon product is washed out with water, allowed to settle (requiring quite a long period), the water is decanted off, and residual water is evaporated at 150°. This yielded solid carbonaceous soots that contain ultradispersed diamond (UDD, 2-15nm), together with amorphous carbon (4-25nm), graphite ribbons (<20nm), and spheres (2-4nm) as investigated by TEM, XRD, SAXS (small angle X-ray scattering) and Auger spectroscopy. Purification was carried out to remove amorphous and graphite-like carbon from the detonation soot by boiling with perchloric acid, and the maximum yield of diamond was identified by X-ray analysis. The best yield was ~7wt% obtained when TNT/RDX was 50/50 and initial pressure in the tank was ~7 atmospheres which optimised the yield of **UDD**. It was also shown that the UDD was thermally transformed into onion-like carbon (**OLC**) particles. [Kuznetsov et al. *Carbon* **32** 873 1994, DOI: 10.1016/0008-6223(94)90044-2.] Annealing studies, in the temperature range of 750°–1900° of detonation diamond (e.g. OLC) in an inert atmosphere (e.g. argon) affected the structure as shown by XRD and HRTEM as well as purified the material further. The IR showed that all the oxygenated groups (mainly OH, COOH and C=O) were removed, and that at the lower temperatures of ~700-800° a higher coverage of the diamond surface with π -bonds resulted. At higher temperatures the powder undergoes a **phase transition** from cubic diamond to graphite. [Chen et al. *Appl Phys Lett* **74** 3651 1999, <http://dx.doi.org/10.1063/1.123211>; Okotrub et al. *J Phys Chem A* **105** 9781 2001, DOI: 10.1021/jp011808o.] Previously, reactions of ND with free radicals lead to diamond alkylation [Nakamura et al. *Chem Commun* 900 2003, DOI: 10.1039/B211807F]; and arylation with diazonium salts [Yeap et al. *Langmuir* **25** 185 2009, DOI: 10.1021/la8029787; Liang et al. *ACS Nano* **3** 2288 2009, DOI: 10.1021/nn900339s] resulted in low yields of **functionalized nanodiamond**. However, with ND annealing at ~800°, the larger coverage of diamond surface with π -bonds allowed A. Kreuger and coworkers [Jarre et al. *Chem Commun* **47** 544 2011, DOI: 10.1039/C0CC02931A] to achieve stable covalent C-C bonding of aromatic groups using Diels-Alder reactions onto surface annealed ND. The key reaction involved the addition of *o*-quinodimethane [acting as diene, generated *in situ* from *o*-bis(bromomethyl)benzene and substituted benzenes] to the π -bonds of ND in the presence of **18-crown-6** and KI in refluxing toluene for 72 hours. The substituent on the benzene ring can be **functionalised** further, e.g. with an SO₃H group (formed by sulfonation of the benzene ring) which can be converted into an SH group (PPh₃, I₂, *C₆H₆ reflux, 24 hours), and which can be made to react with *N*-dye substituted maleimide to form a condensed sulfide. The dye being fluorescent Oregon Green 48 which provided fluorescent diamond particles. The applications of such functionalisations are similar to those of functionalised fullerenes and CNTs including fluorescence labeling, drug delivery, magnetic sensing, electrochemistry, and composite materials among others.

SOLVENT RESISTANT NANO FILTERS (SRNF)

During the past decade and a half considerable effort has been expended in developing filters for separating mixtures down to the molecular level. An alternative name to SRNF is organic solvent nanofiltration (**OSN**). This shows promise as an energy- and waste- efficient process. SRNF has been developed particularly for use with organic solvents but it also applies to aqueous media, with the *proviso* that the membranes may be differ-

ent. The driving force is the pressure applied to the membranes which are classified as **UF** (ultrafiltration) with pressures of 1 to 5 bar, **NF** (nanofiltration) with pressures from 5 to 20 bar and **RO** (reverse osmosis) with pressures >10 bar. In UF substances between 2nm and 0.1 μ m are rejected, in NF particles or dissolved molecules smaller than 2nm are removed from solution, and in RO only solvents permeate.

SRNFs are essentially of two types, the organic polymer type or the ceramic type. The former are polymers which need to be cross-linked in order to increase their strength. A variety of polymers have been prepared and studied including the composites: PAN (polyacrylonitrile), PVDF [poly(vinylidene fluoride)], PI [polyimide (Matrimide) and (Lenzing P84)], PEI [poly(etherimide)], PA (polyamide), PAH [poly(amide hydrazide)], PSf (polysulfone), PES [poly(ethersulfone)], PEEK (polyether ether ketone), SPEEK (sulfonated PEEK), PPESK [poly(phthalazinone ether sulfone ketone)], SPPEK (sulfonated PPESK), CA (cellulose acetate), and PBI (polybenzimidazole). The polymers are supported by an ultra-thin (submicron) barrier (TFC, thin film composite) on top of a chemically different polymer support. The TFCs are made of a variety of substances including PDMS (polydimethylsiloxane), PEI (polyethyleneimine), PAA (polyacrylic acid), PPz (polyphosphazene) among others. The TFCs cause the membranes to be quite strong, as the membranes need to be used and reused over again.

Ceramic membranes, which are much stronger than polymer membranes, are commonly made of Al, Si, Ti or Zr oxides and mixed oxides. The top layers of the membrane are prepared by sol-gel synthesis which converts a colloidal or polymeric solution of inorganic precursors (e.g. alkoxides or salts) to a gelatinous product. Viscous binders are usually added prior to layering followed by controlled calcination and finally sintering to form the ceramic membrane. The process can be controlled to provide membranes with specific MWCOs (molecular weight cut-offs) and which can sustain defined solvents, temperatures and pressures.

Unfortunately at present not many of either type of membranes are available commercially, but undoubtedly many more will appear on the market in the near future. Commercially available SRNF include the *SelRO membranes* (Koch Membrane Systems, USA), *Starmem membranes* (Membrane Extraction Technologies, UK), *SolSep membranes* (SolSep, The Netherlands), and *Desal-5 and Desal-5-DK membranes* (GE/Osmomics, USA) designed for aqueous applications, as well as for SRNF.

SRNFs have found applications in food chemistry, catalysis, chiral separations, petrochemical industry and pharmaceutical manufacturing. The preparation, practical considerations, and theoretical transport mechanisms of SRNFs have been critically reviewed by P. Vandezande, L.E.M. Gevers and I.F.J. Vankelecom [*Chem Soc Rev* **37** 365 2008, DOI: 10.1039/B610848M].

NANO METALS AND METAL DERIVATIVES

Generally the methods for producing such nanoparticles include attrition in which the particles are ground, e.g. in a ball mill, to a fine size; pyrolysis where the precursor is vapourised and forced through an orifice at high pressure and temperature as well as using aids such as ultrasonic nozzles for spraying; by radiolytic methods with radiation chemistry; and hydrothermal synthesis.

Cadmium selenide nanoparticles [1306-24-7] **CdSe**, **M 191.4, m 1268°**; **Cadmium sulfide** [1306-23-6] **CdS**, **M 144.5, m 1750°**; and **Cadmium telluride** [1306-25-8] **CdTe**, **M 240.0, m 1041°**. For details of crystalline forms see Chapter 4 in 'Inorganic Compounds'. CdSe, CdS and CdTe nanoparticles with particles that are smaller than 10nm exhibit *quantum confinement*, in which case the electrons are restricted in a very small volume and have specific properties. Such nanoparticles are CdSe, CdS and CdTe *quantum nanodots* which can fluoresce strongly with high quantum yields.

Preparation of monodispersed CdSe semiconductor nanocrystallites [capped with trioctylphosphine (TOP see [4731-53-7] in 'Miscellaneous As, B, P, Si, S Se and Te Compounds' Chapter 3) and trioctylphosphine oxide (TOPO, see in 'Miscellaneous As, B, P, Si, S, Se and Te Compounds' Chapter 3)]: The method of Bawendi and coworkers [Murray, Norris and Bawendi *J Am Chem Soc* **115** 8706 1993, DOI: 10.1021/ja00072a025] is described here. In a vessel containing TOPO (50g) which is dried and degassed by heating at ~200°/1mm for 20min and flushing with argon is then kept at 300° under 1mm pressure of Ar. To this reaction vessel is added a mixture of Me₂Cd (1ml, 13.35mmol, see [506-82-1] below) and TOP (25ml) prepared separately in a dry-box, and a mixture of 1.0M TOPSe (10ml, 10mmol) stock solution in TOP (15ml) together all at once through a syringe with vigorous stirring. A deep-yellow orange solution develops and light absorption at 440-460nm results while the temperature drops (to ~180°), and is then restored gradually to 230-260°. Aliquots (~10ml) are withdrawn at 5-10min intervals and their light absorptions are monitored for crystal growth. Best quality cryst-

allites are formed over a period of a few hours of steady growth by regulating the growth temperature depending on size distribution as estimated from the absorption spectra; the temperature being raised if growth appeared to stop. As the desired absorption characteristics are obtained, the portions (aliquots) of the growth solution that are withdrawn through a cannula are stored in appropriate vials. Thus a series of nanocrystallites (II to IV) with $\sim 15\text{\AA}$ to 115\AA diameters is obtained. These are **purified** by cooling the 10ml aliquots to $\sim 60^\circ$, slightly above the m.p. of TOP and adding anhydrous MeOH (20ml) whereby the crystallites flocculate and are separated from the supernatant by centrifugation. The flocculant is then dispersed in anhydrous 1-butanol (25ml) followed by centrifugation which results in an optically clear solution of microcrystallites with a grey precipitate of reaction byproducts (shown to be Cd and Se by X-ray analyses) which are discarded. Addition of anhydrous MeOH (25ml) to the supernatant causes the crystallites to flocculate again and removes excess TOP and TOPO. Final rinsing with dry MeOH (50mL) and subsequent drying *in vacuo* produces $\sim 300\text{mg}$ of free flowing TOP/TOPOcapped CdSe nanocrystallites. This powder is readily dispersed in a variety of alkanes, long-chain alcohols, aromatics, chlorinated solvents and bases such as amine, pyridines as well as furans and phosphines. **Size-selective precipitation** can be achieved by dispersion in 1-butanol till optically clear, adding dry MeOH dropwise until opalescence persists on stirring or sonication, then separation of flocculant by centrifugation provides a precipitate of the *largest* crystallites in the sample. This process is repeated until no further sharpening of optical absorption is observed. This size selection can also be carried out in a variety of solvent/non-solvent pair such as pyridine/hexane or $\text{CHCl}_3/\text{MeOH}$. Similarly, **CdS semiconductor nanocrystallites** and **CdTe semiconductor nanocrystallites** can be obtained. Surface exchange, e.g. replacing the TOP/TOPO caps by pyridine, can be achieved by heating the TOP/TOPOcapped nanocrystallites ($\sim 50\text{mg}$) to $\sim 60^\circ$ with pyridine ($\sim 5\text{--}10\text{ml}$) which gradually disperses the crystallites in the solvent, and is then treated with excess of hexane and the pyridine capped crystallites are isolated by centrifugation. This hexane precipitation is repeated a number of times to produce crystallites that will disperse readily in pyridine, MeOH and aromatic solvents, but no longer in hexane. These nearly monodispersed II–IV semiconductor nanocrystallites ranging from $\sim 12\text{\AA}$ to 115\AA were characterised by TEM (Transmission Electron Microscopy) and X-ray diffraction spectra which provide a self-consistent description of the nanocrystallite structures. The physical properties have been described and discussed in detail by Bawendi and coworkers [Murray et al. *Science* **270** 1335 1995, DOI: 10.1126/science.270.5240.1335; Murray et al *Annu Rev Mater Sci* **30** 545 2000, DOI: 10.1146/annurev.matsci.30.1.545].

A similar protocol was used by Alivisatos and coworkers [Owen et al. *J Am Chem Soc* **130** 12279 2008, DOI: 10.1021/ja804414f] for preparing nanocrystalites of **CdSe capped with ODPa** (*n*-octadecylphosphonic acid) in the presence of TOPO with $\sim 4\text{nm}$ diameter, fluorescence max = 580nm , fluorescence *fwhm* $\sim 30\text{nm}$ after 7min at 315° . **CdSe/ZnS-ODPA capped crystallites** were prepared in a closed system by adding, *via* syringe Et_2Zn (1.0g, 0.49nmol Zn, see [557-20-0] below) to a mixture of TOPO (10.0g, 25.86mmol) and ODPa (0.197g, 1.2 equiv/Zn) [previously heated at $100^\circ/30\text{min}$ with vigorous stirring under Ar, then at 275° (measured with a thermocouple)] and the temperature was lowered to 250° . Dry CdSe-ODPA-capped, prepared as before, dissolved in TOP (1.0g, 2.70mmol) was then injected into the mixture under Ar, the temperature was set to 235° , followed immediately by a solution of $(\text{Me}_3\text{Si})_2\text{S}$ (1.0g, 0.246mmol S, 1 part, see [3385-94-2] in 'Miscellaneous As, B, P, Si, S, Se and Te Compounds,' Chapter 3) in TOP (100 parts) using a syringe pump (at 0.015ml/min), and allowed to cool slowly to $<100^\circ$, and the cleaning process was started. This involved injecting dry degassed Me_2CO (50ml) under argon in a glovebox into the mixture; a milky precipitate was formed, collected by centrifugation, and redissolved in CHCl_3 (15ml) to give a dark solution which readily formed a layer of foam on shaking. *n*-Octylamine (5ml) was then added followed by Me_2CO (50ml) and the white milky precipitate was removed by centrifugation leaving a thin layer of nanocrystals and a clear or cloudy supernatant. The film of nanocrystals was dissolved in hexane (15ml) and Me_2CO added until solids start to separate, which will redissolve on strong shaking or sonication but addition of more Me_2CO causes precipitation of solids, and this is repeated until no more solids separate. The solids are removed by centrifugation, discarded, and the deeply coloured supernatant was treated further with Me_2CO to precipitate the nanocrystals which were isolated by centrifugation. This step was repeated a second time to **produce CdSe/ZnS-ODPA** capped nanocrystalites which exhibited variable fluorescence quantum efficiencies (24 to 55%), and are freely soluble in hexane, toluene, CHCl_3 and other non-polar solvents. Their ^1H and ^{31}P NMR did not have sharp signals characteristic of non-bound ligands. The caps can be exchanged with MeSiCl , *n*-tridecyltrimethylammonium chloride, *S*-trimethylsilyl-2,5,8,11-tetraoxatridecane-13-thiol in toluene at $\sim 25^\circ$ to give the bis trimethylsilyl (and related) ester of ODPa and the nanocrystallites with the respective de-trimethylsilyl ligand. The caps were characterised by their ESI-MS and XPS.

Stiegerwald and coworkers [*J Am Chem Soc* **110** 3046 1988, DOI: 10.1021/ja00218a008.] have isolated and derivatised nanometer-sized semiconductor cluster molecules of CdSe by *arrested precipitation* in reverse micelles to give bare semiconductor lattices which can be modified. They prepared surfactant-stabilised CdSe nanoparticles thus: a typical AOT/H₂O/heptane microemulsion was first prepared by dissolving the dried surfactant bis(2-ethylhexyl)sulfosuccinate (AOT, 33.3g, 75mmol, see [577-11-7] below) in heptane (1.3L) and deoxygenated H₂O (4.3ml) and magnetically stirred until homogeneous (with $W = 3.2$; W was defined as the molar ratio of H₂O/AOT). Then Cd²⁺ solution (1.12ml, of stock 1.0M Cd(ClO₄)₂ 2H₂O in deoxygenated H₂O) was added with stirring until optically homogeneous (emulsion with $W = 4.0$), and was followed by injecting (Me₃Si)₂Se (210μl, 0.93mmol) in heptane (50ml) rapidly *via* syringe. As the semiconductor particles were formed, the emulsion developed colour with absorption spectra shifting to its final persistent form with time; from seconds (for large W , e.g. ~ 5.4 , red) to minutes (for small W , e.g. ~ 2 , blue). The surfactant-stabilised colloids collected were treated in three ways: (a) for further surface modification (see below), (b) evaporation to dryness or (c) for flocculation. (b) Produced coloured surfactant particles with embedded CdSe which can be dissolved in hydrocarbon solvents to give optically homogeneous solutions that are quite stable in air or room light, although some deterioration was observed in solution after ~ 8 –10 months. If the solvent was not removed as in (c) flocculation occurs within hours, but addition of large amounts of H₂O resulted in immediate flocculation and the precipitate did not re-dissolve in organic or inorganic solvents. In typical further capping preparations these authors used material in (a) and coated the surfactant-stabilised CdSe particles with Cd²⁺ by adding 0.5ml of above stock 1.0M Cd(ClO₄)₂ 2H₂O in deoxygenated H₂O, stirring till homogeneous then excess of *phenyl trimethylsilyl selenide* (350μl, 1.5mmol, see [33861-17-5] below) in heptane (50ml) was injected *via* syringe when the cloudy mixture precipitated a coloured solid which was collected by centrifugation or filtration, washed with petroleum ether and dried to give a free-flowing powder of crystallites (~ 260 mg). The yields of *phenyl capped crystallites* from preparation ($0 < W < 10$) were $\sim 55\%$ based on input mass, or $\sim 75\%$ based on Cd as limiting reagent. These should be completely soluble in pyridine; if not, the pyridine solution was filtered to remove any flocculated colloidal material from proper molecular crystallites which were then re-precipitated with petroleum ether, and the *purification* process was repeated until complete solubility in pyridine was achieved. The amount of insoluble flocculated material was an indication of the extent of capping. The particles were shown to be of molecular size (10–100 Å).

Preparation and purification of CdSe and CdTe Quantum Nano-Dots (QDs) by Chemical Aerosol Flow Synthesis (CAFS) devised by Didenko & Suslick [*J Am Chem Soc* **127** 12186 2005, DOI: 10.1021/ja054124t, PMID: 16131177]. In this beautiful new technology various combinations of Cd (e.g. CdO, CdCO, Cd naphthate or acetate) and Se (with TOP to form TOPSe *in situ*) precursors in toluene are placed in an ultrasonic nebulizer which produces an aerosol under argon flow that is allowed to flow through a tube furnace whose temperature can be altered from $\sim 300^\circ$ to $\sim 180^\circ$ where the toluene is rapidly vapourised and the CdSe-capped nano Q-dots are formed (with sizes varying with the furnace temperature) and collected by bubbling through toluene in a quencher trap. The Q-dots are then *purified* by precipitation with dry MeOH, redissolved in CHCl₃ and precipitated again with MeOH. This process is repeated at least four times to remove precursors and solvents. The size of the dots can be fine tuned by careful adjustment of furnace temperature, flowrate and precursors. The CdSe/stearic acid or hexadecylamine (HDA) particles are tunable from 2 to 4nm and the sizes are confirmed by TEM, and XRD, and fluoresce in different colours, e.g. with CAFS at furnace temperatures of 320° , 300° , 280° , 260° , 240° , 220° and 180° the fluorescences of CdSe/hexadecylamine Q-dots were red, orange, light yellow-green, green, deep green, light blue and deeper blue respectively. Experimental details of these reactions can be found in the above reference of Didenko and Suslick. Surfactant systems that proved successful include TOP/TOPO, in the presence or absence of octadecylamine, steric acid, hexadecylamine (HDA) or oleic acid (OLA). Similar procedures can be used successfully with S for *CdS Q-dots* (by using thiourea) and Te for *CdTe Q-dots* (by using TOPTe). Highly fluorescent *CdTe Q-dot* were produced by CAFS from a mixture of CdO/octadecylphosphonic acid (ODPA), octadecene and TOPTe in toluene which had quantum yields of fluorescence of 40% with fwhm ~ 30 nm. When formed in the gas phase these high quality CdTe nanoparticles could be deposited directly on surfaces to *produce solar cells, LEDs* or other *Q-dot composites*. The above authors state that CAFS can be extended to the preparation of nanostructured metals, metal oxides and other materials. and can be used for larger quantities. It is superior to the earlier batch process.

Highly luminescent capped colloidal CdS quantum dots with efficient NIR electroluminescence in light emitting diodes have been described by I.D.W. Samuel and coworkers [Bansal et al. *J Phys Chem C* **120** 1871

2016, DOI: 10.1021/acs.jpcc.5b09109]. They injected, *via* syringe, a solution of cadmium bis(octanethiol) (480mg, see preparation below) in TOP (15.4ml) under N₂ to hot TOPO (24g, heated at 120° under a 50mbar vacuum with stirring for 30 minutes then brought to 220° under N₂ for an additional 30 minutes) whereby the temperature dropped to 180°, then the temperature is raised within 5-10min to a stable temperature of 220°. At this point 5ml aliquots are withdrawn after 30, 60, 75, 90, 120, 150, 180 and 240 minutes (as the CdS QDs are being formed) and the reactions stopped by cooling the aliquots to ~20° with external cold H₂O. Each aliquot is poured into Me₂CO (40ml), kept at -20° for 20 minutes and the QDs are collected by centrifugation at 4000rpm for 20 minutes. The pelleted QDs are **purified** by dissolving them in CHCl₃ (200μl) and for duplicate analyses half portions (100μl) are treated with Me₂CO (1.2ml) and the precipitated QDs are obtained by centrifugation at 14,000rpm for 10 minutes, and the dissolution/precipitation repeated twice. The final pellets are dissolved in CHCl₃ (0.5ml) for optical and structural characterisation. In the same vain, but with some modification, CdS and cadmium bis(octanethiol) with TOP, diphenylphosphine (DPP), octadecene (ODE), and/or oleic acid (OLA), produce capped CdS QDs with these ligand. **Purification of the CdS QDs** is generally as stated above, but in some cases (e.g. with ODE) they were purified by pouring into Me₂CO (~40ml, as above) then a volume of MeCN (2-5ml) was added as soon as the solution became cloudy, centrifuged at 4000rpm for 20 minutes, the pellet was re-dissolved in CHCl₃ (0.2-0.4ml) and transferred into Eppendorfs. To each CHCl₃ volume of 0.2ml, Me₂CO/MeCN (1.2ml, 1:1 v/v) was added and the mixture was centrifuged at 14,000rpm for 10 minutes. The pellet was washed again with Me₂CO/MeCN (1.2ml, 1:1 v/v), centrifuged at 14,000rpm for 10 minutes, and re-suspended (dissolved) in CHCl₃ (0.5ml) and characterised. For the special procedure that was reported for the **fabrication of Thin Films and Devices** for analyses by see I.D.W. Samuel and coworkers (above). The **highest photoluminescence** (quantum yield 68.7 — 3.7%, PL_{max} 676 ± 12nm) was found in the **DPP capped CdS QD** in TOPO solvent, and quantum yield 34.7 ± 3.1%, PL_{max} 622 ± 17 in **OLA/DPP capped CdS QD** in neat thin films in the NIR range. deMello and coworkers [*Lab Chip* **11** 1221 2011, DOI: 10.1039/C0LC00507j] reported a versatile capillary-based droplet **microfluidic** reactor for the synthesis of **CdSe oleate NP** by highly controlled nanoparticle growth up to temperatures of 250°. This droplet reactor was very stable during one day's continuous use under constant flow-rates and temperatures producing well defined particles. It was successfully applied to the preparation of **metal (Ag) and metal oxide (TiO₂) NPs**.

Skrabalak and Brutchey [*Chem Mater* **28** 1003 2016, DOI: 10.1021/acs.chemmater.6b00472] recently reviewed all the **Continuous Flow Routes to Colloidal Inorganic Nanoparticles**, and included **microfluidic reactors** [see Laboratory on a Chip/microfluidics in Chapter 1, and gold and platinum nanoparticles below].

CdSe quantum dots have been implemented in a wide range of applications including solar cells. When CdSe nanocrystals are molecularly linked with TiO₂, the films formed are capable of harvesting light energy and are used for making **Quantum Dot solar cells** [Robel et al. *J Am Chem Soc* **128** 2385 2006, DOI:10.1021/ja056494n]. CdSe can be modified by producing two phase materials, e.g. with ZnS coating, whose surface can be further modified with e.g. mercaptoacetic acid to *confer* solubility [for review on 'CdSe nanocrystal based chem-/bio- sensors' see Somers et al. *Chem Soc Rev* **36** 579 2007, DOI: 10.1039/B517613C]. Other applications include light emitting diodes [for 'Light-emitting diodes made from CdSe nanocrystals and a semiconducting polymer' see Colvin et al. [*Nature* **370** (6488): 354 1994, DOI:10.1038/370354a0]; and M Krüger and coworkers reported that the improved efficiency of bulk heterojunction hybrid solar cells can be achieved by utilising **CdSe quantum dot-graphene nanocomposites**. [Eck et al. *Phys. Chem. Chem. Phys.* **16** 12251 2014, DOI: 10.1039/C4CP01566e.]

Cobalt [7440-48-4] Co, M **58.9**, m **1495°**, b **2870°**, d ²⁰ **8.71**, resistivity **6.24 μΩcm/20°**, Brinell hardness **125**. **Cobalt nanoparticles** for use in **magnetic fluids (MFs)** are available commercially with an average particle size of 10-12nm. It is in the form of a black powder, and is prepared by a **thermolysis process** using a strict formula. Thus, to dicobalt octacarbonyl {Co₂(CO)₈, 17.1g, 100mmol, see [10210-68-1]} under flowing argon is added Al(C₈H₁₇)₃ (4.4ml, 10mmol) [Co:Al = 10.1] dissolved in toluene (300ml) in one portion, and the mixture is stirred and heated at 110° under a reflux condenser for 18 hours. The reaction is carried out in an efficient fume cupboard because **POISONOUS** carbon monoxide evolution occurs. The colour of the solution changes to dark brown and a black precipitate separates from the clear solution. The mixture is cooled to 20°, and still under argon a further amount of Al(C₈H₁₇)₃ (1.2ml, ~30% of the initial quantity) is added, the temperature is raised to 110° and is heated for 3 hours more. The mixture is cooled to 20°, and stirred further for 16 hours. Smooth oxidation is carried out by bubbling (use a capillary) air through the mixture for 6 hours. The product is stirred overnight and the precipitated Co is allowed to settle during 2 hours. The supernatant is

decanted and the solid is isolated in **toluene wet form**, or by drying *in vacuo* to give an air stable **Co nanopowder** (5.2g); both of which can be handled under laboratory conditions. The particle size of the Co, as determined by TEM and HRTEM is $\sim 10 \pm 1.1$ nm (see further, however). The particle size is sensitive to the thermolysis conditions and to the $\text{Al}(\text{alkyl})_3$. With $\text{alkyl} = \text{Me, Et or } n\text{-octyl (C}_8\text{H}_{17})$ and a Co:Al ratio of 10:1 as above, the nanoparticle sizes are 3-4, 5-8 or ~ 10 nm respectively. Moreover, with $\text{Al}(\text{C}_8\text{H}_{17})_3$ and a Co:Al ratio of 1:2 the particle size was always ~ 5.4 nm.

Peptisation of Co nanoparticles: The Co particles are suspended in toluene and peptised by adding 2 ml of Korantin SH (BASF, *N*-oleylsarcosine + AOT, an excellent emulsifying, solubilising corrosion inhibitor), oleic acid, LP-4 (a fatty acid condensation polymer, or AOT (sodium dioctylsulfosuccinate, see [577-11-7] below) as surfactants and sonicated in an ultrasound bath for 10 minutes to give a clear dark brown **Co-MF**. After drying a sample *in vacuo* elemental analysis indicated the presence 67.8% of Co and 0.98% of Al. These MFs were found to be stable for more than 5 months [by X-ray absorption near edge structure (XANES) measurements] under air. It should be pointed out that if the $\text{Al}(\text{C}_8\text{H}_{17})_3$ is washed away from the Co particles immediately after the synthesis, the magnetic properties are markedly reduced after exposure to air. The chemical nature of the analysed nanoparticles shows that the stability of the local electronic and geometric structure is significantly affected by the surfactant used. Peptised Co nanoparticles can also be dispersed in kerosene or mineral oil as precursors for MFs.

Cobalt nanoparticles (surface modified with L-cysteine ethyl ester) ethanol wet are available as ~ 10 nm black powder (wet with EtOH) which is best used within 3 months. The material can be easily transferred into stable aqueous suspensions, and can be used as a starting material for surface modifications (e.g. dextran coating) or the preparation of magnetic polymer microspheres.

Carbon coated cobalt nanoparticles (magnetic cobalt) with particle size > 50 nm (TEM $\geq 99\%$, magnetization is > 150 emu/g, mass saturation), $6.24 \mu\Omega \text{ cm}/20^\circ$, [7440-48-4] M 58.9, [MDL number MFCD00010935] are commercially available. The carbon shell (< 8 wt%) has *ca* three graphitic layers covering the cobalt particle which can be functionalised, using the SDS/diazotisation reactions of Dyke & Tour [*Nano Lett* **3** 1215 2003, DOI: 10.1021/nl034537x, see functionalised SWCNTs above] which consequently place a variety of *p*-substituted (e.g. with Cl, NO_2 , etc) aryl groups on the carbon surface. The functionalised particles are collected *via* their magnetic properties and used for whatever purpose they are intended. [Grass et al. *Angew Chem Int Ed* **46** 4909 2007, DOI: 10.1002/anie.200700613].

Copper zinc iron oxide, 98.5% trace metal basis (zinc copper ferrite, copper zinc ferrite) [66402-68-4, 228402-49-1] $\text{CuZnFe}_4\text{O}_4$, [MDL number MFCD06200729, PubChem CID 24883132] M 416.3 d₄²⁵ 5.5g/ml. It is a nano powder with particle size < 100 nm (BET). The actual particle size is < 50 nm (XRD) and is commercially available.

Gold nanoparticles stabilised by gum Arabic (GA-AuNP). AuNPs have been widely used for a variety of purposes and water soluble composites with gum Arabic (GA) are commercially available. Gum Arabic [9000-01-5] from *Acacia* gum is a branched polymer of galactose, rhamnose, arabinose and glucuronic acid as the Ca, Mg and K salts with $M \sim 250,000$, and is a non-toxic and physiologically compatible substance. **GA-AuNP** has been developed by Kannan, Katti and coworkers [Kattumuri et al. *Small* **3** 333 2007, DOI: 10.1002/smll.200600427]. It is prepared simply by mixing NaAuCl_4 with an aqueous solution of a non-toxic phosphine amino acid $[\text{P}(\text{CH}_2\text{NHCHCH}_3\text{COOH})_3 \text{ *THPAL}]$, see preparation below] reducing agent in the presence of a 0.2% aqueous solution of gum Arabic. This resulted in an immediate reaction which produced GA-labeled AuNPs in over 98% yield. The aqueous solutions are stable for months. **Note** that if GA is omitted, the reaction is also instantaneous but the solutions agglomerate within 4-6 hours. TEM images showed that the size of **GA-AuNPs** thus produced is 15-20 nm. It is stable in the presence of cysteine, bovine and human serum albumins and does not agglomerate in 25% NaCl solution. Formulations of it can be readily administered site specifically (*intravenously*), for diagnostic imaging (**computer tomography CT**), and is bio-compatible for therapeutic applications in **nanomedicine**. GA-AuNPs are also suitable for **nanoelectronics** which include sensor design, MEMS applications, spin coating, self-assembly and formation of monolayers.

K.V. Katti and coworkers [Raghuraman et al. *J Am Chem Soc* **125** 6955 2003, DOI: 10.1021/ja034682c] prepared the reducing agent ***tris[N-(R or S)-alaninylmethylene]phosphine (*THPAL)** by adding tris(hydroxymethyl)phosphine (0.50g, 4.03 mmol, see [2767-80-8]) in H_2O (5 ml) dropwise to *R*- or *S*-alanine (1.08g, 12 mmol) in H_2O (10 ml) at 25° with stirring under N_2 for 1 hour. Evaporation *in vacuo* gave a white solid which, after washing with MeOH and drying *in vacuo* was pure product (90% yield). The ^1H NMR (300

MHz, D₂O) has δ at 1.38 (d, $J = 6.0$ Hz, 9H, NCH(CH₃)COOH), 3.47 (d, 6H, PCH₂), 3.65 (m, 3H, NCH(CH₃)COOH); the ¹³C NMR (75MHz, D₂O) has δ at 14.84 (s, NCH(CH₃)COOH), 42.65 (d, $J_{p-c} = 12.82$ Hz, PCH₂), 59.35 (d, $J_{p-c} = 5.77$ Hz, NCH(CH₃)COOH), 174.10 (s, NCH(CH₃)COOH); and the ³¹P NMR (121.5MHz, D₂O) has δ at -39.9 (s); and the ESI-MS has calculated for C₁₂H₂₄N₃O₆P 337.3, found m/z 337.6. The X-ray crystal structure analysis of the *S-enantiomer* (4 H₂O) revealed that the alanine portions were packed to form two-dimensional bilayers running parallel to (001). [See also Kannan et al. *J Am Chem Soc* **128** 11342 2006, DOI: 10.1021/ja063280c.]

Gold nanoparticles stabilised by agarose (A-AuNP). K.V. Katti and coworkers [Kattumuri *et al.* **88** 153114 2006, <http://dx.doi.org/10.1063/1.2192573>] prepared this similarly, but replacing GA by agarose. A-AuNPs were used for surface-enhanced Raman spectroscopic detection of DNA nucleosides as well as the other applications stated above.

Gold nanoparticles functionalised with dodecylthiol in solution (particle size 2-4nm DLS, 2w/v% in toluene) Au-SCH₂(CH₂)₁₀CH₃, d²⁵ 0.9308g/cm³, [MDL number MFCD09953506, PubChem CID 24884475]. It has been found useful for the self-assembly of gold monolayers (Au—SAMS) for nanoelectronics. [Fendler et al. *Chem Mater* **13** 3196 2001, DOI: 10.1021/cm010165m] Similar AuNPs with secondary amines, e.g. diocylamine, didodecylamine and dioctadecylamine which are effectively transferred to gold nanoparticles [Solwoda et al. *Langmuir* **30** 6684 2014, DOI: 10.1021/la501135q]. They are useful as a ruler for measuring the thickness of vertically self assembled gold monolayers, e.g. with vertical heights of 0.9, 1.5 and 2.1nm for matrix reference molecules of hexylthiol, dodecylthiol and octadecylthiol respectively [Kelly et al. *Langmuir* **26** 3040 2010, DOI: 10.1021/la9026128].

Gold (~1%) nanoparticles on titanium oxide extrudates (AUROLite, Au/TiO₂), on aluminium oxide extrudates (Au/Al₂O₃), on zinc oxide granulates (Au/ZnO), and on manganese oxide and carbon (Au/MnO_x/C). These are available commercially and used to catalyse some reactions. A 1% AuNP(~2nm) preparation on TiO₂ is used for catalytic purposes in ‘Green’ processes, e.g. in the presence of NaOMe in alcohols as solvents it catalyses the oxidation of aldehydes to esters using air as oxidant at ambient or lower temperatures [Marsden et al. *Green Chem* **10** 168 2008, DOI: 10.1039/B712171G]. It also catalyses the oxidation of CO and H₂ using O₂ or NO as oxidants [Walther et al. *J Catal* **260** 86 2008, DOI: 10.1016/j.jcat.2008.09.003]. Au/Al₂O₃, Au/ZnO and Au/Mn_xO/C are active catalysts for low temperature oxidation of CO and/or of CH₄ [Grisel & Nieuwenhuys *Catal Today* **64** 69 2001, DOI: 10.1016/S0920-5861(00)00510-1; Ma et al. *J Catal* **252** 119 2007, DOI: 10.1016/j.jcat.2007.08.013; see also Freund *Catal Today* **117** 6 2006, DOI: 10.1016/j.cattod.2006.05.043]. The preparations, properties and identification of these and related ‘Gold on support’ substances will be found in these references; and in Hugon et al. *Gold Bull* **41** 127 2008, WOS:000257937100007, ISSN: 0017-1557; Supansomboon et al. *Gold Bull* **41** 296 2008, Steyn et al. *Gold Bull* **41** 318 2008, Murdoch et al. *Nature Chemistry* **3** 489 2011, DOI: 10.1038/nchem.1048; and for a review on *Catalytically active gold on ordered titania supports* see Chen & Goodman [*Chem Soc Rev* **37** 1860 2008, DOI: 10.1039/b707318f].

Murphy and coworkers [Lohse et al. *ACS Nano*, **7** 4135 2013, DOI: 10.1021/nn4005022] constructed a simple **microfluidic reactor** [see Laboratory on a Chip/microfluidics in Chapter 1] which allowed the aqueous gram-scale synthesis of a variety of functionalised **gold nanoparticles** including **gold nanospheres** with controlled core diameters, **gold nanorods** (aspect ratios ~1.5 to 4.0) and **gold nanocubes** which can be tailored to requirements. The reactor can be interfaced with existing purification and monitoring techniques to produce high-throughput functionalisation/purification of gold nanoparticle products. A system which could be applied to the preparation of *monodispersed, hydrophilic functionalised metal nanoparticles* and be an integral part of chemical nanotechnology research laboratories. Marmstadt and coworkers [Lazarus et al. *Lab Chip* **10** 3377 2010, DOI: 10.1039/C0LC00297f] had also described a **microfluidic platform** for the flow-focused synthesis of monodispersed gold nanoparticles (4.38 ± 0.53nm) rapidly and reproducibly using an imidazolium ionic liquid.

Indium-tin oxide nanopowder (ITO) [50926-11-9] M 264.9, surface resistivity of 100Ωcm. Available in particle size <50nm. It is coated on glass for making transparent electrodes for screens (e.g. touch screens). However, graphene electrodes have better transparency and are tougher (see above) [Bae et al. *Nature Nanotechnol* **5** 574 2010, DOI: 10.1038/nnano.2010.132].

Lead sulfide (Galena) [1314-87-0] PbS, M239.3, nanocrystals (NCs), (see PbS in ‘Inorganic Compounds’, Chapter 4). The following synthesis is a literature procedure [for PbS NCs: Hines & Scholes *Adv Mater* **15** 1844 2003, DOI: 10.1002/adma.200305395] as described by R.D. Robinson and coworkers [Zhang et al. *Nano Lett* **11** 5356 2011; DOI: 10.1021/nl202892p] for preparing **6nm PbS NCs**. To a mixture of PbO (0.45g) and oleic acid (20ml, OLA, see [112-80-1] in ‘Aliphatic Compounds’, Chapter 3) which is heated under N₂ at 150° for 15min until clear, the temperature is then decreased to 110°, evacuated to remove any H₂O and then reheated to 130°; was added BTSS-ODE [10ml, of a mixture of bis(trimethylsilyl)sulfide (252ml, BTSS, see [3385-94-2] in ‘Miscellaneous As, B, P, Si, S, Se and Te Compounds’ Chapter 3) in octadecene (12ml, ODE, see [112-88-9] in ‘Aliphatic Compounds’, Chapter 3)] via a syringe in a glovebox and allowed to react for 40 seconds. The mixture is rapidly cooled with H₂O and then EtOH is added to precipitate the NCs which are separated by centrifugation and purified twice by dissolving in hexanes and precipitating with EtOH. The size of PbS NCs can be controlled by changing the reaction temperature and time. PbSe (6.3 ±0.6nm), CdSe with oleate ligand (4.7 ±0.4nm), CdSe with stearate ligand (8.1 ±1nm), CdS with oleate ligand (3.5nm ±0.3nm), CdS with oleate ligand (7.6 ±0.7nm) and CdSe/CdS coreshell with oleate NCs were synthesised in a similar way.

A new procedure for de-capping was discovered where treating with 0.004M (NH₄)₂S in MeOH for 30 seconds, then washing excess salt with MeOH which removed the ligands completely, and converted the NC metal-rich shells into metal sulfides. The now bare metal sulfide NCs were shown to have metal sulfide bonding to form a larger NC film assembly and still retain quantum confinement. This makes these new NCs promising new materials for **electronic** and **photoelectronic applications**. The capped and de-capped NCs were characterised in detail by elemental analysis, FTIR, optical spectroscopy, X-ray photoelectron spectroscopy (XPS), wavelength dispersive X-ray spectroscopy (WDS), small angle X-ray scattering (SAXS), High-Angle Annular Dark-Field (HAADF) imaging and Electron Energy Loss Spectroscopic (EELS) mapping in an aberration-corrected electron microscope. [Zhang et al. *Nano Lett* **11** 5356 2011; DOI: 10.1021/nl202892p.]

In elegant work, Tisdale and coworkers [Wiedman et al. *ACS Nano* **8** 6363 2014, DOI: 10.1021/nn5018654] found that the Pb to S ratio of the precursor during nucleation is a critical parameter that affects growth and monodispersity of PbS—NCs. They succeeded in synthesising highly monodispersed PbS—NCs (of size dispersivity <5%) over a range of sizes and exciton energies in the range 0.70—1.25eV (1000—1800nm), without use of size-selective precipitations. Ensembles close to the homogeneous limit with absorption peak half width at half maximum values (HWHM) as small as 20meV were obtained. Correspondingly narrow photoluminescence emission that have small Stokes shifts and quantum efficiencies of 30-60% were achieved. These PbS—NCs are stable in air for several months and readily self assemble into ordered superlattices.

The synthetic procedures described briefly here are along the lines developed by Tisdale and coworkers (see above). PbS—NCs have been synthesised at high temperature 120° and at the low temperatures of 40° and 80°. All work is done using air-free Schlenk techniques (see Chapter 1) in N₂ atmospheres. For the **120° synthesis** the Pb solution is prepared by adding PbCl₂ (2.50g, 9mmol) to oleylamine (7.5ml), degassed for 10min (Schlenk vacuum line) with stirring to a pressure of ~150mm, then re-pressurised with N₂ and the temperature set at 120° (using a controller). A sulfur solution is prepared by heating a mixture of S (0.040g) with oleylamine to 120° and while stirring, N₂ is bubbled through for 20 minutes to give a clear amber coloured solution, then cooled to ~25° while still under N₂. To the Pb solution at 120°/N₂, is swiftly injected 0.357ml of the above sulfur solution, whereby a nucleation burst takes place with immediate change from colourless to black occurs. The solution is kept at 120°, while ~1ml aliquots (syringe) are removed at various growth times so as to track the reaction progress and provide increasing number of data points per synthesis. The aliquots are then treated as described below. At the end of the synthesis, i.e. when as many aliquots as required are collected, heating is removed and the reaction is quenched by injecting cold hexanes (20ml), and the reaction flask immersed in a water bath. A typical synthesis yield is ~75mg of product (see further for isolation). The process can be scaled up 10-fold with the proviso that the injection of the S solution is as close to instantaneous as possible. This necessitates the reduction of the S solution volumes by concentrating (e.g. to ~5ml) instead of injecting large volumes in the scale up. The procedure is for a 24:1 Pb/S ratio, for other ratios the weight of Pb precursor is altered but that of the S precursor is unaltered. For the **40° and 80° syntheses** the Pb solution is prepared as above at 120° then allowed to cool to the lower temperatures before injecting the S solution. In this case the nucleation burst takes a few seconds longer to occur.

Purification of the PbS—NCs: (a) with alcohols anti-solvents. The above products are set aside overnight when white PbCl₂ precipitates out and is removed by centrifugation (4000rpm/30min). The quantum dots (QDs) are then precipitated from the supernatant by adding alcohol anti-solvents (5ml MeOH and 10ml BuOH), centrifuged as before and the supernatants are discarded. The QDs pellets are dried for several hours before re-

dispersing in hexanes (20ml). If turbid, centrifuge off any PbCl_2 that separates, then perform ligand exchange for better stability of the QDs. This is done by adding oleic acid (2ml) to the hexane dispersion and agitate for 1min, and precipitate the QDs by adding alcohol anti-solvents, collecting the QDs by centrifugation and drying them. The oleic acid exchange and precipitation is repeated once more time before finally dispersing the QDs in the desired solvent.

(b) with oleic acid. The products from synthesis are precipitated by addition of butanol (10ml) and MeOH (5ml). The precipitates are collected by centrifugation as above and re-dispersed in hexanes (10ml) and 100% oleic acid by volume is added, the mixture is agitated for several minutes, and the QDs are precipitated *via* centrifugation as above. The supernatants may contain unreacted S (orange colour). The pellets are allowed to dry for several minutes, and the precipitation with oleic acid is repeated 3-5 times until the supernatant is colourless. The QD pellets are then re-dispersed in hexane and precipitated with BuOH and MeOH to remove unbound oleic acid to the nanocrystalline QDs. Finally the QDs are redispersed in the desired organic solvent and centrifuged if any PbCl_2 is present (as indicated by some turbidity).

The ~1ml *aliquots* isolated at intervals of time are quenched with cold hexanes (5ml). Their stabilities are less robust those of the above QDs, and as before they are kept in the centrifuge tubes overnight, centrifuged the following day to remove PbCl_2 , the QDs are then precipitated with anti-solvents (~3ml BuOH and ~2ml MeOH) and centrifuged. The pellets are dried for several hours, the QDs are redispersed in hexanes and oleic acid (0.5ml) is added. Centrifuge if turbid, and precipitate the QDs with the alcohols, dry the pellets for several hours and finally redisperse in the desired organic solvent. The QDs are now ready for physical evaluation, i.e. absorption spectra from which the HWHM values are obtained for all QDs of the aliquots, by TEM images, solution phase small-angle X-ray scattering (SAXS), X-ray diffraction (XRD), X-ray photoelectron spectroscopy (XPS), wide angle X-ray scattering (WAXS) and obtaining the grazing-incidence small-angle X-ray scattering (GISAXS) patterns.

$\text{Pd}^{(0)}$ EnCatTM is obtained by reducing the $^*\text{EnPd}(\text{OAc})_2$ (0.4g, 0.4mmol/g, see below) in Et_2O (8ml) with formic acid (8ml) at 45° for 2 hours. The mixture is cooled and the solid microcapsules are filtered through a polyethylene frit (20 micron porosity), washed with distilled H_2O (3 x 30ml), Me_2CO (3 x 30ml) and Et_2O (3 x 30ml). The microcapsules are dried (at <0.5mm) at 45° for 5 hours to provide black **$\text{Pd}^{(0)}$ EnCat** microcapsules (0.35g, containing Pd nanoparticles of ~2nm size). It is an efficient catalyst for reductive ring-opening of epoxides to the respective alcohols which occurs in ~99% yields. The reaction requires either $\text{HCOOH}/\text{Et}_3\text{N}$ or H_2 as hydrogen donors, and the catalyst can be recycled ~10 times [Ley et al. *Org Lett* **5** 4665 2003, DOI: 10.1021/ol0358509].

$^*\text{EnPd}(\text{OAc})_2$ [Pd EnCatTM] is prepared from polymethylene polyphenylene di-isocyanate (SUPRASEC 5025, average functionality of 2.7) and $\text{Pd}(\text{OAc})_2$ dissolved in CHCl_3 and then dispersed at 800rpm for 1 minute into an aqueous solution of Na lignosulfonate (Reax 100M), polyvinyl alcohol (Gohsenol GL 03) and the polyoxyethylene ether of butanol (Tergitol XD), using a mechanical overhead stirrer with a rotary blade. The resulting emulsion would have a particle size range of 20-250microns and the mixture is then gently shaken for 16 hours. The solid polyurea microcapsules formed are collected on a polyethylene frit (20 micron porosity), washed with de-ionised H_2O (5 x 50ml), Me_2CO (5 x 50ml), Et_2O (3 x 50ml) and dried at room temperature. Encapsulated Pd catalysts such as **Pd EnCatTM** are available commercially. These particles are defined by their matrix content e.g. 30% or 40%, the latter having the smaller pore size. [See also Bremeyer et al. *Synlett* 1843 2002, DOI: 10.1055/s-2002-34862; Yu et al. *Chem Commun* 678 2003, DOI: 10.1039/B300074P; Vickerstaffe et al. *Org Biomol Chem* **1** 2419 2003, DOI: 10.1039/B305713E.]

Platinum, gold and palladium nanoparticles on spherical polyelectrolyte brushes (SPB). Pt crystals (2-3nm) were deposited on SPB using the procedure of M. Ballauff and coworkers [Schrinner et al. *Science* **323** 617 2009, DOI: 10.1126/science.1166703]. The SPB consists of a polystyrene core (~100nm diameter) onto which a cationic polyelectrolyte ($+\text{H}_3\text{NCH}_2\text{CH}_2\text{OOC}-$)_n is attached [Schrinner et al. *Adv Mater* **20** 1928 2008, DOI: 10.1002/adma.200702421]. A mixture of AuCl_4^- and PtCl_6^{2-} was reduced with NaBH_4 on the surface of the cationic SPB to form alloy nanoparticles ($\text{Au}_{45}\text{Pt}_{55}$ ratio) on the surface. The Au was removed with cyanide and O_2 . Thus a solution of 1.9×10^{-5} M NaCN (4ml) was added dropwise within 25 minutes to a stirred suspension of AuPt-SPB (0.04 weight %) at room temperature under air. High dilution of the cyanide is crucial for avoiding complete dissolution of the NPs and coagulation. Air was bubbled through the solution to remove completely the Au atoms. After 3 hours the colour of the solution was blue due to the formation of pure Pt-NPs. [Schrinner et al. *Macromol Chem Phys* **208** 1542 2007, DOI: 10.1002/macp.200700161.] Detailed examination

of the composite particles by cryo-TEM, wide angle XRS and HR-TEM showed that the Pt-NPs are well-defined faceted single crystals embedded in the SPB-chain layer. This composite system has excellent colloidal stability, and high catalytic activity with turn over numbers of as high as $1580 \pm 50 \text{ sec}^{-1}$ for the reduction of *p*-nitrophenol to *p*-aminophenol using NaBH_4 , which is among the higher ever observed for this reaction.

Similar **Palladium (SPB)** nanoparticles encapsulated in spherical polyelectrolyte brushes and Core-Shell microgels have been prepared in much the same way by Ballauff and coworkers [Mei et al. *Chem Mater* **19** 1062 2007, DOI: 10.1021/cm062554s]. The metal in a variety of composites exhibits powerful catalytic activities [as judged by the rates of catalytic reduction of *p*-nitrophenol with NaBH_4 to start the reaction, that are exceedingly fast (see above)] and the rates depended on the composite material and its shape. Comparison of the catalytic activities of these composites depended on the metal and showed that they increase in the order $\text{Au} < \text{Pt} < \text{Pd}$ [see Schrinner et al. *Macromol Chem Phys* **208** 1542 2007, DOI: 10.1002/macp.200700161]. [See also Zaera and coworkers in Lee et al. *Nature Materials* **8** 132 2009, DOI: 10.1038/nmat2371; and for a review see H. Birch *RSC Chemistry World* **6** (3) 29 2009.]

Microfluidic (see Laboratory on a Chip/microfluidics in Chapter 1) **continuous platforms** have been devised by Riche, Roberts and coworkers [*Nat Commun* **7** 10780 2016, DOI: 10.1038/ncomms10780, PMCID: PMC4766398] whereby three dimensional droplet generating devices with robust flow invariant behaviour in which a droplet generator produces droplet volumes spanning four orders of magnitude can be used. With a parallel network, the platforms can be made to synthesise **platinum nanoparticles** reproducibly using an ionic liquid solvent, and after recycling an ionic solvent the yield of the reaction was doubled compared with similar batch syntheses.

Rhodium nanoparticles entrapped in Rh/AlO(OH) matrix (~5wt% Rh loading). The catalyst is prepared from a mixture of $\text{RhCl}_3 \cdot x\text{H}_2\text{O}$, $\text{Al}(\text{secBuO})_3$ and 2-butanol [compare with Ru/AlO(OH) below] at 100° for 3 hours, aging in air for 1 day, filtering, washing the solid with Me_2CO and drying in air at 25° . It is estimated by high resolution transmission electron microscopy (HRTEM) and X-ray diffraction that the Rh particle sizes in the matrix are 2.5-3.0nm. It is an efficient catalyst for the hydrogenation of arenes, e.g. benzene, naphthalene, quinoline, at low pressures of H_2 (~1 atmosphere) and room temperature in yields approaching 100% and with high turn over, with *ca* 1 mol% of Rh in the catalyst. [Park et al. *Chem Commun* 5667 2005, DOI: 10.1039/B511577A].

Ruthenium (nanoparticles) in aluminium oxide/hydroxide (~2.5wt% Ru loading). In this catalyst, Ru is encapsulated in an aluminium oxy-hydroxide matrix where the metal has been shown by energy dispersive X-ray analysis (EDX) and X-ray photoelectron spectroscopy (XPS) to exist mainly as Rh(0) in the matrix. It is prepared in a one-pot synthesis through nanoparticle generation and gelation. Thus $\text{RuCl}_3 \cdot x\text{H}_2\text{O}$ (52mg, 0.25mmol) and $\text{Al}(\text{secBuO})_3$ (2.5g, 10mmol) in EtOH (1.2ml, 20mmol) under a reflux condenser are heated, with stirring, at 100° for 1 hour, forming a black suspension to which H_2O (2.0ml) is rapidly added. The mixture is stirred for a further 30 minutes at 100° , filtered, the black solid is washed with Me_2CO and dried in air at 25° to give the catalyst as a grey powder (0.74g, 2.5wt% of Ru). It is used as a recyclable catalyst for efficient oxidant-free alcohol dehydrogenation see Chapter 5, Catalysts-Part 1. [Kim, Park and Park *Org Lett* **8** 2543 2006, DOI: 10.1021/ol060750z.]

Ruthenium nanoparticles stabilised in polymer ligands. A general procedure is used for making these colloids (agglomerates) which involve introducing ruthenium (1,5-cyclooctadiene)(1,3,5-cyclooctatetraene) (158mg, 0.5mmol, $[\text{Ru}(\text{COD})(\text{COT})]$ see [127382-91-6]) in a Fischer-Parr bottle and left *in vacuo* for 30 minutes, then THF (25ml, degassed by freeze-pump cycles) is added and the yellow solution is cooled to -80° , after which the ligand (0.1mmol) in THF (~60ml) is introduced to the bottle. The bottle is pressurised under 3 bars of H_2 and allowed to warm to room temperature. After ~20 hours a brown-black suspension is obtained. Complete decomposition of the Ru complex is checked by eliminating the H_2 (e.g. by blowing N_2 through) and taking a 3ml aliquot through a small Al_2O_3 column when the filtrate should be colourless. The volume of the solution is then reduced to *ca* 15ml, pentane (50ml) is added, the resulting mixture is cooled to -80° at which temperature a brown precipitate is formed. This is filtered off (or centrifuged), washed with pentane and dried *in vacuo*. The ligands used are PVP (poly vinylpyrrolidone), *n*-alkyl (C8, C12 and C16) amines and *n*-alkyl (C8, C12 and C16) thiols in varying amounts (from 0.2 to 1 equivalent) giving Ru percentages of ~55 to 80%. The colloids are characterised by microanalysis, IR after CO absorption, high resolution EM and wide-angle X-ray scattering. In THF and in a polymer matrix (Ru/polymer ~5%) crystalline particles of uniform mean size

(1.1nm) and agglomerated particles (1.7nm) are thus obtained in PVP and cellulose acetate. The reaction with various concentrations of alkyl amines and alkyl thiols lead to agglomerate particles or particles dispersed in solution both displaying mean size of 2-3nm. In the case with the amine ligands the particles are generally elongated with the tendency of forming worm or rod like structures at high amine concentrations, whereas ^{13}C NMR studies show that the stabilised particles are not fluxional, but thiol groups are oxidised to disulfide groups probably by the Ru surface. Because of their chemically reactive groups they are potentially very useful in nanotechnology. [Pan et al. *J Am Chem Soc* **123** 7584 2001, DOI: 10.1021/ja003961m]. In a similar synthesis ruthenium nanoparticles are stabilised by organosilane fragments, by using *n*-octylsilane, giving particles of ~2.3nm diameter which is a very narrow distribution of size. They were characterised by TEM, as well as solid state ^{13}C MAS NMR. [Pelzer et al. *Chem Mater* **16** 4937 2004, DOI: 10.1021/cm049086b.]

Silver acetate [563-63-3] see also entry in Chapter 5, 'Catalysis-Part 1'. Among the current intense activities in the synthesis of hybrid materials, where an inorganic phase is present as nanometer-sized particles dispersed throughout an organic matrix, is the preparation of a *ca* 190nm uniform film of exceptionally reflective and surface conductive silver 'mirror' on a flexible poly(amic) matrix. The Ag-containing resin solutions are made first by dissolving AgOAc in a small volume of dimethylacetamide (DMAc, [127-19-5]) by adding 1.35 equivalents of 1,1,1-trifluoro-2,4-pentanedione (TFAH). As AgOAc is not very soluble in DMAc, addition of TFAH converts the acetate salt into the much more soluble AgTFAH complex. The poly(amic) matrix is prepared by co-polymerising 3,3',4,4'-benzophenonetetracarboxylic anhydride (BTDA, [2421-28-5]) with 4,4'-oxydianiline (APE, 4,4'-diaminodiphenyl ether, [101-80-4]), and making a 15%w/w solution in DMAc by stirring for 5 hours to give a viscosity of 1.7-1.8 dL/g at 35°. The poly(amic) solution is added to the Ag solution to give the desired Ag to BTDA/APE polymer ratio of 10.7 to 13.0% Ag, cast into films (onto soda lime glass plates) to give cured films of 20-25 mm thickness. Air (10% relative humidity) is slowly flowed over the film for 18 hours, then thermally cured in a forced air oven with a cycle involving heating at 135° for 20 minutes, holding for 1 hour, heating at 300° for 4 hours, and holding at 300° for 7 hours. This process reduces Ag(I) to the metal and releases all the volatiles. Uniform nanometer thin films of Ag metal (see above) are thus formed in the polymer matrix. Characterisation by X-ray diffraction, transmission (TEM) and scanning (SEM) microscopy, tapping mode atomic force microscopy, X-ray photoelectron microscopy (XPS), and conductivity, reflectivity, thermal and mechanical measurements showed that the films have excellent properties, and are stable in air below 325°. This '**film-to-film**' method minimises the amount of Ag used. The applications for these Ag-copolymer silvered films with exceptional reflectivity, thermal, and surface conductivity are immense. They include the fabrication of lightweight optical mirrors, sunshields (e.g. in space telescopes), radiofrequency antennas for management of electromagnetic signals in space, Ag coated tubing to deter catheter-induced urethritis, elastomeric devices, concentration of solar energy, flexible conductive patterned surfaces and tapes. [Southward et al. *Chem Mater* **11** 501 1999, DOI: 10.1021/cm981014v.]

Silver nanoparticles on spherical electrolyte brushes (SPB). Compare these with the above Pt-NPs on SPB. They were also developed by M. Ballauff and coworkers [Lu et al. *J Phys Chem C* **111** 7676 2007, DOI: 10.1021/jp070973m]. In this case Ag nanoparticles were synthesised onto the polystyrene (PS) core—poly(acrylic acid) (PAA) SPB particles *in situ* by **photoemulsion polymerisation**. The PS core with a thin layer of photo initiator {2-[*p*-(2-hydroxy-2-methylpropiophenone)]-ethylene glycol-methacrylate} was made by standard emulsion polymerisation procedures also developed by M. Ballauff and coworkers [Guo et al. *Macromolecules* **32** 6043 1999, DOI: 10.1021/ma990609o]. The PS-PAA-Ag composite particles were prepared *in situ* by mixing diluted PS core solution (1 wt %) with a defined quantity of functional monomer silver acrylate (30 mol% of the amount of styrene), degassing and refilling with N_2 (5 times). Then the mixture was irradiated under N_2 with UV-VIS radiation at ~25° for 90 minutes while being vigorously stirred to keep the mixture homogeneous. Coagulum was removed by filtration through glass wool and the Ag nanocomposite particles were washed by dialysis against purified H_2O using a cellulose nitrate membrane (100nm pore size). DLS showed that the polyelectrolyte brushes were on the PS core; TEM, Cryo-TEM, and wide-angle XRS revealed that the AgNPs in the brushes were crystalline with a diameter of $3 \pm 1.2\text{nm}$. This **Ag-composite** is stable and has very good catalytic activity. Reduction of *p*-nitrophenol using NaBH_4 is very fast, albeit slightly slower than with Pt or Pd immobilised on related carriers.

Silver nanoparticles [AgNP], on gum Arabic, gelatin or starch. Available as yellow-brownish liquids. They are water soluble in the form of 5-10nm spheres (10-15nm for starch) and have UV with λ_{max} at 405-410nm.

The gum Arabic (GA), gelatin and starch stabilise these particles and can be used for sensor design applications, for *in vitro* as well as *in vivo* antimicrobial and antifungal purposes. Among the useful AgNPs are the commercially available **decanethiol-functionalised silver nanoparticles Ag-SCH₂(CH₂)₈CH₃**, in 3-7nm size (TEM) as a 0.1w/v & in hexane with 0.075-0.125% loading, **d²⁵ 0.672g/cm³**, [MDL number MFCD08705344]. Also available are **PVP(polyvinylpyrrolidone)-functionalised silver nanoparticles** with 10nm average size as 0.02mg/ml solutions in H₂O. Apart from their anti-microbial and anti-fungal properties, the AgNPs have optical colour properties that depend on their shape and size brought about by their attachments. Their strong coupling with the wavelengths of incident light has been used to produce highly sufficient thermal absorbers and ultrabright reporter molecules with nanoscale *antennae* which could amplify the strengths of local magnetic fields.

Silver Nanostrings: Nanostrings are a way of enhancing the electrical connection between two nanoelectrodes. The formation of necklace-like silver strings can be achieved as follows: Mix thoroughly Triton X-100 [92046-34-9] [4-(C₈H₁₇)C₆H₄(OCH₂CH₂)_{n-10}OH] (2ml, of 0.5 x 10⁻²M) with AgNO₃ (0.2ml of 10⁻²M) and ascorbic acid AA (5ml of 5.68 x 10⁻²M) in a plastic or quartz cuvette and expose to UV light (from a germicidal 15W lamp) for 10min to generate a green sol. [A more powerful lamp requires less exposure time.] Ag₂SO₄ or AgClO₄ give similar reactions. The green sol can also be prepared in gelatin (2ml, 0.001%) with AgNO₃ (0.2ml of 10⁻²M), and ascorbic acid AA (20ml of 5.68 x 10⁻²M) under UV light for 20-30 minutes, and the cuvettes in all cases are kept at a distance of 3cm from the UV lamp. These concentrations and conditions are critical. In the absence of ascorbic acid (AA) no green sol was formed, at lower amounts than stated a yellow sol of silver is formed, and at very high concentrations of AA (e.g. >1.7 x 10⁻¹M, with TX-100 or gelatin) the solution becomes turbid and forms larger yellow to black particles. Slightly longer UV exposure times are not deleterious. At high concentrations of TX-100 (>10⁻²M) or gelatin (>0.002%) the linear aggregation of silver sol is inhibited and a pink silver particle (~50nm) sol is produced. A very low concentration of TX-100 or gelatin produced a yellow sol which is not stable and precipitates black silver particles on standing for ~24hrs. The UV spectrum of the green sol has bands at ~400nm (high absorbance) and a band at ~630nm of lower intensity. The latter band is **indicative of nanostring formation** (solutions of other colour did not possess this band) which is confirmed by studies of fluorescence quenching of 1-aminonaphthalene, TEM (transmission electron microscopy), SERS (Surface Enhanced Raman Scattering) spectral effects (e.g. enhancement of the Raman signal of pyridine) and TAFM (Tapping Mode Atomic Force Microscopy). The Ag-nanostrings, which have similarities with DNA strings, are composed of particles as small as 3-5nm covered with 0.1-0.2mm thick layers of micelle or polymer. **Note** that SERS studies predict that linear aggregates of Ag will be produced in a green coloured sol. [Pal et al. *J Mater Sci Lett* **18** 1391 1999, DOI: 10.1023/A:1006659021058.]

Silver-tin alloy nanoparticles [MDL number MFCD00801108, PubChem Substance ID 24885373] **Ag-Sn₂₅, M 118.3**. These particles are available in <150nm in size and contain 3.5% Ag with the formula Ag_{0.035}Sn_{0.965}. They have a surface area >5.4 m²/g. For the *Synthesis of tin, silver and their alloy nanoparticles for lead-free interconnect applications* see Hongjin Jiang: PhD thesis April 2008, School of Chemistry and Biochemistry, Georgia Institute of Technology, GA, USA, <https://smartechn.gatech.edu/bitstream/handle/1853/22636/Jiang>, and is available in public domain.

Stannic oxide nanopowder [18582-10-5] **Sn(IV)O₂** [MDL number MFCD00011244, PubChem Substance ID 24874714], **M 150.7, d²⁵ 6.95g/cm³**. It is available in particle size of <100nm, and is used for polishing metal or glass, used in coloured glass, enamels, as a mordant for colouring fabrics and in battery manufacture. It is toxic and should not be inhaled. The oxide is prepared by precipitation from tin acetate solution in AcOH, upon addition of 12.5% aqueous ammonia solution with stirring to cool it. The precipitated sol is centrifuged down, washed twice with de-ionised H₂O, then dried at 120° for 15 hours. The total impurity content in the original acetate used did not exceed 1wt%. For *Studies of thermal stability of nanocrystalline SnO₂, ZrO₂, and SiC for semiconductor and thermocatalytic gas sensors and comparison of long-term stability of the sensors' signal with the thermal stability of the nanopowders* see Pavelco et al. [*Russ J Electrochem* **45**(4) 470 2009, DOI: 10.1134/S1023193509040181].

Zinc oxide nanoparticles [10139-47-6] **ZnO, M 81.4**. The preparation of ZnO-nanoparticles can be carried out as follows: A solution of Zn(CH₃CO₂)₂·2H₂O (0.10g) in isoPrOH (25ml) is added to isoPrOH (125ml) in a bath at 65° and chilled, then 0.050M NaOH (15ml in chilled isoPrOH) is added rapidly with stirring *via* a pipette. After addition is complete aliquots are withdrawn (e.g. after 1, 3, 5, 10, 15, 30, 60 and 120 minutes) and

the UV spectra are measured. A series of spectra between 303 to 375nm are thus obtained each having a steep rise in absorption at the shortest wavelength. From the x-axis intercepts of the tangents to the steep rise in absorptions, the average radii of the nanoparticles can be calculated. The longer the sampling time, the larger the nanoparticle sizes that are formed. [For Growth kinetics and modeling of ZnO nanoparticles see Hale et al. *J Chem Educ* **82** 775 2005, DOI: 10.1021/ed082p775.] It should be possible to collect the nanoparticles by centrifugation for 10min, wash three times with de-ionised H₂O followed by two washes with anhydrous MeOH (all by centrifugation), and dried in an oven at 100° [as described by Hasanpoor et al. who used a microwave method *Procedia Material Science* **11** 320 2015, DOI: 10.1016/j.mspro.2015.11.101].

In a **room temperature synthesis of ZnO nanoparticles** Bagabas, Alshammari, Aboud and Kosslick [*Nano Lett* **8** 516 2013, DOI: 10.1186/1556-276X-8-516.] prepared nanoparticles which were effective catalysts for the photochemical destruction of cyanide, a very poisonous ingredient and/or product in the chelating agents, pharmaceutical, electroplating and mining industries. At 25°, cyclohexylamine (60mmoles, see [108-91-8] in 'Alicyclic compounds', Chapter 3) in H₂O (20ml) was added to a solution of Zn(NO₃)₂·6H₂O (30mmoles) in H₂O (60ml) with magnetic stirring followed by more H₂O (80ml), which formed a white precipitate that was stirred for 4 days. The precipitate was filtered (F-size fritted filter) off, washed with Milli-Q H₂O (100ml) and dried for 1 day *in vacuo*. It was then suspended in Milli-Q H₂O (300ml) to remove further impurities, magnetically stirred for 1 day, filtered off again and dried at 25° *in vacuo* to give **pure oxide** (2.43g 89.7% yield). Inductively coupled plasma (ICP) elemental analysis of this uncalcined oxide showed that it had the formula ZnO·0.5H₂O. This was calcined by raising the temperature at the rate of 1°/min up to 500° in air and maintained at this temperature for 3hrs, and was designated ZnO_w.

A similar reaction carried out in EtOH medium gave 2.572g (98.1% yield) of ZnO·0.33H₂O (by CIP) and after similar calcination was designated ZnO_E. Uncalcined and calcined ZnO_w and ZnO_E were subjected to various analyses including: surface areas (BET-SA) and pore size distribution, diffuse reflectance infrared Fourier transformation (DRIFT), FTIR, X-ray diffraction (XRD), morphology studies with field-emission scanning electron microscopy equipped with an energy-dispersive X-ray (EDX), high-resolution transmission electron microscopy (HRTEM), solid state UV-Vis and UV/Vis/NIR spectroscopy.

Analytical studies showed that ZnO_E and ZnO_w are nanocrystallites respectively with: Ave area 7.5 and 12.41 m²/g (BET-SA); pore size 0.02 and 0.05 cm³/g; Ave crystallite 28.8nm (15.6nm uncalcined) and 28.8nm (33.9nm uncalcined); 0.281nm (0.263nm uncalcined) and 0.263nm (0.262nm uncalcined) for d-spacing from HRTEM; 0.281nm (0.260nm uncalcined) and 0.260nm (0.260nm uncalcined) for d-spacing in bulk ZnO; 100hkl(002hkl uncalcined) and 002hkl(002hkl uncalcined) for Miller indices assignment.

The catalytic photodecompositions of cyanide kinetics were performed in aqueous KCN (100ppm, 300ml) made to pH 8.5 with aqueous NH₃ at 25° by UV light irradiation at 365nm (18W lamp) with 0.01, 0.02, 0.03, 0.05, 0.07 or 0.09wt% of suspended calcined ZnO and aliquots taken at time intervals (0 to 360min), and the CN⁻ ion concentration determined by volumetric titration with standard AgNO₃. ZnO_E is a more efficient catalyst than ZnO_w by a factor of 1.5 at catalyst concentration of oxide = 0.02%. Increasing the catalyst concentration from 0.01 to 0.09 increased the photolytic decomposition efficiency from 85% to 100% after 180min with a doubling of the *pseudo* first order rate constant. The rate constants at catalyst concentrations of 0.01 and 0.07wt% were 19.2 x 10⁻³ and 42.9 x 10⁻³ min⁻¹ respectively.

M.R. Hoffmann and coworkers [*J Phys Chem* **91** 3789 1987, DOI: 10.1021/j100298a015] had similarly described the preparation of **ZnO quantum sized nanoparticles** by precipitation from Zn(OAc)₂ solution in H₂O, 2-propanol or MeCN with 80% of the stoichiometric amount of NaOH. They characterised the particles in detail by spectroscopy studies. [For 'A comprehensive review of ZnO materials and devices' see Özgür et al. *J Appl Phys* **98** 041301 2005, DOI: 10.1063/1.1992666; and Klingshirn in 'ZnO: Material, Physics and Applications' *Chem Phys Chem* **8** 782 2007, DOI: 10.1002/cphc.200700002; also see ZnO [10139-47-6] in 'Inorganic Compounds', Chapter 4.]

SOME ORGANIC AND METAL-ORGANIC COMPOUNDS USED IN NANO-TECHNOLOGY

1-Adamantane thiol (tricyclo[3.3.1.1^{3,7}]decane-1-thiol) [34301-54-7] C₁₀H₁₆S, M 168.3, m 95-97°, 100-102°, 99-106°, pK ~11.2. This thiol was prepared by various methods namely by formation of 1-adamantylisothiourinium bromide from 1-adamantylbromide followed by alkaline hydrolysis [Geigy A.-G. Belg. Pat 629,370 1963, *Chem Abstr* **60** 9167 1963], by refluxing 1-adamantanol with Lawesson's reagent (see

[19172-47-5]) in toluene [Nishio *JCS Perkin Trans I* 1113 1993, DOI: 10.1039/P19930001113], and by photolysis of *N*-(1-adamanylcabonyloxy)pyridin-2-thione [Barton reaction: Barton et al. *Tetrahedron Lett* **35** 6057 1994, DOI: 10.1016/0040-4039(94)88074-3]. The thiol is generally purified by drying over P₂O₅, dissolving in Skellysolve B (see Aliphatic Compounds, Chapter 3) or *n*-hexane, passing through a column of Alumina in *n*-hexane, and eluting with *n*-hexane/EtOAc (97:3). Store it dry under N₂ or argon. The IR (CCl₄) has ν_{\max} at 2190, 2850, 1450 and 2565 (S-H) cm⁻¹, the ¹H NMR (CCl₄) has τ at 8.06 (s, 9, β and γ H), 8.29 (s, 6, δ H) and 8.57 (s, 1 S-H); and the MS has *m/z* (peak height) 168 (14, M⁺), 136 (12), 135 (100), 93 (15), 79 (17). The *S*-2,4-dinitrophenylsulfenyl derivative gives yellow needles from MeOH with **m 159-160°**. [Tanner & Brownlee *J Canad Chem* **51** 3366 1973, DOI: 10.1139/v73-502]. The *S*-CH₂CO₂Et derivative (from NaNH₂/BrCH₂CO₂Et) has **b 117-118°/0.001mm**, and on hydrolysis (NaOH/EtOH, 5 hours) gave α -(adamant-1-ylthio)acetic acid **m 68-70°** (hexane/pentane). Similarly obtained are α -(adamant-1-ylthio)propionic acid **m 142-144°** (cyclohexane), α -(adamant-1-ylthio)butyric acid **m 113-114°**, α -(adamant-1-ylthio)isovaleric acid **m 134-145°** (cyclohexane), α -(adamant-1-ylthio)caproic acid **m 74-76°**, α -(adamant-1-ylthio)- α -phenylacetic acid **m 122-124°** (cyclohexane/hexane) as well as other acids that are reported [Geigy A.-G. Belg. Pat 629,370 1963, *Chem Abstr* **60** 9167 1963]. 1-Adamantane-thioether is a good novel sialoside protecting group which is also used for coupling with adequately protected sugars [Crich & Li *J Org Chem* **72** 7794 2007, DOI: 10.1021/jo7012912]. 1-Adamantanethiolate molecules have been used in a new patterning technique (microdisplacement printing) to form self-assembled mono-layers (SAMs) on an Au(111) surface for contact printing. The attached thiolate molecular film hinders lateral surface diffusion of the patterning molecules allowing the use of such molecules which are too mobile to pattern by other methods. It is used for SAMs Cat 659452-5G molecular spacer in nanotechnology [*J Org Chem* **72** 7794 2007, DOI: 10.1021/jo7012912; Dameron et al. *Nano Lett* **5** 1834 2005, DOI: 10.1021/nl050981j].

Aerosol-OT [AOT, DOSS, sodium bis(ethylhexyl)sulfosuccinate, sodium ducosate] [577-11-7] C₂₀H₃₇O₇SNa, M 444.6, m 173-179°. The surfactant can be prepared by stirring vigorously di(2-ethylhexyl)malate (1mol, obtained from maleic anhydride and 2-ethylhexyl alcohol) and sodium bisulfite (1.05mols) with H₂O (100 parts to ~250 parts of ester) and heating (preferably under pressure to avoid loss of SO₂) until a homogenous solution is formed. The moisture is removed *in vacuo*, and any residual salts are removed by dissolving in organic solvents such as toluene or petroleum ether, filtering and evaporating to dryness. For variations of this method see A.O. Jaeger [US Patents 2028091 1936 and 2176423 1939 to American Cyanamid & Chem Corp]. For the preparation of reverse micelles of semiconductor nanoparticles, AOT is purified and degassed by dissolving the solid in petroleum ether (b 35-60°), filtering and evaporating to dryness [Stiegerwald et al. *J Am Chem Soc* **110** 3046 1988, DOI: 10.1021/ja00218a008]. AOT is a waxy colourless solid which is stable at pH 7 and below, but hydrolyses slowly at pH >8. It is soluble in CCl₄, light petroleum, xylene, dibutyl phthalate, EtOH, Me₂CO and plant oils. Its solubilities in H₂O (w/v%) are 1.5 (25°), 2.3 (40°), 3.0 (50°) and 5.5 (70°). It is also used as a surfactant wetting agent, in tablet formulations and as a laxative. The *potassium salt* [7491-07-0] C₂₀H₃₇O₇SK, **M 460.7**, is prepared in a similar way and has similar properties. [*Caryl Ind En Chem* **33** 731 1941, DOI: 10.1021/ie50378a011; *Beilstein* **4** III 71; *Fieser* **15** 149.]

Cadmium bis(octanethiol) C₁₆H₃₄S₂Cd, M 398.9. A precursor for making quantum dots, is prepared by addition of 35% aqueous NH₄OH (~10-11ml) to a solution of CdCl₂ (1.83g, 10mmol) in 1:1 v/v EtOH/H₂O (100ml) until the white suspension dissolved to give a clear solution. To this solution is added 1-octanethiol (3.47ml, 20mmol) dropwise when a white precipitate forms, and the mixture is stirred at ~25 for 2hrs. The solid is collected by centrifugation (4000rpm/10min), resuspended in 1:1 v/v EtOH/H₂O (100ml), centrifuged (4000rpm/10min) again, and this washing is repeated twice. The white powder is then dried in air overnight (85% yield, analytically pure for C₁₆H₃₄S₂Cd). Its FTIR (KBr) has ν at 2955 (C-H₃ asym str), 2872 (C-H₃ sym str), 2849 (-CH₂- sym str), 1466 (-CH₂- deform), 1738 (C-H₃ deform), 722 (-CH₂- rock) and 647 (-CH-deform) cm⁻¹. Because of its insolubility the ¹³C (¹H) CPMAS solid state spectrum was measured and gave a duplicate set of 8 signals attributed to a mixture of crystalline and amorphous forms. [Bansal et al. *J Phys Chem C* **120** 1871 2016, DOI: 10.1021/acs.jpcc.5b09109].

Dimethylcadmium [506-82-1] C₂H₆Cd, M 142.5, f.p. -2.4°(corrected), -4.5°, b 105.5°/758mm, 105.7°/760mm (log *p* versus 1/*T* does not give a straight line), n_D^{25} 1.5488. Prepared by siphoning an ethereal solution of MeMgI under N₂ into a flask at -10° with a hopper (design provided by Anderson and Taylor, see reference below) containing dried powdered anhydrous CdCl₂ which was added at a rate (at ~5min intervals) which

allowed gentle reflux. At the end of the reaction the supernatant Et₂O was siphoned off, the residue was washed with dry Et₂O which was removed also by siphoning. The combined Et₂O was siphoned into a fractionation unit (design also provided by Anderson and Taylor, see reference below), and purified by fractionation also under N₂. Extreme precaution should be exercised as the whole system is **highly flammable**. It has a disagreeable odour, highly flammable and should be stored in sealed tubes at <0°. It ignites if dropped on filter paper and can explode spontaneously if heated above 90° in air. It forms dense white fumes when exposed to moist air. Its physiological effects are serious as it causes irritation of the eyes on exposure to the vapour, with distortion of vision which can last a few hours and the eyes can see halos to bright light that may last a day or two. When using CdMe₂ for chemical reactions it is best to use the siphoned Et₂O solution, determine its concentration, and use the required volume without isolating it. [Gilman & Nelson *Recl Trav Chim Pays-Bas* **55** 518 1936, DOI: 10.1002/recl.19360550607; for preparation (including design of equipment) and physical constants see Anderson & Taylor *J Phys Chem* **56** 161 1952, DOI: 10.1021/j150494a001; and for preparation from MeMgBr and CdBr₂ see Krause *Chem Ber* **50** 1813 1917, DOI: 10.1002/cber.19170500292; for the determination of Cd concentration see Vogel's *Textbook of Quantitative Chemical Analysis* by Jeffrey, Bassett, Mendham and Denney Eds, 5th Edition, pp 444 and **541** 1989, Longman Scientific & Technical Ltd UK/John Wiley & Sons Inc, NY, ISBN 0-582-44693-7.]

6-Mercaptohexylferrocene [134029-92-8] C₁₆H₂₂SFe, **M 302.3, b 321-353°**. This thiol is prepared in four steps from ferrocene. Thus a mixture of 6-bromohexanoyl chloride (3.2g, 15mmol, see [22809-37-6]) and anhydrous AlCl₃ (2.0g, 15mmol) in cold (0°) CH₂Cl₂ (40ml), which is stirred (20 minutes) under argon, is added dropwise with stirring to ferrocene (2.79g, 15mmol) in CH₂Cl₂ (40ml) over a period of 20 minutes when the colour of the solution turns purple. After stirring for 2 hours, the stirring rate is increase while H₂O (20ml) is added slowly, and the solvent boils gently. The organic layer is collected after 10 minutes, washed with H₂O until the washings are neutral, dried (MgSO₄), filtered, concentrated *in vacuo*, and the residue is subjected to chromatography on silica-gel (230-400 mesh) with hexane/EtOAc (100:1) as eluent. The second orange band (the first is ferrocene) is collected and gives **6-(ferrocenylcarbonyl)pentyl bromide** (4.5g, 85%) on evaporation. A part of it (1.61g, 7.4mmol) and Zn/Hg amalgam (freshly prepared from 7.4g of granulated Zn and 0.5g of HgCl₂) in H₂O (25ml), 12M HCl (20ml) and toluene (30ml) are stirred vigorously so that the two liquid phases and the solids are in close contact, and are heated under reflux for 16 hours during which time 12M HCl (2 x 5ml) is added. The mixture is cooled to ~25°, the organic layer is collected washed with H₂O, dried (MgSO₄), filtered, concentrated, and the residue is purified through a silica-gel column as before. The yellow-brown band provides the required **6-ferrocenylhexyl bromide** (2.33g, 90%). The bromide (1.11g, 3.2mmol) and thiourea (0.24g, 3.5mmol) in absolute EtOH (20ml) is then boiled under reflux in an atmosphere of argon. After boiling for 18 hours, the solvent is removed *in vacuo*, treated with aqueous KOH (0.20g, 3.5mmol, in 20ml of H₂O) and boiled under reflux in an atmosphere of argon for 2 hours. The orange-yellow oil that settles on cooling is extracted into Et₂O (3 x 50ml), the combined extracts are dried (MgSO₄), concentrated, and chromatographed as before. The first yellow band gives the desired **6-ferrocenylhexanethiol** (0.68g, 62%) on evaporation. It is identified by its ¹H NMR (300 MHz, CDCl₃) which has δ at 1.35 (t, 1H, **SH**), 1.39 (br, 4H, FcCH₂CH₂CH₂), 1.52 (q, 2H, CH₂CH₂CH₂SH), 1.63 (q, 2H, CH₂CH₂SH), 2.34 (t, 2H, FcCH₂), 2.53 (q, 2H, CH₂SH), 4.06 (t, 2H, C₅H₄), 4.07 (t, 2H, C₅H₄), 4.11 (s, 5H, C₅H₅); and EI-MS *m/z* at 302 (M⁺). The second yellow band contains the corresponding **disulfide** which can be reduced and used. [Creager & Rowe *J Electroanal Chem (Lausanne Switz)* **370** 203 1994, DOI: 10.1016/0022-0728(93)03173-M; cf Yu et al. *J Org Chem* **66** 2937 2001, DOI: 10.1021/jo001283g.] This and other related ferrocenyl-alkane thiols have been synthesised and characterised electrochemically in solution and in self-assembled monolayer (SAM) films on gold electrodes. The affinities of these on the electrode surface have been described in detail. These redox active thiols can be used for selective electrochemical sensing and preparing a new type of thiol self-assembled sensing devices (SAMs). [Creager & Rowe *J Electroanal Chem (Lausanne Switz)* **370** 203 1994, DOI: 10.1016/0022-0728(93)03173-M.]

Gold nanoparticles covered fully with 6-ferrocenylhexanethiolate ligands, with average composition Au₂₂₅(6-ferrocenylhexanethiolate)₄₂, were prepared, and had a unique combination of adsorption properties on Pt electrodes which were studied comprehensively. The adsorbed layers are so robust that sub, mono and multi nanoparticulate layers on the electrodes can be transferred to fresh electrolyte solutions in which they show stable ferrocene voltametry over long time periods. [Stiles et al. *J Am Chem Soc* **130** 1856 2008, DOI: 10.1021/ja074161f.]

NPs for drug delivery to tissues and particularly to brain, i.e. traverse the blood-brain-barrier, include conjugates of dextran, agarose, chitosan polyacetic acid (PLA), poly(lactic-co-glycolytic acid) (PLGA) and NPs with lipophilic moieties, e.g. polyethylene glycol (PEG) to which drugs can be bound have been explored by Fresta and co-workers [Celia et al. *Med Res Rev* **31** (3) 716-756 2011, DOI: 10.1002/med.20201], and shown to be useful devices.

Phenyl trimethylsilyl selenide [33861-17-5] $C_9H_{14}SeSi$, **M 229.3**, **b 93-95°/9-10mm, 92-93°/5mm**. Prepared under positive pressure of N_2 , a solution of diphenyl diselenide (15.7g, 50mmol; Reich et al. *Org Synth* **59** 141 1979, Coll Vol **6** 533 1979, DOI: 10.15227/orgsyn.059.0141) in THF (50ml) is treated with finely cut Na (2.76g, 120mmol) at $\sim 25^\circ$, heated under gentle reflux until the metal dissolves, cooled to $\sim 25^\circ$, then trimethylsilyl chloride (15.2ml, 120mmol) is added slowly to the brown mixture with stirring for 1hr at $\sim 25^\circ$ (orange-yellow solution) followed by refluxing for 30min whereby a copious white precipitate of NaCl is formed. The mixture (containing the NaCl) is evaporated to dryness, and distilled directly *in vacuo* with vigorous stirring to give a colourless liquid which was redistilled to give **phenyl trimethylsilyl selenide** (16.6g, 72%). It has IR (film): ν_{max} 1250, 840 and 739 cm^{-1} , 1H NMR (CCl_4) with δ at 0.34 [s, 9H, $(CH_3)_3Si$], 7.18-7.42 [m, 5H, SeC_6H_5] ppm, ^{13}C NMR ($CDCl_3$) with δ at 1.49 ($(CH_3)_3Si$) and MS (70eV) has $m/e = 230$ (M^+), 215 ($M^+ - CH_3$), 73 [$M^+ - (CH_3)_3Si$]. [Miyoshi et al. *Synthesis* 300 1979, DOI: 10.1055/s-1979-28659]. It is soluble in C_6H_6 , Et_2O , THF and MeCN, but is rather sensitive to moisture and should be handled *via* a syringe using Schlenk-type techniques (see Chapter 1). It can be stored indefinitely under dry argon or N_2 and all reactions should be carried out in efficient fume hoods. [See also Ogawa e-EROS *Encyclopedia of Reagents for Organic Synthesis* published on line on 15 April 2001, DOI: 10.1002/047084289X.rp145.]

Tetra-*n*-propylammonium perruthenate (TPAP, tetrapropyl tetraoxoruthenate) [114615-82-6] $C_{16}H_{28}NO_4Ru$, **M 351.4**, **m 160°(dec)**. This oxidant (at 5 mol%) acts catalytically in oxidising 2-substituted *N*-1, *N*-3-dihydroxy-imidazolidines (1mol) in CH_2Cl_2 at room temperature in 1-12 hours in the presence of *N*-methylmorpholine *N*-oxide (1.1mol) to provide the respective 2-substituted imidazoline-1-nitronyl-3-nitroxide radicals (NNRs) in 44-90% yields. NNRs have found applications in Physics, Chemistry and Nanotechnology for the development of **organic molecular magnets** [Palacio et al. *Phys Rev Lett* **79** 2336 1997, <http://dx.doi.org/10.1103/PhysRevLett.79.2336>] and **single chain magnets** [Caneschi et al. *Angew Chem Int Ed* **40** 1760 2001, DOI: 10.1002/1521-3773(20010504)40:9<1760>]. Gorini et al. *Synlett* 948 2006, DOI: 10.1055/s-2006-939045; for further reading see TPAP entries in Chapter 4, 'Metal-Organic Compounds'; and in Chapter 5, 'Catalysis-Part 1', and for applications see Fieser **14** 302, **16** 325.]

SAFETY ISSUES

Materials of nanometer size have been present on earth almost since its formation ~ 4.6 billion (eon) years ago, as well as in the universe. Recently J. Cami, and coworkers have discovered large clouds in the planetary nebula Tc1 in which C_{60} and C_{70} fullerenes were identified by infra red spectroscopy [*Science* **329** (Issue 5996) 1180 2010, DOI: 10.1126/science.1192035, reviewed by A. Extnance RSC *Chemistry World* **7** (9) 21 2010, and L. Howes RSC *Chemistry World* **8** (1) 34 2011.] During the past 30 years or so the development of nanomaterials and their wide ranging applications have avalanched, and progress is continuing. The safety aspects of nanomaterials have been of concern for many years, and it is generally accepted that conscious efforts have to be made to protect the individual workers in the field, the general public and the environment. Research on safety aspects is slowly gaining pace, and it has already been shown in mice that carbon nanotubes (CNTs) exhibit asbestos-like pathogenicity when inhaled. [K. Donaldson and coworkers: Poland et al. *Nature Nanotechnol* **3** 423 2008, DOI: 10.1038/nnano.2008.111; J.C. Bonner and coworkers: Ryman-Rasmussen et al. *Nature Nanotechnology* **4** 747 2009, DOI: 10.1038/nnano.2009.305; Reviewed by H. Birch RSC *Chemistry World* **6** (12) 24 2009.] The movement of nanomaterials from lungs to lymph nodes to the blood stream and clearance through the kidneys has been demonstrated by J.V. Frangioni and A. Tsuda and their coworkers [Choi et al. *Nature Biotechnol* **28** 1300 2010, DOI: 10.1038/nbt.1696; reviewed by L. Howes RSC *Chemistry World* **7** (12) 24 2010.] More recently W. Wohlleben and coworkers have subjected two thermoplastic materials and two cement materials infused with different nanoparticles (i.e. nanotubes and nanoparticle in polymer filters and in inorganic matrices) to gentle abrasion, high speed sanding and UV radiation. They did not find significant release of nanopowders into the atmosphere in which the sizes of the nanoparticles were very different from

those in control powders. Moreover, rats exposed to the dust fared no worse than those exposed to dust that contained nanocomponents. However, further studies are in progress. [Wohlleben et al. *Small* **7** (16) 2384 2011, DOI: 10: 1002/sml.201002054; reported by K. McAlpine *Chemistry World* **8** (8) 28 2011.]

Safety was rather slack in the early days, but it is being realised that nanomaterials should be treated differently from hazardous chemicals due to their unknown long-term effects. Nano-enabled hoods and nano-enabled-gear for personnel are being used, but only to a small extent. Safety protocols have been written and updated, but nano-regulation is only slowly creeping in [see special report by V. Gill RSC *Chemistry World* **6** (4) 10 2009.] A comprehensive report on nanosafety practices in research laboratories worldwide was compiled by J. Santamaria and coworkers recently, where they detail an up-to-date account of the present situation regarding to safety issues. [Balas et al. *Nature Nanotechnol* **5** 93 2010, DOI: 10.1038/nnano.2010.1; Reviewed by K. McAlpine RSC *Chemistry World* **7** (3) 15 2010.]

See also B. Tomasik, Fundamental Research Institute, http://foundational-research.org/possible_ways_to_promote_compromise, updated Feb 2016; E. Drexler, The Foresight Institute, Background 3, Rev 1 1988 www.foresight.org.a_dialog_on_dangers; Center for Responsible Nanotechnology: www.crnano.org.dangers_of_molecular_manufacturing; www.crnano.org.the_need_for_international_control; www.crnano.org.technical_restrictions_may_make_nanotechnology_safer.

Advice in this regard is that in the manipulation and disposal of nanomaterials the same guidelines for asbestos handling should be used. In addition, further precautions should be exercised which should consider the known toxicity and the chemical nature of the nanomaterials in use.

The following is a list of more recent books that were published on legal, personal, industrial, and environmental safety aspects of nanotechnology:

L.L. Bergeson (Ed), *Nanotechnology: Environmental Law, Policy and Business Considerations*, American Bar Association, ABA Publishing USA, 2010. ISBN 9781604225826.

J-Y. Botero, *Environmental Nanotechnology*, McGraw Hill, 2007. ISBN 97800714777505, 0071477500.

R.E. Hester and R.M. Harrison (Eds), *Nanotechnology: Consequences for Human Health and the Environment*, RSC Publ. 2009. ISBN 9781847559562, 1847559565.

C. Kumar (Ed), *Nanomaterials: Toxicity (Health and Environment Issues)*, WILEY-VCH, Weinheim, 2006. ISBN 9783527313853, 3527313850.

J.R. Lead and E. Smith (Eds), *Environmental and Human Aspects of Nanotechnology*, J. Wiley & Sons, NY, 2009. ISBN 9781405176347.

I. Linkov and J. Steevens, *Nanomaterials: Risks and Benefits*, Springer, 2008. ISBN 9781402094903 (PB), 9781402094907 (HB), 9781402094910 (e-book).

N.A. Montiero-Riviere and C.L. Tran, *Nanotoxicology: Characterisation, Dosing and Health*, Informa Healthcare Publ, 2007. ISBN 9781420024147, 1420045148.

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B. Bhushan (Ed), *Handbook of Nanotechnology 2nd edn*, Springer, Berlin, 2007. ISBN 10: 354029855-x, 13: 9783540298557.

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M.S. Dresselhaus, G. Dresselhaus and P.C. Eklund, *Science of Fullerenes and Carbon Nanotubes*, Academic Press, San Diego, 1996. ISBN: 0-12-221820-5.

J.L. Feather and M.F. Azner, *Nanotechnology: Education and Workforce Development*, CRC Press, Boca Raton, FL, 2011. ISBN 9781420053944.

B.D. Feldman, *Materials Chemistry*, Springer, Dordrecht, 2008. ISBN 978-1-4020-6119-6, 978-1-4020-6120-2 (e-book).

J.H. Grassian, *Nanoscience and Nanotechnology*, J. Wiley & Sons, 2008. ISBN 978047081037.

J.A. Helsen and Y. Missirlis, *Biomaterials: Biological and Medical Physics, Biomedical Engineering*, Springer Verlag, 2010. ISBN: 9783642125317, 9783642125324 (e-book).

C. Hess and R. Schlögl (Eds), *Nanostructured Catalysts*, RSC Publ. Nanoscience & Nanotechnology, 2011. ISBN: 9780854041866.

- G.L. Hornyak and H.F. Tibbals, *Fundamentals of Nanotechnology*, CRC Press, Boca Raton, FL, 2009. ISBN: 978140048032, 1420048031.
- G.L. Hornyak, H.F. Tibbals and J. Dutta, *Introduction to Nanoscience and Nanotechnology*, CRC Press, Boca Raton, FL, 2009. ISBN: 9781420047790, 1420047795.
- M.S. Johal, *Understanding Nanomaterials*, CRC Press, Boca Raton, FL, 2011. ISBN: 9781420073102.
- B.J. Kirby, *Micro- and Nanoscale Fluid Mechanics: Transport in Microfluidic Devices*. Cambridge University Press 2010. ISBN: 978-0-521-11903-0.
- V. Klimov, ed. *Semiconductor and Metal Nanocrystals Synthesis, Electronic, and Optical Properties* Marcel Dekker, NY 2003-2004. Illustrated print ISBN: 082474716x.
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CAS Registry Numbers Index

HOW TO VALIDATE CHEMICAL ABSTRACTS SERVICE REGISTRY NUMBERS

Almost all chemical and biochemical entries in this book have CAS (Chemical Abstract Service) Registry Numbers to identify them and have been entered for each substance. The registry numbers are an efficient way for identifying substances and are independent of the nomenclature of substances. Unlike chemical names which may have more than one synonymous name, there is only one CAS Registry Number for each substance (with only a few exceptions, e.g. where a substance may have another number before purification, or before determination of absolute configuration).

The registry numbers are made up of three sets of digits separated by two hyphens with the general form: $nN_n + \dots + 4N_4 + 3N_3 + 2N_2 + 1N_1 - R$ where the N s are sequential numbers and R is the *check digit*. The first set can be up to eight digits (at this point in time, but not less than two), and the second is made of two digits (zeros are included as a digits) and the third is a single digit which *checks* the validity of the sequential numbers, e.g. 379669-72-4 for *but-3-enylboronic acid* or 83-32-9 for *acenaphthene*. The validity of a registry number can be checked by the following formula (computer generated):

$$\frac{nN_n + \dots + 4N_4 + 3N_3 + 2N_2 + 1N_1}{10} = Q + R/10$$

where the N s include all numbers except the last *check* digit, Q represents an integer (fractions not included) which, although it is disregarded, does give the value of R equal to the *check digit* in a **valid** Registry Number.

The following are three examples demonstrate the application of the formula and the validity of the Registry Number:

1. 1,5-Dihydroxynaphthalene 83-56-7

$$\frac{(4 \times 8) + (3 \times 3) + (2 \times 5) + (1 \times 6)}{10} = \frac{57}{10} = 5 + \frac{7}{10} \quad [\text{Note: the last (check) digit of the CAS number 7 is not included.}]$$

When the value of Q is made to equal 5, the *check* digit R becomes equal to 7; thus the Registry Number is **valid**. In this case the value of Q is in fact the digit(s) (5), before the last *check* digit (7).

2. 4,5-Diamino-6-hydroxypyrimidine hemisulfate 102783-18-6

$$\frac{(8 \times 1) + (7 \times 0) + (6 \times 2) + (5 \times 7) + (4 \times 8) + (3 \times 3) + (2 \times 1) + (1 \times 8)}{10} = \frac{106}{10} = 10 + \frac{6}{10}$$

When the value of Q is made to equal 10, the *check* digit R becomes equal to 6; thus the Registry Number is **valid**. In this case the value of Q is in fact the digit(s) (10), before the last *check* digit (6).

3. 1,4-Bis(pentafluorobenzyl)[C₆₀]fullerene 1260376-31-5

$$\begin{array}{rclclcl}
 (9 \times \mathbf{1}) + (8 \times \mathbf{2}) + (7 \times \mathbf{6}) + (6 \times \mathbf{0}) + (5 \times \mathbf{3}) + (4 \times \mathbf{7}) + (3 \times \mathbf{6}) + (2 \times \mathbf{3}) + (1 \times \mathbf{1}) & = & \mathbf{135} & = & \mathbf{13} & + & \mathbf{5} \\
 \hline
 & & 10 & & 10 & & 10
 \end{array}$$

When the value of *Q* is made to 13, the check digit *R* becomes equal to 5; thus the Registry Number is **valid**. In this case the value of *Q* is in fact the digit(s) (**13**), before the last *check* digit (**5**).

The CAS Registry Handbook (Number Section) and supplements should be consulted for obtaining the substance name and elemental formula corresponding to a Registry Number.

To facilitate the method for locating the purification of a substance, a CAS Registry Number Index of the numbers of the entries with their respective page numbers is included here. This will also provide the reader with a rapid way to see if the purification of a particular substance has been reported in the book.

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